

Level C Data Package Deliverables

Perchlorate



Applied P & Ch Laboratory

Applied P & Ch Laboratory
Wet Analysis Results for Method 314.0

Client Name: GEOFON, Inc.
Project ID: JPL

Project No: 04-4428.10
Service ID: 33391

Anal. Method 314.0
Collected by:

Component Name: Perchlorate
CAS No:

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/29/03	03W3074	µg/L	12	147	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/29/03	03W3074	µg/L	200	1810	
03W3074-MB-01	03W3074-MB-01	Water	05/29/03	05/29/03	05/29/03	03W3074	µg/L	4	<4	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

FORM-3

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 314.0

Client Name: GEOFON, Inc. Contract No: Lab Code: APCL
Case No: SAS No: Service ID: 33391
Project ID: JPL Project No: 04-4428.10 Sample Matrix: Water
Batch No: 03W3074
LCS Filename: - Date Analyzed: 053003 Time Analyzed: 11:43
LCSD Filename: - Date Analyzed: - Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
PERCHLORATE	µg/L	25	0	24.3	97	80-120
# of Out-of-control					0	

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 314.0

Client Name: GEOFON, Inc.	Contract No:	Lab Code: APCL
Case No:	SAS No:	Service ID: 33391
Project ID: JPL	Project No: 04-4428.10	Sample Matrix: Water
	Batch No: 03W3074	
MS Filename: -	Date Analyzed: 053003	Time Analyzed: 16:00
MSD Filename: -	Date Analyzed: 053003	Time Analyzed: 16:19
MS Sample No: MW-7	Sample Lab ID: 03-3444-2	

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
PERCHLORATE	µg/L	50	2.3	51.9	99	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
PERCHLORATE	µg/L	50	53.5	102	3	20	75-125
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-7
Applied P & Ch Laboratory
CCV Recovery for Wet Analysis

Client Name: GEOFON, Inc.

Contract No.:

Lab Code: APCL

Case No:

SAS No.:

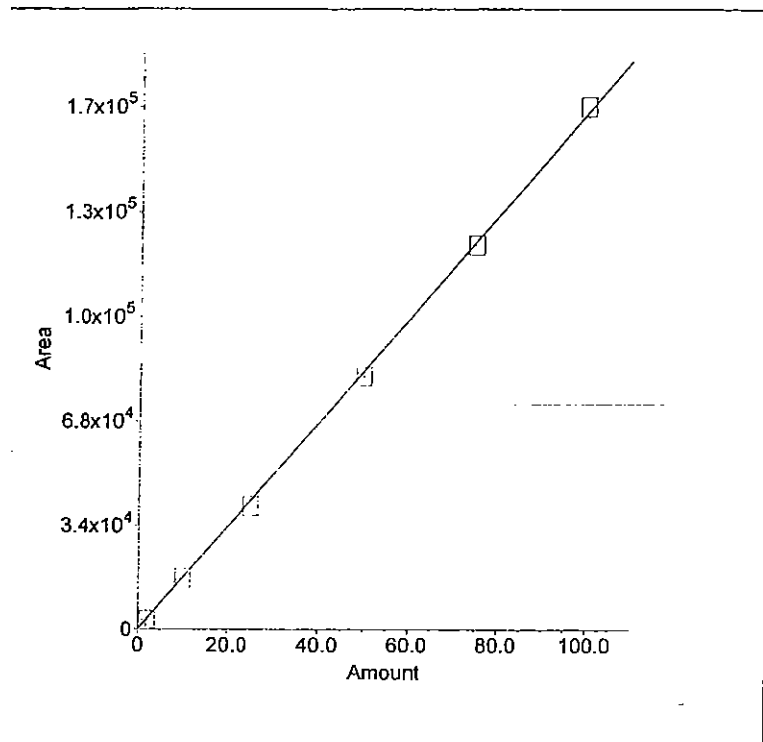
Service ID: 33391

Project ID: JPL

Project No.: 04-4428.10

#	Component Name	Method	Batch No.	Unit	Expected	Test Result	Rec. %	Dev. %	Flag	Control Limit, %	Test Date
1	Perchlorate	314.0	03W3074	µg/L	50	49.7	99	-1	✓	85-115	05/29/2003
	Perchlorate	314.0	03W3074	µg/L	50	52.5	105	5	✓	85-115	05/29/2003
	Perchlorate	314.0	03W3074	µg/L	50	50.8	102	2	✓	85-115	05/29/2003
	Perchlorate	314.0	03W3074	µg/L	50	49.9	100	0	✓	85-115	05/30/2003

1. Component: perchlorate
Standard: External Fit Type: Linear
Origin: Force Calibration: Area
 $r^2 = 0.999492$
Amt = $0.0005893 * \text{Resp} + 0$



Calibration: 7 points, 0, 2, 10, 25, 50, 75, 100 ppb

Analyst C. W
Date 03/12/03
Instrument IC-1c

Line	Sample	Sample Type	Level	Method	Data File	Volume	Dilution
1	Cal blank	Sample		e314-011.met	c:\data\314-011\mb_001.dxd	1	1
2	cal standard 2ppb W7827a	Sample		e314-011.met	c:\data\314-011\std-2pb_002.dxd	1	1
3	cal standard 4ppb W7827b	Sample		e314-011.met	c:\data\314-011\std-4pb_003.dxd	1	1
4	cal standard 10ppb W7827c	Sample		e314-011.met	c:\data\314-011\std-10pb_004.dxd	1	1
5	cal standard 25ppb W7827d	Sample		e314-011.met	c:\data\314-011\std-25pb_005.dxd	1	1
6	cal standard 50ppb W7827e	Sample		e314-011.met	c:\data\314-011\std-50pb_006.dxd	1	1
7	cal standard 75ppb W7827f	Sample		e314-011.met	c:\data\314-011\std-75pb_007.dxd	1	1
8	cal standard 100ppb W7827g	Sample		e314-011.met	c:\data\314-011\std-100pb_008.dxd	1	1
9	ICV 50 ppb w7828a	Sample		e314-011.met	c:\data\314-011\icv-50pb_009.dxd	1	1
10	icb	Sample		e314-011.met	c:\data\314-011\icb_010.dxd	1	1
11	anion 100pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-100_011.dxd	1	1
12	anion 200pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-200_012.dxd	1	1
13	anion 300pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-300_013.dxd	1	1
14	anion 400pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-400_014.dxd	1	1
15	anion 500pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-500_015.dxd	1	1
16	anion 600pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-600_016.dxd	1	1
17	anion 800pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-800_017.dxd	1	1
18	anion 1000pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-1000_018.dxd	1	1
19	anion 400pm each 2pb	Sample		e314-011.met	c:\data\314-011\ipc-2pb_019.dxd	1	1
20	anion 400pm each 4pb	Sample		e314-011.met	c:\data\314-011\ipc-4pb_020.dxd	1	1
21	anion 400pm each 25pb	Sample		e314-011.met	c:\data\314-011\ipc-25pb_021.dxd	1	1
22	ICV 50 ppb	Sample		e314-011.met	c:\data\314-011\ccv-50pb	1	1
23	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-02_023.dxd	1	1
24	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-03_024.dxd	1	1
25	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-04	1	1
26	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-05	1	1
27	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-06	1	1
28	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-07	1	1
29	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-08	1	1
30	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
31	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
32	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
33	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
34	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
35	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
36	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
37	MCT anion 800pm each, 25pbCLO4	Sample		e314-011.met	c:\data\314-011\ipc-25pb	1	1
38	MCT anion 800pm each, 25pbCLO4	Sample		e314-011.met	c:\data\314-011\ipc-25pb	1	1
39	MCT anion 800pm each, 4pbCLO4	Sample		e314-011.met	c:\data\314-011\ipc-4pb	1	1
40	MCT anion 800pm each, 4pbCLO4	Sample		e314-011.met	c:\data\314-011\ipc-4pb	1	1
41	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s01	1	5
42	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s02	1	5
43	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s03	1	5
44	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s04	1	5
45	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s05	1	5
46	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s06	1	5
47	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s07	1	5
48	standard 25ppb W7827d	Sample		e314-011.met	c:\data\314-011\std-25pb	1	1
49	anion 100pm each,4pb CLO4	Sample		e314-011.met	c:\data\314-011\am-100-4pb	1	1
50	anion 200pm each ,4pb CLO4	Sample		e314-011.met	c:\data\314-011\am-200-4pb	1	1
51	anion 300pm each ,4pb CLO4	Sample		e314-011.met	c:\data\314-011\am-300-4pb	1	1
52	anion 100pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\am-100-2pb	1	1
53	anion 200pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\am-200-2pb	1	1
54	anion 300pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\am-300-2pb	1	1
55	1982-01 B S.C 4450us/cm	Sample		e314-011.met	c:\data\314-011\1982-01	1	1
56	1982-01 B S.C 4450us/cm	Sample		e314-011.met	c:\data\314-011\1982-01	1	2
57	1982-02 f=10	Sample		e314-011.met	c:\data\314-011\1982-02_057.dxd	1	10
58		Sample		aastopcl.met		1	1

Line	Sample	Sample Type	Level	Method	Data File	Volume	Dilution	Weight
1	#03w3074kw ipc 25ppb w7759	Sample		e314-011.met	w3074k ipc 25ppb	1	1	1
2	ccv 50ppb w7827e	Sample		e314-011.met	w3074k q01	1	1	1
3	ccb	Sample		e314-011.met	w3074k ccb01	1	1	1
4	lcs 25ppb w7827d	Sample		e314-011.met	w3074k l01	1	1	1
5	LCS 18PPB W7685D	Sample		e314-011.met	w3074k j01	1	1	1
6	ICCS 4ppb w7827b	Sample		e314-011.met	w3074k iccs 4ppb	1	1	1
7	mb	Sample		e314-011.met	w3074k k01	1	1	1
8	3381-01 F=2	Sample		e314-011.met	3381-01	1	2	1
9	3381-06 f=3	Sample		e314-011.met	3381-06	1	3	1
0	3381-07 F=2	Sample		e314-011.met	3381-07	1	2	1
1	3391-01 f=1	Sample		e314-011.met	3391-01	1	1	1
2	3391-02 f=1	Sample		e314-011.met	3391-02	1	1	1
3	ccv 50ppb w7827e	Sample		e314-011.met	w3074k q02	1	1	1
4	ccb	Sample		e314-011.met	w3074k k02	1	1	1
5	3414-01 F=1	Sample		e314-011.met	3414-01	1	1	1
6	3414-02 F=1	Sample		e314-011.met	3414-02_016.dxd	1	1	1
7	3391-01 MD F=3	Sample		e314-011.met	w3074k d01_017.dxd	1	3	1
8	3414-02 MS50PPB F=1 W7428B	Sample		e314-011.met	w3074k m01_018.dxd	1	1	1
9	3414-02 MSD50PPB F=1 W7428B	Sample		e314-011.met	w3074k n01_019.dxd	1	1	1
10	3391-02 F=50	Sample		e314-011.met	3391-02a_020.dxd	1	50	1
11	ccv 50ppb w7827e	Sample		e314-011.met	w3074k q03_021.dxd	1	1	1
12	CCV 50PPB W7827E	Sample		e314-011.met	w3074k q04	1	1	1
13	CCB	Sample		e314-011.met	w3074k k03	1	1	1
14	3444-01 F=1	Sample		e314-011.met	3444-01	1	1	1
15	3444-02 F=1	Sample		e314-011.met	3444-02	1	1	1
16	3444-02 F=100	Sample		e314-011.met	3444-02a	1	100	1
17	3444-03 F=1	Sample		e314-011.met	3444-03a	1	1	1
18	3444-01 MS50PPB F=1 W7428B	Sample		e314-011.met	w3074k m02	1	1	1
19	3444-01 MSD50PPB F=1 W7428B	Sample		e314-011.met	w3074k n02	1	1	1
20	3444-02 MS50PPB F=100 W7428B	Sample		e314-011.met	w3074k m03	1	100	1
21	3444-02 MSD250PPB F=100 W7428B	Sample		e314-011.met	w3074k n03_031.dxd	1	100	1
22	CCV 50PPB W7827E	Sample		e314-011.met	w3074k q05_032.dxd	1	1	1
23	CCB	Sample		e314-011.met	w3074k k04	1	1	1
24	3465-01 F=1	Sample		e314-011.met	3465-01	1	1	1
25	3465-02 F=1	Sample		e314-011.met	3465-02	1	1	1
26	3465-03 F=1	Sample		e314-011.met	3465-03	1	1	1
27	CCV 50PPB W7827E	Sample		e314-011.met	w3074k q06	1	1	1

Analyst Wm Wly
Date 5/29-30/03
Instrument Zc-k

Applied P & Ch Laboratory
 1 Magnolia Ave. Chino CA 91710
 (909) 590-1828 Fax: (909) 590-1498

Conductance (120.1) Worksheet

083

for reference
 Matrix: W
 Constant _____ Calibration STD: 0.0100M KCl

Test Date: 5/28/03 Analyst: mw

SOP: G-45

	Sample ID	Treatment $V/X=f_0$	Dilution $V_f/V_i=f_1$	Temperature $T, ^\circ C$	$C_{25} = C_T f_1 / \{1 - 0.0191(25 - T)\}$		$\rho = 1/C_{25}$ MΩ cm	Note & Anomaly
					$C_T, \mu mhos/cm$	$C_{25}, \mu mhos/cm$		
al.	Lot #:	-	-	25°C				$C_{25} = 1,413 \pm 20$
MB		/ =	/ =					for C ₂₅
1	3381-1	5/28/03	/ =			8160		
2	↓ -6	/ =	/ =			10000		
3	↓ -7	/ =	/ =			8940		
4	3391-1	/ =	/ =			676		
5	↓ -2	/ =	/ =			550		
6	3414-1	/ =	/ =			660		
7	↓ -2	/ =	/ =	↓		501		↓
8	3444-1	/ =	/ =	↑		1300		↓
9	↓ -2	/ =	/ =			593		
10	↓ -3	/ =	/ =	↓		597		↓
11		/ =	/ =					
12		/ =	/ =					
13		/ =	/ =					
14		/ =	/ =					
15		/ =	/ =					
16		/ =	/ =					
17		/ =	/ =					
18		/ =	/ =					
19		/ =	/ =					
20		/ =	/ =					
Dup.		/ =	/ =					

Level C Data Package Deliverables

Wet Chemistry



Applied P & Ch Laboratory

Applied P & Ch Laboratory
Wet Analysis Results for Method SM2320B

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method SM2320B
Project ID: JPL Service ID: 33391 Collected by:

Component Name: Bicarbonate
CAS No:

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	06/02/03	03W3113	mg/L	2	115	
03-3391-2	MW-16	Water	05/27/03	05/27/03	06/02/03	03W3113	mg/L	2	115	
03W3113-MB-01	03W3113-MB-01	Water	06/02/03	06/02/03	06/02/03	03W3113	mg/L	2	< 2	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method SM2320B

Client Name: GEOFON, Inc.
Project ID: JPL

Project No: 04-4428.10
Service ID: 33391

Anal. Method SM2320B
Collected by:

Component Name: Carbonate
CAS No:

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	06/02/03	03W3113	mg-CaCO ₃ /L	2	<2	U
03-3391-2	MW-16	Water	05/27/03	05/27/03	06/02/03	03W3113	mg-CaCO ₃ /L	2	<2	U
03W3113-MB-01	03W3113-MB-01	Water	06/02/03	06/02/03	06/02/03	03W3113	mg-CaCO ₃ /L	2	<2	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 9040B

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 9040B
Project ID: JPL Service ID: 33391 Collected by:

Component Name: pH
CAS No: 10-29-7

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/27/03	03W3048	pH unit	0.01	7.00	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/27/03	03W3048	pH unit	0.01	7.18	
03W3048-MB-01	03W3048-MB-01	Water	05/27/03	05/27/03	05/27/03	03W3048	pH unit	0.01	6.82	

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 160.1

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 160.1
Project ID: JPL Service ID: 33391 Collected by:

Component Name: Solids, Total Dissolved (TDS)

CAS No: 10-33-3

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/29/03	03W3071	mg/L	10	529	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/29/03	03W3071	mg/L	10	347	
03W3071-MB-01	03W3071-MB-01	Water	05/29/03	05/29/03	05/29/03	03W3071	mg/L	10	<10	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 7196

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 7196
Project ID: JPL Service ID: 33391 Collected by:

Component Name: Chromium (VI)
CAS No: 1333-82-0

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/27/03	03W3046	mg/L	0.01	0.024	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/27/03	03W3046	mg/L	0.01	<0.01	U
03W3046-MB-01	03W3046-MB-01	Water	05/27/03	05/27/03	05/27/03	03W3046	mg/L	0.01	<0.01	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 300.0

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 300.0
Project ID: JPL Service ID: 33391 Collected by:

Component Name: Chloride Cl^-
CAS No: 16887-00-6

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	1.6	52.8	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	1	33.5	
03W3042-MB-01	03W3042-MB-01	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	0.2	<0.2	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 300.0

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 300.0
Project ID: JPL Service ID: 33391 Collected by:

Component Name: Nitrate as N
CAS No: 14797-55-8

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	0.32	9.4	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	0.2	9.5	
03W3042-MB-01	03W3042-MB-01	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	0.04	<0.04	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

Applied P & Ch Laboratory
Wet Analysis Results for Method 300.0

Client Name: GEOFON, Inc.
Project ID: JPL

Project No: 04-4428.10
Service ID: 33391

Anal. Method 300.0
Collected by:

Component Name: Sulfate SO_4^{--}
CAS No: 14808-79-8

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3391-1	MW-13	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	4	80.7	
03-3391-2	MW-16	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	2.5	33.7	
03W3042-MB-01	03W3042-MB-01	Water	05/27/03	05/27/03	05/27/03	03W3042	mg/L	0.5	<0.5	U

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

B - Less than RL (PQL, EQL or CRDL), but greater than MDL.

FORM-3

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method SM2320B

Client Name: GEOFON, Inc.	Contract No:	Lab Code: APCL
Case No:	SAS No:	Service ID: 33391
Project ID: JPL	Project No: 04-4428.10	Sample Matrix: Water
	Batch No: 03W3113	
LCS Filename: -	Date Analyzed: 060203	Time Analyzed: 11:45
LCSD Filename: -	Date Analyzed: 060203	Time Analyzed: 11:45

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
BICARBONATE	mg/L	100	0	101	101	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, % RPD REC
BICARBONATE	mg/L	100	99.6	100	1	20 80-120
# of Out-of-control				0	0	

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 300.0

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 33391

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W3042

LCS Filename: -

Date Analyzed: 052703

Time Analyzed: 12:12

LCSD Filename: -

Date Analyzed: 052703

Time Analyzed: 12:24

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHLORIDE CL ⁻	mg/L	4.0	0	4.06	102	80-120
NITRATE AS N	mg/L	1.5	0	1.55	103	80-120
SULFATE SO ₄ ⁻	mg/L	15	0	15.3	102	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE CL ⁻	mg/L	4.0	4.05	101	1	20	80-120
NITRATE AS N	mg/L	1.5	1.54	103	0	20	80-120
SULFATE SO ₄ ⁻	mg/L	15	15.2	101	1	25	80-120
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 300.0

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 33391

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W3042

MS Filename: -

Date Analyzed: 052703

Time Analyzed: 16:39

MSD Filename: -

Date Analyzed: 052703

Time Analyzed: 16:52

MS Sample No: AE113

Sample Lab ID: 03-3370-13

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHLORIDE CL ⁻	mg/L	8000	4420	12600	102	75-125
NITRATE AS N	mg/L	3000	46.9	3030	99	75-125
SULFATE SO ₄ ⁻	mg/L	30000	1690	31300	99	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE CL ⁻	mg/L	8000	12500	101	1	20	75-125
NITRATE AS N	mg/L	3000	2960	97	2	20	75-125
SULFATE SO ₄ ⁻	mg/L	30000	30900	97	2	25	75-125
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 160.1

Client Name: GEOFON, Inc.	Contract No:	Lab Code: APCL
Case No:	SAS No:	Service ID: 33391
Project ID: JPL	Project No: 04-4428.10	Sample Matrix: Water
	Batch No: 03W3071	
LCS Filename: -	Date Analyzed: 052903	Time Analyzed: 09:36
LCSD Filename: -	Date Analyzed: 052903	Time Analyzed: 09:36

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	Unspiked	LCS	96	88-108
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, % RPD REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	391	98	2	20 88-108
# of Out-of-control				0	0	

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 160.1

Client Name:	GEOFON, Inc.	Contract No:		Lab Code:	APCL
Case No:		SAS No:		Service ID:	33391
Project ID:	JPL	Project No:	04-4428.10	Sample Matrix:	Water
		Batch No:	03W3071		
MS Filename:	-	Date Analyzed:	052903	Time Analyzed:	09:36
MSD Filename:	-	Date Analyzed:	052903	Time Analyzed:	09:36
MS Sample No:	AE420	Sample Lab ID:	03-3389-9		

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	4140	4450	78 *	80-119
# of Out-of-control					1	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	4500	90	14	20	80-119
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 7196

Client Name: GEOFON, Inc.	Contract No:	Lab Code: APCL
Case No:	SAS No:	Service ID: 33391
Project ID: JPL	Project No: 04-4428.10	Sample Matrix: Water
	Batch No: 03W3046	
LCS Filename: -	Date Analyzed: 052703	Time Analyzed: 14:37
LCSD Filename: -	Date Analyzed: 052703	Time Analyzed: 14:37

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHROMIUM (VI)	mg/L	0.25	0	0.231	92	80-115
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHROMIUM (VI)	mg/L	0.25	0.249	100	8	19	80-115
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 7196

Client Name:	GEOFON, Inc.	Contract No:		Lab Code:	APCL
Case No:		SAS No:		Service ID:	33391
Project ID:	JPL	Project No:	04-4428.10	Sample Matrix:	Water
		Batch No:	03W3046		
MS Filename:	-	Date Analyzed:	052703	Time Analyzed:	14:37
MSD Filename:	-	Date Analyzed:	052703	Time Analyzed:	14:37
MS Sample No:	MW-13	Sample Lab ID:	03-3391-1		

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHROMIUM (VI)	mg/L	0.25	0.024	0.245	88	78-115
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHROMIUM (VI)	mg/L	0.25	0.255	92	4	19	78-115
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

6A
INITIAL CALIBRATION DATA

Lab Name: Applied P & Ch Lab Contract: _____

Analysis: Chromium (VI) Calibration Date: 01/29/2003

Concentration (mg/L)	0.000	0.0125	0.050	0.125	0.250	0.50
Absorbance	0.000	0.006	0.041	0.109	0.214	0.415

$A = 0.000 + 0.836C$

A=Absorbance

C=Concentration (mg/L)

$r = 0.9997$

Wet Chemistry QC Report B
Duplicate Results

Matrix: Water

APCL Service ID: 03-3391

Analysis	Batch ID	Analysis Date	Sample Name	Unit	Result	Duplicate Result	RPD %	RPD Control limit
Alkalinity	03W3113	06/02/2003	03-3444-1	mg-CaCO ₃ /L	284	282	1	20
PH	03W3048	05/27/2003	MW-13	PH UNITS	7.00	7.03	0	20

Note: N/A = Not applicable; NR: Not requested; NC= Not Calculated; ND: Not detected.

FORM-7
Applied P & Ch Laboratory
CCV Recovery for Wet Analysis

Client Name: GEOFON, Inc.

Contract No.:

Lab Code: APCL

Case No:

SAS No.:

Service ID: 33391

Project ID: JPL

Project No.: 04-4428.10

#	Component Name	Method	Batch No.	Unit	Expected	Test Result	Rec. %	Dev. %	Flag	Control Limit, %	Test Date
1	Chloride Cl^-	300.0	03W3042	mg/L	4.0	4.06	102	2	✓	90-110	05/27/2003
	NITRATE as N-NO_3^- , BY	300.0	03W3042	mg/L	1.5	1.54	103	3	✓	90-110	05/27/2003
	SULFATE SO_4^{--} , BY I	300.0	03W3042	mg/L	15	15.2	101	1	✓	90-110	05/27/2003
	Chloride Cl^-	300.0	03W3042	mg/L	4.0	4.06	101	1	✓	90-110	05/27/2003
	NITRATE as N-NO_3^- , BY	300.0	03W3042	mg/L	1.5	1.54	103	3	✓	90-110	05/27/2003
	SULFATE SO_4^{--} , BY I	300.0	03W3042	mg/L	15	15.2	101	1	✓	90-110	05/27/2003
	Chloride Cl^-	300.0	03W3042	mg/L	4.0	4.03	101	1	✓	90-110	05/27/2003
	NITRATE as N-NO_3^- , BY	300.0	03W3042	mg/L	1.5	1.55	103	3	✓	90-110	05/27/2003
	SULFATE SO_4^{--} , BY I	300.0	03W3042	mg/L	15	15.2	101	1	✓	90-110	05/27/2003
2	Chromium (VI)	7196	03W3046	mg/L	0.25	0.252	101	1	✓	90-110	05/27/2003
	Chromium (VI)	7196	03W3046	mg/L	0.25	0.255	102	2	✓	90-110	05/27/2003

DIONEX METHOD PARAMETERS - E300-063.MET

Method Comment: APCL EPA 300 ANALYSIS DX-100
Column ID: Dionex AS4A-SC
Analyst ID: David

System Parameters

System Name: DX-100.

