

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: 32767

Collected by:

Component Name: Bicarbonate

CAS No:

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	< 2	U
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	181	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	319	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	286	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	220	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/18/03	03W2446	mg/L	2	98.4	
03W2446-MB-01	03W2446-MB-01	Water	04/18/03	04/18/03	04/18/03	03W2446	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: 32767

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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/18/03	03W2446	mg-CaCO ₃ /L	2	<2	U
03W2446-MB-01	03W2446-MB-01	Water	04/18/03	04/18/03	04/18/03	03W2446	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 9040

Client Name: GEOFON, Inc.
Project ID: JPL

Project No: 04-4428.10
Service ID: 32767

Anal. Method 9040
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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	6.04	
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	6.88	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	7.28	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	7.44	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	7.23	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	7.71	
03W2443-MB-01	03W2443-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2443	pH unit	0.01	6.85	

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: 32767 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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	<10	U
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	637	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	755	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	651	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	413	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/21/03	03W2463	mg/L	10	503	
03W2463-MB-01	03W2463-MB-01	Water	04/21/03	04/21/03	04/21/03	03W2463	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 ID: JPL

Project No: 04-4428.10
Service ID: 32767

Anal. Method 7196
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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2441	mg/L	0.01	<0.01	U
03W2441-MB-01	03W2441-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2441	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 314.0

Client Name: GEOFON, Inc.
Project ID: JPL

Project No: 04-4428.10
Service ID: 32767

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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	< 4	U
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	3.6	B
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	2.9	B
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	2.9	B
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	2.1	B
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2440	µg/L	4	2.7	B
03W2440-MB-01	03W2440-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2440	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.

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: 32767

Anal. Method 300.0
Collected by:

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

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.25	0.29	
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	4	110	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	4	127	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	4	102	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	1.6	52.7	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	1.6	68.5	
03W2435-MB-01	03W2435-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2435	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 ID: JPL

Project No: 04-4428.10
Service ID: 32767

Anal. Method 300.0
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
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.05	0.079	
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.8	13.2	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.8	5.3	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.8	8.8	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.32	7.1	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.32	6.8	
03W2435-MB-01	03W2435-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2435	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: 32767

Anal. Method 300.0
Collected by:

Component Name: Sulfate
CAS No: 14808-79-8

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-2767-1	EB-1-4/17/03	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	0.63	0.70	
03-2767-2	MW-21-1	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	10	181	
03-2767-3	MW-21-2	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	10	144	
03-2767-4	MW-21-3	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	10	122	
03-2767-5	MW-21-4	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	4	65.2	
03-2767-6	MW-21-5	Water	04/17/03	04/17/03	04/17/03	03W2435	mg/L	4	103	
03W2435-MB-01	03W2435-MB-01	Water	04/17/03	04/17/03	04/17/03	03W2435	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 300.0

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2435

LCS Filename: -

Date Analyzed: 041703

Time Analyzed: 12:32

LCSD Filename: -

Date Analyzed: 041703

Time Analyzed: 12:45

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHLORIDE	mg/L	4.0	0	3.91	98	80-120
NITRATE AS N	mg/L	1.5	0	1.49	99	80-120
SULFATE	mg/L	15	0	14.7	98	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE	mg/L	4.0	3.91	98	0	20	80-120
NITRATE AS N	mg/L	1.5	1.49	99	0	20	80-120
SULFATE	mg/L	15	14.7	98	0	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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2435

MS Filename: -

Date Analyzed: 041703

Time Analyzed: 16:01

MSD Filename: -

Date Analyzed: 041703

Time Analyzed: 16:59

MS Sample No: MW-21-1

Sample Lab ID: 03-2767-2

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHLORIDE	mg/L	160	110	269	99	75-125
NITRATE AS N	mg/L	60.0	13.2	72.3	99	75-125
SULFATE	mg/L	600	181	777	99	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE	mg/L	160	273	102	3	20	75-125
NITRATE AS N	mg/L	60.0	72.7	99	0	20	75-125
SULFATE	mg/L	600	781	100	1	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 314.0

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2440

LCS Filename: -

Date Analyzed: 041703

Time Analyzed: 15:30

LCSD Filename: -

Date Analyzed: -

Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
PERCHLORATE	µg/L	25	0	24.1	96	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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2440

MS Filename: -

Date Analyzed: 041703

Time Analyzed: 20:47

MSD Filename: -

Date Analyzed: 041703

Time Analyzed: 21:06

MS Sample No: MW-21-4

Sample Lab ID: 03-2767-5

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
PERCHLORATE	µg/L	50.0	2.1	54.0	104	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
PERCHLORATE	µg/L	50.0	53.4	103	1	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-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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2463

LCS Filename: -

Date Analyzed: 042103

Time Analyzed: 09:57

LCSD Filename: -

Date Analyzed: 042103

Time Analyzed: 09:57

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	0	391	98	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	395	99	1	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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2463

MS Filename: -

Date Analyzed: 042103

Time Analyzed: 09:57

MSD Filename: -

Date Analyzed: 042103

Time Analyzed: 09:57

MS Sample No: MW-21-5

Sample Lab ID: 03-2767-6

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	503	886	96	80-119
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, % RPD REC	
						RPD	REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	893	98	2	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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2441

LCS Filename: -

Date Analyzed: 041703

Time Analyzed: 15:04

LCSD Filename: -

Date Analyzed: 041703

Time Analyzed: 15:04

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHROMIUM (VI)	mg/L	0.25	0	0.233	93	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	0.231	92	1	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: 32767

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2441

MS Filename: -

Date Analyzed: 041703

Time Analyzed: 15:04

MSD Filename: -

Date Analyzed: 041703

Time Analyzed: 15:04

MS Sample No: MW-21-1

Sample Lab ID: 03-2767-2

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHROMIUM (VI)	mg/L	0.25	0	0.242	97	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	0.239	96	1	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: 03-2767

Analysis: Chromium (VI) Calibration Date: 1/29/03

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

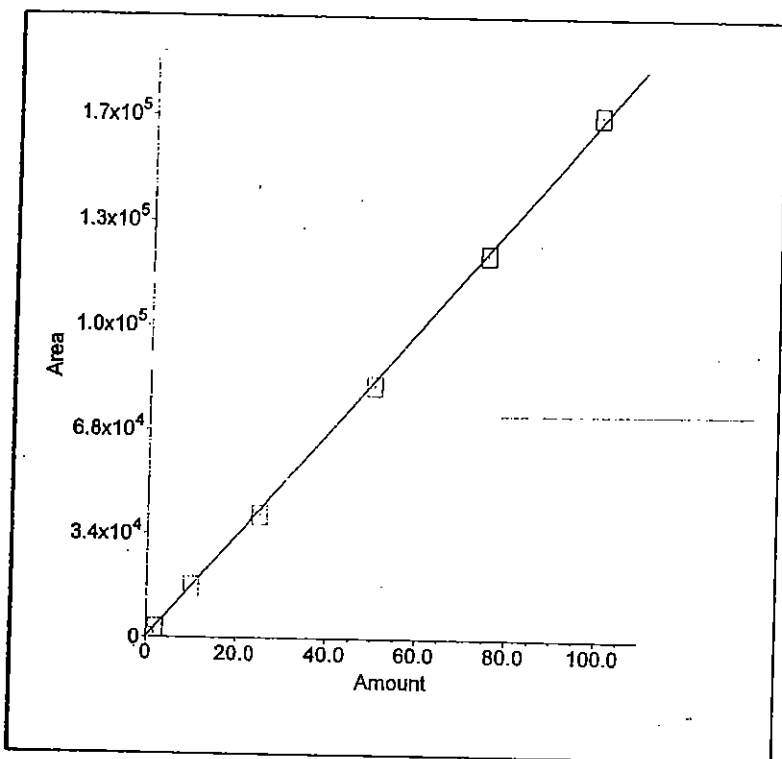
Matrix: Water

APCL Service ID: 03-2767

Analysis	Batch ID	Analysis Date	Sample Name	Unit	Result	Duplicate Result	RPD %	RPD Control limit
Bicarbonate	03W2446	4/18/03	03-2755-1	mg/L	207.0	209.5	1	20
pH	03W2443	4/17/03	MW-21-5	pH unit	7.71	7.73	0	20

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

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



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

Analyst C. W
Date 03/12/03
Instrument LC-10

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\W1991Q01.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 Retention Time 3.45
 Reference Comp. Nitrate-N Window Size 0.20 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 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 Retention Time 3.87
 Reference Comp. Nitrate-N Window Size 0.25 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 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 Retention Time 6.38
 Reference Comp. Phosphate-P Window Size 0.60 min.
 Amount = $K0 + K1 \cdot \text{Area}$
 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

