

Level C Data Package Deliverables

Metals



Applied P & Ch Laboratory

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03
Sample ID: **03M1458-MB-01**
Sample Type: Method Blank

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03M1458-MB-01
Sample Matrix: Water
Collection Date: 05/13/2003
Collected by:
Received Date: 05/13/2003
Moisture %: -

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	< 5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	117	B	P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	< 50	U	P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	< 100	U	P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	64.7	B	P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	1040	B	P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-1
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Sample ID: **EB-12-5/8/03**
Sample Type: Field Sample

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	< 5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	< 200	U	P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	31.4	B	P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	20.7	B	P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	93.5	B	P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	1240	B	P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-2
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Sample ID: MW-22-1
Sample Type: Field Sample

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	< 5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	108000		P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	115		P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	39100		P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	3110		P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	30700		P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03
Sample ID: **MW-22-2**
Sample Type: Field Sample

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-3
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	<5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	35000		P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	81.3		P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	16800		P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	2360		P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	30400		P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-4
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Sample ID: **MW-22-3**
Sample Type: Field Sample

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	<5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	29500		P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	130		P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	14500		P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	2360		P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	36800		P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-5
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Sample ID: **MW-22-4**
Sample Type: Field Sample

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	< 5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	29400		P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	53.8		P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	8450		P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	1700		P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	26900		P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

Applied P & Ch Laboratory
Metal Analysis Results

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130
Lab Sample ID: 03-3130-6
Sample Matrix: Water

Collection Date: 05/08/2003
Collected by:
Received Date: 05/08/2003
Moisture %: -

Sample ID: MW-22-5
Sample Type: Field Sample

Element Name	CAS No	Unit	RL	Result	C	M	Q	Batch	D-Date	A-Date	DF	Method
ARSENIC	7440-38-2	µg/L	5	< 5	U	F		03M1458E	05/13/03	05/13/03	1	200.9
CALCIUM	7440-70-2	µg/L	200	4580		P		03M1452L	05/12/03	05/12/03	1	200.7
IRON	7439-89-6	µg/L	50	57.5		P		03M1452L	05/12/03	05/12/03	1	200.7
MAGNESIUM	7439-95-4	µg/L	100	625		P		03M1452L	05/12/03	05/12/03	1	200.7
POTASSIUM	7440-09-7	µg/L	400	1130		P		03M1452L	05/12/03	05/12/03	1	200.7
SODIUM	7440-23-5	µg/L	2000	70200		P		03M1452L	05/12/03	05/12/03	1	200.7

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

Note: RL: PQL (EQL) or CRDL D-Date: Digestion Date; A-Date: Analysis Date; DF: Dilution Factor

C Qualifier: U - Not Detected or less than IDL

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

Q Qualifier: N - Spike recovery out of control

* - Duplicate analysis out of control

W - Post digestion spike for GFAA out of control

E - Serial dilution difference out of control

M Qualifier: P - ICP

A - FLAA

F - GFAA

CV - Cold Vapor

FORM-2A Metal
Applied P & Ch Laboratory
Initial and Continuing Calibration Verification