Number of Detectors.....	1
Run Time (minutes).....	10.00
Sampling Rate (seconds).....	0.20

```

Detector 1 Type..... COND
Detector 1 real time plot scale maximum (uS )..... 30.000
                                         minimum..... -3.000
Detector 1 Output Equivalent to 1 Volt (in uS ) ..... 30.00
Detector 1 ACI Analog Input Connection ..... DET1
Save Data File..... Yes
Data File Name: C:\DX\DATA\03W1991\W1991001.D10

```

```
-- DETECTOR 1 PARAMETERS --
```

Report Options

```

Create ASCII Report File..... No
Print Report..... Yes
Print All Components..... Yes
Print Components Found..... No
Print Missing Components..... No
Print All Peaks..... No
Print Unknown Peaks..... No
Print Chromatogram..... Yes
Autoscale Chromatogram Maximum..... No
Autoscale Chromatogram Minimum..... No
Fill Peaks with Color ..... No
Draw Grid Lines on Chromatogram..... No
Show Component Fraction Numbers..... No
Label with Peak Number..... No
Label with Retention Times on Chromatogram..... Yes
Label with Component Name..... Yes
Format File Name: C:\DX\METHOD\DEFAULT.PRF

```

Integration Parameters

Starting Peak Width (seconds).....	10.0
Peak Threshold	2.000
Peak Area Reject.....	1000
Area Reject for Reference Peaks.....	1000

Data Events

Time	Description
0.13	Start peak detection
10.00	Stop peak detection

Calibration Parameters

Number Of Levels for Calibration.....	6
Force Calibration Curve Through Origin.....	No
Calibration Fit Type.....	Linear
Replace Or Average Calibrations.....	Replace
External or Internal Calibration.....	External
Calculate Unknowns by Area or Height.....	Area
Default Sample Volume.....	1.0
Default Dilution Factor.....	1.0
Default Response Factor for Unknown Peaks.....	0.0
Calibration Standard Volume	1.0
Internal Standard Amount in Samples	1.0
Amount Units	ppm

Component # 4 Bromide
 Reference Comp. Nitrate-N
 Retention Time 3.45
 Window Size 0.20 min.
 Amount = K0 + K1*Area
 K0 = 4.58974E-002
 K1 = 4.83279E-006

Level	Amount	Area	Height
1	7.50000E-002	13830	1488
2	1.50000E+000	298206	30100
3	3.00000E+000	591234	61776
4	6.00000E+000	1219933	128845
5	7.50000E+000	1559887	166594
6	0.00000E+000	0	0

Component # 5 Nitrate-N
 Reference Comp. Nitrate-N
 Retention Time 3.87
 Window Size 0.25 min.
 Amount = K0 + K1*Area
 K0 = 4.24689E-002
 K1 = 8.05553E-007

Level	Amount	Area	Height
1	3.75000E-002	40157	3802
2	7.50000E-001	849179	77129
3	1.50000E+000	1713421	152776
4	3.00000E+000	3610927	313707
5	3.75000E+000	4688990	396441
6	0.00000E+000	0	0

Component # 6 Phosphate-P
 Reference Comp. Phosphate-P
 Retention Time 6.38
 Window Size 0.60 min.
 Amount = K0 + K1*Area
 K0 = 8.68926E-002
 K1 = 2.12227E-006

Level	Amount	Area	Height
1	7.50000E-002	24783	1450
2	1.50000E+000	642376	38579
3	3.00000E+000	1301126	79971
4	6.00000E+000	2756481	168994
5	7.50000E+000	3546397	217521
6	0.00000E+000	0	0

Component # 1 Fluoride Retention Time 1.32
 Reference Comp. Fluoride Window Size 0.15 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 $K0 = -9.62851\text{E-}004$
 $K1 = 1.37614\text{E-}006$

Level	Amount	Area	Height
1	2.50000E-002	28534	2732
2	5.00000E-001	373164	44629
3	1.00000E+000	707646	82595
4	2.00000E+000	1435865	173007
5	2.50000E+000	1837162	220914
6	0.00000E+000	0	0

Component # 2 Chloride Retention Time 1.97
 Reference Comp. Chloride Window Size 0.15 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 $K0 = 1.28188\text{E-}001$
 $K1 = 1.95287\text{E-}006$

Level	Amount	Area	Height
1	1.00000E-001	51206	7044
2	2.00000E+000	909455	126181
3	4.00000E+000	1856586	261681
4	8.00000E+000	3987563	585791
5	1.00000E+001	5142155	754321
6	0.00000E+000	0	0

Component # 3 Nitrite-N Retention Time 2.33
 Reference Comp. Chloride Window Size 0.15 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 $K0 = 2.38085\text{E-}002$
 $K1 = 9.76240\text{E-}007$

Level	Amount	Area	Height
1	3.75000E-002	30884	3582
2	7.50000E-001	734006	79701
3	1.50000E+000	1468106	162005
4	3.00000E+000	3021523	336616
5	3.75000E+000	3856614	429219
6	0.00000E+000	0	0

Component # 7 Sulfate Retention Time 7.92
 Reference Comp. Sulfate Window Size 0.90 min.
 Amount = K0 + K1*Area
 K0 = 5.32283E-001
 K1 = 2.53252E-006

Level	Amount	Area	Height
1	3.76000E-001	129999	6524
2	7.50000E+000	2598757	138579
3	1.50000E+001	5330209	287851
4	3.00000E+001	11507107	615917
5	3.75000E+001	14859049	776426
6	0.00000E+000	0	0

Timed Events File: C:\DX\METHOD\W761CAL.TE

Step	Time	Description
Init		ACI Autosmp OFF
Init		ACI pump st ON
Init		ACI inject OFF
Init		ACI auto zer OFF
Init		ACI TTL 1 OFF
Init		ACI TTL 2 OFF
Init		ACI TTL 3 OFF
Init		ACI TTL 4 OFF
Init		ACI OFF
Init		ACI OFF
1	0.0	ACI Autosmp ON
1	0.0	ACI auto zer ON
2	2.5	ACI Autosmp OFF
2	2.5	ACI inject ON
2	2.5	ACI TTL 1 ON
2	2.5	Start Sampling

Component: Fluoride

Fit Type: Linear

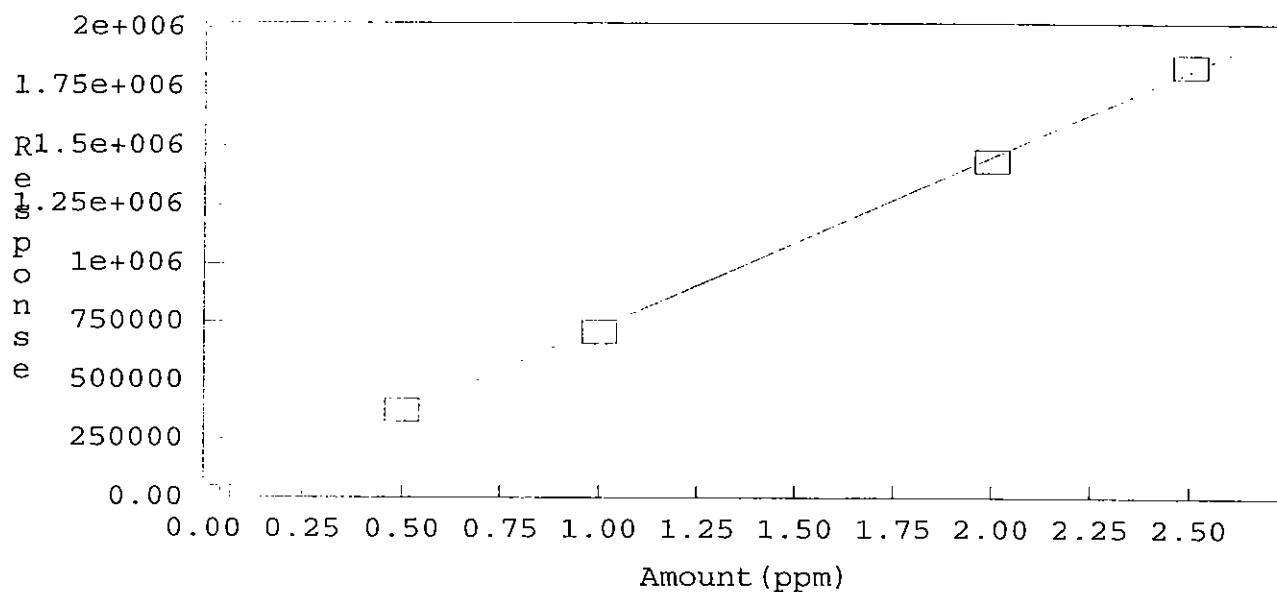
$r^2 = 0.999552$

Amt = Resp * $1.376e-006$ + -0.000962

Resp = Amt * $7.267e+005$ + 699.7

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Chloride

Fit Type: Linear

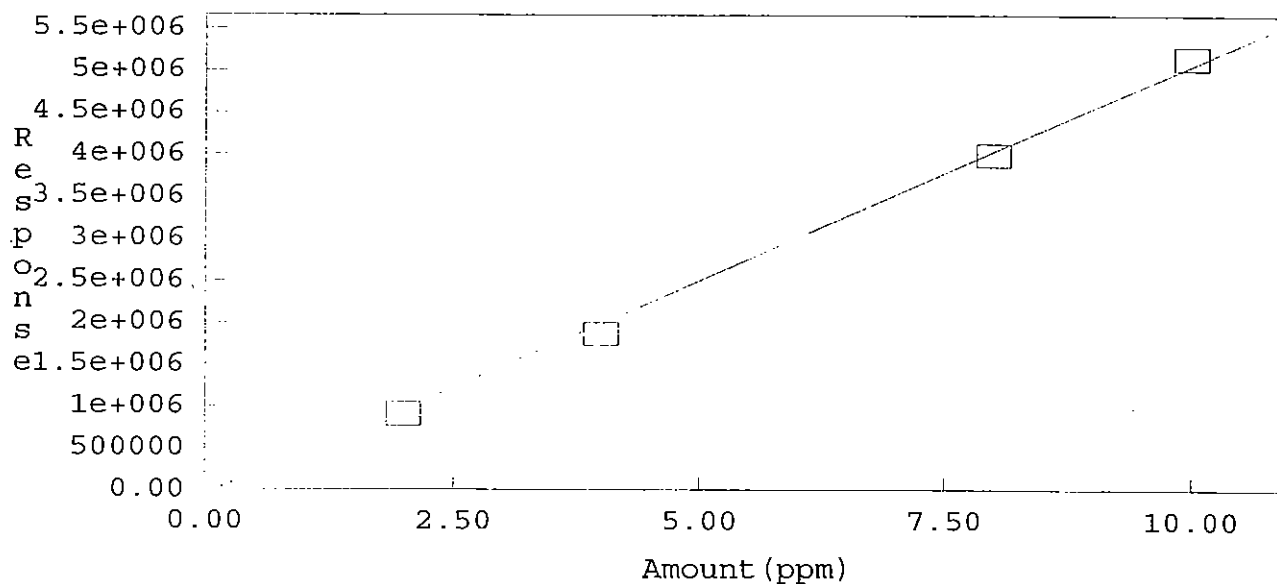
$r^2 = 0.998409$

Amt = Resp * $1.953e-006$ + 0.1282

Resp = Amt * $5.121e+005$ + $-6.564e+00$

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Nitrite-N

Fit Type: Linear

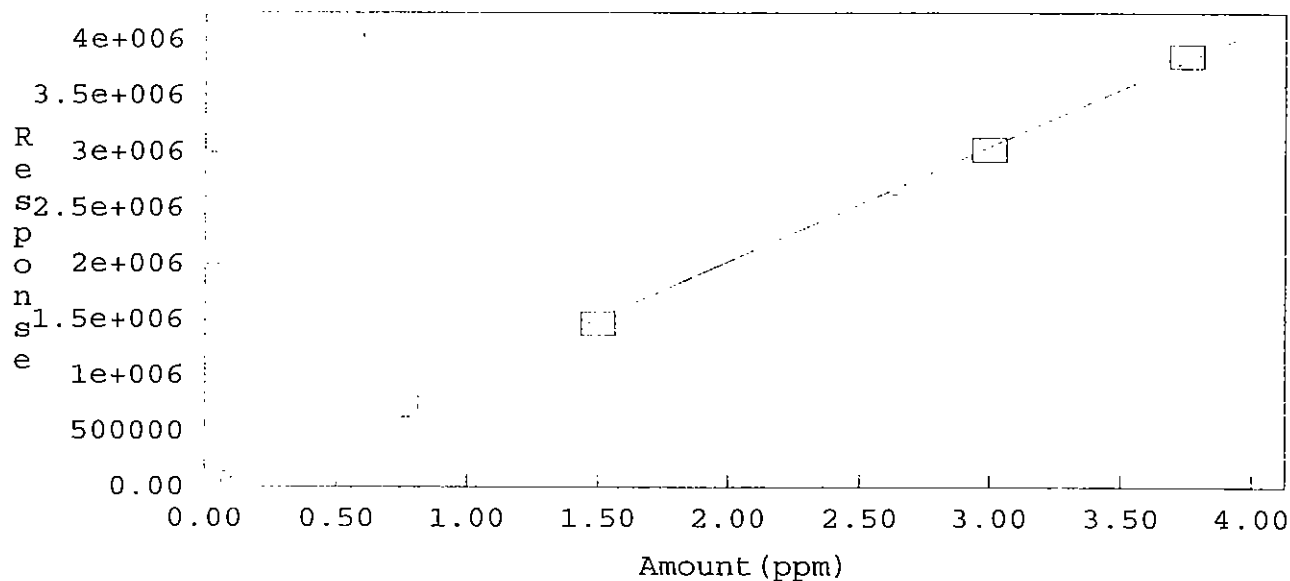
$r^2 = 0.999594$

Amt = Resp * $9.762e-007$ + 0.02381

Resp = Amt * $1.024e+006$ + -2.439e+00

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Bromide

Fit Type: Linear

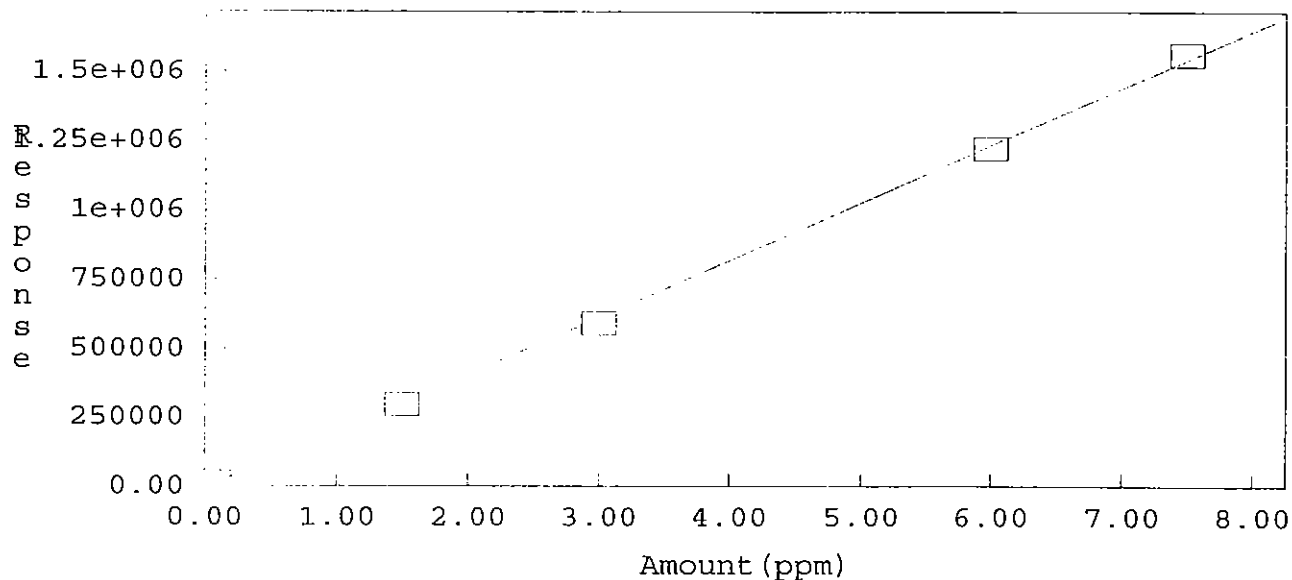
$r^2 = 0.999518$

Amt = Resp * $4.833e-006$ + 0.0459

Resp = Amt * $2.069e+005$ + -9497

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Nitrate-N

Fit Type: Linear

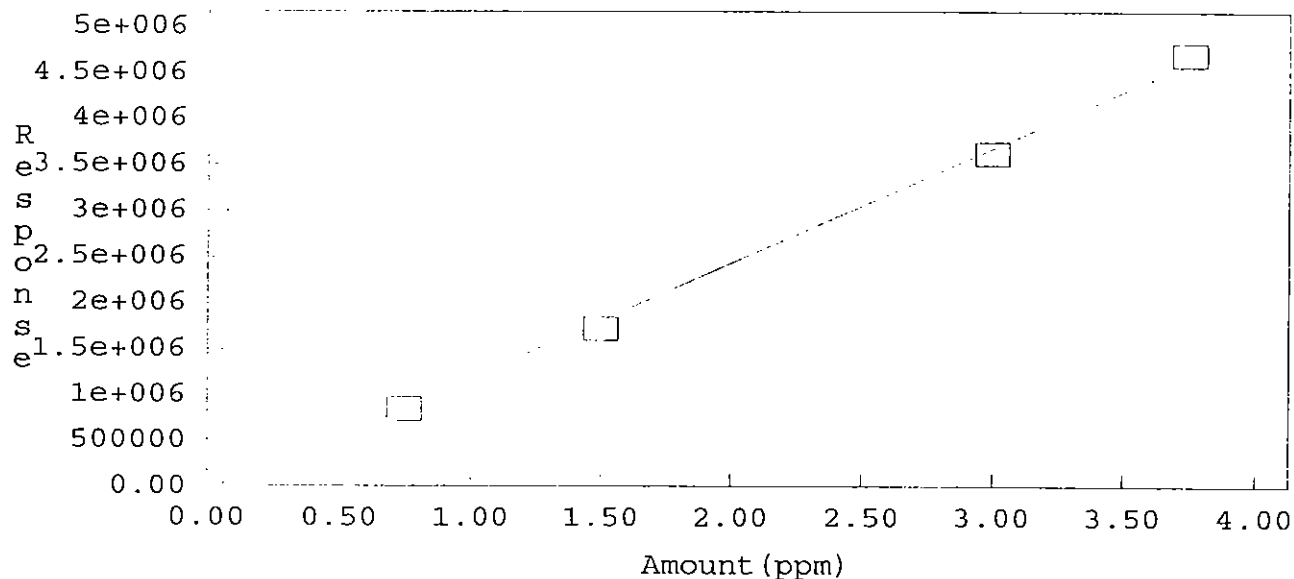
$r^2 = 0.998618$

Amt = Resp * $8.056e-007$ + 0.04247

Resp = Amt * $1.241e+006$ + $-5.272e+00$

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Phosphate-P

Fit Type: Linear

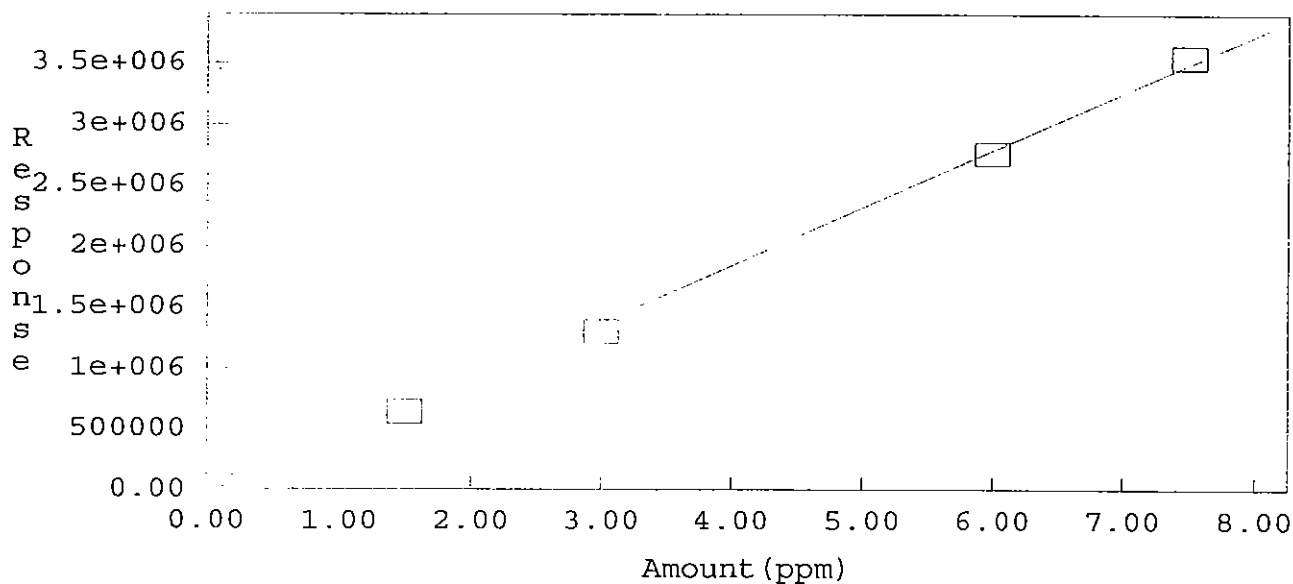
$r^2 = 0.998898$

Amt = Resp * $2.122e-006$ + 0.08689

Resp = Amt * $4.712e+005$ + $-4.094e+00$

Standardization: External

Calibration: Area



Method: C:\DX\METHOD\E300-063.MET

Component: Sulfate

Fit Type: Linear

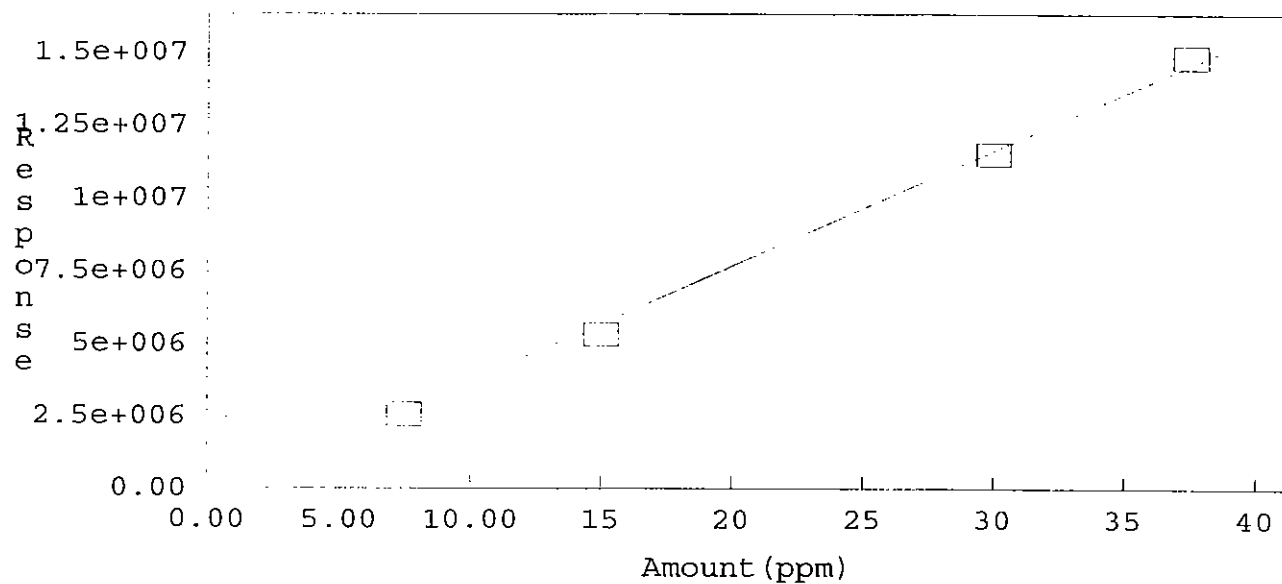
$r^2 = 0.998245$

Amt = Resp * $2.533\text{e-}006$ + 0.5323

Resp = Amt * $3.949\text{e+}005$ + $-2.102\text{e+}00$

Standardization: External

Calibration: Area



DIONEX SCHEDULE - C:\DX\SCHEDULE\E300-063.SCH

Inj#	Sample Name	Method	Data File	Vol.	Dil.	Int.Std.
1	autocal1r	..\E300-063	..\W7767Q01.D01	1	1	1
2	autocal2r	..\E300-063	..\W7767Q01.D02	1	1	1
3	autocal3r	..\E300-063	..\W7767Q01.D03	1	1	1
4	autocal4r	..\E300-063	..\W7767Q01.D04	1	1	1
5	autocal5r	..\E300-063	..\W7767Q01.D05	1	1	1
6	autocal6r	..\E300-063	..\W7767Q01.D06	1	1	1
7	icv-w7768-100X	..\E300-063	..\W7768Q01.D07	1	1	1
8	icb	..\E300-063	..\W7767Q01.D08	1	1	1

Comment:

Analyst DRDate 3/21/03Instrument J

DIONEX SCHEDULE - C:\DX\SCHEDULE\03W3042.SCH

Inj#	Sample Name	Method	Data File	Vol.	Dil.	Int.Std.
1	##03W3042, W CCVW79	..\E300-063	..\W3042Q01.D01	1	1	1
2	MB RW1413	..\E300-063	..\W3042K01.D02	1	1	1
3	LCS W7934-100X	..\E300-063	..\W3042L01.D03	1	1	1
4	LCSD W7934-100X	..\E300-063	..\W3042J01.D04	1	1	1
5	3370-1 F=400	..\E300-063	..\3370-101.D05	1	400	1
6	3370-2 F=400	..\E300-063	..\3370-201.D06	1	400	1
7	3370-3 F=400	..\E300-063	..\3370-301.D07	1	400	1
8	3370-11 F=100	..\E300-063	..\33701101.D08	1	100	1
9	3370-12 F=400	..\E300-063	..\33701201.D09	1	400	1
10	3370-13 F=1000	..\E300-063	..\33701301.D10	1	1000	1
11	3370-14 F=1.25	..\E300-063	..\33701401.D11	1	1.25	1
12	CCV2W7933-100X	..\E300-063	..\W3042Q01.D12	1	1	1
13	MB RW1412	..\E300-063	..\W3042K11.D13	1	1	1
14	\$3370-13 MS F=2000	..\E300-063	..\W3042M01.D14	1	2000	1
15	\$3370-13 MSD F=2000	..\E300-063	..\W3042N01.D15	1	2000	1
16	3360-1 F=8	..\E300-063	..\3360-101.D16	1	8	1
17	3360-7 F=4	..\E300-063	..\3360-701.D17	1	4	1
18	3391-1 F=8	..\E300-063	..\3391-101.D18	1	8	1
19	3391-2 F=5	..\E300-063	..\3391-201.D19	1	5	1
20	CCV3W7933-100X	..\E300-063	..\W3042Q01.D20	1	1	1
21		..\STOP.MET		1	1	1

Comment:

LCS/LCSD LOT # W7934

MS/MSD LOT # W7933

ELUENT LOT # W7868

ANALYTICAL METHOD 9056/E300 MATRIX WAnalyst DCDate 7/27/03Instrument I

160 Magnolia Ave. Chino CA 91710

tel: (909) 590-1828 Fax: (909) 590-1498

atch # 03W3071 Matrix W

Solid Analysis ((160.1, 160.2, 160.3) Worksheet

Method: 160.1 Balance No. _____

Date: 5/29/07 Analyst: Dr

EPA 160.1 TDS - Total Dissolved (filterable) Solids - Dry for 1hr. or more at 180 °C

EPA 160.2 TSS - Total Suspended (nonfilterable) Solids - Dry for 1hr. or more at 103-105 °C

EPA 160.3 TS - Total Solids - Dry for 1hr. or more at 103-105 °C

Other method (specify):

Result = $10^4 \times \Delta W \times f_1 / V$

SOP: G-81

#	Analysis Type	Sample ID (STD Lot #)	Treatment Ratio $V_1/\bar{X}=f_1$	Volume V, mL	W ₁ g	W ₂ 1st, g	W ₂ 2nd, g	$\Delta W = W_2 - W_1$, g	Results (ppm)	Note
1	Blank	T1116	1 =	100	103.1221	103.1221	103.1224	0.0003	3	GC
2	LCS	T1116	1 =	100	104.3113	104.3496	104.3498	0.0385	385	3
3	Sample-1	3392-2	1 =	100	115.1056	115.1442	115.1440	0.0384	384	GC
4	MS on S-1	3389-9	1 =	50.0	111.7875	112.0102	112.0100	0.2225	4450	R
5	MSD on S-1	✓ - 9	1 =	50.0	103.5388	103.7641	103.7639	0.2251	4502	76
6	Sample-2	3396-1	1 =	100	115.9208	116.2320	116.2317	0.3109	3109	Y9
7	Sample-3	✓ - 2	1 =	100	115.8746	116.1845	116.1842	0.3096	3096	CK
8	Sample-4	✓ - 3	1 =	100	113.1742	113.4346	113.4349	0.2607	2607	25
9	Sample-5	✓ - 4	1 =	100	116.9560	117.2355	117.2356	0.2796	2796	W
10	Sample-6	3395-2	1 =	100	113.0638	113.1108	113.1105	0.0467	467	PR
11	Sample-7	3394-11	1 =	100	117.9022	117.9982	117.9980	0.0958	958	7
12	Sample-8	✓ - 12	1 =	100	116.0616	116.1245	116.1243	0.0627	627	31
13	Sample-9	✓ - 13	1 =	100	111.2759	111.2763	111.2760	0.0001	1	5
14	Sample-10	3390-2	1 =	100	114.7856	114.8790	114.8790	0.0934	942	12
15	LCSD	T1116	1 =	100	117.6075	117.6468	117.6466	0.0391	391	X5
16	Sample-11	3391-1	1 =	100	103.3015	103.3548	103.3544	0.0529	529	L
17	Sample-12	✓ - 2	1 =	100	113.8632	113.8966	113.8979	0.0347	347	6
18	Sample-13	3389-4	1 =	100	115.8629	116.1586	116.1588	0.2959	2959	30
19	Sample-14	✓ - 5	1 =	100	114.1726	114.2306	114.2309	0.0583	583	44
20	Sample-15	✓ - 9	1 =	50.0	99.0867	99.2942	99.2939	0.2075	4144	Y2
21	Sample-16	3413-1	1 =	100	108.5560	108.8817	108.8819	0.3259	3259	1P
22	Sample-17	3414-1	1 =	100	107.7204	107.7469	107.7467	0.0263	263	0
23	Sample-18	✓ - 2	1 =	100	112.3057	112.3336	112.3334	0.0277	277	K
24	Sample-19		1 =							
25	Sample-20		1 =							
26	Mix Dup.	3389-9	1 =	100	116.6468	116.8562	116.8564	0.2096	4192	25

Type	STD Lot #	$C_{STD}(\mu\text{g/mL}) \times V_{STD}(\text{mL}) / X(\text{g or mL}) = T$	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
MS	W- 7618	- - x 1 = 46 ppm	%	85-115 %/80-120 %	PQL(w) 10
MSD	W- 7619	x 1 = 1 ppm	%	" "	PQL(s) 50
CS	W- 7619	x 1 = 1 ppm	%	90-110 %/85-115 %	MDL(w) 4
CSD	W- 7619	x 1 = 1 ppm	%	" "	MDL(s) 20

PCL form 5-127, March 7, 1995. Ver. 3.0

No pencil. Use blue pen for record. Use red pen for correction.

LE: [CUST.DOC.WET]TDS.TEX ROOT-FILE: TDS.ROOT.TEX CONTROL-FILE: TDS.000 1-Page file: TDS1.TEX

Control limits are subjected to change. The updated values are given in the latest version of APCL Technical Handbook Vol. 2

Applied P & Ch Laboratory
13760 Magnolia Ave., Chino CA 91710
Tel: (909) 590-1828 Fax: (909) 590-1488

Balance Daily Calibration Worksheet

Calib. Date	Lab Balance				Note (C)	Digital Balance				Note (D) (C) (AR)	Analytical Balance				Note (D) (C) (AR)	Calib. by
	Balance #	1 g ±0.05g	10 g ±0.1g	200 g ±0.5g		Balance #	1 g ±0.02g	10 g ±0.05g	200 g ±0.10g		Balance #	1 g ±0.0002g	10 g ±0.0005g	200 g ±0.0010g		
5/23/03	A-01	Not in use			✓	B-01	1.00	10.00	200.00	✓✓✓	C-01	1.0000	10.0002	200.0001	✓✓✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.0001	10.0000	200.0000	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	10.01	200.01	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/27/03	A-01	Not in use			✓	B-01	1.00	10.00	200.00	✓✓✓	C-01	1.0000	10.0000	200.0001	✓✓✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.0001	10.0000	200.0000	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	10.01	200.01	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/28/03	A-01	Not in use			✓	B-01	1.00	10.00	200.00	✓✓✓	C-01	1.0000	10.0000	200.0001	✓✓✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.0001	10.0000	200.0000	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	10.01	200.01	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/29/03	A-01	Not in use			✓	B-01	1.00	10.00	200.00	✓✓✓	C-01	1.0000	10.0000	200.0001	✓✓✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.0001	10.0000	200.0000	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	10.01	200.01	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					

Notation: (C) - Cleanliness; (D) - Display; (AR) - Auto Resetting.
No pencil. Use blue pen for record. Use red pen for correction.
APCL form 4-213, March 30, 1995, Ver. 4.0
1. Data File: BAL-CAL1.TEX

Temperature compensation must be performed by the instrument automatically.

Analyst rw

SOP: G-44

Batch # 03W3048 Analysis Date: 5/27/03
 Starting Time: 15:55 Ending Time: _____
 Matrix ☐ Aqueous ☐ Soil

Batch # 03W3061 Analysis Date: 5/28/03
 Starting Time: 16:24 Ending Time: _____
 Matrix ☒ Aqueous ☐ Soil

Standard	4.00	7.00	10.00
Lot #		030660-24	030659-24
Temperature °C		24.4	24.4
pH Reading		7.07	10.07
T-corrected pH		7.00	10.00
Control Limit	±0.05 pH unit		

Standard	4.00	7.00	10.00
Lot #		030660-24	030659-24
Temperature °C		24.0	24.0
pH Reading		6.99	10.00
T-corrected pH		7.00	10.07
Control Limit	±0.05 pH unit		

#	Sample ID	Pre-treat	pH	Note
MB	T1116		6.82	
1	3391-1		7.00	
2	↓ -2		7.18	
3	3421-8		7.22	
4				
5				
6				
7				
8				
9				
10				
11				
12				
13				
14				
15				
16				
17				
18				
19				
20				
	3391-1		7.02	

#	Sample ID	Pre-treat	pH	Note
MB	T1116		6.86	
1	3414-1		6.84	
2	-2		7.00	
3				
4				
5				
6				
7				
8				
9				
10				
11				
12				
13				
14				
15				
16				
17				
18				
19				
20				
	3414-2		7.03	

Laboratory
Ave. Chino CA 91710
590-1828 Fax: (909) 590-1498

Chromium (VI) (7196) Worksheet

Lot #: 03W30486 Matrix: W
Reagent Water 5/17/03

[Holding Time: 24 hours!!]

Test Date: 5/27/03

Analyst: Ba

Test Time: 14236

SOP: G-22

Diphenylcazide solution

Calibration	STD Lot #	$C_{std} \times V_{std} / V_f = C_i$	A_i	$RF_i = A_i / C_i$	Calibration results	Note
STD-1	W-	x / = mg/L			Least Square [RF]=	Cal. Code:
STD-2	W-	x / = mg/L			Average RF=	
STD-3	W-	x / = mg/L			C.C.= (> 0.995)	
STD-4	W-	x / = mg/L			RSD= % (< 15%)	
STD-5	W-	x / = mg/L			Ref. page	
STD-6	W-	x / = mg/L			<u>A=2000+0.826C</u>	

Analysis Type	Sample ID or Lot #	Samp. Amnt X_0 (g or mL)	Dilu./Ext $X/X_0 = f_1$	Treat. Ratio $V/X = f_2$	540 nm A	Concentration $C' = A / RF$	C (Sample) $C = f_1 f_2 C'$	Anomaly Note
CCV	Lot: W- <u>7850</u>	Expected Conc.: x	/ = <u>0.25</u> mg/L		<u>0.211</u>	<u>0.252</u> mg/L	REC. %	90-110 %
Method Blank	Bl. Lot: <u>T1116</u>		$/X_0 = 1$	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</u> ppm	
LCS1	Bl. Lot: <u>1</u>		$/X_0 = 1$	95.0/ =	<u>0.193</u>	mg/L	<u>0.221</u> ppm	
Sample-1 <u>13</u>	<u>3391-1</u>		$/X_0 = 1$	95.0/ =	<u>0.027</u>	<u>0.020</u> mg/L	<u>0.024</u> ppm	
MS on S-1	<u>1</u>		$/X_0 = 1$	95.0/ =	<u>0.205</u>	mg/L	<u>0.245</u> ppm	
MSD on S-1	<u>1</u>		$/X_0 = 1$	95.0/ =	<u>0.213</u>	mg/L	<u>0.255</u> ppm	
Sample 2 <u>16</u>	<u>2</u>		$/X_0 = 1$	95.0/ =	<u>0.005</u>	mg/L	<u>0.006</u> ppm	
Sample 3			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 4			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 5			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 6			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 7			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 8			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 9			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 10			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Blank	Lot: <u>T1116</u>		$/X_0 = 1$	95.0/ =		mg/L	ppm	
LCS2	Bl. Lot: <u>T1116</u>		$/X_0 = 1$	95.0/ =	<u>0.208</u>	mg/L	<u>0.249</u> ppm	
Sample 11			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 12			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 13			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 14			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 15			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 16			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 17			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 18			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 19			$/X_0 = 1$	95.0/ =		mg/L	ppm	
Sample 20			$/X_0 = 1$	95.0/ =		mg/L	ppm	
MTX Dup. <u>0.25 mg/L</u>			$/X_0 = 1$	95.0/ =	<u>0.213</u>	<u>0.255</u> mg/L	ppm	

Type	STD Lot #	$C_{STD}(\mu\text{g/mL}) \times V_{STD}(\text{mL}) / X(\text{g or mL}) = T$	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
MS	W- <u>7850</u>	x / = <u>0.25</u> ppm	%	80-120 %/80-120 %	PQL(w) 0.01
MSD	W- <u>1</u>	x / = ppm	%		PQL(s) 0.05
LCS	W- <u>7759</u>	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W- <u>1</u>	x / = ppm	%		MDL(s) 0.025

APCL form 5-155 May 08, 1996, Ver. 3.1 No pencil. Use blue pen for record. Use red pen for correction.

File:[CUST.DOC.WET]CR6.TEX Root-file: CR6.ROOT.TEX 1-Page-File: CR61.TEX

Control limits are subjected to change. The updated values are given in the latest version of APCL Technical Handbook Vol. 2

Laboratory
Ave. Chino CA 91710
909-1828 Fax: (909) 590-1498
02W1295 Matrix: W

Chromium (VI) (7196) Worksheet

[Holding Time: 24 hours!!]

Test Date: 1/29/03 Analyst: BV

Test Time: _____ SOP: G-22

Lot #: Reagent Water _____ Diphenylcazide solution _____

Calibration	STD Lot #	$C_{std} \times V_{std} / V_f = C_i$	A_i	$RF_i = A_i / C_i$	Calibration results	Note
STD-1	W-7191	x / = 0.000 mg/L	0.000		Least Square [RF]=	Cal. Code:
STD-2	W-	x / = 0.012 mg/L	0.006		Average RF=	A=0.000+0.836C
STD-3	W-	x / = 0.030 mg/L	0.001		C.C.=0.997 (> 0.995)	
STD-4	W-	x / = 0.125 mg/L	0.009		RSD= % (< 15%)	
STD-5	W-	x / = 0.150 mg/L	0.014		Ref. page	
STD-6	W-	x / = 0.50 mg/L	0.015		A=0.003+0.846C	

Analysis Type	Sample ID or Lot #	Samp. Amnt X ₀ (g or mL)	Dilu./Ext X/X ₀ =f ₁	Treat. Ratio V/X=f ₂	540 nm A	Concentration C'=A/RF	C (Sample) C=f ₁ f ₂ C'	Anomaly Note
CCV	Lot: W-7076	Expected Conc.: x	/	= 0.25 mg/L	0.216	0.258 mg/L	REC. %	90-110 %
Method Blank	Bl. Lot: T1115		/X ₀ =	95.0/ =	0.000	mg/L	0.00 ppm	
LCS1	Bl. Lot: 4		/X ₀ =	95.0/ =	0.209	mg/L	0.244 ppm	
Sample-1	1369-1		/X ₀ =	95.0/ =	0.000	mg/L	0.00 ppm	
MS on S-1	6		/X ₀ =	95.0/ =	0.223	mg/L	0.266 ppm	
MSD on S-1	6		/X ₀ =	95.0/ =	0.230	mg/L	0.275 ppm	
Sample 2	12		/X ₀ =	95.0/ =	0.004	mg/L	0.005 ppm	
Sample 3	3		/X ₀ =	95.0/ =	0.002	mg/L	0.002 ppm	
Sample 4	4		/X ₀ =	95.0/ =	0.001	mg/L	0.001 ppm	
Sample 5	5		/X ₀ =	95.0/ =	0.002	mg/L	0.002 ppm	
Sample 6	6		/X ₀ =	95.0/ =	0.004	mg/L	0.005 ppm	
Sample 7			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 8			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 9			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 10			/X ₀ =	95.0/ =		mg/L	ppm	
Blank	Lot:		/X ₀ =	95.0/ =		mg/L	ppm	
LCS2	Bl. Lot: T1115		/X ₀ =	95.0/ =	0.210	mg/L	0.25 ppm	
Sample 11			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 12			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 13			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 14			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 15			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 16			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 17			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 18			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 19			/X ₀ =	95.0/ =		mg/L	ppm	
Sample 20			/X ₀ =	95.0/ =		mg/L	ppm	
MTX Dup.	closing 0.25 mg/L		/X ₀ =	95.0/ =	0.204	mg/L	0.24 ppm	

Type	STD Lot #	$C_{STD}(\mu\text{g/mL}) \times V_{STD}(\text{mL}) / X(\text{g or mL}) = T$	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
MS	W-7076	x / = 0.25 ppm	%	80-120 %/80-120 %	PQL(w) 0.01
MSD	W- v	x / = ppm	%		PQL(s) 0.05
LCS	W-7191	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W- x	x / = ppm	%		MDL(s) 0.025

Alkalinity / OH / CO₃ / HCO₃ (310.1 / SM2320B) Worksheet

Batch # 03W3113 Matrix: W

Titrant H₂SO₄

Lot # W7912

Concentration (C) 0.02574

Test Date: 6/26/03

Analyst: DL

SOP: G-51

#	Sample ID	Dilution V _f /V _i =f _i	Simpl Amt V, mL	H ₂ SO ₄ (mL) by Phen S _A E _A	A	H ₂ SO ₄ (mL) by MR-BCG S _B E _B	B	Phen-Aik., P (in unit of mgCaCO ₃ /L)	Tot. Alk., T	OH ⁻	CO ₃ ²⁻	HCO ₃ ⁻	Note & Anomaly
1	MB: 7.116	1 =	100		0		0	0	0	0	0		
2	1.25	1 =	100				7.90	100.9					
3	1.25	1 =	100				7.80	99.6					
4	3391-1	1 =	100		0		4.50	115.0	0	0	115.0		
5	-2	1 =	100		0		4.50	115.0	0	0	115.0		
6	3413-1	1 =	100		0		9.15	233.8	0	0	233.8		
7	3414-1	1 =	100		0		5.70	145.6	0	0	145.6		
8	-2	1 =	100		0		6.00	153.3	0	0	153.3		
9	3444-1	1 =	100		0		11.0	283.6	0	0	283.6		
10	-2	1 =	100		0		4.81	123.9	0	0	123.9		
11	↓ -3	1 =	100		0		7.60	194.2	0	0	194.2		
12	3444-1	1 =	100		0		8.15	218.5	0	0	218.5		
13	-2	1 =	100		0		6.81	175.0	0	0	175.0		
14	↓ -3	1 =	100		0		8.51	218.5	0	0	218.5		
15		1 =											
16		1 =											
17		1 =											
18		1 =											
19		1 =											
20		1 =											
Dup.	3444-1	1 =	100		0		11.05	282.3	0	0	282.3		
Titration Results OH ⁻ (CaCO ₃ mg/L) CO ₃ ²⁻ (CaCO ₃ mg/L) HCO ₃ ⁻ (CaCO ₃ mg/L) P=0 0 0 T P<T/2 0 2P T-2P P=T/2 0 2P 0 P>T/2 2P-T 2(T-P) 0 P=T T 0 0													

Calculation:

A=S_A-E_AB=S_B-E_BP=50,000 f_i A C / VT=50,000 f_i (A+B) C / V

APCL Form 5-101, Nov. 29, 1999 Ver 3.2

Using blue pen. Correcting by red pen.

File: [CUST.DOC.WET]ALK.TEX

Root-File: [CUST.DOC.WET]ALK-ROOT.TEX

1-Page-File: [CUST.DOC.WET]ALK1.TEX



Applied Physics & Chemistry Laboratory

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Tel. (909) 590-1828 Fax (909) 590-1498

May 23, 2003

GEOFON, Inc.
Attention: Leo Williamson
22632 Golden Spring Dr Ste 270
Diamond Bar CA 91765

Dear Leo,

This package contains samples in our Service ID 03-2843 and your project : 04-4428.10 JPL.
Enclosed please find:

- (1) Original analytical report.
- (2) Original Chain of Custody.
- (3) One diskette containing EDD deliverable.
- (4) One original Level C Data Package Deliverable.

If anything is missing or you have any questions, please feel free to contact me.

Respectfully submitted,

Regina Kirakozova
Associate QA/QC Director
Applied P & Ch Laboratory

Submitted to:

GEOFON, Inc.

Attention: Leo Williamson

22632 Golden Spring Dr Ste 270

Diamond Bar CA 91765

Tel: (909) 396-7662 Fax: (909) 396-1455

APCL Analytical Report

Service ID #: 801-032843

Received: 04/23/03

Collected by: Leo Williamson

Extracted: N/A

Collected on: 04/23/03

Tested: 04/23-29/03

Reported: 05/07/03

Sample Description: Water

Project Description: 04-4428.10 JPL

Analysis of Water Samples

Component Analyzed	Method	Unit	PQL	Analysis Result			
				DUPE-2-2Q03 03-02843-1	EB-4-4/23/03 03-02843-2	MW-14-1 03-02843-3	MW-14-2 03-02843-4
BICARBONATE	SM2320B	mg/L	2	210	< 2	245	234
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	< 2	< 2	< 2	< 2
PH	9040B	pH unit	0.01	7.63	7.00	6.77	7.18
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	669	< 10	731	740
CHROMIUM (VI)	7196	mg/L	0.01	< 0.01	< 0.01	< 0.01	< 0.01
Dilution Factor				1	1	1	1
PERCHLORATE	314.0	µg/L	4	5.4	< 4	2.8J	3.3J
Dilution Factor				20	1.25	20	20
CHLORIDE CL ⁻	300.0	mg/L	0.2	96.7	0.23J	136	111
NITRATE AS N	300.0	mg/L	0.04	13.1	0.088	17.1	14.8
SULFATE SO ₄ ²⁻	300.0	mg/L	0.5	139	0.73	194	179
Dilution Factor				1	1	1	1
ARSENIC	200.9	µg/L	5	< 5	< 5	< 5	< 5
CALCIUM	200.7	µg/L	200	96,500	98.0J	142,000	128,000
IRON	200.7	µg/L	50	19.2J	1.9J	32.4J	5.3J
MAGNESIUM	200.7	µg/L	100	39,800	15.1J	45,000	42,700
POTASSIUM	200.7	µg/L	400	3,630	27.8J	3,130	2,920
SODIUM	200.7	µg/L	2000	36,600	430J	36,200	29,900
VOLATILE ORGANIC COMPOUNDS							
Dilution Factor				1	1	1	1
BENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOCHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMODICHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOFORM	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
N-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
SEC-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
TERT-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
2-BUTANONE	524.2	µg/L	10	< 10	< 10	< 10	< 10
CARBON TETRACHLORIDE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLORODIBROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROFORM	524.2	µg/L	0.5	0.3J	< 0.5	0.4J	0.4J
CHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
2-CHLOROTOLUENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				DUPE-2-2Q03 03-02843-1	EB-4-4/23/03 03-02843-2	MW-14-1 03-02843-3	MW-14-2 03-02843-4
4-CHLOROTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DIBROMO-3-CHLOROPROPANE	524.2	µg/L	1.1 ^(a)	<1.1	<1.1	<1.1	<1.1
1,2-DIBROMOETHANE (EDB)	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
DIBROMOMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,4-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
DICHLORODIFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
2,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
HEXACHLOROBUTADIENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ISOPROPYLBENZENE (CUMENE)	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
P-ISOPROPYLTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
METHYLENE CHLORIDE	524.2	µg/L	1.8 ^(a)	<1.8	<1.8	<1.8	<1.8
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1	<1
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	<10	<10	<10	<10
NAPHTHALENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
N-PROPYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
STYRENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TETRACHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	0.4J	0.5J
TOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRICHLOROETHENE	524.2	µg/L	0.5	0.5	<0.5	1.3	3.7
TRICHLOROFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2,2,3,3-HEXACHLORO-1,2,2,3,3,3-HEXACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3,5-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
VINYL CHLORIDE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
O-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
M/P-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				MW-14-3 03-02843-5	MW-14-4 03-02843-6	MW-14-5 03-02843-7	TB-4-4/23/03 03-02843-8
BICARBONATE	SM2320B	mg/L	2	210	156	129	-
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	< 2	< 2	7.6	-
PH	9040B	pH unit	0.01	7.61	7.98	8.22	-
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	613	300	174	-
CHROMIUM (VI)	7196	mg/L	0.01	< 0.01	< 0.01	< 0.01	-
Dilution Factor				1	1	1	1
PERCHLORATE	314.0	µg/L	4	5.7	2.4J	< 4	-
Dilution Factor				20	4	1.25	1
CHLORIDE CL ⁻	300.0	mg/L	0.2	94.3	37.9	9.9	-
NITRATE AS N	300.0	mg/L	0.04	13.1	10.1	0.15	-
SULFATE SO ₄ ⁻	300.0	mg/L	0.5	136	30.0	16.9	-
Dilution Factor				1	1	1	1
ARSENIC	200.9	µg/L	5	< 5	< 5	< 5	-
CALCIUM	200.7	µg/L	200	97,600	50,300	17,100	-
IRON	200.7	µg/L	50	16.0J	21.2J	55.7	-
MAGNESIUM	200.7	µg/L	100	41,500	16,900	10,800	-
POTASSIUM	200.7	µg/L	400	3,610	2,230	2,150	-
SODIUM	200.7	µg/L	2000	38,100	25,500	28,200	-
VOLATILE ORGANIC COMPOUNDS							
Dilution Factor				1	1	1	1
BENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOCHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMODICHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOFORM	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
BROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
N-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
SEC-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
TERT-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
2-BUTANONE	524.2	µg/L	10	< 10	< 10	< 10	< 10
CARBON TETRACHLORIDE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLORODIBROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
CHLOROFORM	524.2	µg/L	0.5	0.3J	< 0.5	< 0.5	< 0.5
CHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
2-CHLOROTOLUENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
4-CHLOROTOLUENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
1,2-DIBROMO-3-CHLOROPROPANE	524.2	µg/L	1.1 (a)	< 1.1	< 1.1	< 1.1	< 1.1
1,2-DIBROMOETHANE (EDB)	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
DIBROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
1,2-DICHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
1,3-DICHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
1,4-DICHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5
DICHLORODIFLUOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5	< 0.5

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				MW-14-3 03-02843-5	MW-14-4 03-02843-6	MW-14-5 03-02843-7	TB-4-4/23/03 03-02843-8
1,1-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
2,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
HEXACHLOROBUTADIENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ISOPROPYLBENZENE (CUMENE)	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
P-ISOPROPYLTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
METHYLENE CHLORIDE	524.2	µg/L	1.8 ^(a)	<1.8	<1.8	<1.8	<1.8
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1	<1
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	<10	<10	<10	<10
NAPHTHALENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
N-PROPYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
STYRENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TETRACHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRICHLOROETHENE	524.2	µg/L	0.5	0.5J	<0.5	<0.5	<0.5
TRICHLOROFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3,5-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
VINYL CHLORIDE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
O-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
M/P-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5

PQL: Practical Quantitation Limit. MDL: Method Detection Limit. CRDL: Contract Required Detection Limit

N.D.: Not Detected or less than the practical quantitation limit.

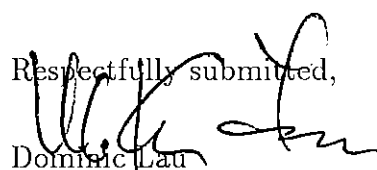
"-": Analysis is not required.

J: Reported between PQL and MDL.