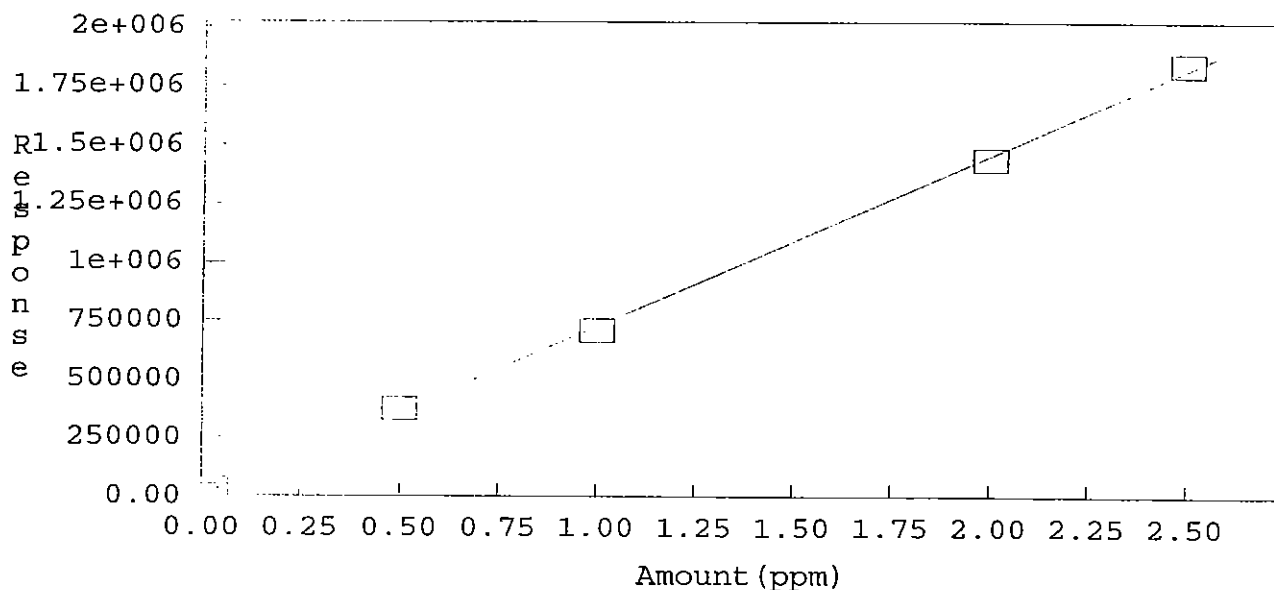
$r^2 = 0.999552$

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

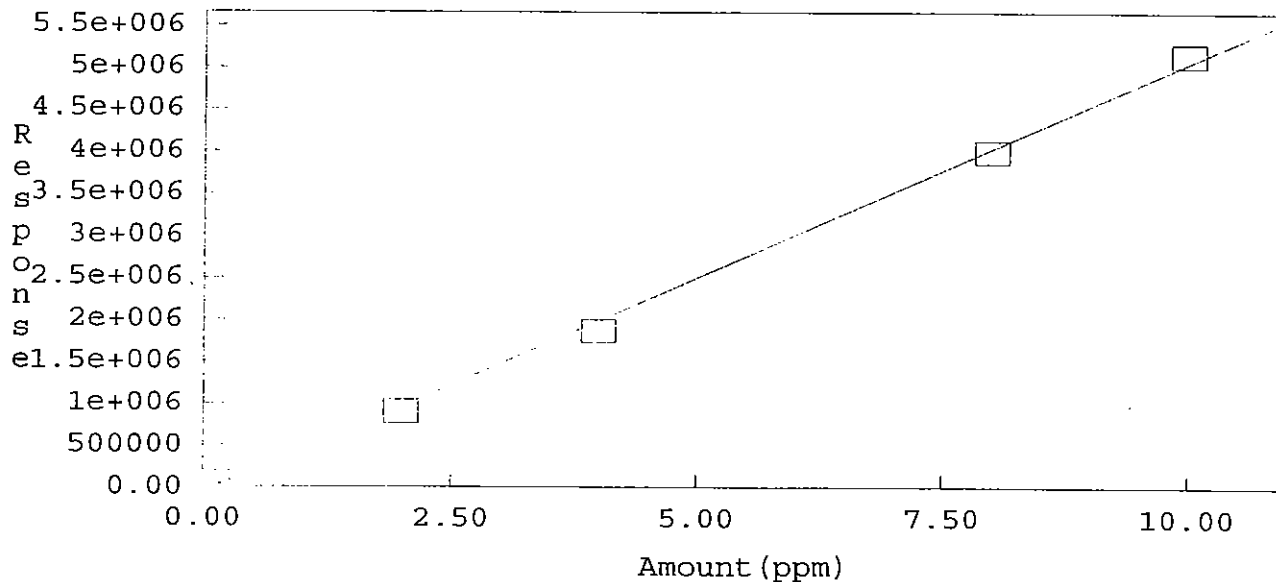
$r^2 = 0.998409$

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

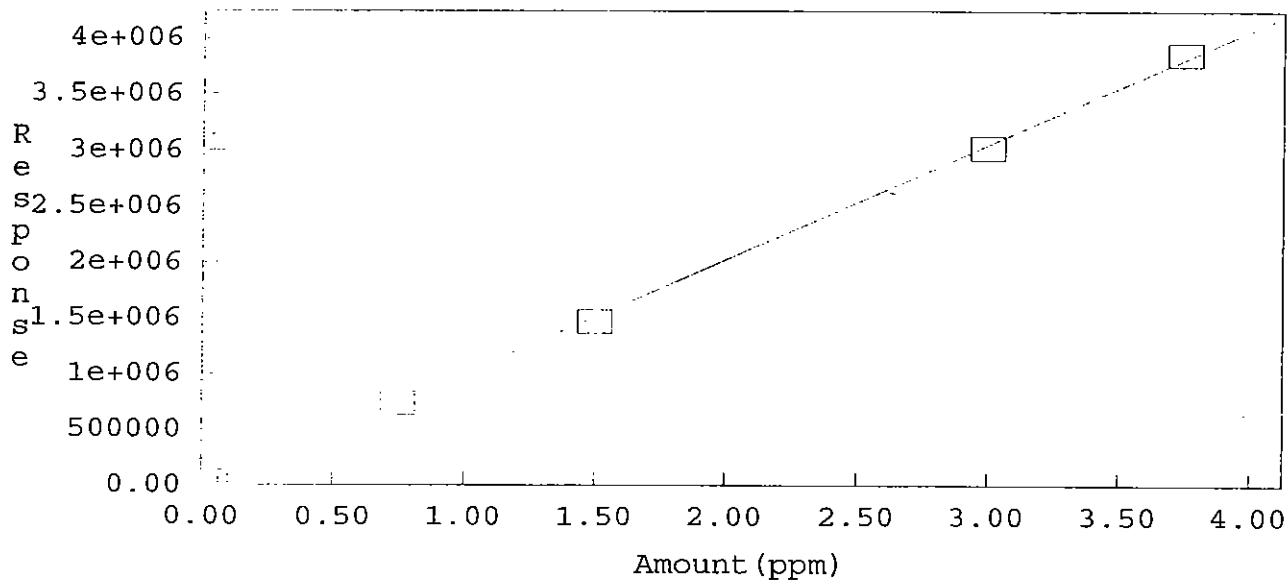
$r^2 = 0.999594$

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

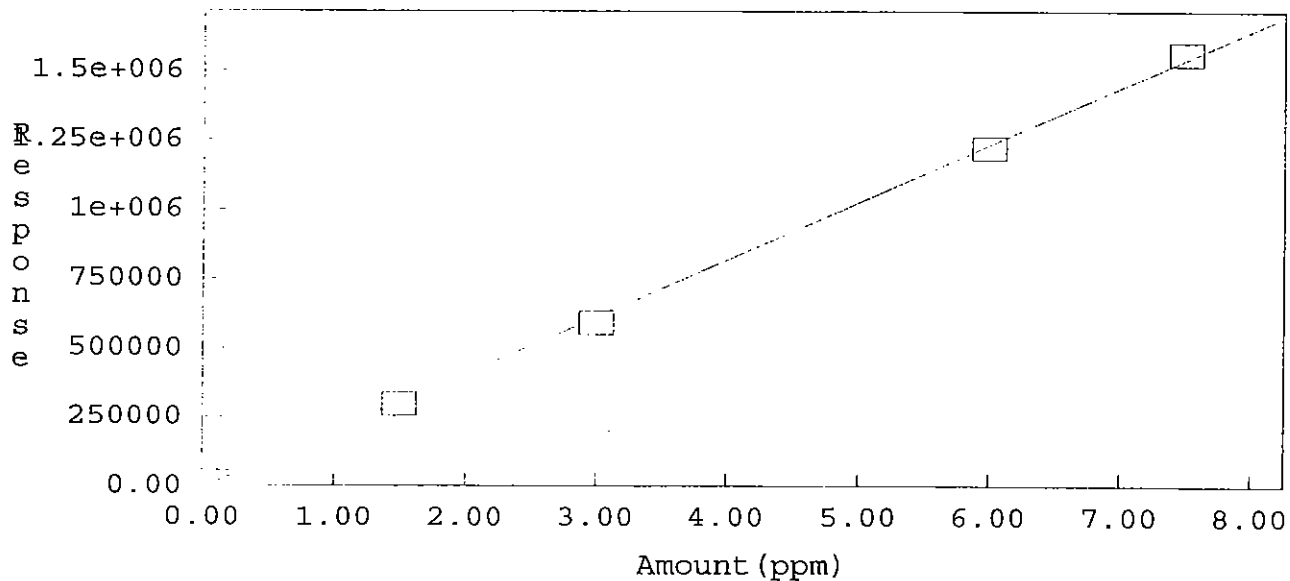
$r^2 = 0.999518$

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

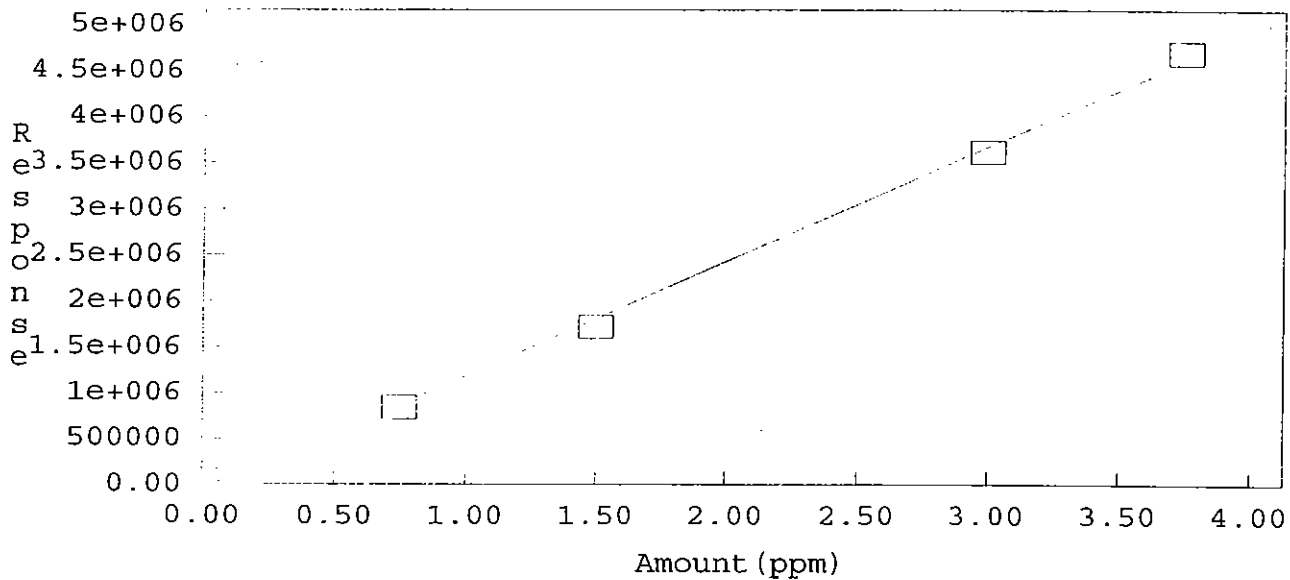
$r^2 = 0.998618$

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

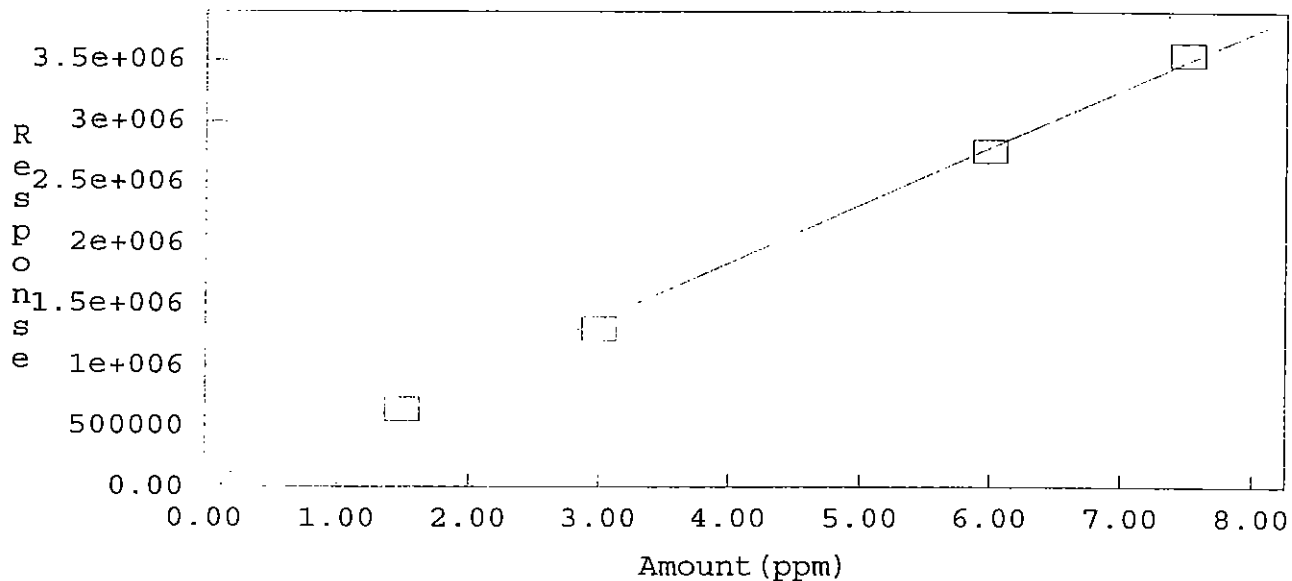
$r^2 = 0.998898$

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

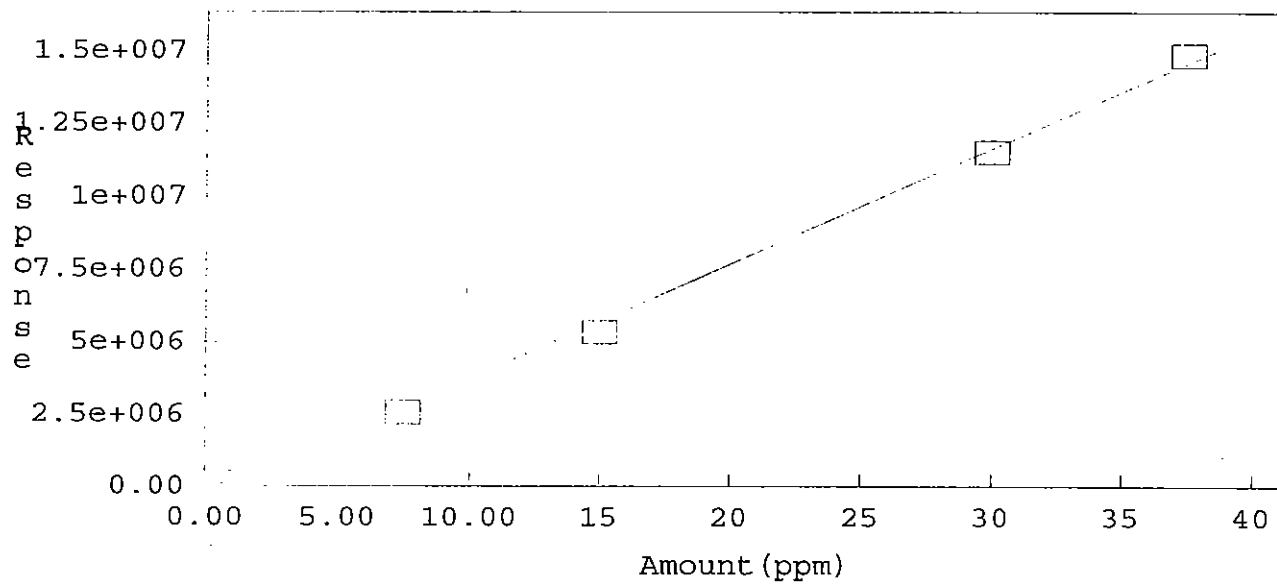
$r^2 = 0.998245$

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



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:

32767

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	300.0	03W2435	mg/L	4.0	3.93	98	-2	✓	90-110	04/17/2003
	NITRATE as N-NO ₃ ⁻ , BY	300.0	03W2435	mg/L	1.5	1.51	101	1	✓	90-110	04/17/2003
	SULFATE SO ₄ ⁻ , BY I	300.0	03W2435	mg/L	15	14.8	99	-1	✓	90-110	04/17/2003
	Chloride	300.0	03W2435	mg/L	4.0	3.90	98	-2	✓	90-110	04/17/2003
	NITRATE as N-NO ₃ ⁻ , BY	300.0	03W2435	mg/L	1.5	1.48	99	-1	✓	90-110	04/17/2003
	SULFATE SO ₄ ⁻ , BY I	300.0	03W2435	mg/L	15	14.7	98	-2	✓	90-110	04/17/2003
	Chloride	300.0	03W2435	mg/L	4.0	3.87	97	-3	✓	90-110	04/17/2003
	NITRATE as N-NO ₃ ⁻ , BY	300.0	03W2435	mg/L	1.5	1.49	99	-1	✓	90-110	04/17/2003
	SULFATE SO ₄ ⁻ , BY I	300.0	03W2435	mg/L	15	14.5	97	-3	✓	90-110	04/17/2003
2	Perchlorate	314.0	03W2440	mg/L	50	51.0	102	2	✓	90-110	04/17/2003
	Perchlorate	314.0	03W2440	mg/L	50	51.3	103	3	✓	90-110	04/17/2003
	Perchlorate	314.0	03W2440	mg/L	50	50.2	100	0	✓	90-110	04/17/2003
3	Chromium (VI)	7196	03W2441	mg/L	0.25	0.252	101	1	✓	90-110	04/17/2003
	Chromium (VI)	7196	03W2441	mg/L	0.25	0.249	100	0	✓	90-110	04/17/2003

Line	Sample	Sample Type	Level	Method	Data File	Volume
1	##03w2440kw ipc 25ppb w7759	Sample		e314-011.met	c:\data\03w2440kw\w2440k ipc 25ppb	1
2	ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2440kw\w2440k q01	1
3	ccb	Sample		e314-011.met	c:\data\03w2440kw\w2440k ccb01	1
4	ics 25ppb w7827d	Sample		e314-011.met	c:\data\03w2440kw\w2440k l01	1
5	LCS 18PPB W7685D	Sample		e314-011.met	c:\data\03w2440kw\w2440k j01	1
6	ICCS 4ppb w7827b	Sample		e314-011.met	c:\data\03w2440kw\w2440k iccs 4ppb_006.dxd	1
7	mb	Sample		e314-011.met	c:\data\03w2440kw\w2440k k01_007.dxd	1
8	2757-05 F=1	Sample		e314-011.met	c:\data\03w2440k\2757-05_008.dxd	1
9	2757-01 F=1	Sample		e314-011.met	c:\data\03w2440k\2757-01_009.dxd	1
10	2757-02 F=1	Sample		e314-011.met	c:\data\03w2440k\2757-02_010.dxd	1
11	2757-03 F=1	Sample		e314-011.met	c:\data\03w2440k\2757-03_011.dxd	1
12	2757-04 F=1	Sample		e314-011.met	c:\data\03w2440k\2757-04	1
13	ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2440kw\w2440k q02	1
14	ccb	Sample		e314-011.met	c:\data\03w2440kw\w2440k k02	1
15	2767-01 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-01	1
16	2767-02 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-02	1
17	2767-03 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-03	1
18	2767-04 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-04	1
19	2767-05 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-05	1
20	2767-06 F=1	Sample		e314-011.met	c:\data\03w2440k\2767-06	1
21	2757-05 MS F=1	Sample		e314-011.met	c:\data\03w2440kw\w2440k m01	1
22	2757-05 MSD F=1	Sample		e314-011.met	c:\data\03w2440kw\w2440k n01	1
23	ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2440kw\w2440k q03	1
24		Sample		aastopcl.met		1

Analyst Wei Wang
Date 04/17/03
Instrument 2C-K

Line	Dilution	Weight	Int. Std.	Comment
1	1	1	1	
2	1	1	1	
3	1	1	1	
4	1	1	1	
5	1	1	1	
6	1	1	1	
7	1	1	1	
8	1	1	1	
9	1	1	1	
10	1	1	1	
11	1	1	1	
12	1	1	1	
13	1	1	1	
14	1	1	1	
15	1	1	1	
16	1	1	1	
17	1	1	1	
18	1	1	1	
19	1	1	1	
20	1	1	1	
21	1	1	1	
22	1	1	1	
23	1	1	1	
24	1	1	1	

Default Method Path: C:\PEAKNET\METHOD

Default Data Path: C:\DATA\03W2323K

Comment:

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 DNDate 3/21/03Instrument J

DIONEX SCHEDULE - C:\DX\SCHEDULE\03W2435.SCH

Inj#	Sample Name	Method	Data File	Vol.	Dil.	Int.Std.
1	##03W2435, W CCVW77	..\E300-063	..\W2435Q01.D01	1	1	1
2	MB RW1407	..\E300-063	..\W2435K01.D02	1	1	1
3	LCS W7768-100X	..\E300-063	..\W2435L01.D03	1	1	1
4	LCSD W7768-100X	..\E300-063	..\W2435J01.D04	1	1	1
5	2755-1 F=1.25	..\E300-063	..\2755-101.D05	1	1.25	1
6	2755-2 F=2	..\E300-063	..\2755-201.D06	1	2	1
7	2757-1 F=800	..\E300-063	..\2757-101.D07	1	800	1
8	2757-2 F=20	..\E300-063	..\2757-201.D08	1	20	1
9	2757-3 F=20	..\E300-063	..\2757-301.D09	1	20	1
10	2757-4 F=20	..\E300-063	..\2757-401.D10	1	20	1
11	2757-5 F=1.25	..\E300-063	..\2757-501.D11	1	1.25	1
12	CCV2W7767-100X	..\E300-063	..\W2435Q01.D12	1	1	1
13	MB RW1407	..\E300-063	..\W2435K01.D13	1	1	1
14	\$2767-2 MS F=40	..\E300-063	..\W2435M01.D14	1	40	1
15	\$2767-2 MSD F=40	..\E300-063	..\W2435N01.D15	1	40	1
16	2767-2 F=20	..\E300-063	..\2767-201.D16	1	20	1
17	\$2767-2 MSD F=40	..\E300-063	..\W2435N01.D17	1	40	1
18	2767-3 F=20	..\E300-063	..\2767-301.D18	1	20	1
19	2767-4 F=20	..\E300-063	..\2767-401.D19	1	20	1
20	2767-5 F=8	..\E300-063	..\2767-501.D20	1	8	1
21	2767-6 F=8	..\E300-063	..\2767-601.D21	1	8	1
22	2767-1 F=1.25	..\E300-063	..\2767-101.D22	1	1.25	1
23	CCV3W7767-100X	..\E300-063	..\W2435Q01.D23	1	1	1
24		..\STOP.MET		1	1	1

Comment:

LCS/LCSD LOT # W7768

MS/MSD LOT # W7767

ELUENT LOT # W7769

ANALYTICAL METHOD 9056/E300 MATRIX WAnalyst PCDate 4/17/03Instrument J

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Batch # 03W2441 Matrix: W

Chromium (VI) (7196) Worksheet

[Holding Time: 24 hours!!]

Test Date: 4/17/03Analyst: h

Lot #: Reagent Water

Diphenylcazide solution

Test Time: 15:04

SOP: G-22

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.= <u>0.997</u> (≥ 0.995)	
STD-4	W-	x / = mg/L			RSD= % ($\leq 15\%$)	
STD-5	W-	x / = mg/L			Ref. page	
STD-6	W-	x / = mg/L			<u>A = 0.000 + 0.836C</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>7853</u>	Expected Conc.: x	/	= <u>0.25</u> mg/L	<u>0.241</u>	<u>0.752</u> mg/L	REC. %	90-110 %
Method Blank	Bl. Lot: <u>T1115</u>		/X ₀ =	95.0/ =	<u>0.000</u>	mg/L	<u>0.00</u> ppm	
LCS1	Bl. Lot: <u>11</u>		/X ₀ =	95.0/ =	<u>0.195</u>	mg/L	<u>0.233</u> ppm	
Sample-1 <u>EB</u>	<u>2767-1</u>		/X ₀ =	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</u> ppm	
MS on S-1	<u>2</u>		/X ₀ =	95.0/ =	<u>0.202</u>	mg/L	<u>0.242</u> ppm	
MSD on S-1	<u>2</u>		/X ₀ =	95.0/ =	<u>0.200</u>	mg/L	<u>0.239</u> ppm	
Sample 2	<u>2</u>		/X ₀ =	95.0/ =	<u>0.001</u>	mg/L	<u>0.001</u> ppm	
Sample 3	<u>3</u>		/X ₀ =	95.0/ =	<u>0.001</u>	mg/L	<u>0.001</u> ppm	
Sample 4	<u>4</u>		/X ₀ =	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</u> ppm	
Sample 5	<u>5</u>		/X ₀ =	95.0/ =	<u>0.001</u>	mg/L	<u>0.001</u> ppm	
Sample 6	<u>6</u>		/X ₀ =	95.0/ =	<u>0.001</u>	mg/L	<u>0.001</u> 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: <u>T1115</u>		/X ₀ =	95.0/ =	<u>0.193</u>	mg/L	<u>0.231</u> 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.	<u>0.05 mg/L</u>		/X ₀ =	95.0/ =	<u>0.208</u>	<u>0.249</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>7757</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>7853</u>	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W- <u>1</u>	x / = ppm	%		MDL(s) 0.025

13760 Magnolia Ave. Chino CA 91710
Tel: (909) 590-1828 Fax: (909) 590-1498

Chromium (VI) (7196) Worksheet

Batch # 03W1295 Matrix: W

[Holding Time: 24 hours!!]

Test Date: 1/24/03 Analyst: B.V.

Lot #: Reagent Water

Diphenylcazide solution

Test Time:

SOP: G-22

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.=0.977 (> 0.995)	
STD-4	W-	x / = 0.15 mg/L	0.109		RSD= % (< 15%)	
STD-5	W-	x / = 0.25 mg/L	0.214		Ref. page	
STD-6	W-	x / = 0.50 mg/L	0.415		A=0.003+0.836C	

Analysis Type	Sample ID or Lot #	Samp. Amt 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	0.26	0.258 mg/L	REC. %	90-110 %
Method Blank	Bl. Lot: T1115		/X ₀ = 1	95.0/ =	0.000	mg/L	0.00 ppm	
LCS1	Bl. Lot: 4		/X ₀ =	95.0/ =	0.204	mg/L	0.204 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 = 1	95.0/ =	0.210	mg/L	0.251 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.			/X ₀ =	95.0/ =	0.204	mg/L	0.204 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- 4	x / = ppm	%		PQL(s) 0.05
LCS	W-7191	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W- 4	x / = ppm	%		MDL(s) 0.025

P & Ch Laboratory

1000 Ave. Chino CA 91710
 (909) 590-1828 Fax: (909) 590-1498

pH (150.1/9045B) Worksheet

0

compensation must be performed by the instrument automatically.

Analyst D.L.