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	ICV 12:49			CCV 13:05			CCV 13:47			CCV 14:30		
		True	Result	%R	True	Result	%R	True	Result	%R	True	Result	%R
1	Aluminum	10000.0	9971.26	99.7	5000.0	4968.25	99.4	5000.0	5032.85	100.7	5000.0	5093.25	101.9
2	Antimony	4000.0	3933.95	98.3	2000.0	1911.54	95.6	2000.0	1981.39	99.1	2000.0	1989.65	99.5
3	Arsenic	1000.0	992.16	99.2	500.0	496.86	99.4	500.0	503.27	100.7	500.0	510.76	102.2
4	Barium	10000.0	9983.63	99.8	5000.0	4598.13	92.0	5000.0	4998.30	100.0	5000.0	4980.00	99.6
5	Beryllium	1000.0	975.87	97.6	500.0	463.44	92.7	500.0	488.99	97.8	500.0	487.32	97.5
6	Cadmium	2000.0	2000.64	100.0	1000.0	953.29	95.3	1000.0	995.82	99.6	1000.0	1003.45	100.3
7	Calcium	100000.0	97268.01	97.3	50000.0	48119.15	96.2	50000.0	50239.37	100.5	50000.0	50757.96	101.5
8	Chromium	1000.0	1005.01	100.5	500.0	464.10	92.8	500.0	496.19	99.2	500.0	495.53	99.1
9	Cobalt	4000.0	4023.16	100.6	2000.0	1879.05	94.0	2000.0	1986.95	99.3	2000.0	1972.56	98.6
10	Copper	4000.0	4031.21	100.8	2000.0	1870.91	93.5	2000.0	1954.75	97.7	2000.0	1961.78	98.1
11	Iron	10000.0	10026.26	100.3	5000.0	4767.28	95.3	5000.0	4969.23	99.4	5000.0	4966.22	99.3
12	Lead	1000.0	980.22	98.0	500.0	477.15	95.4	500.0	501.76	100.4	500.0	509.28	101.9
13	Magnesium	50000.0	50688.56	101.4	25000.0	23165.49	92.7	25000.0	24804.51	99.2	25000.0	24777.11	99.1
14	Manganese	4000.0	4010.00	100.3	2000.0	1854.90	92.7	2000.0	1984.66	99.2	2000.0	1959.68	98.0
15	Nickel	4000.0	4038.54	101.0	2000.0	1913.00	95.7	2000.0	1994.38	99.7	2000.0	1996.29	99.8
16	Potassium	30000.0	28900.23	96.3	15000.0	14904.84	99.4	15000.0	15526.90	103.5	15000.0	15606.02	104.0
17	Selenium	1000.0	982.48	98.2	500.0	483.08	96.6	500.0	501.21	100.2	500.0	510.78	102.2
18	Silver	2000.0	2027.10	101.4	1000.0	927.66	92.8	1000.0	980.31	98.0	1000.0	987.77	98.8
19	Sodium	200000.0	195310.22	97.7	100000.0	97679.50	97.7	100000.0	107540.84	107.5	100000.0	106394.91	106.4
20	Thallium	1000.0	983.05	98.3	500.0	475.36	95.1	500.0	507.20	101.4	500.0	509.38	101.9
21	Vanadium	4000.0	4011.74	100.3	2000.0	1879.82	94.0	2000.0	1992.04	99.6	2000.0	1989.58	99.5
22	Zinc	4000.0	4031.40	100.8	2000.0	1945.57	97.3	2000.0	1982.54	99.1	2000.0	2002.68	100.1
23	Molybdenum	4000.0	4026.90	100.7	2000.0	1885.93	94.3	2000.0	2006.03	100.3	2000.0	2024.31	101.2

(a) ICV Control Limit 95-105%; For Hg, 90-110%.

(b) CCV Control Limit 90-110%; For Hg, 80-120%.

FORM-2A Metal
Applied P & Ch Laboratory
Initial and Continuing Calibration Verification