Listed Dilution Factors (DF) are relative to the method default DF. All unlisted DFs are 1.0

^(a) MDL reported.

Respectfully submitted,


 Dominic Lau
 Laboratory Director
 Applied P & Ch Laboratory

Level C Data Package Deliverables

General Information

Project: 04-4428.10 JPL

APCL Service ID: 03-2843



Applied P & Ch Laboratory

13760 Magnolia Ave. Chino, CA 91710

Telephone (909)590-1828

Fax (909)590-1498

Case Narrative

Project: JPL/04-4428.10

For GEOFON, Inc.

APCL Service No: 03-2843

1. Sample Identification

The sample identifications are listed in the following table:

GEOFON, Inc. Sample ID	APCL Sample ID
MW-14-5	03-02843-7
MW-14-4	03-02843-6
MW-14-3	03-02843-5
MW-14-2	03-02843-4
MW-14-1	03-02843-3
TB-4-4/23/03	03-02843-8
EB-4-4/23/03	03-02843-2
DUPE-2-2Q03	03-02843-1

2. Analytical Methodology

Samples are analyzed by EPA methods

524.2 (Volatile Organic Compounds),
7196 (Chromium (VI)),
314.0 (Perchlorate, low level),
300.0 (Anions by IC),
SM2320B (Carbonate),
SM2320B (Bicarbonate),
9040B (pH),
160.1 (Solids, Total Dissolved (TDS)),
200.7 (Metals by ICP),
200.9 (Arsenic, As, by GFAA),

3. Holding Time

All samples were extracted, digested and analyzed within the holding times defined by the appropriate EPA methods of the analyses.

4. Preservation

All samples were preserved and stored according to the appropriate EPA methods.

5. Tele-log

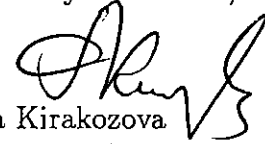
None

6. Anomaly

None

"I certify that these data are technically accurate, complete, and in compliance with the terms and conditions of the contract, for other than the conditions detailed above. Release of the data contained in the hardcopy data package and its electronic data deliverable submitted on diskette had been authorized by the Laboratory Manager or her/his designee, as verified by the following signature."

Respectfully submitted,

A handwritten signature in black ink, appearing to read 'Regina Kirakozova', written over the printed name.

Regina Kirakozova
Associate QA/QC Director
Applied P & Ch Laboratory



IN CORPORATE
DIAMOND BAR, CA 91765 • (909) 396-7662 • FAX (909) 396-1455

CHAIN-OF-CUSTODY RECORD

LABORATORY COPY

MW-14 0021

GEOPON, LAB COORDINATOR

LABORATORY SERVICE ID

LABORATORY CONTACT

MAIL REPORT (COMPANY NAME)

BRAD SHOYEE

LAB COORDINATOR'S PHONE

LAB COORDINATOR'S FAX

LABORATORY PHONE

LABORATORY FAX

REMITTANT NAME

PROJECT NAME

PROJECT LOCATION

PROJECT NUMBER

LABORATORY ADDRESS

CITY, STATE AND ZIP CODE

ADDRESS

PROJECT CONTACT

PROJECT PHONE NUMBER

PROJECT FAX

CITY, STATE AND ZIP CODE

ADDRESS

PROJECT ADDRESS

CITY, STATE AND ZIP CODE

CLIENT

CITY, STATE AND ZIP CODE

ADDRESS

PROJECT MANAGER

PROJECT MANAGER'S PHONE

PROJECT MANAGER'S FAX

CITY, STATE AND ZIP CODE

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CITY, STATE AND ZIP CODE

ADDRESS

Sample Receiving Checklist

APCL ServiceID:

2843

Client Name/Project:

Geslon JPL

1. Sample Arrival

Date/Time Received

4/23/03 1515

Date/Time Opened

4/23/03 1515

By (name):

Kenny Chan

Custody Transfer:

☐ Client☐ Golden State☐ UPS☐ US Mail☐ FedEx☐ APCL Empl:Adam Wood

2. Chain-of-Custody (CoC)

☒ With Samples?☐ Faxed?☐ Client has Copy?☐ Signed, dated? By:☒ Project ID?☐ Analyses Clear?☐ Hold Samples?

on Hold

Received

☒ CoC/Docs Zip-Locked under lid?☐ Compos. #:☒ #Samples OK?☐ Discrepancies?☐ Client notified?☐ Response (attach docs):

3. Shipping Container/Cooler

☒ Cooler Used? # of1

Cooled by:

☒ Ice☐ Blue Ice☐ Dry Ice☐ None

Temp °C

4.0

(Cooler temperature measured from temp blank if present, otherwise measured from the cooler).

Cooler Custody Seal?

☐ Absent☐ Intact☐ Tampered?

4. Sample Preservation

☐ pH <2☐ pH >12

If Not, pH =

Preserved by:

☐ Client☐ APCL☐ Third Party

5. Holding-time Requirements

☐ pH 24hr☐ BACT 6/24hr☒ Cr^{VI} 24hr☐ NO₃⁻ 48hr☐ BOD 48hr☐ Cl₂ ASAP☐ Turbidity 48hr☐ DO ASAP☐ Fe(II) ASAP☐ HT Expired?☐ Client notified?

6. Sample Container Condition

☒ Intact?☐ Broken?☐ Documented?

Number:

Type:

☒ plastic☒ glass☐ Tube: brass/SS☐ Tedlar Bag☒ Quantity OK?☒ Leaking?☐ Anomaly?☒ Caps tight?☒ Air Bubbles?☐ Anomaly?

Labels:

☒ Unique ID?☒ Date/Time☐ Preserved?

7. Turn Around Time

☐ RUSH TAT:☒ Std (7-10 days)☐ Not Marked

8. Sample Matrix

☐ Drinking H₂O☒ Other Liq☐ Soil☐ Wipe☐ Polymer☐ Air☐ Other:☐ Ground H₂O☐ Sludge☐ Filter☐ Oil/Petro☐ Paint☐ W. Water☐ Extract☐ Unknown

9. Pre-Login Check List Completed & OK?

☒ ALL OK? (if not, attach docs)☐ Client Contact? (Name:) Date/Time:

Received/Checked by:

Kenny Chan

Date: 23 Apr 2003

Time: 7:42 a.m.

*HT: Samples must be analyzed for results to reflect total concentrations. Results generated outside required of holding times are considered minimal values and may be used to define waste as hazardous but not as non-hazardous.

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

Tel: (909) 590-1828 Fax: (909) 590-1498

Sample Login: Check List

03-02843 (0470_ 131) (2202777_ 131)

04/23/03

Part 1: General Information

☐ Company Information	Name:	<i>GEOFON, Inc.</i>
	Address:	<i>22632 Golden Spring Dr Ste 270 ,Diamond Bar ,CA 91765</i>
☐ Project Information	Project Description:	<i>JPL</i>
	Project #:	<i>04-4428.10</i>
☐ Billing Information	P.O. #:	
	Bill Address:	<i>22632 Golden Spring Dr Ste 270 ,Diamond Bar ,CA 91765</i>
	Lab Project ID:	
	Client Database #:	<i>3</i>
☐ Receiving Information	Who Received Sample?	<i>Kenny Chan</i>
	Receiving Date/Time:	<i>04/23/03 1515</i>
	COC No.	
☐ Shipping Information	Shipping Company	<i>APCL pick up</i>
	Packing Information:	<i>Cooler/Ice Chester</i>
	Cooler Temperature:	<i>4.0 °C</i>
☐ Container Information	Container Provider:	<i>Client</i>
☐ Sampling Information	Sampling Person:	
	Sampling Company:	<i>Client</i>
☐ Turn-Around-Time Option:		<i>Rush 5 working day(s)</i>
☐ QC Option:		<i>NEESA C</i>
☐ Disposal Option:		<i>Not specify</i>

Part 2: Sample Information

Seq. #	Sample ID (on COC)	Sample Sub-ID	APCL Sample ID	Matrix	Cont- tainer	Preser- vative	Vol, ml Am. g	# of Replica	Condition G, L, B	Collected mmddyy	Hold ?	Composite Group	TAT Days	
1	MW-14-5	VOC	03-02843-7- α	W	V	C	40	3	G	042303	N	0	7	☐
	MW-14-5	Metal	03-02843-7- β	W	P	N	500	1	G	042303	N	0	7	☐
	MW-14-5	300	03-02843-7- γ	W	P		1000	1	G	042303	N	0	7	☐
2	MW-14-4	VOC	03-02843-6- α	W	V	C	40	3	G	042303	N	0	7	☐
	MW-14-4	Metal	03-02843-6- β	W	P	N	500	1	G	042303	N	0	7	☐
	MW-14-4	300	03-02843-6- γ	W	P		1000	1	G	042303	N	0	7	☐
3	MW-14-3	VOC	03-02843-5- α	W	V	C	40	3	G	042303	N	0	7	☐
	MW-14-3	Metal	03-02843-5- β	W	P	N	500	1	G	042303	N	0	7	☐
	MW-14-3	300	03-02843-5- γ	W	P		1000	1	G	042303	N	0	7	☐
4	MW-14-2	VOC	03-02843-4- α	W	V	C	40	3	G	042303	N	0	7	☐
	MW-14-2	Metal	03-02843-4- β	W	P	N	500	1	G	042303	N	0	7	☐
	MW-14-2	300	03-02843-4- γ	W	P		1000	1	G	042303	N	0	7	☐
5	MW-14-1	VOC	03-02843-3- α	W	V	C	40	3	G	042303	N	0	7	☐
	MW-14-1	Metal	03-02843-3- β	W	P	N	500	1	G	042303	N	0	7	☐
	MW-14-1	300	03-02843-3- γ	W	P		1000	1	G	042303	N	0	7	☐
6	TB-4-4/23/03	VOC	03-02843-8	W	V	C	40	2	G	042303	N	0	7	☐
7	EB-4-4/23/03	VOC	03-02843-2- α	W	V	C	40	3	G	042303	N	0	7	☐
	EB-4-4/23/03	Metal	03-02843-2- β	W	P	N	500	1	G	042303	N	0	7	☐
	EB-4-4/23/03	300	03-02843-2- γ	W	P		1000	1	G	042303	N	0	7	☐
8	DUPE-2-2Q03	VOC	03-02843-1- α	W	V	C	40	3	G	042303	N	0	7	☐
	DUPE-2-2Q03	Metal	03-02843-1- β	W	P	N	500	1	G	042303	N	0	7	☐
	DUPE-2-2Q03	300	03-02843-1- γ	W	P		1000	1	G	042303	N	0	7	☐

Part 3: Analysis Information

Test Items:

- ☐ 524.2 Volatile Organic Compounds
- ☐ 7196A Chromium (VI)
- ☐ 314.0/300.0 Perchlorate, low level
- ☐ 300.0 Chloride Cl^- by IC
- ☐ 300.0 Sulfate (SO_4^{--}), by IC
- ☐ 300.0/SM4500NOM Nitrate (NO_3^-) as N by IC
- ☐ SM2320B Carbonate
- ☐ SM2320B Bicarbonate
- ☐ 9040B/150.1 pH
- ☐ 160.1 Solids, Total Dissolved (TDS)
- ☐ 200.7/6010B Sodium, Na, by ICP
- ☐ 200.7/6010B Calcium, Ca, by ICP

Level C Data Package Deliverables

Volatile Organics



Applied P & Ch Laboratory

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/28/2003
Project ID: JPL	Service ID: 32843	Collected by:
Sample ID: 03G2233-MB-01	Lab Sample ID: 03G2233-MB-01	Received Date: 04/28/2003
Sample Type: Method Blank	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: G2233K02	Prep. No: -	Anal. Time: 17:37
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	< 0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	< 0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	< 0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	< 0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	< 0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	< 0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	< 0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	< 0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	< 0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	< 10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	< 0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	< 0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	< 0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	< 0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	< 0.5	U
16	CHLOROMETHANE	74-87-3	µg/L	0.5	< 0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	< 0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	< 0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1	< 1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	< 0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	< 0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	< 0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	< 0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	< 0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	< 0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	< 0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	< 0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	< 0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	< 0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	< 0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	< 0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	< 0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	< 0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	< 0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	< 0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	< 0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	< 0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	< 0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	< 0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	4-BROMO-FLUOROBENZENE (BFB)	460-00-4		70-129	115	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	91	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	94	
4	TOLUENE-D8	2037-26-5		73-129	107	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	91	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	88	
3	FLUOROBENZENE	462-06-6		50-200	94	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

Qualifier: U - Not Detected or less than MDL

E - Exceed calibration range

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

B - A positive value was found in the method blank
D - Diluted

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233K02.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M

Sample : f=1
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

3208

Acq. Time : Apr 28 17:37 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Apr 28 17:57 2003
 Print Time : Tue Apr 29 09:51 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.10	9.03	0.007	96	70	685.618	10.00		Dev(Min)	
47	47 Cl-benzene-d5, I2	12.75	12.66	0.007	82	119	180.325	10.00		0.06	
62	62 1,4-DCB-d4 150 15	15.24	15.15	0.005	152	150	143.101	10.00		0.09	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.52	7.44	0.006	111	113	457.593	18.83		%Recovery	
29	29 1,2-di-Cl-ethane-	8.10	8.02	0.005	65	102	204.750	18.08		18.8	94.16%
55	55 toluene-d8(S2)	11.23	11.15	0.005	100	99	729.771	21.31		18.1	90.38%
70	70 4-Br-1-F-Bz (S3)	13.98	13.90	0.005	174	95	293.390	22.95		21.3	106.54%
										22.9	114.73%

Target Compounds

<<< 11 : ISTD ID = 1 >>>											
111 111 isopropyl alcoho	4.31	4.27	0.004	45	43	2.547	8.23			8.2	1
119 119 methyl acetate	5.04	5.06	-0.001	43	74	1.033	0.39			0.4	56
91 91 2-butanone MEKx10	6.83	6.83	0.000	43	72	3.133	0.49			0.5	74
201 201 Ethyl acetate x2	7.35	7.31	0.004	43	61	2.981	0.33			0.3	34
117 117 Iso-butyl alcoho	7.35	7.31	0.004	43	42	2.981	1.63			1.6	1
48 48 112-tri-Cl-Et	11.24	11.03	0.024	97	83	22.049	2.47			2.5	4
<<< 12 : ISTD ID = 47 >>>											
49 49 1,3-di-Cl-propane	11.23	11.29	-0.005	76	78	8.515	0.72			0.7	82
<<< 13 : ISTD ID = 62 >>>											
87 87 124-tri-Cl-Bz	17.33	17.24	0.006	180	182	1.111	0.69			0.7	76

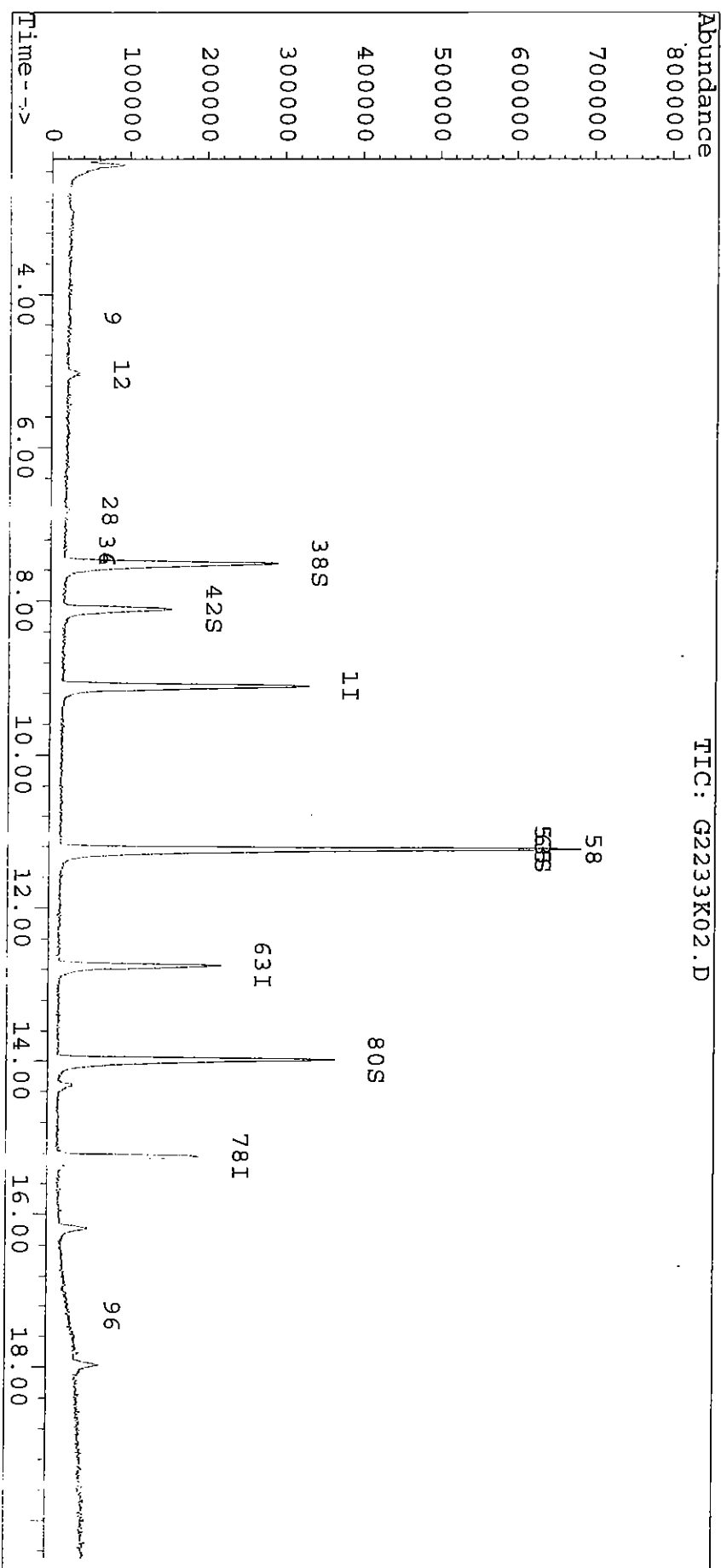
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

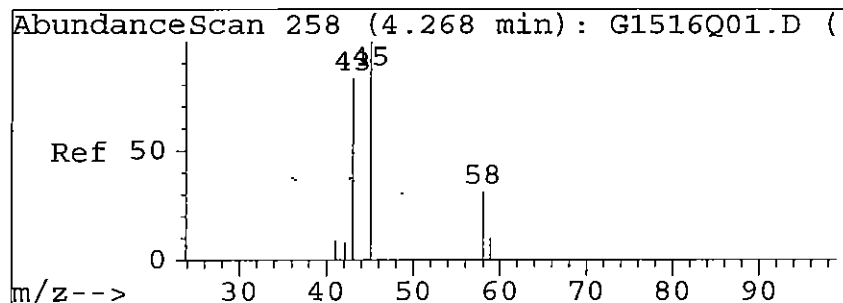
Data File : C:\HPCHEM\1\DATA\03G2233\G2233K02.D
Acq On : 28 Apr 03 5:37 pm
Sample : f=1

Misc :
Quant Time: Apr 28 17:57 2003

Vial: 25
Operator: Eddie
Inst : GCMS-G
Multiplr: 1.00
Quant Results File: quant.res

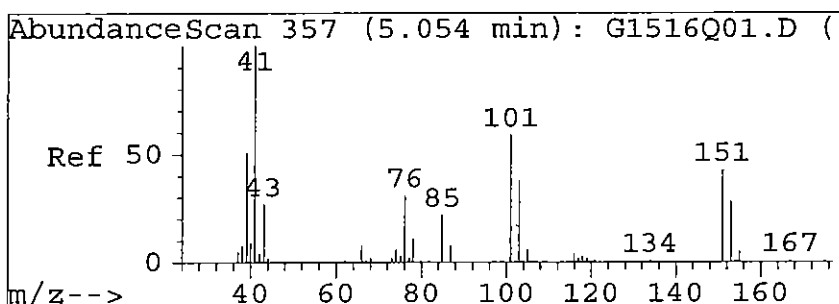
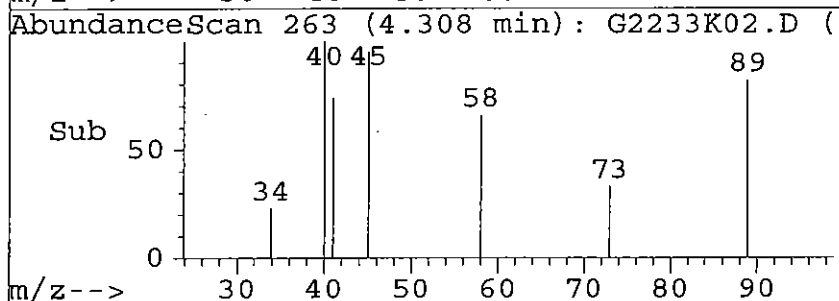
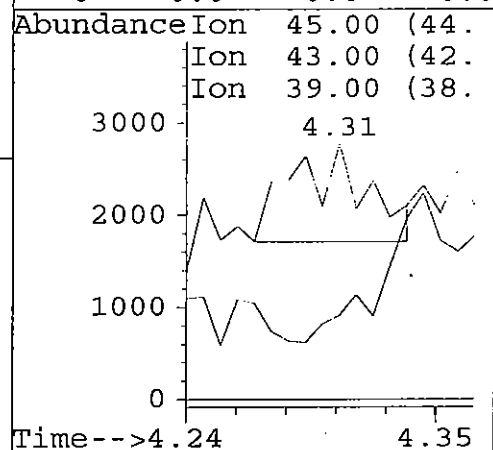
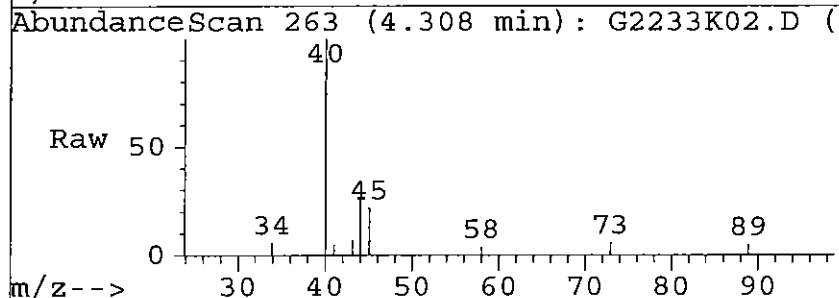
Method : C:\HPCHEM\1\METHODS\E524G003.M
Title : **Applied P & Ch Lab** EPA 524.2
Last Update : Mon Jan 13 10:38:23 2003
Response via : Multiple Level Calibration





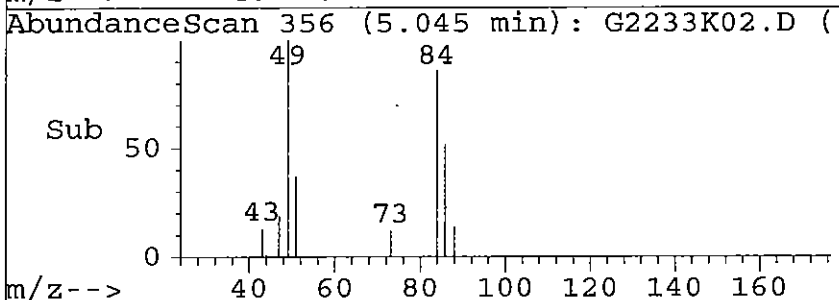
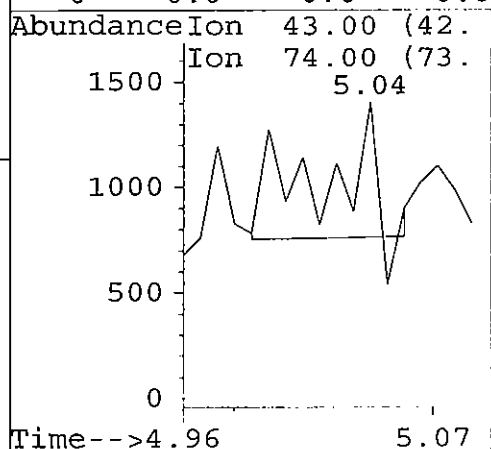
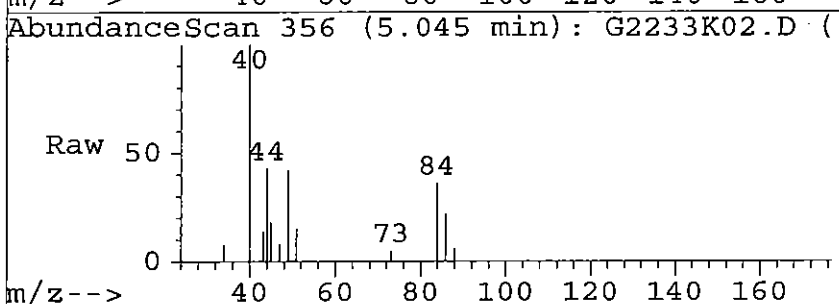
#9
111 isopropyl alcohol x10
Concen: 8.23 ppb
RT: 4.31 min Scan# 263
Delta R.T. 0.04 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

Tgt Ion:45 Resp: 2547
Ion Ratio Lower Upper
45 100
43 21.9 395.5 593.2#
39 0.0 28.3 42.4#
0 0.0 0.0 0.0

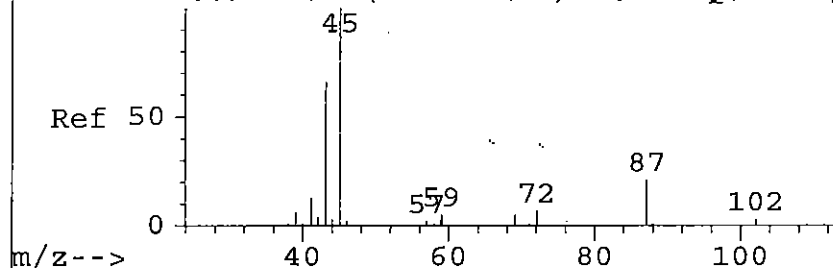


#12
119 methyl acetate
Concen: 0.39 ppb
RT: 5.04 min Scan# 356
Delta R.T. -0.01 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

Tgt Ion:43 Resp: 1033
Ion Ratio Lower Upper
43 100
74 0.0 0.6 40.6#
0 0.0 0.0 0.0
0 0.0 0.0 0.0



AbundanceScan 581 (6.830 min): G1516Q01.D (



#28

91 2-butanone MEKx10

Concen: 0.49 ppb

RT: 6.83 min Scan# 581

Delta R.T. -0.01 min

Lab File: G2233K02.D

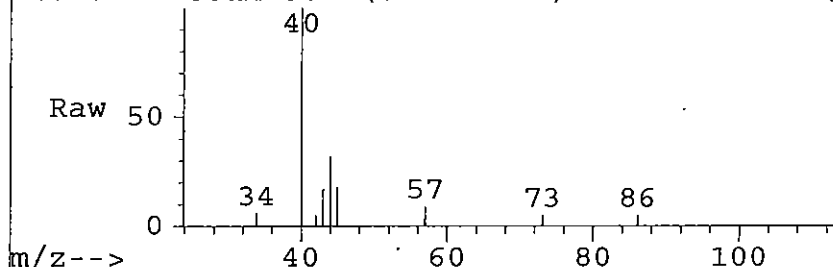
Acq: 28 Apr 03 5:37 pm

Tgt Ion:43 Resp: 3133

Ion Ratio Lower Upper

43	100		
72	0.0	7.7	11.6#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

AbundanceScan 581 (6.826 min): G2233K02.D (



AbundanceIon 43.00 (42.