SOP: G-

Batch # <u>03W2443</u> Analysis Date: <u>4.17.03</u>		Batch # <u>03W2445</u> Analysis Date: <u>4.18.03</u>	
Starting Time: <u>16:32</u> Ending Time: _____		Starting Time: <u>08:19</u> Ending Time: _____	
Matrix ☒ Aqueous ☐ Soil		Matrix ☒ Aqueous ☐ Soil	
Standard	4.00	7.00	10.00
Standard	4.00	7.00	10.00
Temperature °C		<u>21.20</u>	<u>030619-24</u>
Temperature °C		<u>22.0</u>	<u>21.2</u>
pH Reading		<u>7.03</u>	<u>10.02</u>
pH Reading		<u>7.02</u>	<u>10.01</u>
T-corrected pH		<u>7.01</u>	<u>10.03</u>
T-corrected pH		<u>7.01</u>	<u>10.03</u>
Control Limit	±0.05 pH unit		Control Limit
Control Limit	±0.05 pH unit		Control Limit

#	Sample ID	Pre-treat	pH	Note	#	Sample ID	Pre-treat	pH	Note
MB	<u>T1115</u>		<u>6.81</u>		MB	<u>T1115</u>	<u>4/18/03</u>	<u>6.84</u>	<u>1:1</u>
1	<u>2767-1</u>		<u>6.04</u>		1	<u>-</u>		<u>5.68</u>	<u>↓</u>
2	<u>-2</u>		<u>6.88</u>		2				
3	<u>-3</u>		<u>7.28</u>		3				
4	<u>-4</u>		<u>7.44</u>		4				
5	<u>-5</u>		<u>7.23</u>		5				
6	<u>-6</u>		<u>7.71</u>		6				
7					7				
8					8				
9					9				
10					10				
11					11				
12					12				
13			<u>4.17.03</u>		13				
14			<u>22</u>		14				
15					15				
16					16				
17					17				
18					18				
19					19				
20					20				
Dup.	<u>2767-6</u>		<u>7.73</u>		Dup.	<u>-</u>		<u>5.75</u>	<u>1:1</u>

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Solid Analysis (160.1, 160.2, 160.3) Worksheet

Batch # 03W2463 Matrix WMethod: 160.1

Balance No. _____

Date: 4/21/03Analyst: DK

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 = $10^6 \times \Delta W \times f_1 / V$

SOP: G-81

#	Analysis Type	Sample ID (STD Lot #)	Treatment Ratio $V_f/X=f_1$	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	T1115	1 =	100	103.2185	103.1186	103.1187	0.0002	2	GC
2	LCS	T1115	1 =	100	115.7069	115.7462	115.7462	0.0391	391	HK
3	Sample-1	2738-1	1 =	100	105.3119	105.3117	105.3118	0.0000	0	12
4	MS on S-1	2767-6	1 =	100	103.9436	104.0323	104.0322	0.0886	886	14
5	MSD on S-1	✓-6	1 =	100	105.8925	105.9866	105.9868	0.0893	893	19
6	Sample-2	2787-1	1 =	20.0	112.9023	113.1379	113.1377	0.2354	11770	K
7	Sample-3	-2	1 =	100	117.8955	117.9654	117.9652	0.0697	697	7
8	Sample-4	-3	1 =	100	115.1073	115.1761	115.1762	0.0689	689	G
9	Sample-5	-4	1 =	100	115.8739	116.0195	116.0194	0.1464	1464	CK
10	Sample-6	-5	1 =	100	98.0942	98.0943	98.0942	0.0000	0	YI
11	Sample-7	2767-1	1 =	100	113.0698	113.0697	113.0698	0.0000	0	PR
12	Sample-8	-2	1 =	100	115.3407	115.4045	115.4044	0.0637	637	J
13	Sample-9	-3	1 =	100	112.3016	112.3773	112.3771	0.0757	757	K
14	Sample-10	-4	1 =	100	117.6052	117.6704	117.6703	0.0651	651	XJ
15	LCSD	T1115	1 =	100	115.1311	115.1708	115.1706	0.0395	395	10
16	Sample-11	2787-5	1 =	100	116.5582	116.5996	116.5995	0.0413	413	101
17	Sample-12	-6	1 =	100	107.3030	107.3531	107.3533	0.0503	503	L
18	Sample-13	2786-1	1 =	10.0	116.9497	116.9888	116.9886	0.0389	389	N
19	Sample-14		1 =						778	
20	Sample-15		1 =							
21	Sample-16		1 =							
22	Sample-17		1 =							
23	Sample-18		1 =							
24	Sample-19		1 =							
25	Sample-20		1 =							
26	Mtx Dup.		1 =							

STD Lot #	CSTD(μ s/mL) $\times$ VSTD(mL) / X(s or mL) = T	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
W- 7618	x / = 400 ppm	%	85-115 %/80-120 %	PQL(w) 10
W- ✓	x / = ppm	%		PQL(s) 50
W- 7619	x / = ppm	%	90-110 %/85-115 %	MDL(w) 4
W- ✓	x / = ppm	%		MDL(s) 20

Form 5-127, March 7, 1995. Ver. 3.0

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

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

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

Alkalinity/OH/CO₃/HCO₃ (310.1/SM2320B) WorksheetsBatch # 03W2446 Matrix: WTitrant H₂SO₄ Lot # W7777Concentration (C) 0.02535N: Test Date: 4/18/03Analyst: SL

SOP: G-51

#	Sample ID	Dilution V ₁ /V ₂ =f ₁	Smpl Amt V ₁ , mL	H ₂ SO ₄ (mL) by Pnhl S _A E _A	A	H ₂ SO ₄ (mL) by MR-BCG S _B E _B	B	Pnhl-Alk., P (in unit of mgCaCO ₃ /L)	To. Alk., T	OH ⁻	CO ₃ ²⁻	HCO ₃ ⁻	Note & Anomaly
1	MB: T1118	/ =	100		0		0	0	0	0	0	0	
2	21.55-1	/ =	25		4.05		207.0	0	0	0	207.0		
3	-2	/ =	1		3.90		199.3	0	0	0	199.3		
4	21.57-1	/ =	50		5.40		138.0	0	0	0	138.0		
5	-2	/ =	25		5.40		238.0	0	0	0	238.0		
6	-3	/ =	1		5.00		255.5	0	0	0	255.5		
7	-4	/ =	1		6.60		337.3	0	0	0	337.3		
8	-5	/ =	1		0		0	0	0	0	0		
9	27.67-1	/ =	1		0		0	0	0	0	0		
10	-2	/ =	1		3.55		181.4	0	0	0	181.4		
11	-3	/ =	1		6.25		319.4	0	0	0	319.4		
12	-4	/ =	1		5.60		286.2	0	0	0	286.2		
13	-5	/ =	1		4.30		219.7	0	0	0	219.7		
14	-6	/ =	1		3.85		98.4	0	0	0	98.4		
15		/ =	1										
16		/ =	1										
17		/ =	1										
18		/ =	1										
19	LCS	/ =	100		2.95		101.6						
20	LCSO	/ =	100		8.00		102.2						
Dup.	27.55-1	/ =	25		4.10		209.5	0	0	0	209.5		

Calculation:

A = S_A - E_AB = S_B - E_BP = 50,000 f₁ A C / VT = 50,000 f₁ (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

Balance Daily Calibration Worksheet

Weight Set S/N: 12006

Calib. Date	Lab Balance					Digital Balance					Analytical Balance					Calib. by
	Balance #	1 g ±0.05g	10 g ±0.1g	200 g ±0.5g	Note (C)	Balance #	1 g ±0.02g	10 g ±0.05g	200 g ±0.10g	Note (D) (C) (AR)	Balance #	1 g ±0.0002g	10 g ±0.0005g	200 g ±0.0010g	Note (D) (C) (AR)	
4/13/03	A-01	Net	in use		✓	B-01	1.00	10.00	200.01	✓ ✓ ✓	C-01	1.0000	200.0003	200.0001	✓ ✓ ✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓ ✓ ✓	C-02	1.0000	200.0001	200.0000	✓ ✓ ✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.00	200.01	✓ ✓ ✓	C-					
	A-04					B-07	1.00	10.00	199.99	✓ ✓ ✓	C-					
	A-					B-					C-					
4/16/03	A-01	Net	in use		✓	B-01	1.00	10.00	200.01	✓ ✓ ✓	C-01	1.0000	10.00	200.0000	✓ ✓ ✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓ ✓ ✓	C-02	1.0000	10.02	199.9999	✓ ✓ ✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.01	200.00	✓ ✓ ✓	C-					
	A-04					B-07	1.00	10.00	199.99	✓ ✓ ✓	C-					
	A-					B-					C-					
4/17/03	A-01	Net	in use		✓	B-01	1.00	10.00	200.01	✓ ✓ ✓	C-01	1.0000	10.00	200.0001	✓ ✓ ✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓ ✓ ✓	C-02	1.0000	10.01	200.0000	✓ ✓ ✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.01	200.00	✓ ✓ ✓	C-					
	A-04					B-07	1.00	10.00	199.99	✓ ✓ ✓	C-					
	A-					B-					C-					
4/18/03	A-01	Net	in use		✓	B-01	1.00	10.00	200.01	✓ ✓ ✓	C-01	1.0000	10.00	200.0001	✓ ✓ ✓	1/2
	A-02					B-05	1.00	9.99	200.00	✓ ✓ ✓	C-02	1.0000	10.01	200.0000	✓ ✓ ✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.01	200.00	✓ ✓ ✓	C-					
	A-04					B-07	1.00	10.00	199.99	✓ ✓ ✓	C-					
	A-					B-					C-					

Notation: (C) - Cleanliness; (D) - Display; (AR) - Auto Rereading.
 APCI, form 4-21a, March 30, 1995, Ver. 4.0 No pencil. Use blue pen for record. Use red pen for correction.
 File: CUST.DOC\LAB\BAL-CAL-TEX Root-File: BAL-CAL-ROOT-TEX 1-Page-File: BAL-CAL-1-TEX

Balance Daily Calibration Worksheet

Weight Set S/N: 12006

067

3134

Calib. Date	Lab Balance					Digital Balance					Analytical Balance					Calib. by
	Balance #	1 g ±0.05g	10 g ±0.1g	200 g ±0.5g	Note (C)	Balance #	1 g ±0.02g	10 g ±0.05g	200 g ±0.10g	Note (D) (C) (AR)	Balance #	1 g ±0.0002g	10 g ±0.0005g	200 g ±0.0010g	Note (D) (C) (AR)	
4/21/03	A-01	Net in Wt			✓	B-01	1.00	9.99	200.01	✓✓✓	C-01	1.000	9.9999	200.0001	✓✓✓	✓
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.000	9.9999	200.0000	✓✓✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.01	200.00	✓✓✓	C-03					
	A-04					B-07	1.00	10.00	199.99	✓✓✓	C-04					
	A-					B-					C-					
4/22/03	A-01	Net in Wt			✓	B-01	1.00	10.00	200.01	✓✓✓	C-01	1.000	9.9999	200.0001	✓✓✓	✓
	A-02					B-05	1.00	9.99	200.00	✓✓✓	C-02	1.000	9.9999	200.0000	✓✓✓	
	A-03	1.00	10.00	200.01	✓	B-06	1.00	10.00	200.00	✓✓✓	C-03					
	A-04					B-07	1.00	10.00	199.99	✓✓✓	C-04					
	A-					B-					C-					
4/24/03	A-01	Net in Wt			✓	B-01	1.00	10.00	200.00	✓✓✓	C-01	1.000	9.9999	200.0000	✓✓✓	✓
	A-02					B-05	1.00	10.01	200.01	✓✓✓	C-02	1.000	10.0000	200.0002	✓✓✓	
	A-03	1.00	10.00	200.00		B-06	1.00	10.00	200.00	✓✓✓	C-03					
	A-04					B-07	1.00	9.99	199.99	✓✓✓	C-04					
	A-					B-					C-					

Notation: (C) - Cleanliness; (D) - Display; (AR) - Auto Rerzeroing.
APCL form 4-213, March 30, 1995, Ver. 4.0 No pencil. Use blue pen for record. Use red pen for correction.
File: [CUST.DOC]LABAL.CAL.TEX Root-File: BAL.CAL.ROOT.TEX 1-Page-File: BAL.CAL.1.TEX



Applied Physics & Chemistry Laboratory

13760 Magnolia Ave. Chino CA 91710

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

June 9, 2003

GEOFON, Inc.

Attention: Leo Williamson

22632 Golden Spring Dr Ste 270

Diamond Bar CA 91765

Dear Leo Williamson,

This package contains samples in our Service ID 03-3130 and your project : 04-4428.10 JPL GW Mon-2Q03.

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

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Submitted to:

GEOFON, Inc.

Attention: Leo W. Williamson

22632 Golden Spring Dr Ste 270

Diamond Bar CA 91765

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

APCL Analytical Report

Service ID #: 801-033130

Received: 05/08/03

Collected by: Leo Williamson

Extracted: N/A

Collected on: 05/08/03

Tested: 05/08-20/03

Reported: 05/30/03

Sample Description: Water from MW-22 (N. of Bl 180)

Project Description: 04-4428.10 JPL GW Mon-2Q03

Analysis of Water Samples

Component Analyzed	Method	Unit	PQL	Analysis Result		
				EB-12-5/8/03 03-03130-1	MW-22-1 03-03130-2	MW-22-2 03-03130-3
BICARBONATE	SM2320B	mg/L	2	< 2	271	148
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	< 2	< 2	2.6
PH	9040B	pH unit	0.01	7.31	7.64	8.21
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	< 10	646	287
CHROMIUM (VI)	7196	mg/L	0.01	< 0.01	< 0.01	< 0.01
Dilution Factor				1	1	1
PERCHLORATE	314.0	µg/L	4	< 4	3.2J	1.6J
Dilution Factor				1.25	12.5	5
CHLORIDE CL ⁻	300.0	mg/L	0.2	0.27	107	34.7
NITRATE AS N	300.0	mg/L	0.04	0.087	9.1	6.8
SULFATE SO ₄ ⁻²	300.0	mg/L	0.5	0.69	138	29.7
Dilution Factor				1	1	1
ARSENIC	200.9	µg/L	5	< 5	< 5	< 5
CALCIUM	200.7	µg/L	200	< 200	108,000	35,000
IRON	200.7	µg/L	50	31.4J	115	81.3
MAGNESIUM	200.7	µg/L	100	20.7J	39,100	16,800
POTASSIUM	200.7	µg/L	400	93.5J	3,110	2,360
SODIUM	200.7	µg/L	2000	1,240J	30,700	30,400
VOLATILE ORGANIC COMPOUNDS						
Dilution Factor				1	1	1
BENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
BROMOBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
BROMOCHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
BROMODICHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
BROMOFORM	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
BROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
2-BUTANONE	524.2	µg/L	10	< 10	< 10	< 10
N-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
SEC-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
TERT-BUTYLBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CARBON TETRACHLORIDE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CHLOROBENZENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CHLORODIBROMOMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CHLOROETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CHLOROFORM	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
CHLOROMETHANE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5
2-CHLOROTOLUENE	524.2	µg/L	0.5	< 0.5	< 0.5	< 0.5

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result		
				EB-12-5/8/03 03-03130-1	MW-22-1 03-03130-2	MW-22-2 03-03130-3
4-CHLOROTOLUENE	524.2	µg/L	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
1,2-DIBROMOETHANE (EDB)	524.2	µg/L	0.5	<0.5	<0.5	<0.5
DIBROMOMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,3-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,4-DICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
DICHLORODIFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
CIS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TRANS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,3-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
2,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
CIS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TRANS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
ETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
HEXACHLOROBUTADIENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
ISOPROPYLBENZENE (CUMENE)	524.2	µg/L	0.5	<0.5	<0.5	<0.5
P-ISOPROPYLTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	5J	3J	5J
METHYLENE CHLORIDE	524.2	µg/L	1.8 (a)	<1.8	2.0	<1.8
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1
NAPHTHALENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
N-PROPYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
STYRENE	524.2	µg/L	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
1,1,2,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TETRACHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2,4-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1,1-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1,2-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TRICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
TRICHLOROFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2,4-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,3,5-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
VINYL CHLORIDE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
O-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
M/P-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				MW-22-3 03-03130-4	MW-22-4 03-03130-5	MW-22-5 03-03130-6	TB-12-5/8/03 03-03130-7
BICARBONATE	SM2320B	mg/L	2	130	114	49.9	-
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	5.2	2.6	76.6	-
PH	9040B	pH unit	0.01	8.18	8.07	9.23	-
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	287	213	208	-
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	1.8J	<4	<4	-
Dilution Factor				5	2	2	1
CHLORIDE CL ⁻	300.0	mg/L	0.2	33.9	11.8	8.1	-
NITRATE AS N	300.0	mg/L	0.04	6.9	4.1	0.11	-
SULFATE SO ₄ ⁻	300.0	mg/L	0.5	34.6	7.6	22.0	-
Dilution Factor				1	1	1	1
ARSENIC	200.9	µg/L	5	<5	<5	<5	-
CALCIUM	200.7	µg/L	200	29,500	29,400	4,580	-
IRON	200.7	µg/L	50	130	53.8	57.5	-
MAGNESIUM	200.7	µg/L	100	14,500	8,450	625	-
POTASSIUM	200.7	µg/L	400	2,360	1,700	1,130	-
SODIUM	200.7	µg/L	2000	36,800	26,900	70,200	-
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
2-BUTANONE	524.2	µg/L	10	<10	<10	<10	<10
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
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	<0.5	<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-22-3 03-03130-4	MW-22-4 03-03130-5	MW-22-5 03-03130-6	TB-12-5/8/03 03-03130-7
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
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	6J	9J	5J	5J
METHYLENE CHLORIDE	524.2	µg/L	1.8 ^(a)	<1.8	<1.8	<1.8	3.6
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1	<1
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	<0.5	<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
1,1,2,2,3,3-HEXACHLORO-1,2-DIFLUOROETHANE	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.

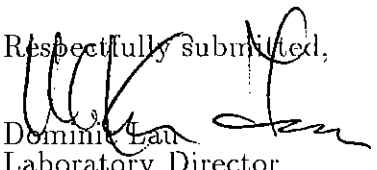
"-": Analysis is not required.

J: Reported between PQL and MDL.

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

^(a) MDL reported.

Respectfully submitted,


 Dominick Lau
 Laboratory Director
 Applied P & Ch Laboratory

Level C Data Package Deliverables

General Information

Project: 04-4428.10 JPL GW Mon-2Q03

APCL Service ID: 03-3130



Applied Earth & Ch Laboratory

13760 Magnolia Ave. Chino, CA 91710

Telephone (909)590-1828

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Case Narrative

Project: JPL / 04-4428.10

For GEOFON, Inc.

APCL Service No: 03-3130

1. Sample Identification

The sample identifications are listed in the following table:

GEOFON, Inc. Sample ID	APCL Sample ID
MW-22-5	03-03130-6
MW-22-4	03-03130-5
MW-22-3	03-03130-4
MW-22-2	03-03130-3
MW-22-1	03-03130-2
TB-12-5/8/03	03-03130-7
EB-12-5/8/03	03-03130-1

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),
206.2 (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

(1) SW8260B:

Methylene Chloride in the amount of 4.7 ug/L was detected in the Method Blank of batch 03G2534. Methylene Chloride was also detected in the associated field samples, due to lab contamination.

(2) Arsenic - 200.9:

Batch 03M1458E included SDG 03-3082 and 03-3130 from the same project. MS/MSD for Arsenic was performed only on the sample MW-11-3, from SDG 03-3082; and it was not analyzed on the sample MW-22-1, from SDG 03-3130.