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	CCV 15:04			CCV 15:41			CCV 16:51			CCV 17:21		
		True	Result	%R	True	Result	%R	True	Result	%R	True	Result	%R
1	Aluminum	5000.0	5020.90	100.4	5000.0	4826.65	96.5	5000.0	4812.73	96.3	5000.0	5115.32	102.3
2	Antimony	2000.0	1861.92	93.1	2000.0	1874.85	93.7	2000.0	1991.76	99.6	2000.0	2057.32	102.9
3	Arsenic	500.0	485.72	97.1	500.0	477.65	95.5	500.0	536.62	107.3	500.0	517.60	103.5
4	Barium	5000.0	4847.02	96.9	5000.0	4684.24	93.7	5000.0	4532.10	90.6	5000.0	5035.18	100.7
5	Beryllium	500.0	476.69	95.3	500.0	462.96	92.6	500.0	454.20	90.8	500.0	490.76	98.2
6	Cadmium	1000.0	937.19	93.7	1000.0	927.84	92.8	1000.0	984.86	98.5	1000.0	1006.97	100.7
7	Calcium	50000.0	48197.91	96.4	50000.0	46396.52	92.8	50000.0	50277.04	100.6	50000.0	50681.31	101.4
8	Chromium	500.0	457.94	91.6	500.0	453.70	90.7	500.0	469.85	94.0	500.0	498.37	99.7
9	Cobalt	2000.0	1925.45	96.3	2000.0	1871.10	93.6	2000.0	1903.41	95.2	2000.0	2033.06	101.7
10	Copper	2000.0	1926.89	96.3	2000.0	1870.35	93.5	2000.0	1899.53	95.0	2000.0	1995.48	99.8
11	Iron	5000.0	4853.37	97.1	5000.0	4702.96	94.1	5000.0	4682.53	93.7	5000.0	5064.92	101.3
12	Lead	500.0	482.53	96.5	500.0	468.97	93.8	500.0	511.30	102.3	500.0	512.98	102.6
13	Magnesium	25000.0	24130.69	96.5	25000.0	23313.06	93.3	25000.0	23309.07	93.2	25000.0	25211.54	100.8
14	Manganese	2000.0	1902.17	95.1	2000.0	1847.41	92.4	2000.0	1845.72	92.3	2000.0	2027.06	101.4
15	Nickel	2000.0	1965.23	98.3	2000.0	1910.48	95.5	2000.0	1949.68	97.5	2000.0	2036.48	101.8
16	Potassium	15000.0	15465.96	103.1	15000.0	14932.53	99.6	15000.0	14622.86	97.5	15000.0	15057.02	100.4
17	Selenium	500.0	489.04	97.8	500.0	494.11	98.8	500.0	544.37	108.9	500.0	511.98	102.4
18	Silver	1000.0	974.31	97.4	1000.0	943.97	94.4	1000.0	930.92	93.1	1000.0	986.90	98.7
19	Sodium	100000.0	102101.83	102.1	100000.0	98824.82	98.8	100000.0	92495.88	92.5	100000.0	102369.23	102.4
20	Thallium	500.0	476.86	95.4	500.0	482.95	96.6	500.0	491.20	98.2	500.0	510.91	102.2
21	Vanadium	2000.0	1950.97	97.5	2000.0	1891.11	94.6	2000.0	1875.19	93.8	2000.0	2025.70	101.3
22	Zinc	2000.0	1989.42	99.5	2000.0	1933.75	96.7	2000.0	2013.24	100.7	2000.0	2033.11	101.7
23	Molybdenum	2000.0	1910.53	95.5	2000.0	1877.52	93.9	2000.0	1912.50	95.6	2000.0	1998.55	99.9

(a) ICV Control Limit 95-105%; For Hg, 90-110%.

(b) CCV Control Limit 90-110%; For Hg, 80-120%.

FORM-2A Metal
Applied P & Ch Laboratory
Initial and Continuing Calibration Verification

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Lab Code: APCL
Project Name: JPL GW Mon-2Q03	Service ID: 033130	Sequence No.: 03M1458E
	Instrument: GFAA-E	Method: 200.9
Batch No.(s): 03M1458		

Analysis Date: 05/13/03

Concentration Units: UG/L

#	Analyte	ICV 11:10			CCV 12:27			CCV 13:42			CCV 14:59		
		True	Result	%R	True	Result	%R	True	Result	%R	True	Result	%R
1	Arsenic	50.0	51.40	102.8	50.0	56.60	113.2	50.0	55.60	111.2	50.0	64.60	129.2

(a) ICV Control Limit 95-105%; For Hg, 90-110%.