Ion 72.00 (71.

6.83

2000

1500

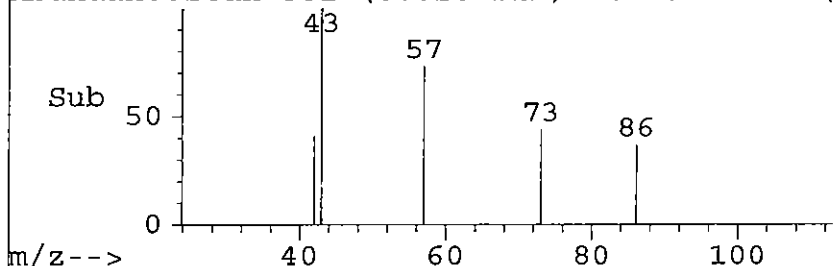
1000

500

0

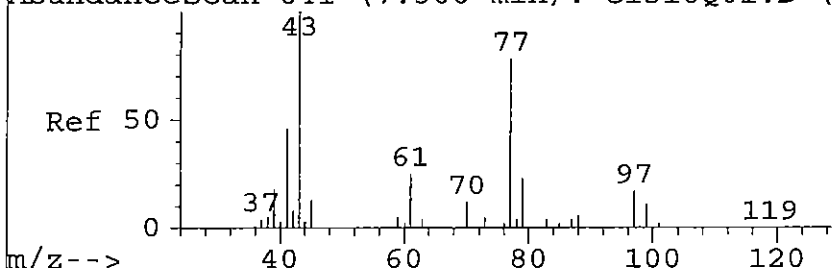
m/z-->

AbundanceScan 581 (6.826 min): G2233K02.D (



Time-->6.72 6.88

AbundanceScan 641 (7.306 min): G1516Q01.D (



#34

201 Ethyl acetate x2

Concen: 0.33 ppb

RT: 7.35 min Scan# 647

Delta R.T. 0.04 min

Lab File: G2233K02.D

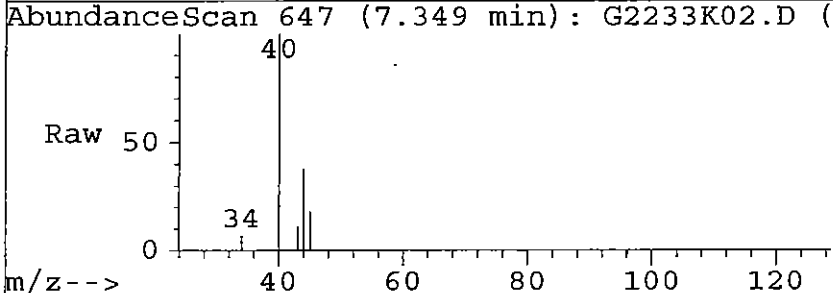
Acq: 28 Apr 03 5:37 pm

Tgt Ion:43 Resp: 2981

Ion Ratio Lower Upper

43	100		
61	0.0	33.7	50.6#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

AbundanceScan 647 (7.349 min): G2233K02.D (



AbundanceIon 43.00 (42.

Ion 61.00 (60.

7.35

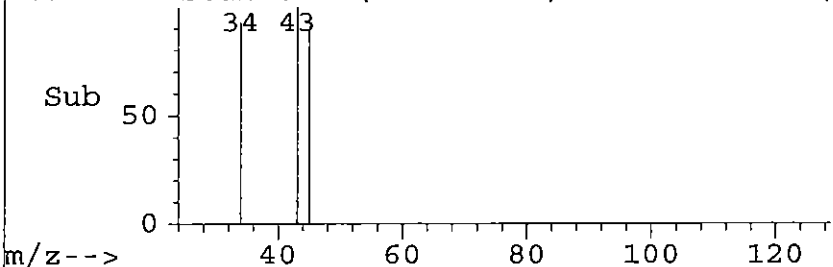
1000

500

0

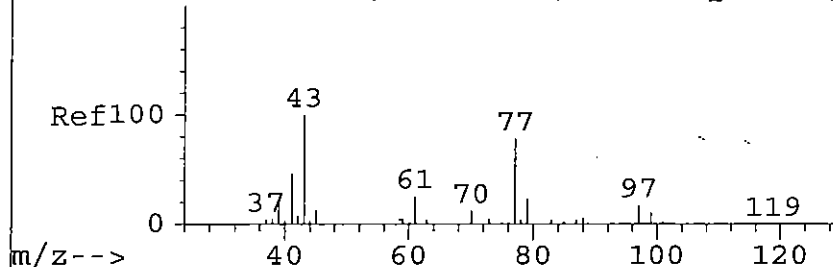
m/z-->

AbundanceScan 647 (7.349 min): G2233K02.D (

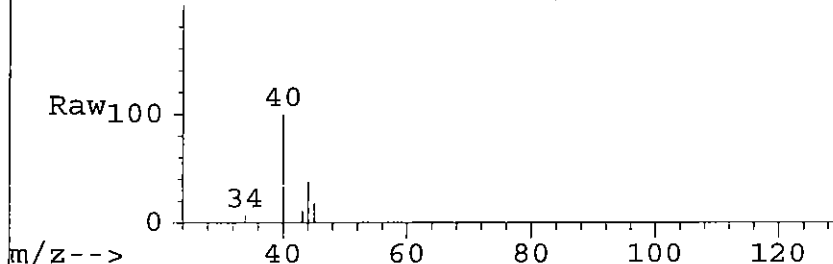


Time-->7.25 7.38

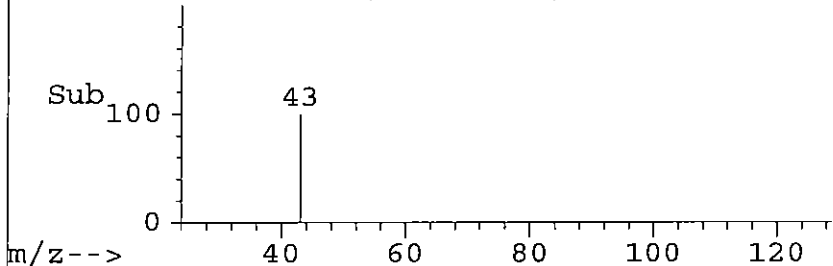
AbundanceScan 641 (7.306 min): G1516Q01.D (



AbundanceScan 647 (7.349 min): G2233K02.D (

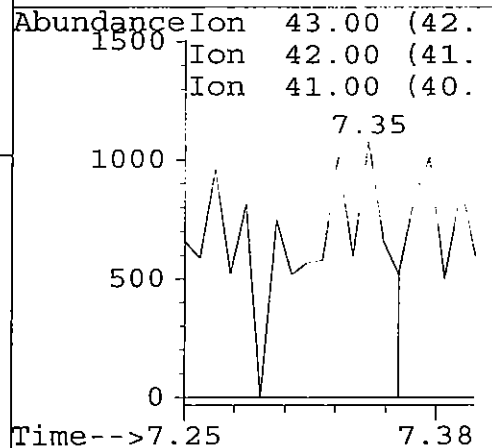


AbundanceScan 647 (7.349 min): G2233K02.D (

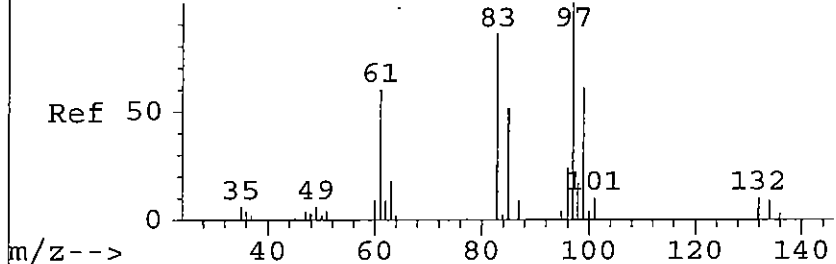


#36
117 Iso-butyl alcohol X10
Concen: 1.63 ppb
RT: 7.35 min Scan# 647
Delta R.T. 0.04 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

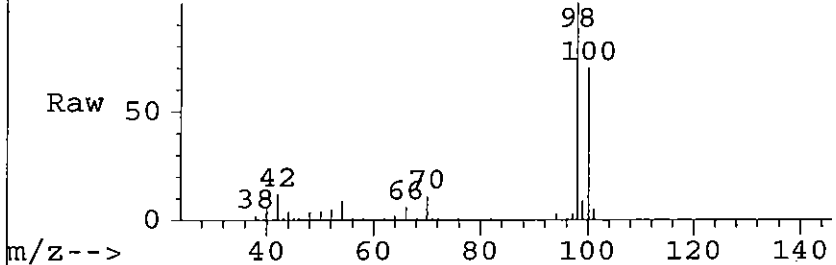
Tgt Ion:	43	Resp:	2981
Ion	Ratio	Lower	Upper
43	100		
42	0.0	21.1	31.6#
41	0.0	161.8	242.7#
0	0.0	0.0	0.0



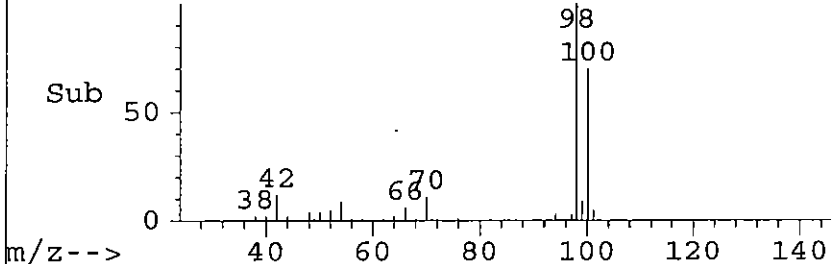
AbundanceScan 1111 (11.036 min): G1516Q01.D



AbundanceScan 1139 (11.245 min): G2233K02.D

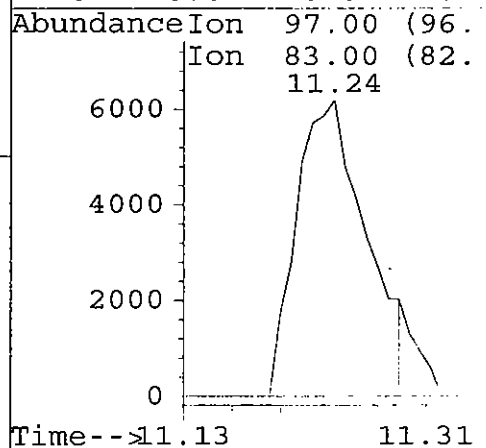


AbundanceScan 1139 (11.245 min): G2233K02.D

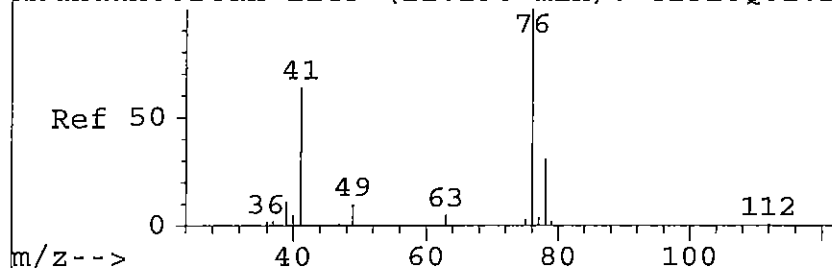


#58
48 112-tri-Cl-Et 97 83
Concen: 2.47 ppb
RT: 11.24 min Scan# 1139
Delta R.T. 0.22 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

Tgt Ion:	97	Resp:	22049
Ion	Ratio	Lower	Upper
97	100		
83	0.0	46.0	137.9#
0	0.0	0.0	0.0
0	0.0	0.0	0.0



AbundanceScan 1143 (11.290 min): G1516Q01.D

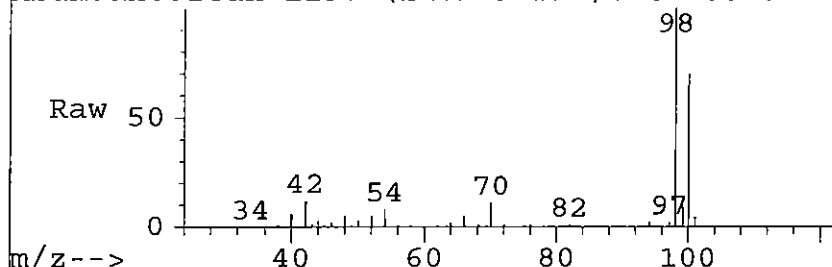


#65

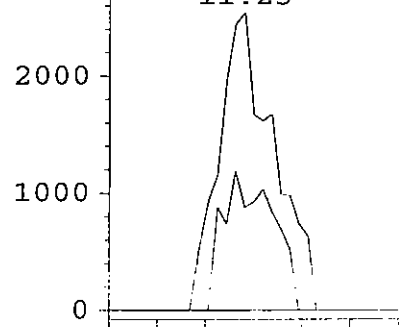
49 1,3-di-cl-propane 76 78
Concen: 0.72 ppb
RT: 11.23 min Scan# 1137
Delta R.T. -0.06 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

Tgt Ion:	76	Resp:	8515
Ion	Ratio	Lower	Upper
76	100		
78	43.1	26.5	39.8#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

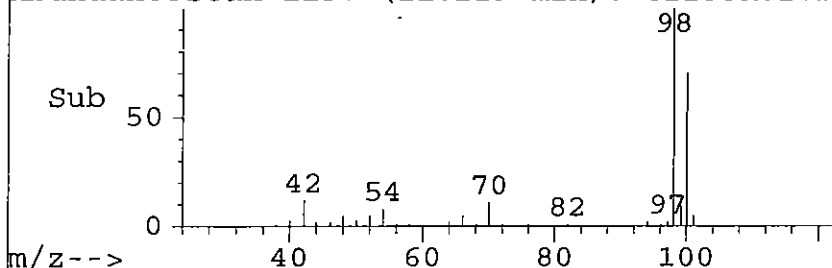
AbundanceScan 1137 (11.229 min): G2233K02.D



AbundanceIon 76.00 (75.
3000 Ion 78.00 (77.
11.23

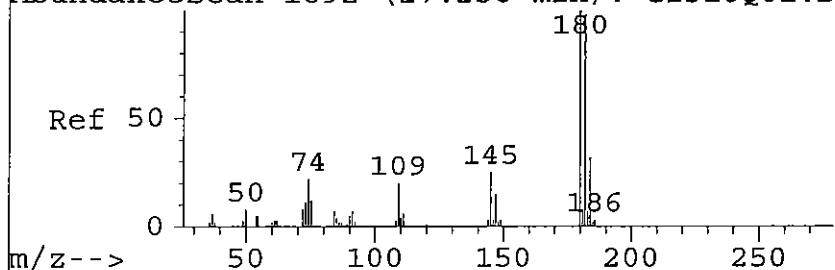


AbundanceScan 1137 (11.229 min): G2233K02.D



Time-->11.11 11.32

AbundanceScan 1892 (17.236 min): G1516Q01.D

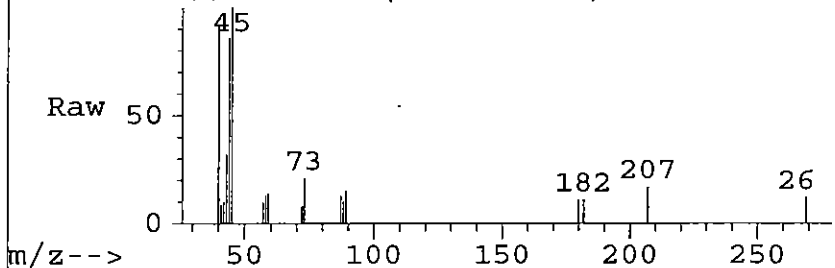


#96

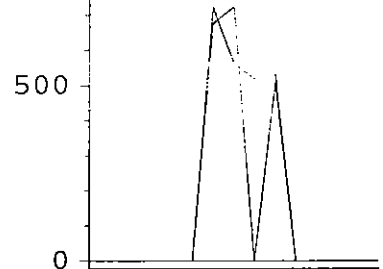
87 124-tri-Cl-Bz 180 182
Concen: 0.69 ppb
RT: 17.33 min Scan# 1907
Delta R.T. 0.08 min
Lab File: G2233K02.D
Acq: 28 Apr 03 5:37 pm

Tgt Ion:	180	Resp:	1111
Ion	Ratio	Lower	Upper
180	100		
182	81.6	47.8	143.3
184	0.0	15.1	45.1#
0	0.0	0.0	0.0

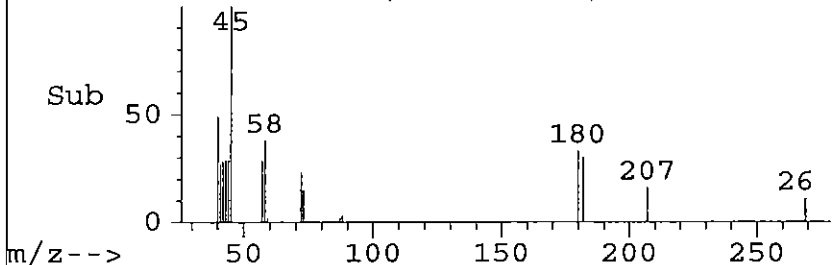
AbundanceScan 1907 (17.328 min): G2233K02.D



AbundanceIon 180.00 (179
1000 Ion 182.00 (181
Ion 184.00 (183
17.33



AbundanceScan 1907 (17.328 min): G2233K02.D



Time-->17.28 17.37

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: DUPE-2-2Q03	Lab Sample ID: 03-2843-1	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: 2843-01	Prep. No: -	Anal. Time: 21:55
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	0.3	J
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	0.5	
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	103	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	88	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	89	
4	TOLUENE-D8	2037-26-5		73-129	100	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	105	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	99	
3	FLUOROBENZENE	462-06-6		50-200	104	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

E - Exceed calibration range

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

B - A positive value was found in the method blank

D - Diluted

Data Filename: C:\HPCHEM\1\DATA\03G2233\2843-01.D Sample : f=1 9
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 21:55 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Apr 28 22:16 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 09:47 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.08	9.03	0.005	96	70	759.953	10.00		0.05	
47	47 Cl-benzene-d5, I2	12.73	12.66	0.006	82	119	208.715	10.00		0.07	
62	62 1,4-DCB-d4 150 15	15.23	15.15	0.005	152	150	161.113	10.00		0.07	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.48	7.44	0.003	111	113	479.998	17.81	17.8	89.03%	
29	29 1,2-di-Cl-ethane-	8.07	8.02	0.003	65	102	220.128	17.53	17.5	87.66%	
55	55 toluene-d8(S2)	11.21	11.15	0.004	100	99	754.943	19.89	19.9	99.44%	
70	70 4-Br-1-F-Bz (S3)	13.96	13.90	0.004	174	95	296.702	20.61	20.6	103.06%	

Target Compounds	<<< I1 : ISTD ID = 1 >>>	Qvalue
111 111 isopropyl alcoho	4.26 4.27 0.000 45 43 1.256 3.66	3.7 1
119 119 methyl acetate	5.06 5.06 0.000 43 74 1.437 0.42	0.4 56
25 25 chloroform	7.29 7.27 0.003 83 85 12.016 0.30	0.3 87
117 117 Iso-butyl alcoho	7.31 7.31 0.000 43 42 2.068 1.02	1.0 1
40 40 trichloroethene	9.60 9.54 0.006 130 132 10.429 0.51	0.5 90
93 93 2-Hexanone x5	11.48 11.48 0.000 43 58 1.580 0.37	0.4 24
48 48 112-tri-Cl-Et	11.23 11.03 0.022 97 83 25.302 2.56	2.6 4
<<< I2 : ISTD ID = 47 >>>		
49 49 1,3-di-Cl-propane	11.21 11.29 -0.006 76 78 8.856 0.65	0.6 66

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

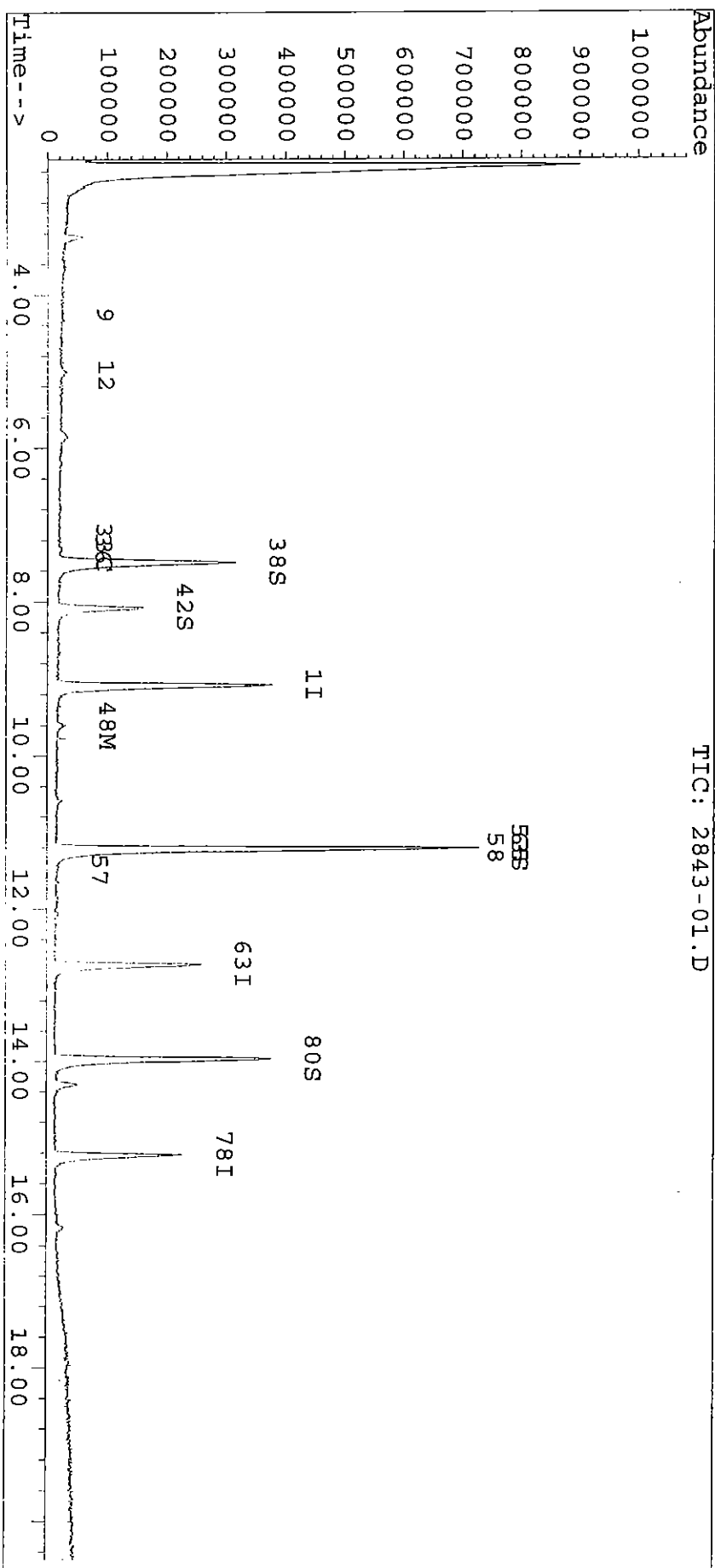
Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2233\2843-01.D
 Acq On : 28 Apr 03 9:55 pm
 Sample : f=1 9
 Misc :

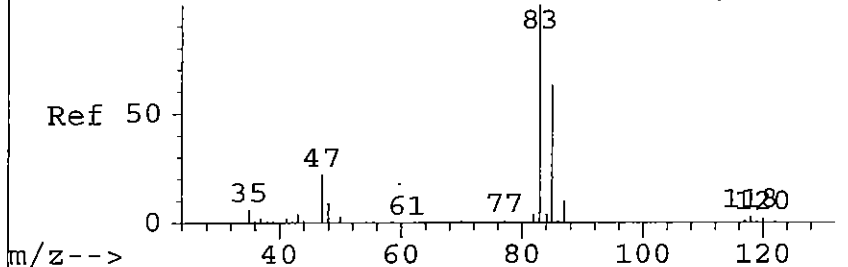
Quant Time: Apr 28 22:16 2003

Vial: 21
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Sch Lab** EPA 524.2
 Last Update : Mon Jan 13 10:38:23 2003
 Response via : Multiple Level Calibration