"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,



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



INCOPIATE
22632 GOLDEN SPRINGS DR., SUITE 270
DIAMOND BAR, CA 91765 • (909) 396-7662 • FAX (909) 396-1455

CHAIN-OF-CUSTODY RECORD

LABORATORY COPY

MW-22 0031

GEOFON LAB COORDINATOR		LAB COORDINATOR'S PHONE		LAB COORDINATOR'S FAX		LABORATORY SERVICE ID		LABORATORY CONTACT		MAIL REPORT (COMPANY NAME)	
Brad Shojee		(909) 396-7662		(909) 396-1455		-		Kenny Chan		GEOFON, INC.	
PROJECT NAME:		PROJECT LOCATION		PROJECT NUMBER		LABORATORY PHONE		LABORATORY FAX		RECIPIENT NAME	
JPL-BW MON-2003		MW-22 (N. of B1 180)		04-442810		(909) 396-1828		(909) 396-1498		Leo W. Williamson	
PROJECT CONTACT		PROJECT PHONE NUMBER		PROJECT FAX		LABORATORY ADDRESS		LABORATORY ADDRESS		ADDRESS	
Leo W. Williamson		(714) 920-8729		(909) 396-1455		13700 Magnolia Ave.		13700 Magnolia Ave.		22632 Golden Springs Dr. #270	
PROJECT ADDRESS		CITY, STATE AND ZIP CODE		CLIENT		CITY, STATE AND ZIP CODE		CITY, STATE AND ZIP CODE		CITY, STATE AND ZIP CODE	
4800 Oak Lane Dr.		Pasadena, CA		US NAVY SWD/V		Ching CA.		Ching CA.		Diamond Bar, CA. 91765	
PROJECT MANAGER		PROJECT MANAGER'S PHONE		PROJECT MANAGER'S FAX		PROJECT MANAGER'S FAX		PROJECT MANAGER'S FAX		PROJECT MANAGER'S FAX	
Asrar Fakhrean		(909) 396-7662		(909) 396-1455		(909) 396-1455		(909) 396-1455		(909) 396-1455	
Item	Sample Identifier	Matrix	Date	Time	Preserved	# of Cont.	OC Level	TAT	Comments		
1	MW-22-5	H ₂ O	5/8/03	800	NONE	3+1+	III	NORMAL	X	X	MINERALS: Na/K/Ca/As/Mg/Fe
2	MW-22-4			840	NONE	1+1			X	X	
3	MW-22-3			925					X	X	
4	MW-22-2			1000					X	X	
5	MW-22-1			1105					X	X	
6											
7	TB-12-5/8/03	H ₂ O	-	-	HCl	2	III	NORMAL	X	X	
8	EB-12-5/8/03		5/8/03	850	NONE	3+1+			X	X	
9											
10											
SAMPLES COLLECTED BY: Leo W. Williamson		COURIER AND AIR BILL NUMBER		RECEIVED BY: S. Shojee		DATE: 5-9-03		TIME: 12:05		COOLER TEMPERATURE UPON RECEIPT	
RELINQUISHED BY: S. Shojee				RECEIVED BY: S. Shojee		DATE: 5/8/03		TIME: 1330		SAMPLE'S CONDITION UPON RECEIPT	
X.S. Shojee				X.S. Shojee							

Sample Receiving Checklist

APCL ServiceID: **3130** Client Name/Project: Geon JPL

1. Sample Arrival

Date/Time Received 5/8/03 1330 Date/Time Opened 5/8/03 1330 By (name): Venus Chan
Custody Transfer: ☐ Client ☐ Golden State ☐ UPS ☐ US Mail ☐ FedEx ☒ APCL Empl: Scott B.

2. Chain-of-Custody (CoC)

☒ With Samples? ☐ Faxed? ☐ Client has Copy? ☐ Signed, dated? By: _____
☐ Project ID? ☒ Analyses Clear? ☐ Hold Samples? #on Hold _____ # Received 6
☒ CoC/Docs Zip-Locked under lid? ☐ Compos.#: _____ ☒ #Samples OK? _____
☐ Discrepancies? ☐ Client notified? ☐ Response (attach docs): _____

3. Shipping Container/Cooler

☒ Cooler Used? # of 1 Cooled by: ☒ Ice ☐ Blue Ice ☐ Dry Ice ☐ None
Temp °C 40
(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: [Signature] Date: 8 May 2003 Time: 7:41 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

03-03130 (0470_ 141) (2202777_ 141)

05/08/03

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>05/08/03 1330</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-22-5	VOC	03-03130-6- α	W	V	C	40	3	G	050803	N	0	7 ☐
	MW-22-5	Metal	03-03130-6- β	W	P	N	500	1	G	050803	N	0	7 ☐
	MW-22-5	300	03-03130-6- γ	W	P		1000	1	G	050803	N	0	7 ☐
2	MW-22-4	VOC	03-03130-5- α	W	V	C	40	3	G	050803	N	0	7 ☐
	MW-22-4	Metal	03-03130-5- β	W	P	N	500	1	G	050803	N	0	7 ☐
	MW-22-4	300	03-03130-5- γ	W	P		1000	1	G	050803	N	0	7 ☐
3	MW-22-3	VOC	03-03130-4- α	W	V	C	40	3	G	050803	N	0	7 ☐
	MW-22-3	Metal	03-03130-4- β	W	P	N	500	1	G	050803	N	0	7 ☐
	MW-22-3	300	03-03130-4- γ	W	P		1000	1	G	050803	N	0	7 ☐
4	MW-22-2	VOC	03-03130-3- α	W	V	C	40	3	G	050803	N	0	7 ☐
	MW-22-2	Metal	03-03130-3- β	W	P	N	500	1	G	050803	N	0	7 ☐
	MW-22-2	300	03-03130-3- γ	W	P		1000	1	G	050803	N	0	7 ☐
5	MW-22-1	VOC	03-03130-2- α	W	V	C	40	6	G	050803	N	0	7 ☐
	MW-22-1	Metal	03-03130-2- β	W	P	N	500	2	G	050803	N	0	7 ☐
	MW-22-1	300	03-03130-2- γ	W	P		1000	2	G	050803	N	0	7 ☐
6	TB-12-5/8/03	VOC	03-03130-7	W	V	C	40	2	G	050803	N	0	7 ☐
7	EB-12-5/8/03	VOC	03-03130-1- α	W	V	C	40	3	G	050803	N	0	7 ☐
	EB-12-5/8/03	Metal	03-03130-1- β	W	P	N	500	1	G	050803	N	0	7 ☐
	EB-12-5/8/03	300	03-03130-1- γ	W	P		1000	1	G	050803	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/SM4500NOR 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
- ☐ 200.7/6010B Potassium, K, by ICP
- ☐ 200.7/6010B Magnesium, Mg, by ICP
- ☐ 200.7/6010B Iron, Fe, 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: 05/19/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: 03G2534-MB-01	Lab Sample ID: 03G2534-MB-01	Received Date: 05/19/2003
Sample Type: Method Blank	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: G2534K01	Prep. No: -	Anal. Time: 20:12
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8	4.7	
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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)	70-129	107
2	1,2-DICHLOROETHANE-D4	70-129	98
3	DIBROMOFLUOROMETHANE	70-122	104
4	TOLUENE-D8	73-129	107
# of out-of-control			0

Internal Standard

		Control Limit, %	IS Rec.%
1	CHLOROBENZENE-D5	50-200	85
2	1,4-DICHLOROBENZENE-D4	50-200	89
3	FLUOROBENZENE	50-200	93
# 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

Applied P & Ch Laboratory
Organic Analysis Results for Method 524.2

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: EB-12-5/8/03	Lab Sample ID: 03-3130-1	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-01	Prep. No: -	Anal. Time: 21:10
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	5	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 (a)	<1.8	U
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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-TETRACHLORO-1,2-DIFLUOROETHANE	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

70-129

98

70-129

96

70-122

99

73-129

103

of out-of-control

0

Internal Standard

Control Limit, %

IS Rec.%

1	CHLOROBENZENE-D5	3114-55-4
2	1,4-DICHLOROBENZENE-D4	3855-82-1
3	FLUOROBENZENE	462-06-6

50-200

85

50-200

93

50-200

93

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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: MW-22-1	Lab Sample ID: 03-3130-2	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-02	Prep. No: -	Anal. Time: 21:39
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	3	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 (a)	2.0	B
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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-TETRACHLORO-1,2,2,2-TRIFLUOROETHANE	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	98	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	92	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	99	
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	92	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	101	
3	FLUOROBENZENE	462-06-6		50-200	100	
# 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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: MW-22-2	Lab Sample ID: 03-3130-3	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-03	Prep. No: -	Anal. Time: 22:08
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	2-BUTANONE	78-93-3	µg/L	10	< 10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	< 0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	< 0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	< 0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	5	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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-TETRACHLORO-1,2-DIFLUOROETHANE	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

70-129

90

70-129

87

70-122

93

73-129

98

of out-of-control

0

Internal Standard

Control Limit, %

IS Rec. %

1	CHLOROBENZENE-D5	3114-55-4
2	1,4-DICHLOROBENZENE-D4	3855-82-1
3	FLUOROBENZENE	462-06-6

50-200

94

50-200

109

50-200

107

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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: MW-22-3	Lab Sample ID: 03-3130-4	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-04	Prep. No: -	Anal. Time: 22:37
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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
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19	1,2-DIBROMO-3-CHLOROPROPANE	96-12-8	µg/L	1.1 (a)	<1.1	U
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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
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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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	6	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 (a)	<1.8	U
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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	112TRICHLORO-122TRIFLUOROETHANE	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	98	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	105	
4	TOLUENE-D8	2037-26-5		73-129	111	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	90	
2	1,4-DICHLOROETHANE-D4	3855-82-1		50-200	105	
3	FLUOROBENZENE	462-06-6		50-200	98	
# 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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: MW-22-4	Lab Sample ID: 03-3130-5	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-05	Prep. No: -	Anal. Time: 23:06
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	9	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	<1.8	U
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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	98	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	91	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	97	
4	TOLUENE-D8	2037-26-5		73-129	102	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	97	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	105	
3	FLUOROBENZENE	462-06-6		50-200	107	
# 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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: MW-22-5	Lab Sample ID: 03-3130-6	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/19/03	Anal. Date: 05/19/03
Data File Name: 3130-06	Prep. No: -	Anal. Time: 23:35
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	5	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 (a)	<1.8	U
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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

70-129

110

70-129

101

70-122

106

73-129

114

of out-of-control

0

Internal Standard

Control Limit, %

IS Rec.%

1	CHLOROBENZENE-D5	3114-55-4
2	1,4-DICHLOROBENZENE-D4	3855-82-1
3	FLUOROBENZENE	462-06-6

50-200

83

50-200

96

50-200

95

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: 05/08/2003
Project ID: JPL GW Mon-2Q03	Service ID: 33130	Collected by:
Sample ID: TB-12-5/8/03	Lab Sample ID: 03-3130-7	Received Date: 05/08/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: G
Batch No: 03G2534	Prep. Date: 05/20/03	Anal. Date: 05/20/03
Data File Name: 3130-07	Prep. No: -	Anal. Time: 00:04
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	2-BUTANONE	78-93-3	µg/L	10	<10	U
8	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
9	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
10	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	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	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	5	J
42	METHYLENE CHLORIDE	75-09-2	µg/L	1.8 ^(a)	3.6	B
43	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	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

70-129

94

70-129

89

70-122

96

73-129

101

of out-of-control

0

Internal Standard

Control Limit, %

IS Rec.%

1	CHLOROBENZENE-D5	3114-55-4
2	1,4-DICHLOROBENZENE-D4	3855-82-1
3	FLUOROBENZENE	462-06-6

50-200

97

50-200

109

50-200

107

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.
 Case No:
 Project ID: JPL GW Mon-2Q03

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03G2534

Lab Code: APCL
 SDG Number: 033130
 Sample Matrix: Water

#	Client Sample No	Lab Sample ID	S1 % #	S2 % #	S3 % #	S4 % #	TOT OUT
1	03G2534-LCS-01	03G2534-LCS-01	88	105	102	99	0
2	MW-22-1MS	03-3130-2MS	87	97	95	94	0
3	MW-22-1MSD	03-3130-2MSD	96	111	107	103	0
4	03G2534-MB-01	03G2534-MB-01	107	98	104	107	0
5	EB-12-5/8/03	03-3130-1	98	96	99	103	0
6	MW-22-1	03-3130-2	98	92	99	104	0
7	MW-22-2	03-3130-3	90	87	93	98	0
8	MW-22-3	03-3130-4	101	98	105	111	0
9	MW-22-4	03-3130-5	98	91	97	102	0
10	MW-22-5	03-3130-6	110	101	106	114	0
11	TB-12-5/8/03	03-3130-7	94	89	96	101	0
12							
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.
 Case No:
 Project ID: JPL GW Mon-2Q03

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03G2534

Lab Code: APCL
 Service ID: 33130
 Sample Matrix: Water

LCS Filename: G2534L01
 LCSD Filename: -

Date Analyzed: 051903
 Date Analyzed: -
 Time Analyzed: 17:18
 Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
BENZENE	µg/L	20	0	20.1	101	65-120
CHLOROBENZENE	µg/L	20	0	20.4	102	65-134
1,1-DICHLOROETHENE	µg/L	20	0	20.0	100	65-127
TOLUENE	µg/L	20	0	19.2	96	65-134
TRICHLOROETHENE	µg/L	20	0	20.0	100	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: _____

FORM-3A

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 524.2

Client Name: GEOFON, Inc.
 Case No:
 Project ID: JPL GW Mon-2Q03

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03G2534

Lab Code: APCL
 Service ID: 33130
 Sample Matrix: Water

MS Filename: G2534M01
 MSD Filename: G2534N01
 MS Sample No: MW-22-1

Date Analyzed: 051903
 Date Analyzed: 051903
 Sample Lab ID: 03-3130-2

Time Analyzed: 17:47
 Time Analyzed: 18:16

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
BENZENE	µg/L	20	0	18.7	94	65-121
CHLOROBENZENE	µg/L	20	0	19.2	96	65-134
1,1-DICHLOROETHENE	µg/L	20	0	18.7	94	65-127
TOLUENE	µg/L	20	0	18.0	90	65-134
TRICHLOROETHENE	µg/L	20	0	19.1	96	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	20.5	103	9	28	65-121
CHLOROBENZENE	µg/L	20	21.4	107	11	35	65-134
1,1-DICHLOROETHENE	µg/L	20	20.3	102	8	31	65-127
TOLUENE	µg/L	20	19.8	99	10	35	65-134
TRICHLOROETHENE	µg/L	20	21.6	108	12	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: _____

FORM-4A

Applied P & Ch Laboratory

Method Blank Summary for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 33130

Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10

Analysis Date: 05/19/03

Sample Matrix: Water

Analysis Time: 20:12

Sample ID: 03G2534-MB-01

Batch No: 03G2534

Instrument ID: GC/MS: G

Lab Sample ID: 03G2534-MB-01

Data File Name: G2534K01

GC Column: DB-VEX

Heated Purge: (Y/N) N

Column ID: 0.45 mm

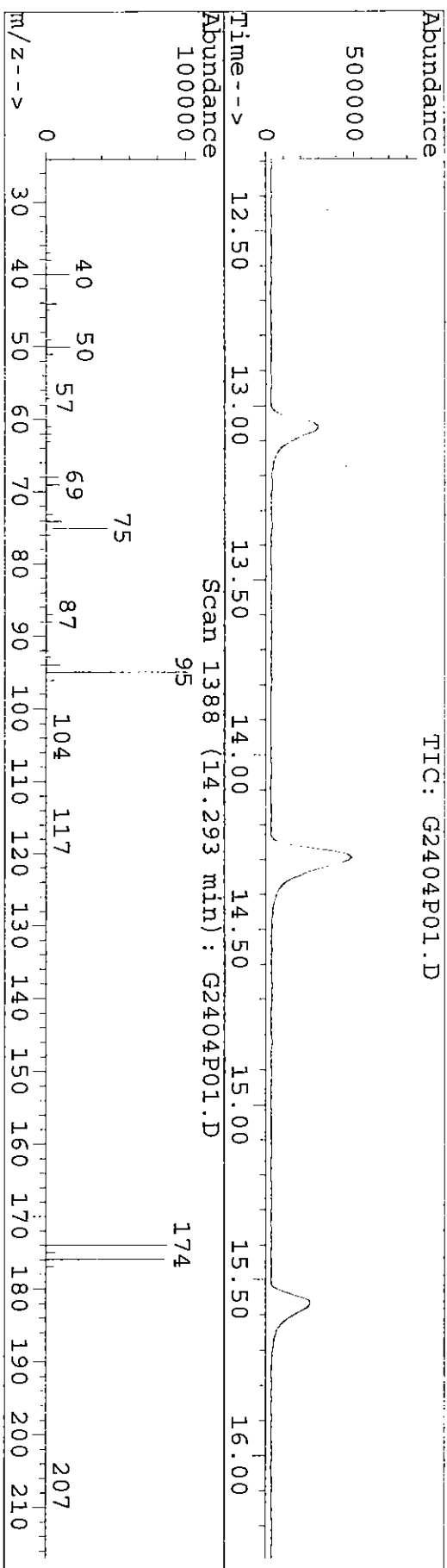
This Method Blank applies to the following samples and QC samples:

#	Client Sample No	Lab Sample ID	Sample Type	Data Filename	Analysis Date	Analysis Time
1	03G2534-LCS-01	03G2534-LCS-01	Lab Control Spike	G2534L01	05/19/03	17:18
2	MW-22-1MS	03-3130-2MS	Matrix Spike	G2534M01	05/19/03	17:47
3	MW-22-1MSD	03-3130-2MSD	Matrix Spike Duplicate	G2534N01	05/19/03	18:16
4	EB-12-5/8/03	03-3130-1	Field Sample	3130-01	05/19/03	21:10
5	MW-22-1	03-3130-2	Field Sample	3130-02	05/19/03	21:39
6	MW-22-2	03-3130-3	Field Sample	3130-03	05/19/03	22:08
7	MW-22-3	03-3130-4	Field Sample	3130-04	05/19/03	22:37
8	MW-22-4	03-3130-5	Field Sample	3130-05	05/19/03	23:06
9	MW-22-5	03-3130-6	Field Sample	3130-06	05/19/03	23:35
10	TB-12-5/8/03	03-3130-7	Field Sample	3130-07	05/20/03	00:04
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						
23						
24						
25						

Data File : C:\HPCHEM\1\DATA\03G2404\G2404P01.D
 Acq On : 15 May 03 10:18 am
 Sample : ##03g2404, w
 Misc :

Vial: 18
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & ch Lab** EPA 524.2



Peak Apex is scan: 1388

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
50	95	15	40	16.7	17208	PASS
75	95	30	60	43.0	44184	PASS
95	95	100	100	100.0	102784	PASS
96	95	5	9	6.1	6301	PASS
173	174	0	2	0.0	0	PASS
174	95	50	100	84.8	87112	PASS
175	174	5	9	7.7	6735	PASS
176	174	95	101	97.9	85240	PASS
177	176	5	9	6.8	5764	PASS

5A
VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name : APPLIED P & CH LAB Contract: _____
 Lab Code: _____ Case No.: _____ SAS No.: _____ SDG No.: 03-3130
 Lab File ID: G2404 P01 BFB Injection Date: 05/15/2003
 Instrument ID: GCMS-G BFB Injection Time: 1018
 GC Column: VOCOL ID: 0.32 (mm) Heated Purge: (Y/N) N

m/e	ION ABUNDANCE CRITERIA	%RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	16.7
75	30.0 - 60.0% of mass 95	43.0
95	Base peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.1
173	Less than 2.0% of mass 174	0.0 (0.0)1
174	50.0 - 100.0% of mass 95	84.8
175	5.0 - 9.0% of mass 174	6.6 (7.7)1
176	95.0 - 101.0% of mass 174	82.9 (97.9)1
177	5.0 - 9.0% of mass 176	5.6 (6.8)2