(b) CCV Control Limit 90-110%; For Hg, 80-120%.

FORM-2A Metal
Applied P & Ch Laboratory
Initial and Continuing Calibration Verification

Client Name:	GEOFON, Inc.	Project No:	04-4428.10	Lab Code:	APCL
Project Name:	JPL GW Mon-2Q03	Service ID:	033130	Sequence No.:	03M1458E
		Instrument:	GFAA-E	Method:	200.9
Batch No.(s):	03M1458				

Analysis Date: 05/13/03

Concentration Units: UG/L

#	Analyte	CCV 15:15			CCV 16:06								
		True	Result	%R	True	Result	%R	True	Result	%R	True	Result	%R
1	Arsenic	50.0	51.00	102.0	50.0	49.70	99.4						

(a) ICV Control Limit 95-105%; For Hg, 90-110%.

(b) CCV Control Limit 90-110%; For Hg, 80-120%.

FORM-2B Metal
Applied P & Ch Laboratory
CRDL Standard For AA and ICP

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	True	12:57 Found	R%	Time Found	R%
1	Aluminum	200.0	119.50	59.7		
2	Antimony	20.0	20.27	101.3		
3	Arsenic	20.0	18.22	91.1		
4	Barium	10.0	7.25	72.5		
5	Beryllium	4.0	3.58	89.4		
6	Cadmium	5.0	3.78	75.5		
7	Calcium	1000.0	875.42	87.5		
8	Chromium	10.0	6.89	68.9		
9	Cobalt	20.0	13.23	66.1		
10	Copper	10.0	8.36	83.6		
11	Iron	50.0	39.88	79.8		
12	Lead	10.0	7.41	74.1		
13	Magnesium		29.34			
14	Manganese	10.0	6.26	62.6		
15	Nickel	20.0	14.67	73.4		
16	Potassium		71.92			
17	Selenium	10.0	10.51	105.1		
18	Silver	10.0	8.25	82.5		
19	Sodium		1103.41			
20	Thallium	10.0	9.05	90.5		
21	Vanadium	10.0	7.09	70.9		
22	Zinc	20.0	17.89	89.4		
23	Molybdenum	15.0	11.91	79.4		

FORM-3 Metal
Applied P & Ch Laboratory
Metal ICB/CCB Summary

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	ICB Result	12:54 C	CCB Result	13:12 C	CCB Result	13:56 C	CCB Result	14:43 C	CCB Result	15:12 C
1	Aluminum	6.80	B	9.24	B	5.60	U	5.60	U	5.60	U
2	Antimony	2.70	U	2.70	U	3.60	B	2.70	U	2.85	B
3	Arsenic	1.90	U	1.90	U	1.90	U	1.90	U	-2.67	B
4	Barium	1.20	U	1.20	U	1.20	U	1.20	U	1.20	U
5	Beryllium	0.06	U	0.06	B	0.06	U	0.06	U	0.06	U
6	Cadmium	0.24	B	0.32	B	0.13	B	0.15	B	-0.18	B
7	Calcium	70.73	B	81.17	B	136.33	B	59.00	U	158.98	B
8	Chromium	0.14	U	-0.22	B	0.14	U	0.14	U	0.14	U
9	Cobalt	0.49	U	0.49	U	0.49	U	0.49	U	0.49	U
10	Copper	1.70	U	1.70	U	1.70	U	1.70	U	1.70	U
11	Iron	1.50	U	1.50	U	1.50	U	1.50	U	1.50	U
12	Lead	1.30	U	1.30	U	1.30	U	1.30	U	-2.18	B
13	Magnesium	9.70	U	14.65	B	9.70	U	9.70	U	19.93	B
14	Manganese	0.16	U	0.27	B	0.26	B	0.45	B	0.66	B
15	Nickel	-0.59	B	-0.46	B	0.46	U	0.46	U	0.46	U
16	Potassium	62.13	B	70.47	B	75.05	B	74.01	B	87.28	B
17	Selenium	2.30	U	2.30	U	2.30	U	2.30	U	2.30	U
18	Silver	-2.21	B	0.61	U	0.61	U	-1.20	B	0.84	B
19	Sodium	979.83	B	985.41	B	772.32	B	985.80	B	1087.80	B
20	Thallium	1.60	U	1.60	U	1.60	U	1.60	U	1.60	U
21	Vanadium	0.35	B	0.55	B	-0.27	B	0.13	U	0.13	U
22	Zinc	2.20	U	2.20	U	2.20	U	2.20	U	2.20	U
23	Molybdenum	0.32	U	0.45	B	0.89	B	0.34	B	0.78	B