AbundanceScan 635 (7.258 min): G1516Q01.D (

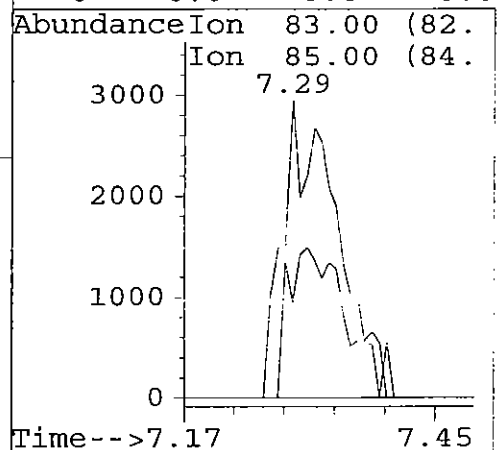
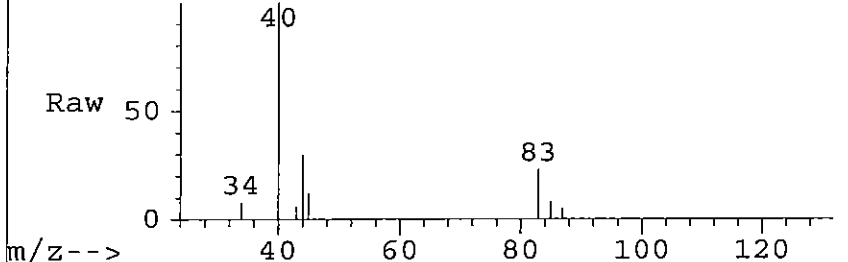


#33

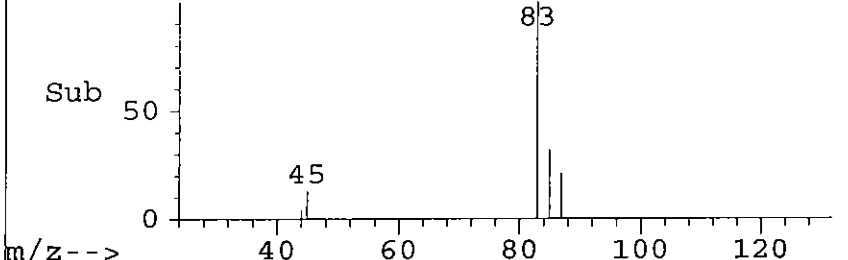
25 chloroform 83 85
 Concen: 0.30 ppb
 RT: 7.29 min Scan# 640
 Delta R.T. 0.02 min
 Lab File: 2843-01.D
 Acq: 28 Apr 03 9:55 pm

Tgt	Ion:83	Resp:	12016
Ion	Ratio	Lower	Upper
83	100		
85	55.5	32.8	98.3
0	0.0	0.0	0.0
0	0.0	0.0	0.0

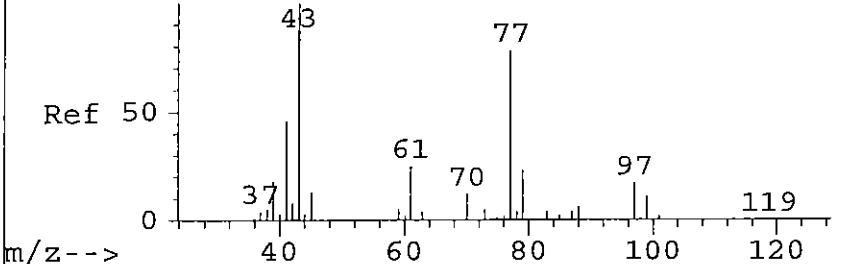
AbundanceScan 640 (7.293 min): 2843-01.D (*)



AbundanceScan 640 (7.293 min): 2843-01.D (-



AbundanceScan 641 (7.306 min): G1516Q01.D (

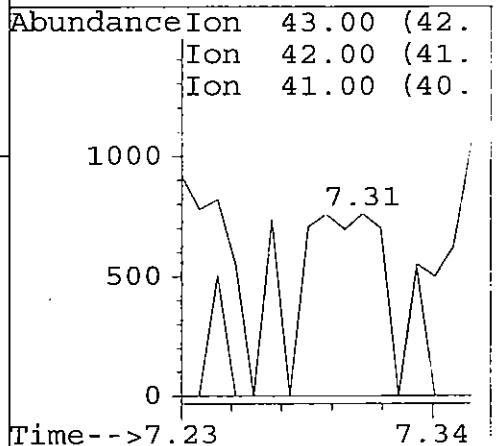
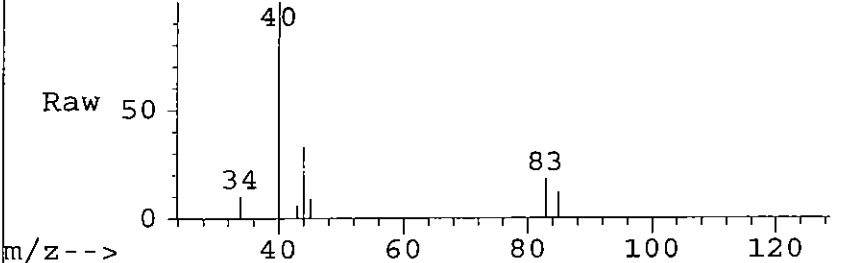


#36

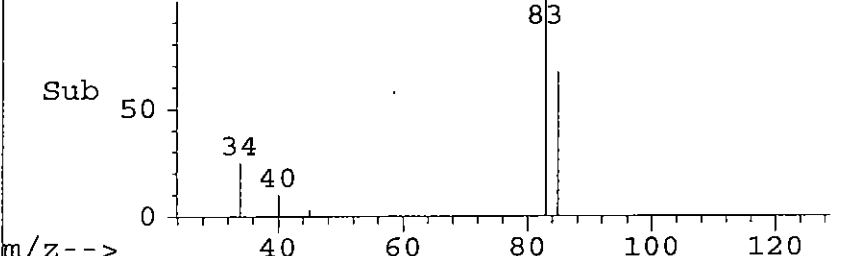
117 Iso-butyl alcohol X10
 Concen: 1.02 ppb
 RT: 7.31 min Scan# 642
 Delta R.T. -0.00 min
 Lab File: 2843-01.D
 Acq: 28 Apr 03 9:55 pm

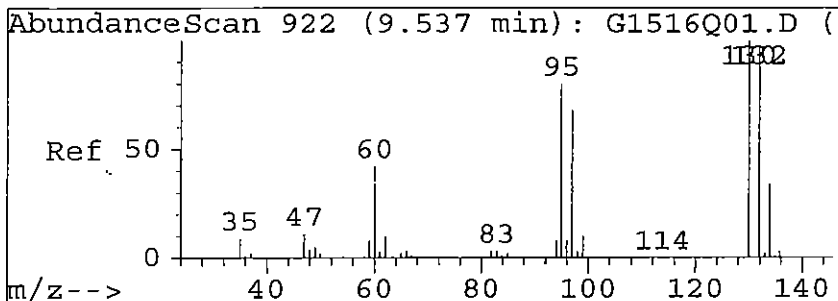
Tgt	Ion:43	Resp:	2068
	Ratio	Lower	Upper
43	100		
42	12.3	21.1	31.6#
41	0.0	161.8	242.7#
0	0.0	0.0	0.0

AbundanceScan 642 (7.309 min): 2843-01.D (*)



AbundanceScan 642 (7.309 min): 2843-01.D (-





#48

40 trichloroethene 130 132

Concen: 0.51 ppb

RT: 9.60 min Scan# 931

Delta R.T. 0.06 min

Lab File: 2843-01.D

Acq: 28 Apr 03 9:55 pm

Tgt Ion:130 Resp: 10429

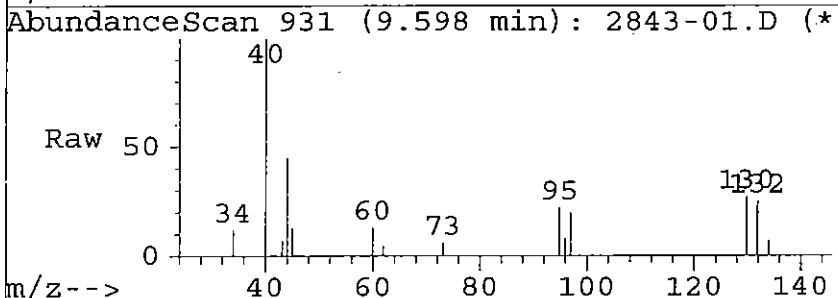
Ion Ratio Lower Upper

130 100

132 88.0 48.8 146.3

0 0.0 0.0 0.0

0 0.0 0.0 0.0



AbundanceIon 130.00 (129

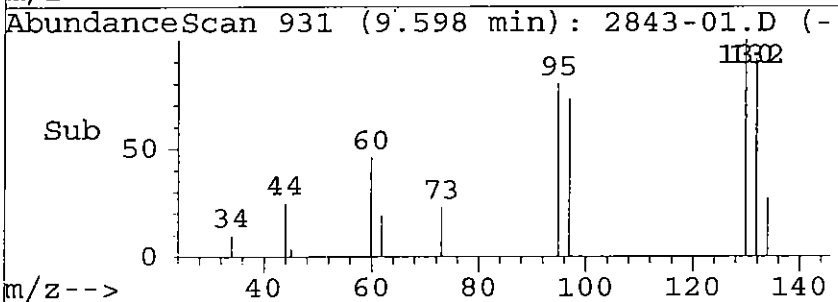
Ion 132.00 (131

3000 9.60

2000

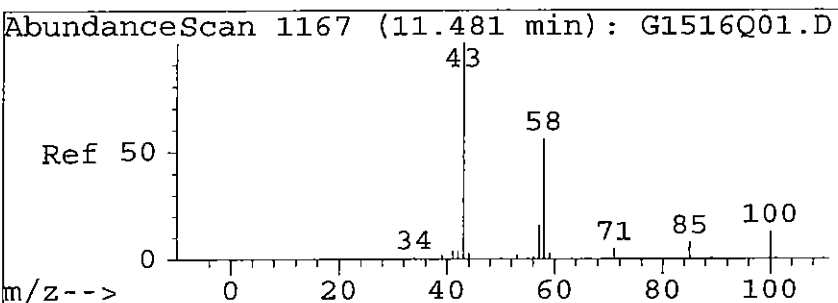
1000

0



Time-->9.48

9.72



#57

93 2-Hexanone x5

Concen: 0.37 ppb

RT: 11.48 min Scan# 1169

Delta R.T. 0.00 min

Lab File: 2843-01.D

Acq: 28 Apr 03 9:55 pm

Tgt Ion:43 Resp: 1580

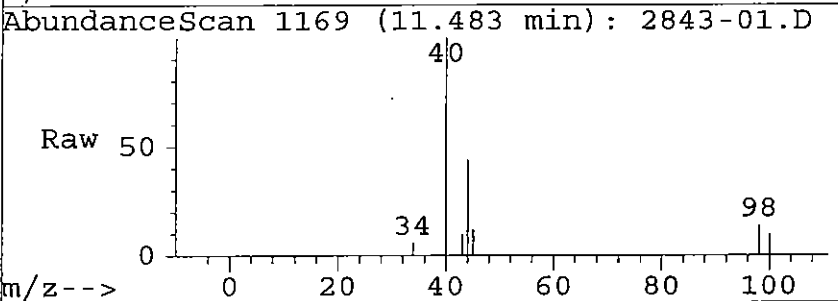
Ion Ratio Lower Upper

43 100

58 0.0 44.5 66.7#

0 0.0 0.0 0.0

0 0.0 0.0 0.0



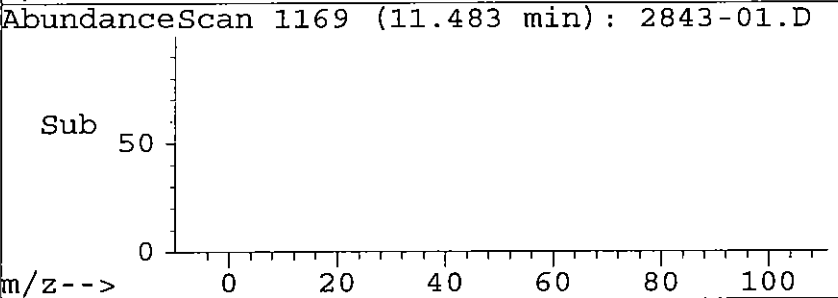
AbundanceIon 43.00 (42.

1000 Ion 58.00 (57.

11.48

500

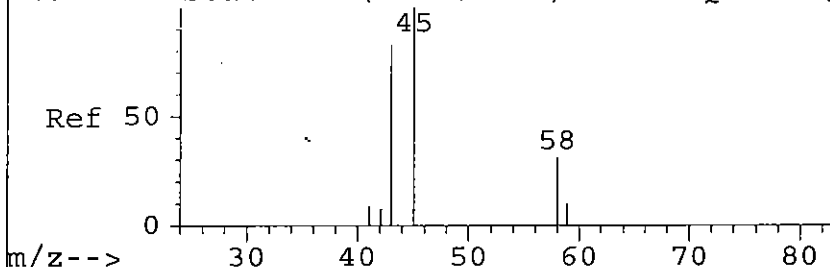
0



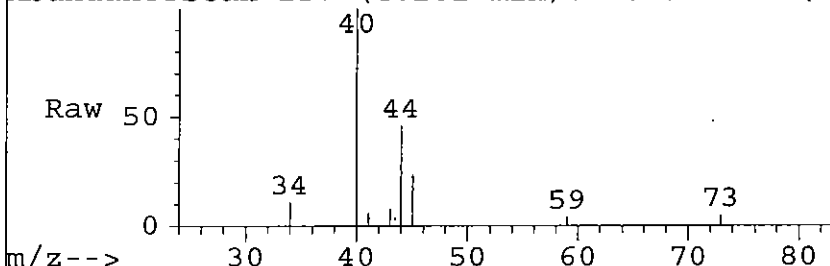
Time-->11.46

11.53

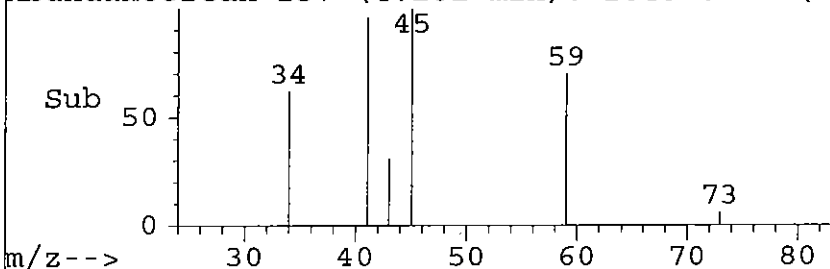
AbundanceScan 258 (4.268 min): G1516Q01.D (



AbundanceScan 257 (4.261 min): 2843-01.D (*)



AbundanceScan 257 (4.261 min): 2843-01.D (-

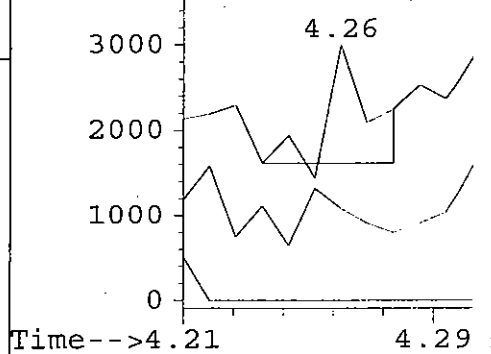


#9

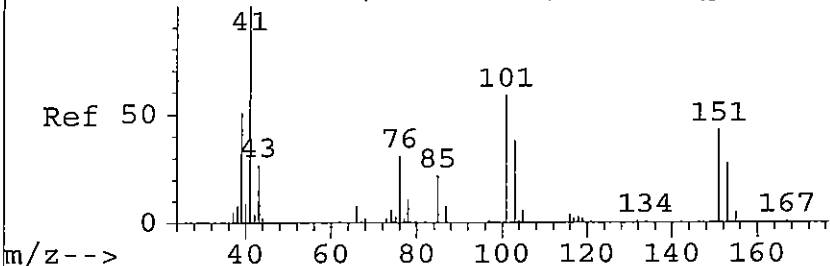
111 isopropyl alcohol x10
Concen: 3.66 ppb
RT: 4.26 min Scan# 257
Delta R.T. -0.01 min
Lab File: 2843-01.D
Acq: 28 Apr 03 9:55 pm

Tgt Ion:	45	Resp:	1256
Ion Ratio	Lower	Upper	
45	100		
43	58.1	395.5	593.2#
39	0.0	28.3	42.4#
0	0.0	0.0	0.0

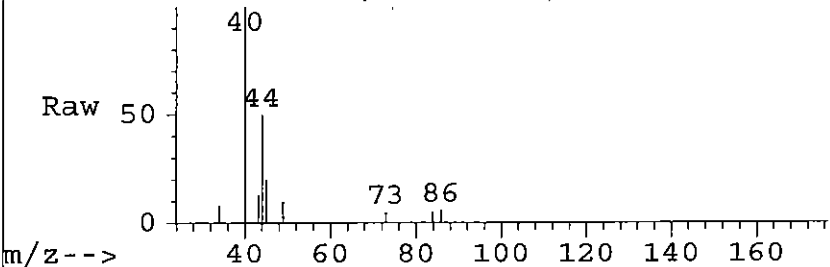
AbundanceIon 45.00 (44.
4000 Ion 43.00 (42.
Ion 39.00 (38.



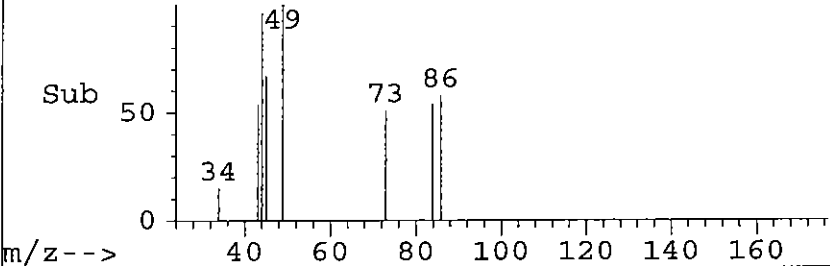
AbundanceScan 357 (5.054 min): G1516Q01.D (



AbundanceScan 358 (5.060 min): 2843-01.D (*)



AbundanceScan 358 (5.060 min): 2843-01.D (-

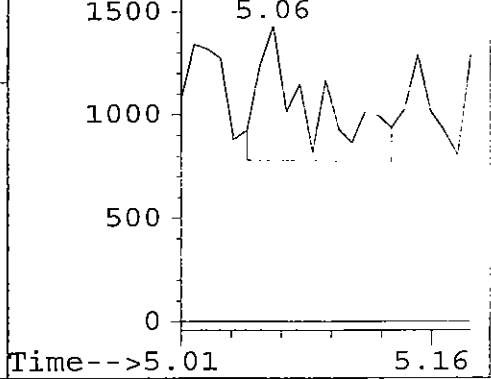


#12

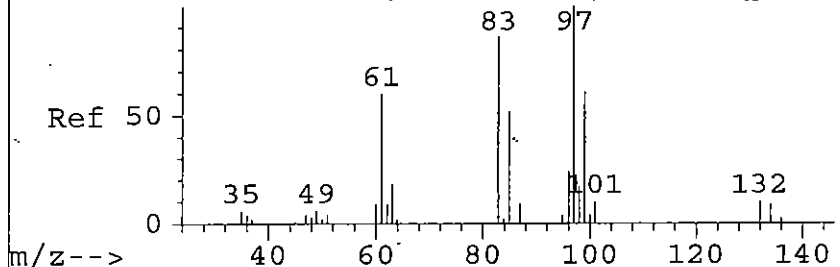
119 methyl acetate
Concen: 0.42 ppb
RT: 5.06 min Scan# 358
Delta R.T. 0.00 min
Lab File: 2843-01.D
Acq: 28 Apr 03 9:55 pm

Tgt Ion:	43	Resp:	1437
Ion Ratio	Lower	Upper	
43	100		
74	0.0	0.6	40.6#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

AbundanceIon 43.00 (42.
Ion 74.00 (73.



AbundanceScan 1111 (11.036 min): G1516Q01.D

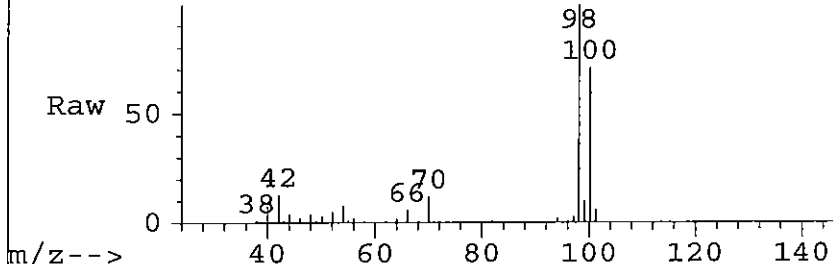


#58

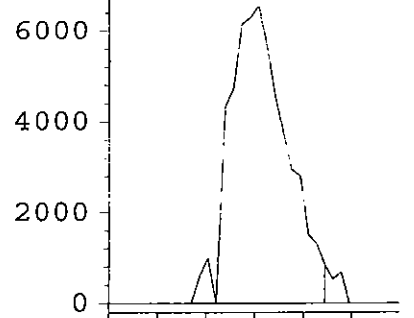
48 112-tri-Cl-Et 97 83
 Concen: 2.56 ppb
 RT: 11.23 min Scan# 1137
 Delta R.T. 0.20 min
 Lab File: 2843-01.D
 Acq: 28 Apr 03 9:55 pm

Tgt Ion:	97	Resp:	25302
Ion Ratio	Lower	Upper	
97	100		
83	0.0	46.0	137.9#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

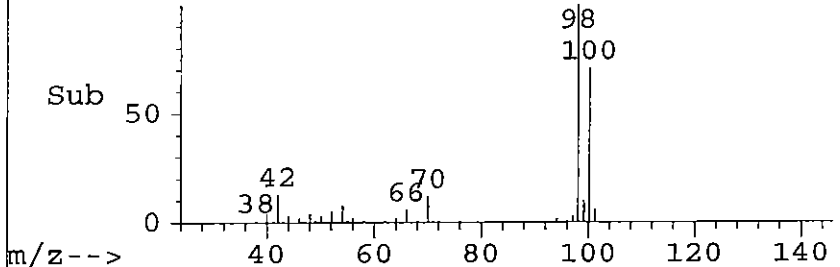
AbundanceScan 1137 (11.229 min): 2843-01.D



AbundanceIon 97.00 (96.
 Ion 83.00 (82.
 11.23

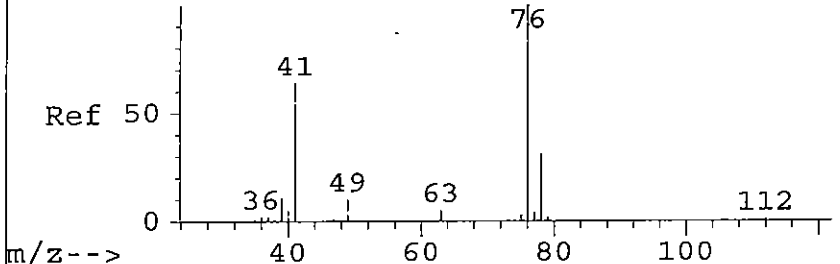


AbundanceScan 1137 (11.229 min): 2843-01.D



Time-->11.09 11.32

AbundanceScan 1143 (11.290 min): G1516Q01.D

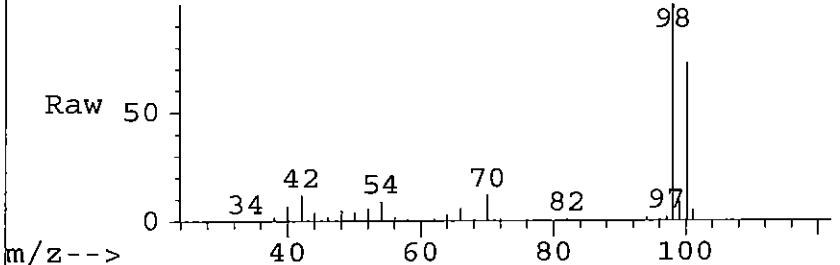


#65

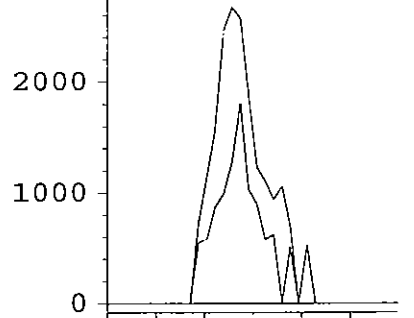
49 1,3-di-cl-propane 76 78
 Concen: 0.65 ppb
 RT: 11.21 min Scan# 1135
 Delta R.T. -0.08 min
 Lab File: 2843-01.D
 Acq: 28 Apr 03 9:55 pm

Tgt Ion:	76	Resp:	8856
Ion Ratio	Lower	Upper	
76	100		
78	52.2	26.5	39.8#
0	0.0	0.0	0.0
0	0.0	0.0	0.0

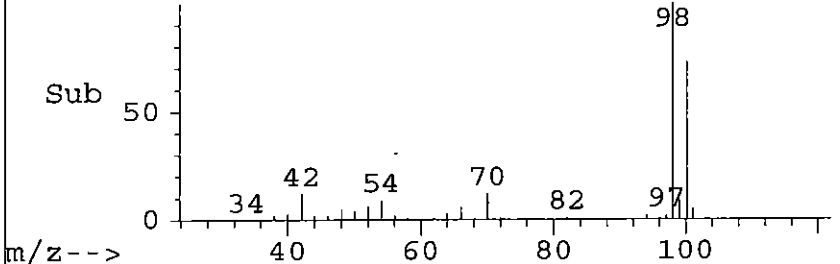
AbundanceScan 1135 (11.214 min): 2843-01.D



AbundanceIon 76.00 (75.
 Ion 78.00 (77.
 11.21



AbundanceScan 1135 (11.214 min): 2843-01.D



Time-->11.09 11.33

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: EB-4-4/23/03	Lab Sample ID: 03-2843-2	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: 2843-02	Prep. No: -	Anal. Time: 22:24
Methanol Vol. -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	<0.5	U
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U

Surrogates

		Control Limit, %	Surro. Rec.%
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	70-129	104
2	1,2-DICHLOROETHANE-D4	70-129	90
3	DIBROMOFLUOROMETHANE	70-122	91
4	TOLUENE-D8	73-129	101
#	of out-of-control		0

Internal Standard

		Control Limit, %	IS Rec.%
1	CHLOROBENZENE-D5	50-200	107
2	1,4-DICHLOROBENZENE-D4	50-200	103
3	FLUOROBENZENE	50-200	105
#	of out-of-control		0