1-Value is % mass 174

2-Value is % mass 176

This check applies to the following SAMPLES, MS, MSD, BLANKS and STANDARDS:

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01	VSTD003	004-003	004-0003.D	05/15/03	1047
02	VSTD002	004-002	004-0002.D	05/15/03	1145
03	VSTD010	004-0010	004-0010.D	05/15/03	1214
04	VSTD020	004-0020	004-0020.D	05/15/03	1243
05	VSTD040	004-0040	004-0040.D	05/15/03	1312
06	VSTD080	004-0080	004-0080.D	05/15/03	1341
07					
08					
09					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					

INITIAL CALIBRATION SUMMARY

Method File

E524G004

Last Calibration Update

Fri May 16 10:36:26 2003

<u>Level 1 File Name</u>	4-003.D	<u>Level 1 ID</u>	0.3
<u>Level 2 File Name</u>	4-002.D	<u>Level 2 ID</u>	2
<u>Level 3 File Name</u>	4-010.D	<u>Level 3 ID</u>	10
<u>Level 4 File Name</u>	4-020.D	<u>Level 4 ID</u>	20
<u>Level 5 File Name</u>	4-040.D	<u>Level 5 ID</u>	40
<u>Level 6 File Name</u>	4-080.D	<u>Level 6 ID</u>	80
<u>Level 7 File Name</u>	4-020.D	<u>Level 7 ID</u>	cc

<u>Compound Name</u>	<u>Level 1 Response</u>	<u>Level 2 Response</u>	<u>Level 3 Response</u>	<u>Level 4 Response</u>	<u>Level 5 Response</u>	<u>Level 6 Response</u>	<u>Level 7 Response</u>	<u>Coeff X⁰</u>	<u>Coeff X¹ / ave RF</u>	<u>Coeff X²</u>	<u>R² / RSD</u>
1 Fluorobenzene l1 1	815158	799651	824804	746757	764394	689688	-1	-----	-----	-----	-----
3 di-Cl-di-F-methane 85 87	4181	35493	168476	314256	706635	1288384	-1	0.0000	0.2120	0.0000	0.1090
4 Chloromethane 50 52	1706	19850	102587	246113	534218	1069891	-1	-0.0408	0.1954	0.0000	0.9966
9 F114 85 135	4199	44804	220942	386132	819274	1556249	-1	-0.0188	0.2815	0.0000	0.9991
5 vinyl chloride 62 64	3355	33999	162051	300841	621901	1167862	-1	0.0000	0.1938	0.0000	0.1466
6 bromomethane 94 96	4372	26980	129267	267214	519945	1000442	-1	0.0000	0.1724	0.0000	0.0536
7 Chloroethane 64 66	3179	26106	123679	236431	478026	873637	-1	0.0000	0.1527	0.0000	0.0781
8 tri-Cl-F-methane 101 103	8951	64415	295972	594414	1265977	2395096	-1	0.0000	0.3956	0.0000	0.0724
111 isopropyl alcohol x10	3587	6299	29925	70395	111196	226721	-1	0.0000	0.0040	0.0000	0.1113
100 ethyl ether x5	26492	132391	571623	1090032	2124645	3947698	-1	0.0028	0.1423	0.0000	0.9997
102 Acrolein x10	1454	15421	124387	164659	294058	649263	-1	-0.0048	0.0115	0.0000	0.9858
119 methyl acetate	7468	34881	93789	329077	525452	898426	-1	0.0239	0.1624	0.0000	0.9860
104 Carbon disulfide	15360	108670	470593	924294	1876377	3448920	-1	0.0000	0.6226	0.0000	0.0560
103 Acrylonitrile x10	4460	38141	123788	403095	766254	1455017	-1	-0.0461	0.0268	0.0000	0.9965
95 Acetone x10	18003	42021	146218	266351	522927	928171	-1	0.0183	0.0166	0.0000	0.9999
108 F-113	7993	56604	258951	515229	1098171	2025833	-1	0.0000	0.3443	0.0000	0.0590
13 11-dichloroethene 61 96	8587	60316	284348	535921	11112083	2068506	-1	0.0000	0.3617	0.0000	0.0355
101 Acetonitrile x10	2831	10077	31282	102405	196047	363830	-1	-0.0063	0.0067	0.0000	0.9966
109 Iodomethane	6929	75147	336586	628909	1205092	2300947	-1	-0.0067	0.4140	0.0000	0.9991
113 Tert butyl alcohol x10	1373	18588	77974	138129	265783	388696	-1	0.0000	0.0092	0.0000	0.1787
18 methylene chloride 49 84	-1	48402	184549	355095	696481	1289109	-1	0.0000	0.2451	0.0000	0.1330

112 Allyl chloride	13441	77142	331252	615063	1148799	1977528	-1	0.0468	0.3564	0.0000	0.9986
200 Nitro methane x10	6304	66437	325452	650702	1254351	2397871	-1	0.0000	0.0391	0.0000	0.1719
10 t-Bu-Me-ether 73 57	11529	91189	366763	722425	1502372	2824413	-1	0.0000	0.4955	0.0000	0.0864
19 t-12-di-Cl-ethene 96 61	5258	44435	215839	423127	799696	1549462	-1	0.0000	0.2634	0.0000	0.0969
98 Vinyl acetate x5	3645	167151	292003	1342007	2982123	5784036	-1	-0.2666	0.2134	0.0000	0.9933
21 11-dichloroethane 63 83	11126	88915	412703	833014	1647004	3000180	-1	0.0000	0.5252	0.0000	0.0765
91 2-butanone MEKx10	25733	153111	642126	1432839	2827847	5285771	-1	0.0000	0.0938	0.0000	0.0952
115 Di isoprop ether	40820	232542	1079994	2060781	4081017	7613956	-1	0.0000	1.4212	0.0000	0.0923
22 c-12-di-Cl-ethene 96 61	6716	44823	222557	432230	852613	1623067	-1	0.0000	0.2812	0.0000	0.0324
23 22-Dichloropropane 77 97	11379	71432	324674	629042	1268689	2353196	-1	0.0000	0.4280	0.0000	0.0585
24 Br-Cl-methane 128 130	619	19664	94361	180264	365219	680793	-1	-0.0056	0.1234	0.0000	0.9997
25 chloroform 83 85	15134	81585	381866	741343	1478585	2772452	-1	0.0000	0.5124	0.0000	0.1068
201 Ethyl acetate x2	2846	46100	136924	355515	703484	1425799	-1	-0.0563	0.1306	0.0000	0.9958
116 ETBE	33538	144136	644550	1245123	2437411	4486393	-1	0.0049	0.8099	0.0000	0.9998
117 Iso-butyl alcohol X10	940	62706	135151	357401	710558	1430613	-1	-0.0353	0.0259	0.0000	0.9953
26 tetrahydrofuranx5	257	8307	43847	84254	161549	290888	-1	0.0000	0.0107	0.0000	0.0325
27 Di-Br-F-Methane (S1) 111 1	8284	60816	289627	568312	1126216	2075916	-1	0.0000	0.3713	0.0000	0.0331
34 111-tri-Cl-ethane 97 99	8942	65770	311389	599971	1237882	2397744	-1	0.0000	0.3993	0.0000	0.0616
30 12-dichloroethane 64 62	1802	33229	153871	310815	631494	1211777	-1	-0.0194	0.2197	0.0000	0.9990
35 11-Di-Cl-propene 75 110	8689	58001	275836	532605	1078521	2009125	-1	0.0000	0.3543	0.0000	0.0301
29 1,2-di-Cl-ethane-d4 [Surf] 10	4884	26465	134448	261823	505671	963545	-1	0.0000	0.1688	0.0000	0.0341
36 benzene 78 52	20191	142907	668867	1302042	2578473	4754791	-1	0.0000	0.8512	0.0000	0.0359
37 CC14 117 119	8483	58348	273728	543038	1134691	2073848	-1	0.0000	0.3590	0.0000	0.0461
97 thiophene	9443	75171	331986	649163	1291816	2373848	-1	0.0000	0.4243	0.0000	0.0680
118 TAME	2423	114119	445234	894867	1715454	3200762	-1	-0.0058	0.5780	0.0000	0.9994
39 12-di-Cl-propane 63 76	5331	41840	206799	390600	777542	1446554	-1	0.0000	0.2514	0.0000	0.0677
40 trichloroethene 130 132	7533	51222	233771	454200	926225	1695738	-1	0.0000	0.3044	0.0000	0.0393
96 Me-methacrylate	1437	15341	47596	161499	295494	591170	-1	-0.0168	0.1076	0.0000	0.9949
42 Br-di-Cl-methane 83 85	12659	54887	257854	508851	988624	1842908	-1	-0.0042	0.3328	0.0000	0.9996
41 dibromomethane 174 172	618	19288	93298	195812	390733	738622	-1	-0.0102	0.1342	0.0000	0.9994
Compound	Level 1 Response	Level 2 Response	Level 3 Response	Level 4 Response	Level 5 Response	Level 6 Response	Level 7 Response	Coeff X ⁰	Coeff X ¹ / ave RF	Coeff X ²	R ² / RSD
45 c-13-di-Cl-propene 75 110	7615	48884	239463	477602	948616	1759634	-1	-----	-----	-----	-----
55 toluene-d8(S2) 100 99	13666	83281	390496	768293	1521774	2828221	-1	0.0000	0.5038	0.0000	0.0376
92 2-ClEt-VI-ether10	5311	41678	205749	413282	830636	1543091	-1	0.0000	0.0259	0.0000	0.0904
56 toluene 91 92	24063	127484	612856	1179183	2400670	4445368	-1	0.0000	0.8174	0.0000	0.1033

107 Et methacrylate	3479	23955	53825	215372	441923	820108	-1	-0.0241	0.1510	0.0000	0.9952
93 2-Hexanone x5	1510	42848	144809	430950	860345	1584358	-1	-0.0326	0.0581	0.0000	0.9977
48 112-tri-Cl-Et 97 83	906	19429	109523	184224	367637	694437	-1	-0.0023	0.1252	0.0000	0.9994
58 1,2-di-br-ethane 107 109	1824	20769	99010	199115	395372	752280	-1	-0.0087	0.1362	0.0000	0.9993
51 di-Br-Cl-methane 129 127	7347	33435	155803	299945	605212	1137349	-1	-0.0082	0.2056	0.0000	0.9995
46 t-13-di-cl-propene 75 110	7615	48884	239463	477602	948616	1759634	-1	0.0000	0.3094	0.0000	0.0348
105 1-Chlorohexane	11922	55414	207900	393247	824539	1487258	-1	-0.0003	0.2692	0.0000	0.9998
47 Cl-benzene-d5, 12	231880	227963	249245	222899	243736	235423	-1	0.0000	1.0000	0.0000	0.0000
54 MIBK	4406	1842	61603	163289	329191	585241	-1	0.0022	0.3172	0.0000	0.9942
49 1,3-di-cl-propane 76 78	5042	34635	178128	334360	648825	1196619	-1	0.0000	0.7083	0.0000	0.0687
59 tetra-Cl-ethene 166 168	5645	43932	206527	398332	812967	1516368	-1	0.0000	0.8560	0.0000	0.0716
Compound											
Name	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	Level 7	Coeff	Coeff	Coeff	R²/RSD
	Response	Response	Response	Response	Response	Response	Response	X¹0	X¹1/ave RF	X¹2	
60 chlorobenzene	112 77	11129	80876	394322	752942	1478617	2729365	-1	0.0000	-----	-----
61 1112-tetra-Cl-Et 131 133	4438	35454	177813	344569	695345	1340863	-1	0.0000	0.7212	0.0000	0.0707
64 ethylbenzene 91 106	21645	144618	668986	1270086	2566878	4842788	-1	0.0000	2.8368	0.0000	0.0896
65 m/p-Xylenes x2	30844	222047	1015495	1972002	3905278	7205695	-1	0.0000	2.1361	0.0000	0.0886
99 1-4-di-Cl-butane	6740	36572	153006	301374	572112	1066373	-1	0.0798	0.5616	0.0000	0.9979
52 bromoform	173 175	2520	15462	69675	136091	278788	522738	-1	0.0000	0.3083	0.0000
66 styrene 104 78	15564	84384	382712	759242	1511886	2814241	-1	0.1396	1.4904	0.0000	0.9987
67 o-xylene 91 106	15170	107821	489526	954644	1857499	3377446	-1	0.0000	2.0583	0.0000	0.1015
68 1122-Tetra-Cl-Et 83 85	2369	17006	81453	170717	330326	601304	-1	0.0000	0.3469	0.0000	0.0735
110 t-1,4-dichloro-2-butene	598	5155	17689	36488	58046	107382	-1	0.0000	0.0781	0.0000	0.2649
106 Cl-benzyl	1709	34984	130453	246745	493146	894786	-1	0.0686	0.4730	0.0000	0.9980
62 1,4-DCB-d4 150 152 13	220209	211064	218269	198057	195846	179706	-1	0.0000	1.0000	0.0000	0.0000
69 123-tri-Cl-Pr 110 97	247	4417	25007	46613	92169	179941	-1	0.0000	0.1159	0.0000	0.0641
70 4-Br-1-F-Bz (S3) 174 95	7551	36349	170967	334015	639559	1173651	-1	0.0000	0.8241	0.0000	0.0360
71 isopropylbenzene 105 120	20338	154289	681700	1327629	2705126	5006864	-1	0.0000	3.3574	0.0000	0.0661
72 bromobenzene 156 158	3634	32102	153148	304357	590583	1107164	-1	0.0000	0.7174	0.0000	0.1196
73 n-propylbenzene 120 78	5459	46356	210844	394872	801684	1463801	-1	0.0000	0.9881	0.0000	0.0916
74 2-Cl-Tl 126 128	3366	25894	120864	249899	481903	908178	-1	0.0000	0.5924	0.0000	0.0839
75 4-Cl-Tl 126 128	4268	41895	196931	372201	722805	1334882	-1	0.0000	0.8886	0.0000	0.1380
76 135-tri-Me-Bz 105 120	17066	117538	539476	1018963	2059638	3828683	-1	0.0000	2.6173	0.0000	0.0399
79 tert-butylbenzene 119 91	16192	139975	610164	1182229	2412602	4386919	-1	0.0000	2.9464	0.0000	0.1001
78 124-tri-Me-Bz 105 120	13993	99021	441266	843153	1722397	3314335	-1	0.0000	2.1864	0.0000	0.0560
80 13-di-Cl-Bz 146 148	8334	52974	229141	454607	873397	1637769	-1	0.0000	1.1613	0.0000	0.0711

82 14-di-Cl-Bz	146 148	9119	77324	360564	688087	1350534	2427479	-1	0.0000	1.6689	0.0000	0.0921
81 sec-butylbenzene	105 134	27562	188712	808141	1585357	3307792	5902646	-1	0.0000	4.1126	0.0000	0.0619
77 4-iso-Pr-toluene	119 134	19882	146089	661221	1200767	2437020	4387114	-1	0.0000	3.1156	0.0000	0.0554
84 12-di-Cl-benzene	146 148	6500	51953	234009	454651	862171	1667156	-1	0.0000	1.1158	0.0000	0.0757
85 n-butylbenzene	91 134	17225	125422	538366	1017966	2160614	4022017	-1	0.0000	2.6951	0.0000	0.0677
86 12-diBr-3-Cl-Pra	157 155	-1	1829	14905	28995	56245	114486	-1	-0.0147	0.0803	0.0000	0.9979
87 124-tri-Cl-Bz	180 182	2922	30767	135221	262726	578192	1051545	-1	-0.0553	0.7381	0.0000	0.9992
88 naphthalene	128 129	1954	26649	107459	235161	598470	1033706	-1	-0.0995	0.7367	0.0000	0.9955
90 123-tri-Cl-Bz	180 182	1279	22090	101362	206041	447433	779205	-1	-0.0222	0.5495	0.0000	0.9986
89 hx-Cl-butadiene	225 260	4633	32362	134569	257067	538819	952000	-1	0.0000	0.6806	0.0000	0.0759

Response Factor Report GCMS-G

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Initial Calibration

Calibration Files

0.3 =4-003.D 2 =4-002.D 10 =4-010.D
 20 =4-020.D 40 =4-040.D 80 =4-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD	
1) I 1 Fluorobenzene I1	0.171	0.222	0.204	0.210	0.231	0.234	0.212	10.90	
2) 3 di-Cl-di-F-metha	0.070	0.124	0.124	0.165	0.175	0.194	0.142	31.72#	0.996
3) P 4 Chloromethane	0.172	0.280	0.268	0.259	0.268	0.282	0.255	16.33	0.999
4) 9 F114 85 135	0.137	0.213	0.196	0.201	0.203	0.212	0.194	14.66	
5) C 5 vinyl chloride	0.179	0.169	0.157	0.179	0.170	0.181	0.172	5.36	
6) 6 bromomethane	0.130	0.163	0.150	0.158	0.156	0.158	0.153	7.81	
7) 7 Chloroethane	0.366	0.403	0.359	0.398	0.414	0.434	0.396	7.24	
8) 8 tri-Cl-F-methane	0.004	0.004	0.005	0.005	0.004	0.004	0.004#	11.13	
9) 111 isopropyl alcoh	0.217	0.166	0.139	0.146	0.139	0.143	0.158	19.19	1.000
10) 100 ethyl ether x5	0.010	0.015	0.011	0.011	0.010	0.012	0.011#	19.61	0.986
11) 102 Acrolein x10	0.218	0.114	0.220	0.172	0.163	0.163	0.177	24.90	0.986
12) 119 methyl acetate	0.628	0.679	0.571	0.619	0.614	0.625	0.623	5.60	
13) 104 Carbon disulfid	0.024	0.015	0.027	0.025	0.026	0.026	0.023#	20.79	0.996
14) 103 Acrylonitrilex1	0.026	0.018	0.018	0.018	0.017	0.017	0.019#	20.91	1.000
15) 95 Acetone x10	0.327	0.354	0.314	0.345	0.359	0.367	0.344	5.90	
16) 108 F-113	0.351	0.377	0.345	0.359	0.364	0.375	0.362	3.55	0.996
17) M,C 13 11-dichloroethen	0.012	0.006	0.004	0.007	0.006	0.007	0.007#	36.67	0.996
18) 101 Acetonitrilex1	0.283	0.470	0.408	0.421	0.394	0.417	0.399	15.58	
19) 109 Iodomethane	0.012	0.009	0.009	0.009	0.009	0.007	0.009#	17.87	
20) 113 Tert butyl alco	0.303	0.224	0.238	0.228	0.234	0.245	0.245	13.30	
21) 18 methylene chlori	0.550	0.482	0.402	0.412	0.376	0.358	0.430	16.86	0.998
22) 112 Allyl chloride	0.026	0.042	0.039	0.044	0.041	0.043	0.039#	17.19	
23) 200 Nitro methane x	0.471	0.570	0.445	0.484	0.491	0.512	0.496	8.64	
24) 10 t-Bu-Me-ether	0.215	0.278	0.262	0.283	0.262	0.281	0.263	9.69	
25) 19 t-12-di-Cl-ethen	0.030	0.209	0.071	0.180	0.195	0.210	0.149	52.56	0.993
26) 98 Vinyl acetate x5	0.455	0.556	0.500	0.558	0.539	0.544	0.525	7.65	
27) P 21 11-dichloroethan	0.105	0.096	0.078	0.096	0.092	0.096	0.094	9.52	
28) 91 2-butanone MEKx1									