FORM-3 Metal
Applied P & Ch Laboratory
Metal ICB/CCB Summary

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	CCB Result	15:45 C	CCB Result	16:54 C	CCB Result	17:24 C	CCB Result	Time C	CCB Result	Time C
1	Aluminum	5.60	U	5.60	U	30.40	B				
2	Antimony	2.70	U	2.70	U	2.70	U				
3	Arsenic	1.90	U	1.90	U	1.90	U				
4	Barium	1.62	B	1.20	U	1.62	B				
5	Beryllium	0.20	B	0.09	B	0.16	B				
6	Cadmium	0.19	B	0.38	B	0.30	B				
7	Calcium	62.81	B	59.00	U	227.84					
8	Chromium	0.26	B	0.14	U	0.14	U				
9	Cobalt	0.49	U	0.73	B	0.52	B				
10	Copper	1.70	U	1.70	U	1.70	U				
11	Iron	1.50	U	1.50	U	19.77	B				
12	Lead	1.30	U	1.30	U	-2.18	B				
13	Magnesium	16.17	B	9.70	U	25.97	B				
14	Manganese	1.18	B	0.54	B	1.06	B				
15	Nickel	0.46	U	0.46	U	0.46	U				
16	Potassium	72.47	B	115.69	B	110.28	B				
17	Selenium	3.81	B	2.73	B	2.30	U				
18	Silver	-1.59	B	0.91	B	0.61	U				
19	Sodium	973.29	B	1265.55	B	1314.67	B				
20	Thallium	4.33	B	3.16	B	3.47	B				
21	Vanadium	1.04	B	0.13	U	0.13	U				
22	Zinc	2.20	U	2.20	U	2.20	U				
23	Molybdenum	2.34	B	2.30	B	2.04	B				

FORM-3 Metal
Applied P & Ch Laboratory
Metal ICB/CCB Summary

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: GFAA-E

Lab Code: APCL
Sequence No.: 03M1458E
Method: 200.9

Batch No.(s): 03M1458

Analysis Date: 05/13/03

Concentration Units: UG/L

#	Analyte	ICB	11:17	CCB	12:34	CCB	13:49	CCB	15:06	CCB	15:21
		Result	C	Result	C	Result	C	Result	C	Result	C
1	Arsenic	2.10	U	2.10	U	2.10	U	2.10	U	2.10	U

FORM-3 Metal
Applied P & Ch Laboratory
Metal ICB/CCB Summary

Client Name:	GEOFON, Inc.	Project No:	04-4428.10	Lab Code:	APCL
Project Name:	JPL GW Mon-2Q03	Service ID:	033130	Sequence No.:	03M1458E
		Instrument:	GFAA-E	Method:	200.9
Batch No.(s):	03M1458				