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

E - Exceed calibration range

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: MW-14-1	Lab Sample ID: 03-2843-3	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: 2843-03	Prep. No: -	Anal. Time: 22:52
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	0.4	J
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	0.4	J
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	1.3	
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	113	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	98	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	97	
4	TOLUENE-D8	2037-26-5		73-129	109	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	98	
2	1,4-DICHLOROETHANE-D4	3855-82-1		50-200	91	
3	FLUOROBENZENE	462-06-6		50-200	94	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

E - Exceed calibration range

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: MW-14-2	Lab Sample ID: 03-2843-4	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: 2843-04	Prep. No: -	Anal. Time: 23:20
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	0.4	J
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	0.5	J
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	3.7	
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	101	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	88	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	87	
4	TOLUENE-D8	2037-26-5		73-129	98	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	108	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	104	
3	FLUOROBENZENE	462-06-6		50-200	106	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

E - Exceed calibration range

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: MW-14-3	Lab Sample ID: 03-2843-5	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/28/03	Anal. Date: 04/28/03
Data File Name: 2843-05	Prep. No: -	Anal. Time: 23:49
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	0.3	J
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	0.5	J
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	108	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	92	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	97	
4	TOLUENE-D8	2037-26-5		73-129	104	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	104	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	99	
3	FLUOROBENZENE	462-06-6		50-200	101	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

E - Exceed calibration range

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
	Lab Sample ID: 03-2843-6	Received Date: 04/23/2003
Sample ID: MW-14-4	Sample Matrix: Water	Moisture %: -
Sample Type: Field Sample	Prep. Method: 5030	Instrument ID: GC/MS: G
Anal. Method: 524.2	Prep. Date: 04/29/03	Anal. Date: 04/29/03
Batch No: 03G2233	Prep. No: -	Anal. Time: 00:17
Data File Name: 2843-06	Sample Amount: 25 mL	Dilution Factor: 1
Methanol Vol: -		
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	<0.5	U
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	115	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	95	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	97	
4	TOLUENE-D8	2037-26-5		73-129	109	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	101	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	96	
3	FLUOROBENZENE	462-06-6		50-200	101	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

E - Exceed calibration range

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: MW-14-5	Lab Sample ID: 03-2843-7	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/29/03	Anal. Date: 04/29/03
Data File Name: 2843-07	Prep. No: -	Anal. Time: 00:45
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	<0.5	U
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	105	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	86	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	89	
4	TOLUENE-D8	2037-26-5		73-129	99	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	108	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	103	
3	FLUOROBENZENE	462-06-6		50-200	108	
# of out-of-control					0	

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

E - Exceed calibration range

B - A positive value was found in the method blank

D - Diluted

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/23/2003
Project ID: JPL	Service ID: 32843	Collected by: Leo Williamson
Sample ID: TB-4-4/23/03	Lab Sample ID: 03-2843-8	Received Date: 04/23/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2233	Prep. Date: 04/29/03	Anal. Date: 04/29/03
Data File Name: 2843-08	Prep. No: -	Anal. Time: 01:14
Methanol Vol: -	Sample Amount: 25 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	<10	U
11	CARBON TETRACHLORIDE	56-23-5	µg/L	0.5	<0.5	U
12	CHLOROBENZENE	108-90-7	µg/L	0.5	<0.5	U
13	CHLORODIBROMOMETHANE	124-48-1	µg/L	0.5	<0.5	U
14	CHLOROETHANE	75-00-3	µg/L	0.5	<0.5	U
15	CHLOROFORM	67-66-3	µg/L	0.5	<0.5	U
16	CHLOROMETHANE	74-87-3	µg/L	0.5	<0.5	U
17	2-CHLOROTOLUENE	95-49-8	µg/L	0.5	<0.5	U
18	4-CHLOROTOLUENE	106-43-4	µg/L	0.5	<0.5	U
19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U

Surrogates

Control Limit, %

Surro. Rec.%

1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4
2	1,2-DICHLOROETHANE-D4	17060-07-0
3	DIBROMOFLUOROMETHANE	1868-53-7
4	TOLUENE-D8	2037-26-5
# of out-of-control		0

Control Limit, %

IS Rec.%

Internal Standard

1	CHLOROBENZENE-D5	3114-55-4
2	1,4-DICHLOROBENZENE-D4	3855-82-1
3	FLUOROBENZENE	462-06-6
# of out-of-control		0

Not Detected is shown as PQL, with dilution and moisture corrected if applicable.

^(a)MDL reported.

Qualifier: U - Not Detected or less than MDL

E - Exceed calibration range

J - Less than RL (PQL, EQL or CRDL), but greater than MDL, or an estimated result (e.g. for TIC)

B - A positive value was found in the method blank

D - Diluted

FORM-2A

Applied P & Ch Laboratory

Surrogate Recovery Summary for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

SDG Number: 032843

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2233

#	Client Sample No	Lab Sample ID	S1 % #	S2 % #	S3 % #	S4 % #	TOT OUT
1	03G2233-LCS-01	03G2233-LCS-01	112	87	99	110	0
2	MW-20-3MS	03-2866-5MS	105	86	99	107	0
3	MW-20-3MSD	03-2866-5MSD	108	83	96	108	0
4	03G2233-MB-01	03G2233-MB-01	115	91	94	107	0
5	DUPE-2-2Q03	03-2843-1	103	88	89	100	0
6	EB-4-4/23/03	03-2843-2	104	90	91	101	0
7	MW-14-1	03-2843-3	113	98	97	109	0
8	MW-14-2	03-2843-4	101	88	87	98	0
9	MW-14-3	03-2843-5	108	92	97	104	0
10	MW-14-4	03-2843-6	115	95	97	109	0
11	MW-14-5	03-2843-7	105	86	89	99	0
12	TB-4-4/23/03	03-2843-8	107	89	88	101	0
13							
14							
15							
16							
17							
18							
19							
20							
21							
22							
23							
24							
25							

QC Control Limit

S1 = 1-BROMO-4-FLUOROBENZENE (4-BROMOFL

70-129

S2 = 1,2-DICHLOROETHANE-D4

70-129

S3 = DIBROMOFLUOROMETHANE

70-122

S4 = TOLUENE-D8

73-129

Column to be used to flag recovery values:

* - Values outside of contract required QC Limits

D - Surrogate diluted out

I - Matrix Interference

FORM-3A

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32843

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2233

LCS Filename: G2233L01

Date Analyzed: 042803

Time Analyzed: 15:19

LCSD Filename: -

Date Analyzed: -

Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
BENZENE	µg/L	20	0	21.8	109	65-120
CHLOROBENZENE	µg/L	20	0	23.2	116	65-134
1,1-DICHLOROETHENE	µg/L	20	0	21.8	109	65-127
TOLUENE	µg/L	20	0	21.2	106	65-134
TRICHLOROETHENE	µg/L	20	0	21.8	109	67-122
# of Out-of-control					0	

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233L01.D

Sample : F=1

Method : C:\HPCHEM\1\METHODS\E524G003.M

Inst. : GCMS-G

Acq. Time : Apr 28 15:19 2003

RF via : Multiple Level Calibration

Method Update: Tue Apr 29 11:06 2003

Operator: Eddie

Quant. Time : Apr 29 11:18 2003

Multiplr: 1.000000

Print Time : Tue Apr 29 11:19 2003

Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.09	9.03	0.006	96	70	690.083	10.00		0.06	
47	47 Cl-benzene-d5, I2	12.75	12.66	0.007	82	119	189.127	10.00		0.09	
62	62 1,4-DCB-d4 150 15	15.24	15.15	0.006	152	150	154.498	10.00		0.09	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.49	7.44	0.004	111	113	484.829	19.84		19.8	%Recovery
29	29 1,2-di-Cl-ethane-	8.08	8.02	0.004	65	102	199.082	17.46		17.5	87.31%
55	55 toluene-d8 (S2)	11.22	11.15	0.005	100	99	756.236	21.94		21.9	109.69%
70	70 4-Br-1-F-Bz (S3)	13.98	13.90	0.006	174	95	308.803	22.37		22.4	111.85%

Target Compounds

Qvalue

<<< I1 : ISTD ID = 1 >>>											
3	3 di-Cl-di-F-methan	2.60	2.60	0.000	85	87	425.083	21.71		21.7	99
4	4 Chloromethane	2.79	2.78	0.001	50	52	291.426	17.35		17.4	98
9	9 F114 85 135	2.84	2.83	0.001	85	135	497.516	22.58		22.6	94
5	5 vinyl chloride	2.98	2.96	0.002	62	64	308.855	19.40		19.4	97
6	6 bromomethane	3.39	3.37	0.002	94	96	244.354	17.42		17.4	94
7	7 Chloroethane	3.55	3.52	0.004	64	66	251.754	21.35		21.4	100
8	8 tri-Cl-F-methane	4.18	4.14	0.005	101	103	649.857	24.14		24.1	99
111	111 isopropyl alcoho	4.31	4.27	0.005	45	43	40.681	130.65		130.7	86
100	100 ethyl ether x5	4.47	4.44	0.004	59	74	806.844	82.03		82.0	100
102	102 Acrolein x10	4.19	4.15	0.005	56	55	184.404	234.99		235.0	92
119	119 methyl acetate	5.10	5.06	0.005	43	74	236.044	27.81		27.8	98
104	104 Carbon disulfide	5.23	5.18	0.005	76	78	1103.688	19.19		19.2	99
103	103 Acrylonitrile x10	4.93	4.89	0.005	53	52	149.377	90.59		90.6	98
95	95 Acetone x10	4.35	4.31	0.005	43	58	216.038	182.62		182.6	94
108	108 F-113	5.07	5.02	0.005	151	101	544.515	26.32		26.3	98
13	13 11-dichloroethene	4.81	4.76	0.005	61	96	590.028	21.81		21.8	98
101	101 Acetonitrile x10	4.26	4.20	0.006	41	40	46.345	103.96		104.0	87
109	109 Iodomethane	4.85	4.80	0.005	142	127	558.250	25.71		25.7	100
113	113 Tert butyl alcoh	4.90	4.86	0.004	59	57	118.884	199.39		199.4	100

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

EPA 524.2

3239

75
m?
#?

Page 2

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233L01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 28 15:19 2003
 Method Update: Tue Apr 29 11:06 2003
 Quant. Time : Apr 29 11:18 2003
 Print Time : Tue Apr 29 11:19 2003
 Miscellaneous :

Sample : f=1
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

3240

ID	Component Name	R.T.	RT0	DRRT	Q1on	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.43	11.36	0.008	69	99	95.344	8.18	8.2	100	
93	2-Hexanone x5	11.56	11.48	0.009	43	58	230.796	58.88	58.9	94	
48	112-tri-Cl-Et	11.11	11.03	0.010	97	83	138.094	16.71	16.7	99	
58	1,2-di-br-ethane	11.91	11.82	0.009	107	109	143.623	16.28	16.3	99	
51	di-Br-Cl-methane	11.65	11.57	0.009	129	127	197.513	17.78	17.8	98	
46	t-13-di-Cl-propen	10.94	10.87	0.008	75	110	251.962	18.12	18.1	100	
105	1-Chlorohexane	12.72	12.64	0.009	55	93	456.191	26.92	26.9	98	
<<< 12 : ISTD ID = 47 >>>											
54	MIBK	10.59	10.52	0.006	43	58	101.758	17.02	17.0	98	
49	1,3-di-Cl-propane	11.36	11.29	0.006	76	78	245.388	19.85	19.8	97	
59	tetra-Cl-ethene	12.08	12.00	0.006	166	168	445.156	26.75	26.7	99	
60	chlorobenzene	12.77	12.70	0.006	112	77	641.644	23.21	23.2	97	
61	1112-tetra-Cl-Et	12.70	12.62	0.006	131	133	251.624	22.73	22.7	99	
64	ethylbenzene	12.98	12.90	0.006	91	106	1367.932	25.69	25.7	99	
65	m/p-Xylenes x2	13.18	13.10	0.006	91	106	2051.449	50.67	50.7	98	
99	1-4-di-Cl-butane	13.54	13.46	0.006	55	41	227.565	18.65	18.7	99	
52	bromoform	13.30	13.22	0.006	173	175	109.434	19.42	19.4	97	
66	styrene	13.50	13.43	0.006	104	78	660.950	23.12	23.1	99	
67	o-xylene	13.58	13.50	0.006	91	106	947.348	23.78	23.8	99	
68	1122-Tetra-Cl-Et	13.58	13.51	0.006	83	85	136.095	18.61	18.6	97	
110	t-1,4-dichloro-2	13.75	13.67	0.007	89	53	27.942	24.52	24.5	67	
106	Cl-benzyl	15.21	15.12	0.007	91	126	192.644	21.04	21.0	98	
<<< 13 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.72	13.64	0.005	110	97	33.951	18.05	18.0	89	
71	isopropylbenzene	13.93	13.86	0.005	105	120	1441.682	28.40	28.4	100	
72	bromobenzene	14.17	14.10	0.005	156	158	249.664	22.40	22.4	98	
73	n-propylbenzene	14.36	14.29	0.005	120	78	389.376	28.45	28.5	99	
74	2-Cl-Tl	14.46	14.37	0.006	126	128	209.407	23.03	23.0	98	
75	4-Cl-Tl	14.52	14.45	0.005	126	128	302.445	24.33	24.3	94	
76	135-tri-Me-Bz	14.64	14.57	0.005	105	120	1054.677	25.73	25.7	98	
79	tert-butylbenzene	14.91	14.84	0.005	119	91	1179.884	28.04	28.0	99	
78	124-tri-Me-Bz	15.02	14.94	0.005	105	120	854.513	24.52	24.5	97	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233L01.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 15:19 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:18 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:19 2003
 Miscellaneous :

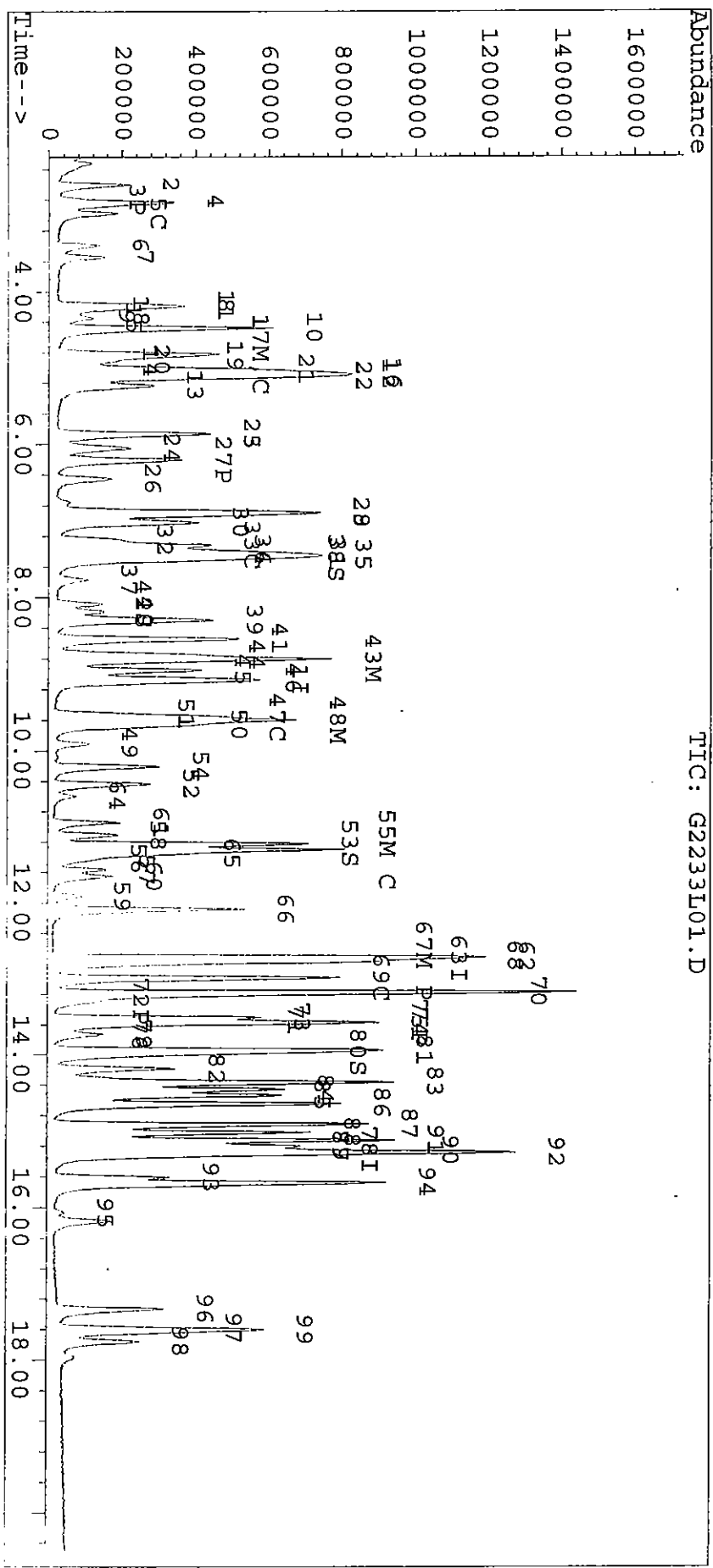
ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.21	15.11	0.006	146	148	375.345	21.14	21.1	91
82	14-di-Cl-Bz	146	15.26	15.19	0.005	146	148	573.822	22.24	22.2	93
81	sec-butylbenzene		15.13	15.04	0.006	105	134	1707.466	27.75	27.7	100
77	4-iso-Pr-toluene		15.30	15.22	0.005	119	134	1216.041	27.39	27.4	100
84	12-di-Cl-benzene		15.61	15.52	0.006	146	148	361.151	21.08	21.1	97
85	n-butylbenzene		15.68	15.60	0.005	91	134	1255.529	26.98	27.0	100
86	12-diBr-3-Cl-Pra		16.07	15.97	0.006	157	155	22.331	17.79	17.8	92
87	124-tri-Cl-Bz		17.33	17.24	0.006	180	182	288.573	21.35	21.3	99
88	naphthalene		17.57	17.49	0.005	128	129	219.109	21.32	21.3	99
90	123-tri-Cl-Bz		17.76	17.68	0.005	180	182	211.332	22.64	22.6	99
89	hx-Cl-butadiene		17.61	17.52	0.005	225	260	314.462	30.99	31.0	95

Phylog

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2233\G2233L01.D Vial: 12
 Acq On : 28 Apr 03 3:19 pm Operator: Eddie
 Sample : f=1 Inst : GCMS-G
 Misc : Multiplier: 1.00
 Quant Time: Apr 29 11:18 2003 Quant Results File: quant.res

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Tue Apr 29 11:06:45 2003
 Response via : Multiple Level Calibration



FORM-3A

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32843

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2233

MS Filename: G2233M01

Date Analyzed: 042803

Time Analyzed: 15:48

MSD Filename: G2233N01

Date Analyzed: 042803

Time Analyzed: 16:17

MS Sample No: MW-20-3

Sample Lab ID: 03-2866-5

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
BENZENE	µg/L	20	0	21.5	108	65-121
CHLOROBENZENE	µg/L	20	0	22.5	113	65-134
1,1-DICHLOROETHENE	µg/L	20	0	21.3	107	65-127
TOLUENE	µg/L	20	0	21.0	105	65-134
TRICHLOROETHENE	µg/L	20	0	21.1	106	65-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
BENZENE	µg/L	20	21.3	107	1	28	65-121
CHLOROBENZENE	µg/L	20	23.8	119	5	35	65-134
1,1-DICHLOROETHENE	µg/L	20	21.2	106	1	31	65-127
TOLUENE	µg/L	20	21.1	106	1	35	65-134
TRICHLOROETHENE	µg/L	20	21.4	107	1	30	65-125
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233M01.D Sample : f=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 15:48 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:27 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:28 2003
 Miscellaneous :

3244

ID	Component Name	R.T.	RT0	DRRT	Q1on	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.11	9.03	0.008	96	70	702.271	10.00		0.07	
47	47 Cl-benzene-d5, I2	12.75	12.66	0.007	82	119	198.407	10.00		0.08	
62	62 1,4-DCB-d4 150 15	15.25	15.15	0.006	152	150	163.321	10.00		0.09	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.51	7.44	0.005	111	113	492.186	19.79	19.8	98.94%	
29	29 1,2-di-Cl-ethane-	8.10	8.02	0.005	65	102	199.130	17.16	17.2	85.81%	
55	55 toluene-d8(S2)	11.23	11.15	0.005	100	99	747.685	21.31	21.3	106.57%	
70	70 4-Br-1-F-Bz (S3)	13.97	13.90	0.005	174	95	306.153	20.98	21.0	104.90%	

Target Compounds

Qvalue

<<< I1 : ISTD ID = 1 >>>											
3	3 di-Cl-di-F-methan	2.61	2.60	0.001	85	87	420.819	21.10	21.1	99	
4	4 Chloromethane	2.80	2.78	0.002	50	52	289.349	16.93	16.9	97	
9	9 F114 85 135	2.84	2.83	0.001	85	135	498.925	22.25	22.2	93	
5	5 vinyl chloride	2.97	2.96	0.001	62	64	302.199	18.65	18.6	100	
6	6 bromomethane	3.39	3.37	0.002	94	96	258.619	18.14	18.1	98	
7	7 Chloroethane	3.55	3.52	0.004	64	66	252.389	21.03	21.0	99	
8	8 tri-Cl-F-methane	4.18	4.14	0.005	101	103	653.032	23.83	23.8	100	
111	111 isopropyl alcoho	4.30	4.27	0.004	45	43	54.278	171.29	171.3	59	
100	100 ethyl ether x5	4.49	4.44	0.005	59	74	805.564	80.48	80.5	99	
102	102 Acrolein x10	4.20	4.15	0.005	56	55	146.113	182.97	183.0	91	
119	119 methyl acetate	5.11	5.06	0.006	43	74	234.595	27.16	27.2	95	
104	104 Carbon disulfide	5.24	5.18	0.006	76	78	1107.564	18.92	18.9	99	
103	103 Acrylonitrilex10	4.94	4.89	0.005	53	52	260.371	155.16	155.2	99	
95	95 Acetone x10	4.35	4.31	0.005	43	58	206.559	170.71	170.7	90	
108	108 F-113	5.07	5.02	0.005	151	101	542.704	25.78	25.8	99	
13	13 11-dichloroethene	4.82	4.76	0.006	61	96	586.494	21.30	21.3	99	
101	101 Acetonitrilex10	4.26	4.20	0.007	41	40	77.522	172.24	172.2	81	
109	109 Iodomethane	4.84	4.80	0.005	142	127	615.349	27.98	28.0	99	
113	113 Tert butyl alcoh	4.91	4.86	0.005	59	57	114.448	188.52	188.5	97	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233M01.D Sample : f=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 15:48 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:27 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:28 2003
 Miscelaneous :

3245

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.02	4.97	0.005	84	49	485.060	17.30	17.3	97	
112	Allyl chloride	5.12	5.07	0.005	41	76	646.442	21.23	21.2	95	?
200	200 Nitro methane x1	5.87	5.82	0.005	61	46	649.107	211.85	211.9	94	#?
10	10 t-Bu-Me-ether	6.06	6.00	0.006	73	57	542.924	18.26	18.3	96	
19	19 t-12-di-Cl-ethene	5.87	5.82	0.005	96	61	429.249	20.34	20.3	95	?
98	98 Vinyl acetate x5	6.47	6.41	0.007	43	86	1020.343	53.72	53.7	98	
21	21 11-dichloroethane	6.22	6.15	0.007	63	83	842.148	22.85	22.9	98	
91	91 2-butanone MEKx10	6.91	6.83	0.008	43	72	1083.791	164.55	164.6	99	?
115	115 Di isoprop ether	6.92	6.85	0.008	45	87	1783.146	19.17	19.2	98	?
22	22 c-12-di-Cl-ethene	7.04	6.97	0.008	96	61	399.002	19.32	19.3	95	
23	23 22-Dichloropropan	7.43	7.35	0.009	77	97	701.128	22.05	22.0	100	
24	24 Br-Cl-methane	7.27	7.18	0.010	128	130	120.298	16.24	16.2	100	
25	25 chloroform	7.33	7.27	0.007	83	85	725.508	19.68	19.7	100	
201	201 Ethyl acetate x2	7.38	7.31	0.008	43	61	233.279	24.85	24.8	77	#?
116	116 ETBE	7.47	7.40	0.008	59	87	997.644	17.61	17.6	99	
117	117 Iso-butyl alcoho	7.38	7.31	0.008	43	42	233.554	124.31	124.3	64	#?
26	26 tetrahydrofuranx5	7.78	7.71	0.008	72	42	56.233	78.14	78.1	90	
34	34 111-tri-Cl-ethane	8.31	8.23	0.009	97	99	687.107	22.37	22.4	99	
30	30 12-dichloroethane	8.19	8.13	0.007	62	64	258.187	17.97	18.0	96	
35	35 11-Di-Cl-propene	8.55	8.48	0.008	75	110	591.923	22.05	22.1	98	
36	36 benzene	8.83	8.75	0.009	78	52	1312.041	21.50	21.5	100	
37	37 CCl4	8.76	8.68	0.009	117	119	552.642	23.51	23.5	99	
97	97 thiophene	8.97	8.89	0.009	84	58	578.945	19.08	19.1	98	
118	118 TAME	9.07	9.00	0.008	73	43	667.118	17.10	17.1	98	
39	39 12-di-Cl-propane	9.57	9.49	0.008	63	76	356.088	19.94	19.9	97	
40	40 trichloroethene	9.62	9.54	0.009	130	132	398.530	21.08	21.1	97	
96	96 Me-methacrylate	9.92	9.85	0.007	69	100	110.986	17.91	17.9	100	
42	42 Br-di-Cl-methane	9.69	9.61	0.008	83	85	443.157	17.79	17.8	97	
41	41 dibromomethane	9.53	9.45	0.008	174	172	151.035	17.53	17.5	96	
45	45 c-13-di-Cl-propen	10.45	10.37	0.009	75	110	395.821	18.02	18.0	98	
92	92 2-ClEt-Vi-ether10	10.22	10.15	0.008	63	43	317.298	84.24	84.2	98	
56	56 toluene	11.31	11.23	0.009	91	92	1202.580	20.97	21.0	100	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233M01.D Sample : F=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 15:48 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:27 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:28 2003
 Miscellaneous :