(#) = Out of Range
 E524G004.M Fri May 16 10:37:12 2003

Response Factor Report GCMS-G

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =4-003.D 2 =4-002.D 10 =4-010.D
 20 =4-020.D 40 =4-040.D 80 =4-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD	R ²
29) 115 Di isoprop ethe	1.669	1.454	1.309	1.380	1.335	1.380	1.421	9.23	
30) 22 c-12-di-Cl-ethen	0.275	0.280	0.270	0.289	0.279	0.294	0.281	3.24	
31) 23 22-Dichloropropa	0.465	0.447	0.394	0.421	0.415	0.426	0.428	5.85	
32) 24 Br-Cl-methane	0.025	0.123	0.114	0.121	0.119	0.123	0.104	37.24	1.000
33) 25 chloroform	0.619	0.510	0.463	0.496	0.484	0.502	0.512	10.68	
34) 201 Ethyl acetate x	1.371	0.901	0.781	0.834	0.797	0.813	0.916	24.75	0.996
35) 116 ETBE	0.039	0.016	0.024	0.023	0.023	0.026	0.022#	52.71	1.000
36) 117 Iso-butyl alcoh	0.004	0.010	0.011	0.011	0.011	0.011	0.011#	3.25	0.995
37) 26 tetrahydrofuranx	0.010	0.011	0.011	0.011	0.011	0.011	0.011#	3.25	
38) S 27 Di-Br-F-Methane	0.380	0.351	0.378	0.381	0.368	0.376	0.371	3.31	
39) 34 111-tri-Cl-ethan	0.366	0.411	0.378	0.402	0.405	0.435	0.399	6.16	
40) 30 12-dichloroethan	0.074	0.208	0.187	0.208	0.207	0.220	0.184	29.91	0.999
41) 35 11-Di-Cl-propene	0.355	0.363	0.334	0.357	0.353	0.364	0.354	3.01	
42) S 29 1,2-di-Cl-ethane	0.165	0.163	0.175	0.175	0.165	0.175	0.169	3.41	
43) M 36 benzene	0.826	0.894	0.811	0.872	0.843	0.862	0.851	3.59	
44) 37 CC14	0.347	0.365	0.332	0.364	0.371	0.376	0.359	4.61	
45) 97 thiophene	0.386	0.470	0.403	0.435	0.422	0.430	0.424	6.80	
46) 118 TAME	0.099	0.714	0.540	0.599	0.561	0.580	0.515	41.29	0.999
47) C 39 12-di-Cl-propane	0.218	0.262	0.251	0.262	0.254	0.262	0.251	6.77	
48) M 40 trichloroethene	0.308	0.320	0.283	0.304	0.303	0.307	0.304	3.93	
49) 96 Me-methacrylate	0.059	0.096	0.058	0.108	0.097	0.107	0.087	26.49	0.994
50) 42 Br-di-Cl-methane	0.518	0.343	0.313	0.341	0.323	0.334	0.362	21.31	1.000
51) 41 dibromomethane	0.025	0.121	0.113	0.131	0.128	0.134	0.109	38.23	0.999
52) 45 c-13-di-Cl-prope	0.311	0.306	0.290	0.320	0.310	0.319	0.309	3.48	
53) S 55 toluene-d8 (S2)	0.521	0.473	0.514	0.498	0.513	0.504	0.504	3.76	
54) 92 2-ClEt-Vi-ether1	0.022	0.026	0.025	0.028	0.027	0.028	0.026#	9.04	
55) M C 56 toluene	0.984	0.797	0.743	0.790	0.785	0.806	0.817	10.33	
56) 107 Et methacrylate	0.142	0.150	0.065	0.144	0.145	0.149	0.132	24.94	0.995
57) 93 2-Hexanone x5	0.012	0.054	0.035	0.058	0.056	0.057	0.045#	40.36	0.998

(#) = Out of Range
 E524G004.M Fri May 16 10:37:17 2003

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =4-003.D 2 =4-002.D 10 =4-010.D
 20 =4-020.D 40 =4-040.D 80 =4-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD	r ²
58) 48 1,12-tri-Cl-Et	0.037	0.121	0.133	0.123	0.120	0.126	0.110	32.76	0.999
59) 58 1,2-di-br-ethane	0.075	0.130	0.120	0.133	0.129	0.136	0.121	19.23	0.999
60) 51 di-Br-Cl-methane	0.300	0.209	0.189	0.201	0.198	0.206	0.217	19.04	1.000
61) 46 t-13-di-Cl-prope	0.311	0.306	0.290	0.320	0.310	0.319	0.309	3.48	
62) 105 1-Chlorohexane	0.488	0.346	0.252	0.263	0.270	0.270	0.315	28.95	1.000
63) I 47 Cl-benzene-d5, I2	0.633	0.040	0.247	0.366	0.338	0.311	0.323	59.49	0.993
64) 54 MIBK	0.725	0.760	0.715	0.750	0.665	0.635	0.708	6.87	
65) 49 1,3-di-Cl-propan	0.811	0.964	0.829	0.894	0.834	0.805	0.856	7.16	
66) 59 tetra-Cl-ethene	1.600	1.774	1.582	1.689	1.517	1.449	1.602	7.29	
67) M P 60 chlorobenzene	0.638	0.778	0.713	0.773	0.713	0.712	0.721	7.07	
68) 61 1,12-tetra-Cl-Et	3.112	3.172	2.684	2.849	2.633	2.571	2.837	8.96	
69) C 64 ethylbenzene	2.217	2.435	2.037	2.212	2.003	1.913	2.136	8.86	
70) 65 m/p-Xylenes x2	0.969	0.802	0.614	0.676	0.587	0.566	0.702	22.19	0.997
71) 99 1-4-di-Cl-butane	0.362	0.339	0.280	0.305	0.286	0.278	0.308	11.37	
72) P 52 bromoform	2.237	1.851	1.535	1.703	1.551	1.494	1.729	16.33	0.998
73) 66 styrene	2.181	2.365	1.964	2.141	1.905	1.793	2.058	10.15	
74) 67 o-xylene	0.341	0.373	0.327	0.383	0.339	0.319	0.347	7.35	
75) P 68 1,122-Tetra-Cl-Et	0.086	0.113	0.071	0.082	0.060	0.057	0.078	26.49	
76) 110 t-1,4-dichloro-	0.246	0.767	0.523	0.553	0.506	0.475	0.512	32.62	0.997
77) 106 Cl-benzyl	0.105	0.115	0.118	0.118	0.118	0.125	0.116	6.41	
78) I 62 1,4-DCB-d4 150 152	0.861	0.783	0.843	0.816	0.816	0.816	0.824	3.60	
79) 69 1,23-tri-Cl-Pr	3.079	3.655	3.123	3.352	3.453	3.483	3.357	6.61	
80) S 70 4-Br-1-F-Bz (S3)	0.550	0.760	0.702	0.768	0.754	0.770	0.717	11.96	
81) 71 isopropylbenzene	0.826	1.098	0.966	0.997	1.023	1.018	0.988	9.16	
82) 72 bromobenzene	0.510	0.613	0.554	0.631	0.615	0.632	0.592	8.39	
83) 73 n-propylbenzene									
84) 74 2-Cl-Tl 126									

(#) = Out of Range
 E524G004.M Fri May 16 10:37:22 2003

Response Factor Report GCMS-G

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =4-003.D 2 =4-002.D 10 =4-010.D
 20 =4-020.D 40 =4-040.D 80 =4-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD
85) 75 4-Cl-Tl 126	0.646	0.992	0.902	0.940	0.923	0.929	0.889	13.80
86) 76 135-tri-Me-Bz	2.583	2.784	2.472	2.572	2.629	2.663	2.617	3.99
87) 79 tert-butylbenzen	2.451	3.316	2.795	2.985	3.080	3.051	2.946	10.01
88) 78 124-tri-Me-Bz	2.118	2.346	2.022	2.129	2.199	2.305	2.186	5.60
89) 80 13-di-Cl-Bz 146	1.262	1.255	1.050	1.148	1.115	1.139	1.161	7.11
90) 82 14-di-Cl-Bz 146	1.380	1.832	1.652	1.737	1.724	1.689	1.669	9.21
91) 81 sec-butylbenzene	4.172	4.470	3.703	4.002	4.222	4.106	4.113	6.19
92) 77 4-iso-Pr-toluene	3.010	3.461	3.029	3.031	3.111	3.052	3.116	5.54
93) 84 12-di-Cl-benzene	0.984	1.231	1.072	1.148	1.101	1.160	1.116	7.57
94) 85 n-butylbenzene	2.607	2.971	2.467	2.570	2.758	2.798	2.695	6.77
95) 86 12-diBr-3-Cl-Pra	0.043	0.068	0.073	0.072	0.080	0.067	20.80	0.448
96) 87 124-tri-Cl-Bz	0.442	0.729	0.620	0.663	0.738	0.731	0.654	17.40
97) 88 naphthalene	0.296	0.631	0.492	0.594	0.764	0.719	0.583	29.17
98) 90 123-tri-Cl-Bz	0.194	0.523	0.464	0.520	0.571	0.542	0.469	29.72
99) 89 hx-Cl-butadiene	0.701	0.767	0.617	0.649	0.688	0.662	0.681	7.59
								0.444
								0.444
								0.444

(#) = Out of Range
 E524G004.M Fri May 16 10:37:26 2003

Continuing Calibration Concentration Summary

Data File G2404Q01.D

Method File E524G004

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
1 Fluorobenzene I1 1	10	10.00	ppb	0.00	649888
3 di-Cl-di-F-methane 85 87	20	20.14	ppb	0.70	277528
4 Chloromethane 50 52	20	21.53	ppb	7.65	246888
9 F114 85 135	20	17.01	ppb	14.97	298895
5 vinyl chloride 62 64	20	21.36	ppb	6.82	269070
6 bromomethane 94 96	20	24.95	ppb	24.76	279586
7 Chloroethane 64 66	20	21.23	ppb	6.16	210692
8 tri-Cl-F-methane 101 103	20	21.76	ppb	8.80	559460
111 isopropyl alcohol x10	200	203.55	ppb	1.78	52983
100 ethyl ether x5	100	107.60	ppb	7.60	996988
102 Acrolein x10	200	139.76	ppb	30.12	100897
119 methyl acetate	20	21.45	ppb	7.23	241873
104 Carbon disulfide	20	18.24	ppb	8.80	738097
103 Acrylonitrile x10	200	214.62	ppb	7.31	344451
95 Acetone x10	200	396.54	ppb	98.27	439971
108 F-113	20	21.41	ppb	7.07	479223
13 11-dichloroethene 61 96	20	20.09	ppb	0.46	472350
101 Acetonitrile x10	200	243.89	ppb	21.95	101324
100 Iodomethane	20	13.22	ppb	33.89	351403
Tert butyl alcohol x10	200	216.00	ppb	8.00	129323
18 methylene chloride 49 84	20	23.61	ppb	18.05	376106
112 Allyl chloride	20	22.42	ppb	12.12	549774
200 Nitro methane x10	200	197.96	ppb	1.02	503531
10 t-Bu-Me-ether 73 57	20	21.00	ppb	4.99	676223
19 t-12-di-Cl-ethene 96 61	20	21.43	ppb	7.15	366814
98 Vinyl acetate x5	100	143.85	ppb	43.85	1821937
21 11-dichloroethane 63 83	20	21.03	ppb	5.16	717926
91 2-butanone MEK x10	200	234.40	ppb	17.20	1429484
115 Di isoprop ether	20	18.37	ppb	8.14	1696951
22 c-12-di-Cl-ethene 96 61	20	20.62	ppb	3.09	376787
23 22-Dichloropropane 77 97	20	21.99	ppb	9.97	611798
24 Br-Cl-methane 128 130	20	21.08	ppb	5.42	165470
25 chloroform 83 85	20	20.12	ppb	0.61	670049
201 Ethyl acetate x2	40	46.95	ppb	17.39	361793
116 ETBE	20	22.19	ppb	10.95	1171113
117 Iso-butyl alcohol X10	200	238.82	ppb	19.41	378345
26 tetrahydrofuran x5	100	105.40	ppb	5.40	73174
27 Di-Br-F-Methane (S1) 111 1	20	21.39	ppb	6.97	516235
34 111-tri-Cl-ethane 97 99	20	21.29	ppb	6.44	552365
30 12-dichloroethane 64 62	20	21.09	ppb	5.45	288522
35 11-Di-Cl-propene 75 110	20	20.00	ppb	0.02	460611
29 1,2-di-Cl-ethane-d4 [Surr] 10	20	22.15	ppb	10.74	242901
36 benzene 78 52	20	20.55	ppb	2.74	1136626
117 117	20	21.01	ppb	5.06	490276

97 thiophene	20	20.54	ppb	2.72	566534
TAME	20	21.35	ppb	6.73	798122
39 12-di-Cl-propane 63 76	20	21.07	ppb	5.34	344200
40 trichloroethene 130 132	20	20.87	ppb	4.36	412821
96 Me-methacrylate	20	21.65	ppb	8.25	140547
42 Br-di-Cl-methane 83 85	20	21.90	ppb	9.50	470944
41 dibromomethane 174 172	20	20.55	ppb	2.73	172546
45 c-13-di-Cl-propene 75 110	20	21.45	ppb	7.23	431214
55 toluene-d8(S2) 100 99	20	20.92	ppb	4.62	685077
92 2-ClEt-Vi-ether10	200	223.46	ppb	11.73	376438

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
56 toluene 91 92	20	19.45	ppb	2.77	1033014
107 Et methacrylate	20	27.67	ppb	38.34	255854
93 2-Hexanone x5	100	146.36	ppb	46.36	531705
48 112-tri-Cl-Et 97 83	20	20.22	ppb	1.10	163013
58 1,2-di-br-ethane 107 109	20	20.86	ppb	4.30	179065
51 di-Br-Cl-methane 129 127	20	21.16	ppb	5.82	277463
46 t-13-di-cl-propene 75 110	20	21.45	ppb	7.23	431214
105 1-Chlorohexane	20	19.65	ppb	1.75	343562
47 Cl-benzene-d5, I2	10	10.00	ppb	0.00	204425
54 MIBK	20	23.65	ppb	18.24	153762
49 1,3-di-cl-propane 76 78	20	20.32	ppb	1.58	294186
59 tetra-Cl-ethene 166 168	20	19.77	ppb	1.13	346025
60 chlorobenzene 112 77	20	20.73	ppb	3.67	678907
112-tetra-Cl-Et 131 133	20	22.02	ppb	10.09	324606
ethylbenzene 91 106	20	19.75	ppb	1.24	1145468

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
65 m/p-Xylenes x2	40	39.48	ppb	1.30	1723939
99 1-4-di-Cl-butane	20	22.55	ppb	12.74	275180
52 bromoform 173 175	20	21.37	ppb	6.85	134679
66 styrene 104 78	20	20.97	ppb	4.85	667454
67 o-xylene 91 106	20	20.91	ppb	4.53	879630
68 1122-Tetra-Cl-Et 83 85	20	21.50	ppb	7.52	152489
110 t-1,4-dichloro-2-butene	20	24.42	ppb	22.08	38966
106 Cl-benzyl	20	25.41	ppb	27.04	259687
62 1,4-DCB-d4 150 152 I3	10	10.00	ppb	0.00	177200
69 123-tri-Cl-Pr 110 97	20	22.77	ppb	13.86	46786
70 4-Br-1-F-Bz (S3) 174 95	20	20.47	ppb	2.34	298875
71 isopropylbenzene 105 120	20	20.08	ppb	0.39	1194531
72 bromobenzene 156 158	20	21.33	ppb	6.63	271114
73 n-propylbenzene 120 78	20	19.94	ppb	0.29	349176
74 2-Cl-TI 126 128	20	21.49	ppb	7.47	225624
75 4-Cl-TI 126 128	20	21.36	ppb	6.82	336390
76 135-tri-Me-Bz 105 120	20	20.52	ppb	2.58	951479
79 tert-butylbenzene 119 91	20	20.34	ppb	1.70	1061912
78 124-tri-Me-Bz 105 120	20	20.47	ppb	2.37	793217
80 13-di-Cl-Bz 146 148	20	21.95	ppb	9.75	451713
82 14-di-Cl-Bz 146 148	20	20.48	ppb	2.39	605588
sec-butylbenzene 105 134	20	18.63	ppb	6.87	1357391

77 4-iso-Pr-toluene	119 134	20	20.27	ppb	1.36	1119193
2-di-Cl-benzene	146 148	20	21.15	ppb	5.73	418081
85 n-butylbenzene	91 134	20	19.11	ppb	4.45	912663
86 12-diBr-3-Cl-Pra	157 155	20	20.75	ppb	3.74	26931
87 124-tri-Cl-Bz	180 182	20	19.47	ppb	2.66	244810
88 naphthalene	128 129	20	18.09	ppb	9.54	218555
90 123-tri-Cl-Bz	180 182	20	19.04	ppb	4.81	181441

Ave.% Dev 9.11

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2404\G2404Q01.D
 Acq On : 16 May 03 11:08 am
 Sample : f=1 ccv/icv/bfb
 Misc :

Vial: 18
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	1.000	1.000	0.0	87	0.00
2	0.212	0.214	-0.7	88	0.00
3 P	0.142	0.190	-33.8#	100	0.00
4	0.255	0.230	9.7	77	0.00
5 C	0.194	0.207	-6.8	89	0.00
6	0.172	0.215	-24.8#	105	0.00
7	0.153	0.162	-6.2	89	0.00
8	0.396	0.430	-8.8	94	0.00
9	0.004	0.004#	-1.8	75	0.00
10	0.158	0.153	3.0	91	0.00
11	0.011	0.008#	32.1#	61	0.00
12	0.177	0.186	-4.9	74	0.00
13	0.623	0.568	8.8	80	0.00
14	0.023	0.027#	-13.0	85	0.00
15	0.019	0.034#	-76.7#	165#	0.00
16	0.344	0.369	-7.1	93	0.00
17 M,C	0.362	0.363	-0.5	88	0.00
18	0.007	0.008#	-12.6	99	0.00
19	0.399	0.270	32.2#	56	0.00
20	0.009	0.010#	-8.0	94	0.00
21	0.245	0.289	-18.1	106	0.00
22	0.430	0.423	1.6	89	0.00
23	0.039	0.039#	1.0	77	0.00
24	0.496	0.520	-5.0	94	0.00
25	0.263	0.282	-7.2	87	0.00
26	0.149	0.280	-88.1#	136	0.00
27 P	0.525	0.552	-5.2	86	0.00