Analysis Date: 05/13/03

Concentration Units: UG/L

#	Analyte	CCB	16:12	CCB	Time	CCB	Time	CCB	Time	CCB	Time
		Result	C	Result	C	Result	C	Result	C	Result	C
1	Arsenic	2.10	U								

FORM-4 Metal
Applied P & Ch Laboratory
ICP Interference Check Sample

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
ICP ID Number: ICP -L

Lab Code: APCL
Sequence No.: 03M1452L

Batch No.(s): 03M1452

Analysis Date: 05/12/03

Concentration Units: UG/L

#	Analyte	Expected		Initial 12:59		%R	Found 13:02		%R
		Sol. A	Sol. AB	Sol. A	Sol. AB		Sol. A	Sol. AB	
1	Aluminum	500000	500000	415714	457650.0	91.5	418470	466762.8	93.4
2	Antimony	0	1000	6	924.7	92.5	-12	967.9	96.8
3	Arsenic	0	1000	-2	930.4	93.0	1	982.9	98.3
4	Barium	0	500	0	475.7	95.1	0	485.9	97.2
5	Beryllium	0	500	1	475.6	95.1	1	474.4	94.9
6	Cadmium	0	1000	0	942.4	94.2	-1	968.5	96.9
7	Calcium	500000	500000	436983	477053.1	95.4	456328	508772.8	101.8
8	Chromium	0	500	7	490.0	98.0	5	492.7	98.5
9	Cobalt	0	500	4	466.0	93.2	4	470.6	94.1
10	Copper	0	500	7	500.2	100.0	7	512.3	102.5
11	Iron	200000	200000	166368	177664.4	88.8	166632	179210.3	89.6
12	Lead	0	1000	3	910.5	91.1	0	930.0	93.0
13	Magnesium	500000	500000	432782	468407.6	93.7	435952	470768.9	94.2
14	Manganese	0	500	-1	477.1	95.4	-1	484.8	97.0
15	Nickel	0	1000	2	914.5	91.4	3	923.2	92.3
16	Potassium	0	0	91	89.4		128	132.9	
17	Selenium	0	1000	9	931.2	93.1	10	971.6	97.2
18	Silver	0	1000	4	994.1	99.4	7	1009.1	100.9
19	Sodium	0	0	898	1080.9		803	1051.3	
20	Thallium	0	1000	-9	869.1	86.9	-2	917.6	91.8
21	Vanadium	0	500	-3	477.5	95.5	-4	484.4	96.9
22	Zinc	0	1000	10	970.1	97.0	11	1003.2	100.3
23	Molybdenum	0	1000	4	921.3	92.1	0	930.9	93.1

FORM-5A Metal

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 200.9

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

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03M1458E

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

MS Filename: -

Date Analyzed: 051303

Time Analyzed: 12:01

MSD Filename: -

Date Analyzed: 051303

Time Analyzed: 12:08

MS Sample No: MW-11-3

Sample Lab ID: 03-3082-4

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
ARSENIC	µg/L	50	0	54.9	110	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
ARSENIC	µg/L	50	55.0	110	0	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-5A Metal

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 200.7

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	Sample Matrix:	Water
		Batch No:	03M1452L		
MS Filename:	-	Date Analyzed:	051203	Time Analyzed:	13:39
MSD Filename:	-	Date Analyzed:	051203	Time Analyzed:	13:42
MS Sample No:	MW-22-1	Sample Lab ID:	03-3130-2		

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CALCIUM	µg/L	20000	108000	133000	125	75-125
IRON	µg/L	1000	115	1070	96	75-125
MAGNESIUM	µg/L	10000	39100	53800	147 *	75-125
POTASSIUM	µg/L	5000	3110	9070	119	75-125
SODIUM	µg/L	40000	30700	77400	117	75-125
# of Out-of-control					1	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CALCIUM	µg/L	20000	126000	90	5	20	75-125
IRON	µg/L	1000	1010	90	6	20	75-125
MAGNESIUM	µg/L	10000	51500	124	4	20	75-125
POTASSIUM	µg/L	5000	8890	116	2	20	75-125
SODIUM	µg/L	40000	73700	108	5	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-5B Metal
Applied P & Ch Laboratory
Post Digest Spike Sample Recovery

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Batch No.: 03M1452
Matrix: WATER
Analysis Date: 05/12/03