3246

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.44	11.36	0.009	69	99	155.802	13.13	13.1	93	
93	2-Hexanone x5	11.57	11.48	0.009	43	58	313.946	78.70	78.7	95	
48	112-tri-Cl-Et	11.11	11.03	0.009	97	83	146.374	17.41	17.4	96	
58	1,2-di-br-ethane	11.91	11.82	0.009	107	109	149.298	16.63	16.6	99	
51	di-Br-Cl-methane	11.65	11.57	0.009	129	127	204.804	18.12	18.1	99	
46	t-13-di-Cl-propen	10.95	10.87	0.009	75	110	253.335	17.90	17.9	99	
105	1-Chlorohexane	12.72	12.64	0.009	55	93	443.431	25.67	25.7	100	
<<< 12 : ISTD ID = 47 >>>											
54	MIBK	10.61	10.52	0.007	43	58	116.256	18.59	18.6	94	
49	1,3-di-Cl-propane	11.37	11.29	0.006	76	78	258.081	19.90	19.9	99	
59	tetra-Cl-ethene	12.09	12.00	0.007	166	168	434.443	24.88	24.9	98	
60	chlorobenzene	12.78	12.70	0.006	112	77	651.486	22.47	22.5	99	
61	112-tetra-Cl-Et	12.71	12.62	0.007	131	133	257.091	22.14	22.1	98	
64	ethylbenzene	12.99	12.90	0.007	91	106	1359.049	24.33	24.3	99	
65	m/p-Xylenes x2	13.19	13.10	0.007	91	106	2030.507	47.81	47.8	98	
99	1-4-di-Cl-butane	13.53	13.46	0.006	55	41	238.436	18.63	18.6	98	
52	bromoform	13.30	13.22	0.007	173	175	114.135	19.30	19.3	95	
66	styrene	13.50	13.43	0.006	104	78	680.671	22.70	22.7	99	
67	o-xylene	13.57	13.50	0.006	91	106	944.219	22.59	22.6	99	
68	1122-Tetra-Cl-Et	13.58	13.51	0.006	83	85	137.991	17.98	18.0	99	
110	t-1,4-dichloro-2	13.74	13.67	0.006	89	53	28.308	23.52	23.5	65	
106	Cl-benzyl	15.21	15.12	0.007	91	126	208.052	21.71	21.7	95	
<<< 13 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.72	13.64	0.005	110	97	36.069	18.14	18.1	87	
71	isopropylbenzene	13.93	13.86	0.005	105	120	1419.781	26.45	26.5	100	
72	bromobenzene	14.18	14.10	0.005	156	158	254.497	21.60	21.6	97	
73	n-propylbenzene	14.36	14.29	0.005	120	78	379.146	26.21	26.2	99	
74	2-Cl-Tl	14.46	14.37	0.006	126	128	223.003	23.20	23.2	95	
75	4-Cl-Tl	14.53	14.45	0.006	126	128	298.544	22.72	22.7	97	
76	135-tri-Me-Bz	14.64	14.57	0.005	105	120	1061.726	24.50	24.5	99	
79	tert-butylbenzene	14.92	14.84	0.006	119	91	1148.581	25.82	25.8	100	
78	124-tri-Me-Bz	15.02	14.94	0.006	105	120	865.549	23.49	23.5	98	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233M01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 28 15:48 2003
 Method Update: Tue Apr 29 11:06 2003
 Quant. Time : Apr 29 11:27 2003
 Print Time : Tue Apr 29 11:28 2003
 Miscellaneous :

Sample : F=1 \$2866-05
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.20	15.11	0.006	146	148	413.711	22.04	22.0	90
82	14-di-Cl-Bz	146	15.27	15.19	0.006	146	148	551.120	20.21	20.2	94
81	sec-butylbenzene		15.13	15.04	0.006	105	134	1699.782	26.13	26.1	100
77	4-iso-Pr-toluene		15.29	15.22	0.005	119	134	1216.594	25.92	25.9	100
84	12-di-Cl-benzene		15.61	15.52	0.006	146	148	362.507	20.02	20.0	97
85	n-butylbenzene		15.68	15.60	0.005	91	134	1236.468	25.14	25.1	100
86	12-diBr-3-Cl-Pra		16.06	15.97	0.005	157	155	22.163	16.70	16.7	89
87	124-tri-Cl-Bz		17.33	17.24	0.006	180	182	285.974	20.05	20.1	97
88	naphthalene		17.57	17.49	0.005	128	129	223.049	20.53	20.5	99
90	123-tri-Cl-Bz		17.76	17.68	0.005	180	182	212.927	21.58	21.6	100
89	hx-Cl-butadiene		17.60	17.52	0.005	225	260	309.148	28.80	28.8	96

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

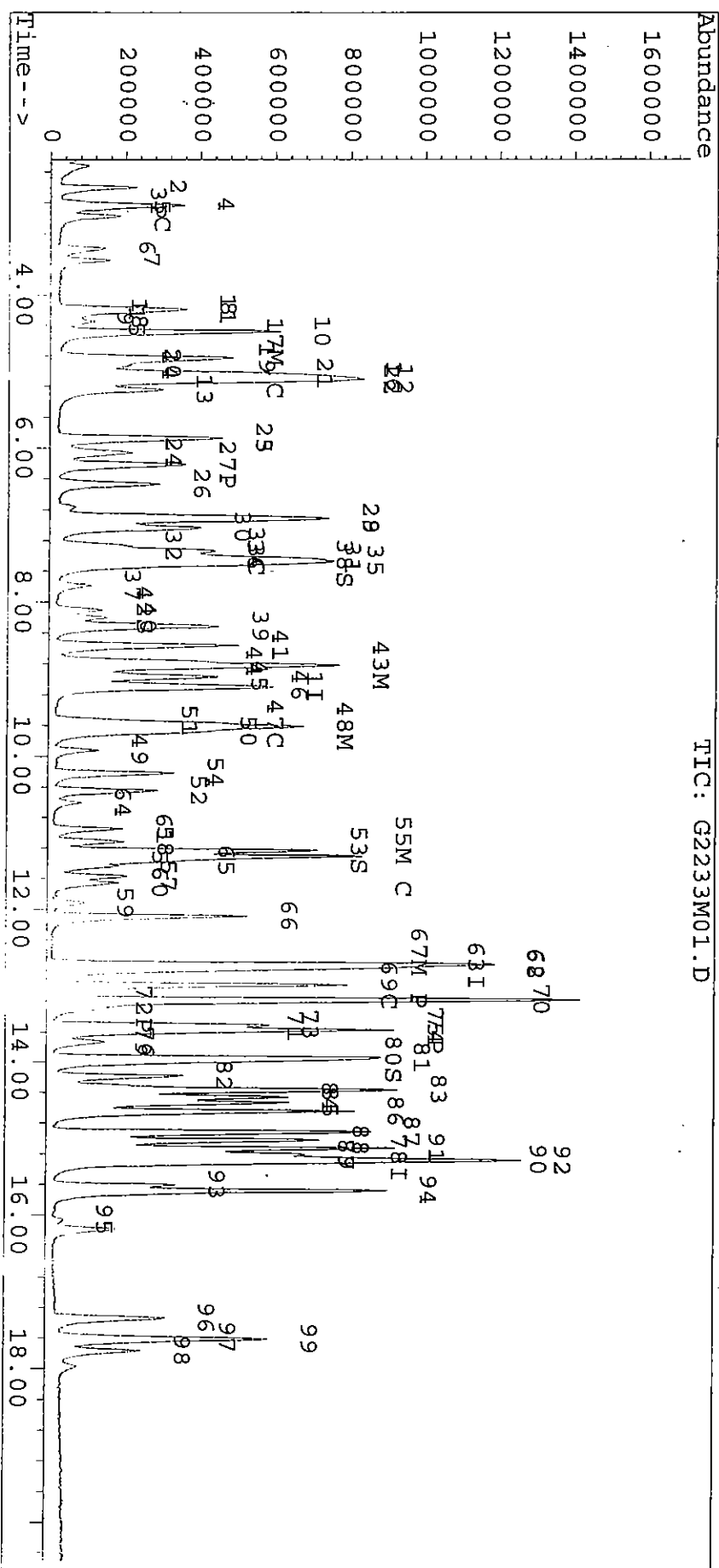
Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2233\G2233M01.D
 Acq On : 28 Apr 03 3:48 pm
 Sample : f=1 \$2866-05
 Misc :
 Quant Time: Apr 29 11:27 2003

Quant Results File: quant.res

Vial: 13
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Tue Apr 29 11:06:45 2003
 Response via : Multiple Level Calibration



Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233N01.D

Sample : f=1 \$2866-05

Method : C:\HPCHEM\1\METHODS\E524G003.M

Inst. : GCMS-G

Acq. Time : Apr 28 16:17 2003

RF via : Multiple Level Calibration

Method Update: Tue Apr 29 11:06 2003

Operator: Eddie

Quant. Time : Apr 29 11:31 2003

Multiplr: 1.000000

Print Time : Tue Apr 29 11:31 2003

Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.10	9.03	0.008	96	70	711.756	10.00		0.07	
47	47 Cl-benzene-d5, I2	12.74	12.66	0.006	82	119	188.117	10.00		0.07	
62	62 1,4-DCB-d4 150 15	15.24	15.15	0.006	152	150	156.706	10.00		0.09	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.52	7.44	0.005	111	113	484.709	19.22		19.2	%Recovery
29	29 1,2-di-Cl-ethane-	8.09	8.02	0.005	65	102	196.000	16.67		16.7	96.10%
55	55 toluene-d8(S2)	11.23	11.15	0.005	100	99	767.056	21.57		21.6	83.34%
70	70 4-Br-1-F-Bz (S3)	13.97	13.90	0.005	174	95	303.394	21.67		21.7	107.87%
											108.34%

Target Compounds

Qvalue

<<< I1 : ISTD ID = 1 >>>											
3	3 di-Cl-di-F-methan	2.61	2.60	0.001	85	87	434.022	21.48		21.5	100
4	4 Chloromethane	2.80	2.78	0.002	50	52	285.903	16.51		16.5	97
9	9 F114 85 135	2.84	2.83	0.001	85	135	493.909	21.73		21.7	96
5	5 vinyl chloride	2.98	2.96	0.002	62	64	316.137	19.25		19.2	98
6	6 bromomethane	3.40	3.37	0.003	94	96	268.925	18.63		18.6	96
7	7 Chloroethane	3.55	3.52	0.004	64	66	257.582	21.18		21.2	99
8	8 tri-Cl-F-methane	4.20	4.14	0.006	101	103	663.029	23.88		23.9	100
111	111 isopropyl alcoho	4.31	4.27	0.005	45	43	48.766	151.85		151.8	78
100	100 ethyl ether x5	4.49	4.44	0.005	59	74	792.239	78.10		78.1	99
102	102 Acrolein x10	4.20	4.15	0.005	56	55	139.193	171.98		172.0	98
119	119 methyl acetate	5.12	5.06	0.007	43	74	194.120	22.23		22.2	95
104	104 Carbon disulfide	5.25	5.18	0.007	76	78	1118.398	18.85		18.8	100
103	103 Acrylonitrile x10	4.95	4.89	0.006	53	52	257.961	151.68		151.7	99
95	95 Acetone x10	4.36	4.31	0.005	43	58	211.444	172.56		172.6	97
108	108 F-113	5.09	5.02	0.007	151	101	547.820	25.68		25.7	98
13	13 11-dichloroethene	4.82	4.76	0.006	61	96	592.320	21.23		21.2	100
101	101 Acetonitrile x10	4.26	4.20	0.006	41	40	68.114	149.04		149.0	94
109	109 Iodomethane	4.86	4.80	0.006	142	127	603.343	27.02		27.0	100
113	113 Tert butyl alcoh	4.90	4.86	0.004	59	57	98.017	159.02		159.0	96

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

?
 #
 m
 4/29/03

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233N01.D Sample : F=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 16:17 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:31 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:32 2003
 Miscellaneous :

3250

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.02	4.97	0.006	84	49	486.222	17.08	17.1	100	
112	Allyl chloride	5.13	5.07	0.006	41	76	651.603	21.11	21.1	95	?
200	200 Nitro methane x1	5.88	5.82	0.007	61	46	645.186	207.77	207.8	100	?
10	10 t-Bu-Me-ether	6.06	6.00	0.007	73	57	530.640	17.62	17.6	97	
19	19 t-12-di-Cl-ethene	5.88	5.82	0.007	96	61	431.065	20.15	20.2	96	?
98	98 Vinyl acetate x5	6.48	6.41	0.008	43	86	1075.035	55.85	55.8	97	
21	21 11-dichloroethane	6.22	6.15	0.008	63	83	836.796	22.40	22.4	98	
91	91 2-butanone MEKx10	6.91	6.83	0.009	43	72	1060.941	158.94	158.9	99	?
115	115 Di isoprop ether	6.92	6.85	0.008	45	87	1739.010	18.44	18.4	99	?
22	22 c-12-di-Cl-ethene	7.05	6.97	0.009	96	61	407.332	19.46	19.5	98	
23	23 22-Dichloropropan	7.43	7.35	0.009	77	97	703.776	21.83	21.8	100	
24	24 Br-Cl-methane	7.28	7.18	0.010	128	130	120.942	16.11	16.1	97	
25	25 chloroform	7.34	7.27	0.008	83	85	730.691	19.55	19.6	99	
201	201 Ethyl acetate x2	7.39	7.31	0.009	43	61	252.779	26.56	26.6	87	?
116	116 ETBE	7.48	7.40	0.009	59	87	983.221	17.12	17.1	97	
117	117 Iso-butyl alcoho	7.39	7.31	0.009	43	42	282.910	148.57	148.6	90	#?
26	26 tetrahydrofuranx5	7.78	7.71	0.008	72	42	54.443	74.64	74.6	89	
34	34 111-tri-Cl-ethane	8.30	8.23	0.008	97	99	689.203	22.14	22.1	100	
30	30 12-dichloroethane	8.20	8.13	0.008	62	64	251.724	17.29	17.3	95	
35	35 11-Di-Cl-propene	8.56	8.48	0.009	75	110	605.962	22.28	22.3	100	
36	36 benzene	8.83	8.75	0.009	78	52	1318.831	21.32	21.3	99	
37	37 CCl4	8.76	8.68	0.009	117	119	567.260	23.81	23.8	100	
97	97 thiophene	8.97	8.89	0.009	84	58	569.162	18.51	18.5	99	
118	118 TAME	9.07	9.00	0.008	73	43	647.881	16.39	16.4	98	
39	39 12-di-Cl-propane	9.57	9.49	0.009	63	76	356.554	19.70	19.7	97	
40	40 trichloroethene	9.62	9.54	0.009	130	132	410.490	21.42	21.4	99	
96	96 Me-methacrylate	9.92	9.85	0.008	69	100	101.354	16.21	16.2	99	
42	42 Br-di-Cl-methane	9.68	9.61	0.008	83	85	436.659	17.30	17.3	97	
41	41 dibromomethane	9.53	9.45	0.009	174	172	153.097	17.54	17.5	98	
45	45 c-13-di-Cl-propen	10.44	10.37	0.008	75	110	393.057	17.66	17.7	97	
92	92 2-ClEt-Vi-ether10	10.21	10.15	0.008	63	43	300.415	78.70	78.7	98	
56	56 toluene	11.30	11.23	0.008	91	92	1225.467	21.08	21.1	99	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233N01.D Sample : f=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 16:17 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:31 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:32 2003
 Miscellaneous :

3251

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.44	11.36	0.008	69	99	156.388	13.00	13.0	99	
93	2-Hexanone x5	11.56	11.48	0.008	43	58	290.667	71.89	71.9	94	
48	112-tri-Cl-Et	11.10	11.03	0.008	97	83	135.976	15.94	15.9	100	
58	1,2-di-br-ethane	11.91	11.82	0.009	107	109	147.030	16.16	16.2	97	
51	di-Br-Cl-methane	11.64	11.57	0.008	129	127	196.834	17.18	17.2	100	
46	t-13-di-Cl-propen	10.94	10.87	0.008	75	110	254.026	17.71	17.7	98	
105	1-Chlorohexane	12.71	12.64	0.008	55	93	455.583	26.03	26.0	98	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.60	10.52	0.007	43	58	114.280	19.29	19.3	97	
49	1,3-di-Cl-propane	11.37	11.29	0.006	76	78	250.256	20.35	20.3	98	
59	tetra-Cl-ethene	12.09	12.00	0.007	166	168	454.542	27.46	27.5	100	
60	chlorobenzene	12.78	12.70	0.006	112	77	653.946	23.79	23.8	100	
61	1112-tetra-Cl-Et	12.70	12.62	0.006	131	133	256.810	23.33	23.3	99	
64	ethylbenzene	12.98	12.90	0.006	91	106	1387.425	26.19	26.2	100	
65	m/p-Xylenes x2	13.18	13.10	0.006	91	106	2066.102	51.31	51.3	97	
99	1-4-di-Cl-butane	13.53	13.46	0.006	55	41	230.043	18.96	19.0	99	
52	bromoform	13.29	13.22	0.006	173	175	109.984	19.63	19.6	98	
66	styrene	13.50	13.43	0.006	104	78	672.321	23.64	23.6	100	
67	o-xylene	13.57	13.50	0.006	91	106	949.746	23.97	24.0	99	
68	1122-Tetra-Cl-Et	13.57	13.51	0.005	83	85	129.921	17.86	17.9	98	
110	t-1,4-dichloro-2	13.76	13.67	0.007	89	53	25.897	22.53	22.5	75	
106	Cl-benzyl	15.20	15.12	0.006	91	126	188.067	20.62	20.6	99	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.72	13.64	0.005	110	97	33.631	17.63	17.6	89	
71	isopropylbenzene	13.93	13.86	0.005	105	120	1435.271	27.87	27.9	99	
72	bromobenzene	14.19	14.10	0.005	156	158	250.958	22.20	22.2	97	
73	n-propylbenzene	14.36	14.29	0.005	120	78	384.949	27.73	27.7	99	
74	2-Cl-Tl	14.46	14.37	0.005	126	128	228.140	24.74	24.7	94	
75	4-Cl-Tl	14.53	14.45	0.005	126	128	285.916	22.68	22.7	99	
76	135-tri-Me-Bz	14.64	14.57	0.005	105	120	1082.843	26.04	26.0	100	
79	tert-butylbenzene	14.92	14.84	0.005	119	91	1183.044	27.72	27.7	98	
78	124-tri-Me-Bz	15.02	14.94	0.005	105	120	903.363	25.56	25.6	98	

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

7/29/03
m

Data Filename: C:\HPCHEM\1\DATA\03G2233\G2233N01.D Sample : f=1 \$2866-05
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 16:17 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 29 11:31 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:32 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.20	15.11	0.006	146	148	411.656	22.86	22.9	91
82	14-di-Cl-Bz	146	15.27	15.19	0.005	146	148	542.143	20.72	20.7	93
81	sec-butylbenzene		15.12	15.04	0.005	105	134	1688.826	27.06	27.1	99
77	4-iso-Pr-toluene		15.29	15.22	0.005	119	134	1239.130	27.51	27.5	98
84	12-di-Cl-benzene		15.60	15.52	0.005	146	148	351.701	20.24	20.2	99
85	n-butylbenzene		15.68	15.60	0.005	91	134	1277.729	27.07	27.1	100
86	12-diBr-3-Cl-Pra		16.07	15.97	0.006	157	155	21.911	17.21	17.2	92
87	124-tri-Cl-Bz		17.32	17.24	0.005	180	182	285.339	20.83	20.8	98
88	naphthalene		17.57	17.49	0.005	128	129	213.287	20.46	20.5	98
90	123-tri-Cl-Bz		17.75	17.68	0.005	180	182	208.185	21.99	22.0	100
89	hx-Cl-butadiene		17.60	17.52	0.005	225	260	321.939	31.28	31.3	95

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= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

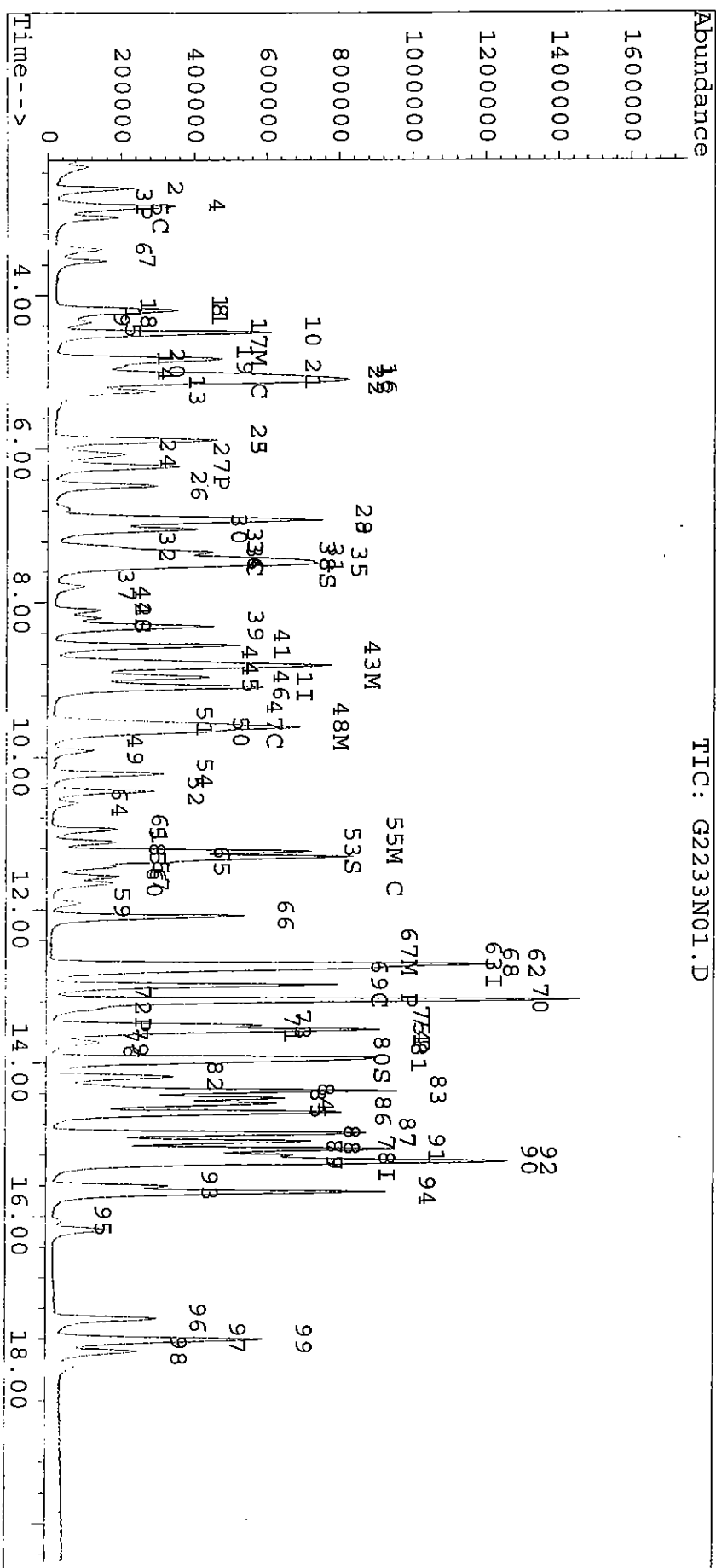
Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2233\G2233N01.D
 Acq On : 28 Apr 03 4:17 pm
 Sample : f=1 \$2866-05
 Misc :
 Quant Time: Apr 29 11:31 2003

Quant Results File: quant.res

Vial: 14
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Tue Apr 29 11:06:45 2003
 Response via : Multiple Level Calibration



Data Filename: C:\HPCHEM\1\DATA\03G2233\2866-05.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 28 20:01 2003 RF via : Multiple Level Calibration
 Method Update: Tue Apr 29 11:06 2003 Operator: Eddie
 Quant. Time : Apr 28 20:21 2003 Multiplr: 1.000000
 Print Time : Tue Apr 29 11:32 2003
 Miscellaneous :

3254

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	1 Fluorobenzene I1	9.09	9.03	0.006	96	70	740.598	10.00		0.06	
47	47 Cl-benzene-d5, I2	12.73	12.66	0.006	82	119	214.196	10.00		0.07	
62	62 1,4-DCB-d4 150 15	15.23	15.15	0.005	152	150	168.157	10.00		0.08	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.49	7.44	0.004	111	113	469.236	17.86		17.9	89.32%
29	29 1,2-di-Cl-ethane-	8.09	8.02	0.004	65	102	227.552	18.60		18.6	92.98%
55	55 toluene-d8(S2)	11.21	11.15	0.004	100	99	767.704	20.75		20.8	103.76%
70	70 4-Br-1-F-Bz (S3)	13.97	13.90	0.005	174	95	317.366	21.12		21.1	105.62%

Target Compounds	<<< I1 : ISTD ID = 1 >>>	Qvalue
111 111 isopropyl alcoho	4.29 4.27 0.003 45 43 1.691 5.06 5.1	1
119 119 methyl acetate	5.04 5.06 -0.001 43 74 0.810 0.36 0.4	56
91 91 2-butanone MEKx10	6.79 6.83 -0.005 43 72 8.853 1.27 1.3	74
117 117 Iso-butyl alcoho	7.32 7.31 0.000 43 42 1.621 0.82 0.8	1
93 93 2-Hexanone x5	11.51 11.48 0.003 43 58 2.201 0.52 0.5	24
48 48 112-tri-Cl-Et	11.21 11.03 0.021 97 83 21.693 2.22 2.2	4
<<< I2 : ISTD ID = 47 >>>		
54 54 MIBK	10.60 10.52 0.006 43 58 26.862 3.51 3.5	90
49 49 1,3-di-cl-propane	11.21 11.29 -0.006 76 78 9.122 0.65 0.7	67

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2233\2866-05.D Vial: 30
 Acq On : 28 Apr 03 8:01 pm Operator: Eddie
 Sample : f=1 Inst : GCMS-G
 Misc : Multiplr: 1.00
 Quant Time: Apr 28 20:21 2003 Quant Results File: quant.res

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Tue Apr 29 11:06:45 2003
 Response via : Multiple Level Calibration