(#) = Out of Range
 G2404Q01 D E524G004.M Fri May 16 12:11:14 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2404\G2404Q01.D
Acq On : 16 May 03 11:08 am
Sample : f=1 ccv/icv/bfb
Misc :

Vial: 18
Operator: Eddie
Inst : GCMS-G
Multiplier: 1.00

6540

Method : C:\HPCHEM\1\METHODS\E524G004.M
Title : **Applied P & Sch Lab** EPA 524.2
Last Update : Fri May 16 10:36:26 2003
Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound	AVGRF	CCRF	%Dev	Area%	Dev(min)
28 91 2-butanone MEKx10	0.094	0.110	-17.2	100	0.00
29 115 Di isoprop ether	1.421	1.306	8.1	82	0.00
30 22 c-12-di-Cl-ethene	0.281	0.290	-3.1	87	0.00
31 23 22-Dichloropropene	0.428	0.471	-10.0	97	0.00
32 24 Br-Cl-methane	0.104	0.127	-22.0#	92	0.00
33 C 25 chloroform	0.512	0.516	-0.6	90	0.00
34 201 Ethyl acetate x2	0.118	0.139	-17.9	102	0.00
35 116 ETBE	0.916	0.901	1.7	94	0.00
36 117 Iso-butyl alcohol X10	0.022	0.029#	-31.8#	106	0.00
37 26 tetrahydrofuranx5	0.011	0.011#	-5.4	87	0.00
38 S 27 Di-Br-F-Methane (S1)	0.371	0.397	-7.0	91	0.00
39 34 111-tri-Cl-ethane	0.399	0.425	-6.4	92	0.00
40 30 12-dichloroethane	0.184	0.222	-20.8#	93	0.00
41 35 11-Di-Cl-propene	0.354	0.354	-0.0	86	0.00
42 S 29 1,2-di-Cl-ethane-d4 [Sur	0.169	0.187	-10.7	93	0.00
43 M 36 benzene	0.851	0.874	-2.7	87	0.00
44 37 CCl4	0.359	0.377	-5.1	90	0.00
45 97 thiophene	0.424	0.436	-2.7	87	0.00
46 118 TAME	0.515	0.614	-19.1	89	0.00
47 C 39 12-di-Cl-propane	0.251	0.265	-5.3	88	0.00
48 M 40 trichloroethene	0.304	0.318	-4.4	91	0.00
49 96 Me-methacrylate	0.087	0.108	-23.7#	87	0.00
50 42 Br-di-Cl-methane	0.362	0.362	-0.1	93	0.00
51 41 dibromomethane	0.109	0.133	-22.2#	88	0.00
52 45 c-13-di-Cl-propene	0.309	0.332	-7.2	90	0.00
53 S 55 toluene-d8 (S2)	0.504	0.527	-4.6	89	0.00
54 92 2-ClEt-Vi-ether10	0.026	0.029#	-11.7	91	0.00

(#) = Out of Range
G2404Q01 D E524G004.M

Fri May 16 12:11:19 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2404\G2404Q01.D
 Acq On : 16 May 03 11:08 am
 Sample : f=1 ccv/icv/bfb
 Misc :

Vial: 18
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 10:36:26 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound	AvgrRF	CCRF	%Dev	Area%	Dev(min)
55 M C 56 toluene	91	9	0.817	0.795	2.8
56 107 Et methacrylate			0.132	0.197	-48.6#
57 93 2-Hexanone x5			0.045	0.082	-80.2#
58 48 112-tri-Cl-Et	97	8	0.110	0.125	-13.9
59 58 1,2-di-br-ethane	107	109	0.121	0.138	-14.3
60 51 di-Br-Cl-methane	129	12	0.217	0.213	1.7
61 46 t-13-di-Cl-propene	75	11	0.309	0.332	-7.2
62 105 1-Chlorohexane			0.315	0.264	16.0
63 I 47 Cl-benzene-d5, I2			1.000	1.000	0.0
64 54 MIBK			0.323	0.376	-16.6
65 49 1,3-di-Cl-propane	76	78	0.708	0.720	-1.6
66 59 tetra-Cl-ethene	166	16	0.856	0.846	1.1
67 M P 60 chlorobenzene	112	7	1.602	1.661	-3.7
68 61 1112-tetra-Cl-Et	131	13	0.721	0.794	-10.1
69 C 64 ethylbenzene	91	10	2.837	2.802	1.2
70 65 m/p-Xylenes x2			2.136	2.108	1.3
71 99 1-4-di-Cl-butane			0.702	0.673	4.2
72 P 52 bromoform	173	17	0.308	0.329	-6.9
73 66 styrene	104	7	1.729	1.633	5.6
74 67 o-xylene	91	10	2.058	2.151	-4.5
75 P 68 1122-Tetra-Cl-Et	83	8	0.347	0.373	-7.5
76 110 t-1,4-dichloro-2-butene			0.078	0.095	-22.1#
77 106 Cl-benzyl			0.512	0.635	-24.1#
78 I 62 1,4-DCB-d4	150	152	1.000	1.000	0.0
79 69 123-tri-Cl-Pr	110	9	0.116	0.132	-13.9

(#) = Out of Range

G2404Q01 D E524G004.M

Fri May 16 12:11:24 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2404\G2404Q01.D
Acq On : 16 May 03 11:08 am
Sample : f=1 ccv/icv/bfb
Misc :

Vial: 18
Operator: Eddie
Inst : GCMS-G
Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
Title : **Applied P &Ch Lab** EPA 524.2
Last Update : Fri May 16 10:36:26 2003
Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

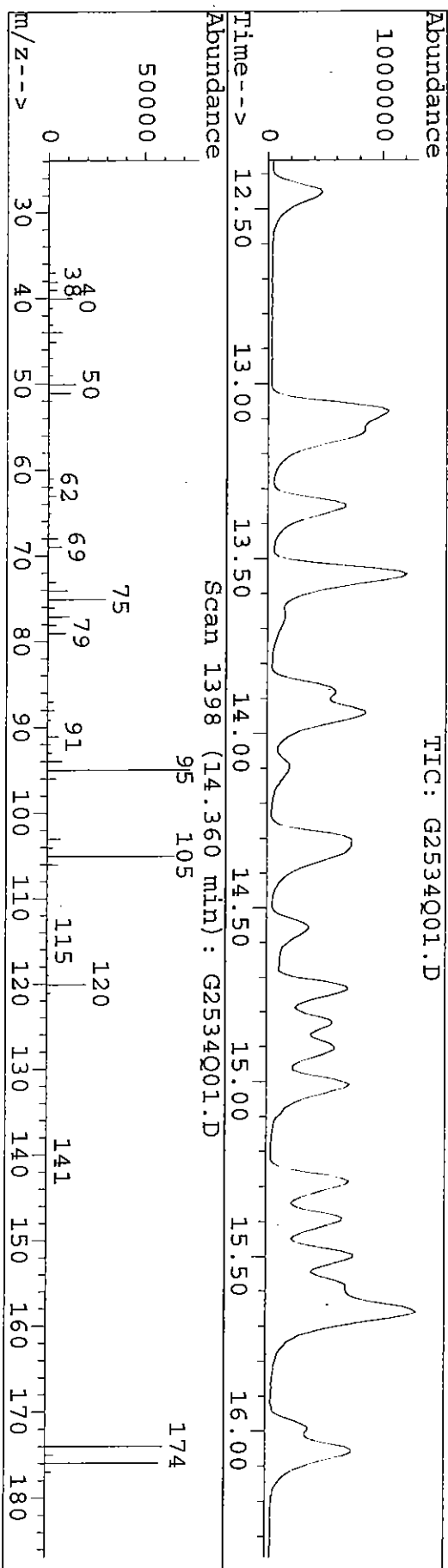
	Compound			AvgRF	CCRF	%Dev	Area%	Dev (min)		
80	S	70	4-Br-1-F-Bz (S3)	174	9	0.824	0.843	-2.3	89	0.00
81		71	isopropylbenzene	105	12	3.357	3.371	-0.4	90	0.00
82		72	bromobenzene	156	15	0.717	0.765	-6.6	89	0.00
83		73	n-propylbenzene	120	7	0.988	0.985	0.3	88	0.00
84		74	2-Cl-Tl	126	128	0.592	0.637	-7.5	90	0.00
85		75	4-Cl-Tl	126	128	0.889	0.949	-6.8	90	0.00
86		76	135-tri-Me-Bz	105	12	2.617	2.685	-2.6	93	0.00
87		79	tert-butylbenzene	119	9	2.946	2.996	-1.7	90	0.00
88		78	124-tri-Me-Bz	105	12	2.186	2.238	-2.4	94	0.00
89		80	13-di-Cl-Bz	146	148	1.161	1.275	-9.8	99	0.00
90		82	14-di-Cl-Bz	146	148	1.669	1.709	-2.4	88	0.00
91		81	sec-butylbenzene	105	13	4.113	3.830	6.9	86	0.00
92		77	4-iso-Pr-toluene	119	13	3.116	3.158	-1.4	93	0.00
93		84	12-di-Cl-benzene	146	14	1.116	1.180	-5.7	92	0.00
94		85	n-butylbenzene	91	13	2.695	2.575	4.4	90	0.00
95		86	12-diBr-3-Cl-Pra	157	15	0.067	0.076	-13.0	93	0.00
96		87	124-tri-Cl-Bz	180	18	0.654	0.691	-5.6	93	0.00
97		88	naphthalene	128	12	0.583	0.617	-5.8	93	0.00
98		90	123-tri-Cl-Bz	180	18	0.469	0.512	-9.1	88	0.00
99		89	hx-Cl-butadiene	225	26	0.681	0.669	1.7	92	0.00

(#) = Out of Range SPC's out = 0 CCC's out = 0
G2404Q01.D E524G004.M Fri May 16 12:11:27 2003

Data File : C:\HPCHEM\1\DATA\03G2534\G2534Q01.D
 Acq On : 19 May 03 4:49 pm
 Sample : f=1 ccv
 Misc :

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2



Peak Apex is scan: 1398

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
50	95	15	40	18.9	14018	PASS
75	95	30	60	40.1	29696	PASS
95	95	100	100	100.0	74024	PASS
96	95	5	9	6.6	4915	PASS
173	174	0	2	0.0	0	PASS
174	95	50	100	82.5	61048	PASS
175	174	5	9	7.3	4481	PASS
176	174	95	101	96.5	58928	PASS
177	176	5	9	5.8	3407	PASS

FORM-5A

Applied P & Ch Laboratory

Volatile Organic Instrument Performance Check for Method 524.2

Bromofluorobenzene (BFB), Part II

Client Name: GEOFON, Inc.
Case No:
Project ID: JPL GW Mon-2Q03

Contract No:
SAS No:
BFB Inj. Date: 05/19/03
BFB Inj. Time: 16:49
Instrument ID: G
Heated Purge: (Y/N) N

Lab Code: APCL
Service ID: 033130
Batch No: 03G2534
Sequence No: 03G2534
GC Column: DB-VEX
Column ID: 0.45 mm

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

#	Client Sample No	Lab Sample ID	Data File Name	Date Analyzed	Time Analyzed
1	03G2534-CCV-01	03G2534-CCV-01	G2534Q01	05/19/03	16:49
2	03G2534-LCS-01	03G2534-LCS-01	G2534L01	05/19/03	17:18
3	MW-22-1MS	03-3130-2MS	G2534M01	05/19/03	17:47
4	MW-22-1MSD	03-3130-2MSD	G2534N01	05/19/03	18:16
5	03G2534-MB-01	03G2534-MB-01	G2534K01	05/19/03	20:12
6	EB-12-5/8/03	03-3130-1	3130-01	05/19/03	21:10
7	MW-22-1	03-3130-2	3130-02	05/19/03	21:39
8	MW-22-2	03-3130-3	3130-03	05/19/03	22:08
9	MW-22-3	03-3130-4	3130-04	05/19/03	22:37
10	MW-22-4	03-3130-5	3130-05	05/19/03	23:06
11	MW-22-5	03-3130-6	3130-06	05/19/03	23:35
12	TB-12-5/8/03	03-3130-7	3130-07	05/20/03	00:04
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					

Continuing Calibration Concentration Summary

Data File G2534Q01.D
Method File E524G004

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
1 Fluorobenzene I1 1	10	10.00	ppb	0.00	696385
3 di-Cl-di-F-methane 85 87	20	20.97	ppb	4.86	309653
4 Chloromethane 50 52	20	20.56	ppb	2.79	251325
9 F114 85 135	20	18.59	ppb	7.07	351251
5 vinyl chloride 62 64	20	20.81	ppb	4.07	280900
6 bromomethane 94 96	20	18.42	ppb	7.91	221134
7 Chloroethane 64 66	20	20.46	ppb	2.32	217609
8 tri-Cl-F-methane 101 103	20	20.64	ppb	3.19	568603
111 isopropyl alcohol x10	200	217.87	ppb	8.93	60767
100 ethyl ether x5	100	99.63	ppb	0.37	989373
102 Acrolein x10	200	157.46	ppb	21.27	122234
119 methyl acetate	20	16.67	ppb	16.63	205204
104 Carbon disulfide	20	18.08	ppb	9.61	783822
103 Acrylonitrile x10	200	202.87	ppb	1.44	347141
95 Acetone x10	200	394.72	ppb	97.36	469337
108 F-113	20	20.71	ppb	3.53	496504
13 11-dichloroethene 61 96	20	19.95	ppb	0.25	502571
101 Acetonitrile x10	200	232.77	ppb	16.39	103421
109 Iodomethane	20	16.60	ppb	16.98	474077
113 Tert butyl alcohol x10	200	187.64	ppb	6.18	120381
18 methylene chloride 49 84	20	20.93	ppb	4.66	357316
112 Allyl chloride	20	21.53	ppb	7.65	566911
200 Nitro methane x10	200	212.62	ppb	6.31	579503
10 t-Bu-Me-ether 73 57	20	19.93	ppb	0.35	687757
19 t-12-di-Cl-ethene 96 61	20	19.92	ppb	0.40	365332
98 Vinyl acetate x5	100	141.46	ppb	41.46	1916784
21 11-dichloroethane 63 83	20	19.95	ppb	0.26	729629
91 2-butanone MEK x10	200	223.85	ppb	11.92	1462810
115 Di isoprop ether	20	18.47	ppb	7.67	1827519
22 o-12-di-Cl-ethene 96 61	20	19.66	ppb	1.71	384942
23 22-Dichloropropane 77 97	20	20.84	ppb	4.18	621068
24 Br-Cl-methane 128 130	20	19.92	ppb	0.39	167328
25 chloroform 83 85	20	18.82	ppb	5.88	671671
201 Ethyl acetate x2	40	47.58	ppb	18.94	393341
116 ETBE	20	20.72	ppb	3.59	1171933
117 Iso-butyl alcohol X10	200	241.27	ppb	20.63	409818
26 tetrahydrofuran x5	100	99.31	ppb	0.69	73877
27 Di-Br-F-Methane (S1) 111 1	20	20.15	ppb	0.77	521115
34 111-tri-Cl-ethane 97 99	20	20.16	ppb	0.81	560579
30 12-dichloroethane 64 62	20	20.04	ppb	0.19	293077
35 11-Di-Cl-propene 75 110	20	19.61	ppb	1.94	483926
29 1,2-di-Cl-ethane-d4 [Surr] 10	20	20.42	ppb	2.10	239972
36 benzene 78 52	20	19.59	ppb	2.05	1161177
37 CCl4 117 119	20	20.01	ppb	0.03	500192

97 thiophene	20	19.48	ppb	2.62	575552
118 TAME	20	19.58	ppb	2.11	784093
39 12-di-Cl-propane 63 76	20	20.73	ppb	3.67	362992
40 trichloroethene 130 132	20	20.21	ppb	1.03	428249
96 Me-methacrylate	20	18.71	ppb	6.43	128592
42 Br-di-Cl-methane 83 85	20	20.28	ppb	1.41	467155
41 dibromomethane 174 172	20	18.96	ppb	5.21	170037
45 c-13-di-Cl-propene 75 110	20	20.52	ppb	2.58	442013
55 toluene-d8(S2) 100 99	20	19.76	ppb	1.21	693153
92 2-ClEt-Vi-ether10	200	205.01	ppb	2.51	370077

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
56 toluene 91 92	20	18.50	ppb	7.48	1053296
107 Et methacrylate	20	25.19	ppb	25.96	248112
93 2-Hexanone x5	100	133.09	ppb	33.09	516042
48 112-tri-Cl-Et 97 83	20	19.07	ppb	4.65	164651
58 1,2-di-br-ethane 107 109	20	19.13	ppb	4.33	175497
51 di-Br-Cl-methane 129 127	20	19.42	ppb	2.92	272277
46 t-13-di-cl-propene 75 110	20	20.52	ppb	2.58	442013
105 1-Chlorohexane	20	19.66	ppb	1.68	368383
47 Cl-benzene-d5, 12	10	10.00	ppb	0.00	217118
54 MIBK	20	21.00	ppb	4.98	145050
49 1,3-di-cl-propane 76 78	20	19.64	ppb	1.78	302095
59 tetra-Cl-ethene 166 168	20	18.79	ppb	6.05	349215
60 chlorobenzene 112 77	20	19.50	ppb	2.52	678038
61 1112-tetra-Cl-Et 131 133	20	20.50	ppb	2.51	321009
64 ethylbenzene 91 106	20	18.87	ppb	5.66	1162099

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
65 m/p-Xylenes x2	40	37.33	ppb	6.69	1731107
99 1-4-di-Cl-butane	20	21.22	ppb	6.10	276088
52 bromoform 173 175	20	18.64	ppb	6.78	124788
66 styrene 104 78	20	19.47	ppb	2.63	660492
67 o-xylene 91 106	20	19.48	ppb	2.58	870690
68 1122-Tetra-Cl-Et 83 85	20	20.88	ppb	4.40	157269
110 t-1,4-dichloro-2-butene	20	18.21	ppb	8.95	30864
106 Cl-benzyl	20	14.06	ppb	29.68	159323
62 1,4-DCB-d4 150 152 13	10	10.00	ppb	0.00	186313
69 123-tri-Cl-Pr 110 97	20	21.95	ppb	9.73	47407
70 4-Br-1-F-Bz (S3) 174 95	20	19.61	ppb	1.94	301114
71 isopropylbenzene 105 120	20	19.92	ppb	0.41	1245976
72 bromobenzene 156 158	20	19.99	ppb	0.03	267246
73 n-propylbenzene 120 78	20	19.25	ppb	3.74	354454
74 2-Cl-Tl 126 128	20	19.21	ppb	3.95	212025
75 4-Cl-Tl 126 128	20	19.99	ppb	0.03	331027
76 135-tri-Me-Bz 105 120	20	19.38	ppb	3.10	945079
79 tert-butylbenzene 119 91	20	19.22	ppb	3.90	1055116
78 124-tri-Me-Bz 105 120	20	20.06	ppb	0.28	816993
80 13-di-Cl-Bz 146 148	20	18.20	ppb	9.01	393743
82 14-di-Cl-Bz 146 148	20	21.23	ppb	6.13	659992
81 sec-butylbenzene 105 134	20	17.62	ppb	11.92	1349801