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9
Instrument: ICP -L

Spike Sample No. : 03-3130-02
Client Sample No.: MW-22-1

Concentration Units: UG/L

#	Analyte	Spiked Sample Result(SSR)	13:45 C	Sample Result(SR)	13:30 C	Spike Added(SA)	% Rec.	Control Limit	Q
1	Aluminum	1852.6040		12.7526	B	2000.00	92.0	75-125	
2	Antimony	463.4429		7.4045	B	500.00	91.2	75-125	
3	Arsenic	489.2482		-0.4512	U	500.00	97.8	75-125	
4	Barium	3734.5156		88.7251		4000.00	91.1	75-125	
5	Beryllium	172.1157		0.1102	B	200.00	86.0	75-125	
6	Cadmium	235.4077		1.0622	B	250.00	93.7	75-125	
7	Calcium	129244.1406		107992.0547		20000.00	106.3		
8	Chromium	904.9807		8.7595		1000.00	89.6	75-125	
9	Cobalt	900.9261		-0.1176	U	1000.00	90.1	75-125	
10	Copper	897.4900		5.4562	B	1000.00	89.2	75-125	
11	Iron	1029.8873		115.1744		1000.00	91.5	75-125	
12	Lead	2819.9768		1.6943	B	3000.00	93.9	75-125	
13	Magnesium	48935.4648		39087.8008		10000.00	98.5	75-125	
14	Manganese	843.7136		27.6662		1000.00	81.6	75-125	
15	Nickel	907.8880		4.4787	B	1000.00	90.3	75-125	
16	Potassium	8431.8789		3111.4026		5000.00	106.4	75-125	
17	Selenium	488.7188		3.0440	B	500.00	97.1	75-125	
18	Silver	892.9365		0.7159	B	1000.00	89.2	75-125	
19	Sodium	69012.2734		30696.1758		40000.00	95.8	75-125	
20	Thallium	479.9632		0.5902	U	500.00	96.0	75-125	
21	Vanadium	1763.5845		2.8254	B	2000.00	88.0	75-125	
22	Zinc	489.8489		17.6908		500.00	94.4	75-125	
23	Molybdenum	1956.8752		3.8302	B	2000.00	97.7	75-125	

FORM-5B Metal
Applied P & Ch Laboratory
Post Digest Spike Sample Recovery

Client Name:	GEOFON, Inc.	Project No:	04-4428.10	Lab Code:	APCL
Project Name:	JPL GW Mon-2Q03	Service ID:	033130	Sequence No.:	03M1458E
		Batch No.:	03M1458	Method:	200.9
Spike Sample No. :	03-3082-04	Matrix:	WATER	Instrument:	GFAA-E
Client Sample No.:	MW-11-3	Analysis Date:	05/13/03		

Concentration Units: **UG/L**

#	Analyte	Spiked Sample	12:14	Sample	11:42	Spike	% Rec.	Control Limit	Q
		Result(SSR)	C	Result(SR)	C	Added(SA)			
1	Arsenic	56.4000		0.9000	U	50.00	112.8	75-125	

FORM-6 Metal
Applied P & Ch Laboratory
Duplicates Verification

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Lab Code: APCL
Project Name: JPL GW Mon-2Q03	Service ID: 033130	Sequence No.: 03M1452L
	Batch No.: 03M1452	Method: 200.9
Spike Sample No. 03-3130-02	Matrix: WATER	Instrument: ICP -L
Client Sample No. MW-22-1	% Solid: 0.00	Analysis Date: 05/12/03