77 4-iso-Pr-toluene	119 134	20	19.78	ppb	1.08	1148361
84 12-di-Cl-benzene	146 148	20	19.97	ppb	0.13	415242
85 n-butylbenzene	91 134	20	19.19	ppb	4.06	963542
86 12-diBr-3-Cl-Pra	157 155	20	20.80	ppb	3.99	28392
87 124-tri-Cl-Bz	180 182	20	19.46	ppb	2.68	257359
88 naphthalene	128 129	20	19.54	ppb	2.29	249689
90 123-tri-Cl-Bz	180 182	20	19.66	ppb	1.68	197180

Ave.% Dev	6.35
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Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2534\G2534Q01.D
 Acq On : 19 May 03 4:49 pm
 Sample : f=1 ccv
 Misc :

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 12:11:07 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound	AVGRF	CCRF	%Dev Area	Dev(min)
1 I 1 Fluorobenzene I1 1	1.000	1.000	0.0	93
2 3 di-Cl-di-F-methane 85 8	0.212	0.222	-4.9	99
3 P 4 Chloromethane 50 5	0.142	0.180	-27.1#	102
4 9 F114 85 135	0.255	0.252	1.0	91
5 C 5 vinyl chloride 62 6	0.194	0.202	-4.1	93
6 6 bromomethane 94 9	0.172	0.159	7.9	83
7 7 Chloroethane 64 66	0.153	0.156	-2.3	92
8 8 tri-Cl-F-methane 101 10	0.396	0.408	-3.2	96
9 111 isopropyl alcohol x10	0.004	0.004#	-8.9	86
10 100 ethyl ether x5	0.158	0.142	10.2	91
11 102 Acrolein x10	0.011	0.009#	23.2#	74
12 119 methyl acetate	0.177	0.147	16.9	62
13 104 Carbon disulfide	0.623	0.563	9.6	85
14 103 Acrylonitrile x10	0.023	0.025#	-6.3	86
15 95 Acetone x10	0.019	0.034#	-75.9#	176#
16 108 F-113	0.344	0.356	-3.5	96
17 M,C 13 11-dichloroethene 61 9	0.362	0.361	0.2	94
18 101 Acetonitrile x10	0.007	0.007#	-7.3	101
19 109 Iodomethane	0.399	0.340	14.7	75
20 113 Tert butyl alcohol x10	0.009	0.009#	6.2	87
21 18 methylene chloride 49 8	0.245	0.257	-4.7	101
22 112 Allyl chloride	0.430	0.407	5.3	92
23 200 Nitro methane x10	0.039	0.042#	-6.3	89
24 10 t-Bu-Me-ether 73 57	0.496	0.494	0.4	95
25 19 t-12-di-Cl-ethene 96 6	0.263	0.262	0.4	86
26 98 Vinyl acetate x5	0.149	0.275	-84.7#	143
27 P 21 11-dichloroethane 63 8	0.525	0.524	0.3	88

(#) = Out of Range

G2534Q01.D E524G004.M Tue May 20 11:18:26 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2534\G2534Q01.D
 Acq On : 19 May 03 4:49 pm
 Sample : f=1 ccv
 Misc :

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 12:11:07 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound	AvgrRF	CCRF	%Dev	Area%	Dev(min)
28	91 2-butanone MEKx10	0.094	0.105	-11.9	102 0.00
29	115 Di isoprop ether	1.421	1.312	7.7	89 0.00
30	22 c-12-di-Cl-ethene	0.281	0.276	1.7	89 0.00
31	23 22-Dichloropropane	0.428	0.446	-4.2	99 0.00
32	24 Br-Cl-methane	0.104	0.120	-15.1	93 0.00
33 C	25 chloroform	0.512	0.482	5.9	91 0.00
34	201 Ethyl acetate x2	0.118	0.141	-19.6	111 0.00
35	116 ETBE	0.916	0.841	8.2	94 0.00
36	117 Iso-butyl alcohol X10	0.022	0.029#	-33.2#	115 0.00
37	26 tetrahydrofuranx5	0.011	0.011#	0.7	88 0.00
38 S	27 Di-Br-F-Methane (S1)	0.371	0.374	-0.8	92 0.00
39	34 111-tri-Cl-ethane	0.399	0.402	-0.8	93 0.00
40	30 12-dichloroethane	0.184	0.210	-14.5	94 0.00
41	35 11-Di-Cl-propene	0.354	0.347	1.9	91 0.00
42 S	29 1,2-di-Cl-ethane-d4 [Sur	0.169	0.172	-2.1	92 0.00
43 M	36 benzene	0.851	0.834	2.1	89 0.00
44	37 CC14	0.359	0.359	-0.0	92 0.00
45	97 thiophene	0.424	0.413	2.6	89 0.00
46	118 TAME	0.515	0.563	-9.2	88 0.00
47 C	39 12-di-Cl-propane	0.251	0.261	-3.7	93 0.00
48 M	40 trichloroethene	0.304	0.307	-1.0	94 0.00
49	96 Me-methacrylate	0.087	0.092	-5.7	80 0.00
50	42 Br-di-Cl-methane	0.362	0.335	7.3	92 0.00
51	41 dibromomethane	0.109	0.122	-12.4	87 0.00
52	45 c-13-di-Cl-propene	0.309	0.317	-2.6	93 0.00
53 S	55 toluene-d8 (S2)	0.504	0.498	1.2	90 0.00
54	92 2-ClEt-Vi-ether10	0.026	0.027#	-2.5	90 0.00

(#) = Out of Range

G2534Q01.D E524G004.M

Tue May 20 11:18:29 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2534\G2534Q01.D
 Acq On : 19 May 03 4:49 pm
 Sample : f=1 ccv
 Misc :

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 12:11:07 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

Compound		AVGRF		CCRF		%Dev Area% Dev(min)	
55 M C	56 toluene	91	9	0.817	0.756	7.5	89 0.00
56	107 Et methacrylate			0.132	0.178	-34.5#	115 0.00
57	93 2-Hexanone x5			0.045	0.074	-63.2#	120 0.00
58	48 112-tri-Cl-Et	97	8	0.110	0.118	-7.3	89 0.00
59	58 1,2-di-br-ethane	107	109	0.121	0.126	-4.5	88 0.00
60	51 di-Br-Cl-methane	129	12	0.217	0.195	10.0	91 0.00
61	46 t-13-di-cl-propene	75	11	0.309	0.317	-2.6	93 0.00
62	105 1-Chlorohexane			0.315	0.264	16.0	94 0.00
63 I	47 Cl-benzene-d5, I2			1.000	1.000	0.0	97 0.00
64	54 MIBK			0.323	0.334	-3.5	89 0.00
65	49 1,3-di-cl-propane	76	78	0.708	0.696	1.8	90 0.00
66	59 tetra-Cl-ethene	166	16	0.856	0.804	6.1	88 0.00
67 M P	60 chlorobenzene	112	7	1.602	1.561	2.5	90 0.00
68	61 1112-tetra-Cl-Et	131	13	0.721	0.739	-2.5	93 0.00
69 C	64 ethylbenzene	91	10	2.837	2.676	5.7	91 0.00
70	65 m/p-Xylenes x2			2.136	1.993	6.7	88 0.00
71	99 1-4-di-Cl-butane			0.702	0.636	9.5	92 0.00
72 P	52 bromoform	173	17	0.308	0.287	6.8	92 0.00
73	66 styrene	104	7	1.729	1.521	12.0	87 0.00
74	67 o-xylene	91	10	2.058	2.005	2.6	91 0.00
75 P	68 1122-Tetra-Cl-Et	83	8	0.347	0.362	-4.4	92 0.00
76	110 t-1,4-dichloro-2-butene			0.078	0.071	9.0	85 0.00
77	106 Cl-benzyl			0.512	0.367	28.3#	65 0.00
78 I	62 1,4-DCB-d4	150	152	1.000	1.000	0.0	94 0.00
79	69 123-tri-Cl-Pr	110	9	0.116	0.127	-9.7	102 0.00

(#) = Out of Range

G2534Q01.D E524G004.M

Tue May 20 11:18:32 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2534\G2534Q01.D
 Acq On : 19 May 03 4:49 pm
 Sample : f=1 ccv
 Misc :

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G004.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Fri May 16 12:11:07 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.050 Min. Rel. Area : 35% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AVGRF	CCRF	%Dev	Area%	Dev(min)
80	S					
81	70 4-Br-1-F-Bz (S3)	174	9	0.824	0.808	1.9 90 0.00
82	71 isopropylbenzene	105	12	3.357	3.344	0.4 94 0.00
83	72 bromobenzene	156	15	0.717	0.717	0.0 88 0.00
84	73 n-propylbenzene	120	7	0.988	0.951	3.7 90 0.00
85	74 2-Cl-Tl	126	128	0.592	0.569	4.0 85 0.00
86	75 4-Cl-Tl	126	128	0.889	0.888	0.0 89 0.00
87	76 135-tri-Me-Bz	105	12	2.617	2.536	3.1 93 0.00
88	79 tert-butylbenzene	119	9	2.946	2.832	3.9 89 0.00
89	78 124-tri-Me-Bz	105	12	2.186	2.193	-0.3 97 0.00
90	80 13-di-Cl-Bz	146	148	1.161	1.057	9.0 87 0.00
91	82 14-di-Cl-Bz	146	148	1.669	1.771	-6.1 96 0.00
92	81 sec-butylbenzene	105	13	4.113	3.622	11.9 85 0.00
93	77 4-iso-Pr-toluene	119	13	3.116	3.082	1.1 96 0.00
94	84 12-di-Cl-benzene	146	14	1.116	1.114	0.1 91 0.00
95	85 n-butylbenzene	91	13	2.695	2.586	4.1 95 0.00
96	86 12-diBr-3-Cl-Pra	157	15	0.067	0.076	-13.3 98 0.00
97	87 124-tri-Cl-Bz	180	18	0.654	0.691	-5.6 98 0.00
98	88 naphthalene	128	12	0.583	0.670	-15.0 106 0.00
99	90 123-tri-Cl-Bz	180	18	0.469	0.529	-12.8 96 0.00
	89 hx-Cl-butadiene	225	26	0.681	0.657	3.4 95 0.00

(#) = Out of Range SPC's out = 0 CCC's out = 0
 G2534Q01.D E524G004.M Tue May 20 11:18:35 2003

FORM-8A

Applied P & Ch Laboratory

Internal Standard Area and RT Summary for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 033130

Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10

Sample Matrix: Water

CCV Data File: G2534Q01

Instrument ID: G

Batch No: 03G2534

#	Client Sample No	Lab Sample ID	Analysis Date & Time	IS-1 Area # RT #	IS-2 Area # RT #	IS-3 Area # RT #
	12 Hour CCV STD		05/19/03 16:49	696385 9.51	217118 13.12	186313 15.63
	CCV Upper Limit			1392770 10.01	434236 13.62	372626 16.13
	CCV Lower Limit			348192 9.01	108559 12.62	93156 15.13
1	03G2534-LCS-01	03G2534-LCS-01	05/19/03 17:18	665707 9.51	201715 13.11	191235 15.61
2	MW-22-1MS	03-3130-2MS	05/19/03 17:47	691053 9.50	213121 13.11	194829 15.63
3	MW-22-1MSD	03-3130-2MSD	05/19/03 18:16	624964 9.50	187075 13.11	178631 15.62
4	03G2534-MB-01	03G2534-MB-01	05/19/03 20:12	648041 9.50	183940 13.11	165027 15.62
5	EB-12-5/8/03	03-3130-1	05/19/03 21:10	649527 9.50	183699 13.10	173342 15.62
6	MW-22-1	03-3130-2	05/19/03 21:39	693146 9.49	199411 13.10	188055 15.60
7	MW-22-2	03-3130-3	05/19/03 22:08	743798 9.50	203470 13.10	203953 15.60
8	MW-22-3	03-3130-4	05/19/03 22:37	681813 9.49	195978 13.10	194874 15.60
9	MW-22-4	03-3130-5	05/19/03 23:06	748362 9.49	211588 13.10	196346 15.61
10	MW-22-5	03-3130-6	05/19/03 23:35	664342 9.49	180402 13.10	178007 15.63
11	TB-12-5/8/03	03-3130-7	05/20/03 00:04	744335 9.49	210234 13.11	203725 15.61
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						

IS-1 = FLUOROBENZENE

IS-2 = CHLOROBENZENE-D5

IS-3 = 1,4-DICHLOROBENZENE-D4

Area Upper Limit = +100% of CCV internal standard area

Area Lower Limit = - 50% of CCV internal standard area

RT Upper Limit = +0.50 minutes of CCV internal standard RT

RT Lower Limit = - 0.50 minutes of CCV internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits

Applied P & Ch Laboratory

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

VOC Analysis General Logbook

Source # 07692404 Batch # 07692404 Matrix: W Date: 5-15-03 Analyst: EddieIS/Surrogate: GC-1514/1515 Methanol(mark-M): _____ PEG (mark-PEG): _____ Defoaming(mark-DF): _____Lab # 1 Ini. Batch ☒ Initial Batch; ☐ Middle Batch; ☐ Final Batch. ☐ Internal Study Datafile Path: _____

	Type	Sample ID	Method	$V/X=f_1$	$V_1/V_i=f_2$	$V_{app}/V_{inj}=f_3$	F	A-#	Datafile	Note	pH
3012	SP	62404P01	E5246	25125 = 1	/ =	/ =	/		62404P01	5-15-03 10:18 AM GC15137	845137
3013	Talib	4-003	004	/ =	/ =	/ =	/		4-003		
3014		4-002		/ =	/ =	/ =	/		4-002		
3015		4-010		/ =	/ =	/ =	/		4-010		
3016		4-020		/ =	/ =	/ =	/		4-020		
3017		4-040		/ =	/ =	/ =	/		4-040		
3018	AK	4-080		/ =	/ =	/ =	/		4-080		
3019	CW	62404R01		/ =	/ =	/ =	/		62404R01	CW/CW 5-16-03 10:08 AM GC15138	
3020	LY	L01		/ =	/ =	/ =	/		L01		
3021	M3	M01		/ =	/ =	/ =	/		M01	3082-04 C2	
3022	M40	N01		/ =	/ =	/ =	/		N01	3082-04 C2	
3023	M13	K01		/ =	/ =	/ =	/		K01		
3024	Sample	3082-01		/ =	/ =	/ =	/		3082-01		C2
3025		02		/ =	/ =	/ =	/		02		
3026		03		/ =	/ =	/ =	/		03		
3027		04		/ =	/ =	/ =	/		04	M3	
3028		05		/ =	/ =	/ =	/		05		
3029		06		/ =	/ =	/ =	/		06		
3030		07		/ =	/ =	/ =	/		07		
3031		3102-01		/ =	/ =	/ =	/		3102-01		
3032		02		/ =	/ =	/ =	/		02		
3033		03		/ =	/ =	/ =	/		03		
3034		04		/ =	/ =	/ =	/		04		
3035		05		/ =	/ =	/ =	/		05		
3036		06		/ =	/ =	/ =	/		06		
3037		07		/ =	/ =	/ =	/		07		
3038		08		/ =	/ =	/ =	/		08		
3039		3115-01		/ =	/ =	/ =	/		3115-01		
3040		02		/ =	/ =	/ =	/		02		
3041		03		/ =	/ =	/ =	/		03		
3042	✓	04	✓	✓	/ =	/ =	✓		04	push 12 min	
3043				/ =	/ =	/ =	/				

Type	Op #	STD Lot #	$C_{std}(ng/\mu L) \times V_{std}(\mu L) / X(g \text{ or } mL) = T$	Op #	STD Lot #	$C_{std}(ng/\mu L) \times V_{std}(\mu L) / X(g \text{ or } mL) = T$
CS/LCSD	3825	GC-15138	202 x 2.5 / X = ppb		GC-	x / X = ppb
MS/MSD	3826/3827	GC-15138	202 x 2.5 / X = ppb		GC-	x / X = ppb

Footnote/Anomaly:

13780 Magnolia Ave. Chino CA 91710
 Tel: (909) 590-1828 Fax: (909) 590-1498

VOC Analysis General Logbook

Sequence # 03612534 Batch # 03612534 Matrix: W Date: 5-19-03 Analyst: Edie

Lot #: IS/Surrogate: GC-1514/15115 Methanol(mark-M): _____ PEG (mark-PEG): _____ Defoaming(mark-DF): _____

☒ Calib. + Ini. Batch ☒ Initial Batch; ☐ Middle Batch; ☐ Final Batch. ☐ Internal Study Datafile Path: _____

Op. #	Type	Sample ID	Method	$V_1/X=f_1$	$V_1/V_i=f_2$	$V_{std}/V_{inj}=f_3$	F	A-#	Datafile	Note	pH
3849	SP	625401	625401	251 25 = 1	/ =	/ =			625401		
3850	CW	201	004	/ =	/ =	/ =			201	5-19-03	
3851	LS	L01		/ =	/ =	/ =			L01	4-49 pm	6.4
3852	MS	M01		/ =	/ =	/ =			M01	3/130-02	6.2
3853	M9D	N01		/ =	/ =	/ =			N01	3/130-02	6.2
3854	MB	K01		/ =	/ =	/ =			K01		
3855	Sample	3115-04A		/ =	/ =	/ =			3115-04A		6.2
3856		3130-01		/ =	/ =	/ =			3130-01	elo	
3857		02		/ =	/ =	/ =			02	6.5	
3858		03		/ =	/ =	/ =			03		
3859		04		/ =	/ =	/ =			04		
3860		05		/ =	/ =	/ =			05		
3861		06		/ =	/ =	/ =			06		
3862		07		/ =	/ =	/ =			07		
3863		3205-01		/ =	/ =	/ =			3205-01		
3864		02		/ =	/ =	/ =			02		
3865		03		/ =	/ =	/ =			03		
3866		04		/ =	/ =	/ =			04		
3867		05		/ =	/ =	/ =			05		
3868		06		/ =	/ =	/ =			06		
3869		07		/ =	/ =	/ =			07		
3870	✓	08	✓	/ =	/ =	/ =	✓		08		
3871				/ =	/ =	/ =					
3872				/ =	/ =	/ =					
3873				/ =	/ =	/ =					
3874				/ =	/ =	/ =					
3875				/ =	/ =	/ =					
3876				/ =	/ =	/ =					
3877				/ =	/ =	/ =					
3878				/ =	/ =	/ =					
3879				/ =	/ =	/ =					
3880				/ =	/ =	/ =					

Type	Op #	STD Lot #	$C_{std}(ng/\mu L) \times V_{std}(\mu L) / X(g \text{ or } mL) = T$	Op #	STD Lot #	$C_{std}(ng/\mu L) \times V_{std}(\mu L) / X(g \text{ or } mL) = T$
LCS/LCSD	385	GC-1513	1.5 /X = ppb		GC-	x /X = ppb
MS/MSD	385/3853	GC-1513	1.5 /X = 20 ppb		GC-	x /X = ppb