Concentration Unit: UG/L

#	Analyte	13:30 Sample(s)	C	13:33 Duplicate	C	RPD(%)	Q
1	Aluminum	12.7526	B	11.9653	B	6.4	
2	Antimony	7.4045	B	3.7321	B	66.0	
3	Arsenic	-0.4512	U	0.6600	U		
4	Barium	88.7251		90.2903		1.7	
5	Beryllium	0.1102	B	0.0176	U	200.0	
6	Cadmium	1.0622	B	0.8392	B	23.5	
7	Calcium	107992.0547		109638.2813		1.5	
8	Chromium	8.7595		8.8449		1.0	
9	Cobalt	-0.1176	U	-0.2775	U		
10	Copper	5.4562	B	5.2974	B	3.0	
11	Iron	115.1744		117.4338		1.9	
12	Lead	1.6943	B	2.1477	B	23.6	
13	Magnesium	39087.8008		40505.9375		3.6	
14	Manganese	27.6662		28.2460		2.1	
15	Nickel	4.4787	B	4.6103	B	2.9	
16	Potassium	3111.4026		3132.7891		0.7	
17	Selenium	3.0440	B	1.7055	U	200.0	
18	Silver	0.7159	B	2.3257	B	105.9	
19	Sodium	30696.1758		32320.1172		5.2	
20	Thallium	0.5902	U	-1.1092	U		
21	Vanadium	2.8254	B	2.4893	B	12.6	
22	Zinc	17.6908		17.6131		0.4	
23	Molybdenum	3.8302	B	2.7063	B	34.4	

FORM-6 Metal
Applied P & Ch Laboratory
Duplicates Verification

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Lab Code: APCL
Project Name: JPL GW Mon-2Q03	Service ID: 033130	Sequence No.: 03M1458E
	Batch No.: 03M1458	Method: 200.9
Spike Sample No. 03-3082-04	Matrix: WATER	Instrument: GFAA-E
Client Sample No. MW-11-3	% Solid: 0.00	Analysis Date: 05/13/03

Concentration Unit: UG/L

#	Analyte	11:42 Sample(s)	C	11:48 Duplicate	C	RPD(%)	Q
1	Arsenic	0.9000	U	0.5000	U		

FORM-7 Metal

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 200.9

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

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03M1458E

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

LCS Filename: -
 LCSD Filename: -

Date Analyzed: 051303
 Date Analyzed: 051303
 Time Analyzed: 11:29
 Time Analyzed: 11:36

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
ARSENIC	µg/L	50	0	56.8	114	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, % RPD REC
ARSENIC	µg/L	50	57.1	114	0	20 80-120
# of Out-of-control				0	0	

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-7 Metal

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 200.7

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

Contract No:
SAS No:
Project No: 04-4428.10
Batch No: 03M1452L

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

LCS Filename: -
LCSD Filename: -

Date Analyzed: 051203
Date Analyzed: 051203
Time Analyzed: 13:24
Time Analyzed: 13:26

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CALCIUM	µg/L	20000	0	21300	107	80-120
IRON	µg/L	1000	0	1020	102	80-120
MAGNESIUM	µg/L	10000	0	9460	95	80-120
POTASSIUM	µg/L	5000	0	5570	111	80-120
SODIUM	µg/L	40000	0	44900	112	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CALCIUM	µg/L	20000	20500	103	4	20	80-120
IRON	µg/L	1000	975	98	4	20	80-120
MAGNESIUM	µg/L	10000	9030	90	5	20	80-120
POTASSIUM	µg/L	5000	5460	109	2	20	80-120
SODIUM	µg/L	40000	42000	105	6	20	80-120
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-9 Metal
Applied P & Ch Laboratory
Serial Dilution

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Batch No.: 03M1452
Matrix: WATER
Analysis Date: 05/12/03

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9
Instrument: ICP -L

Dilution Sample No.: 03-3130-02
Client Sample No.: MW-22-1

Concentration Units: **UG/L**

#	Analyte	Initial Sample Results(I)	13:30 C	Serial Dilut Results(S)	13:37 C	% Diff.	Q
1	Aluminum	12.75	B	73.88	B	479.4	
2	Antimony	7.40	B	22.66	B	206.0	
3	Arsenic	-0.45	U	-6.17	U		
4	Barium	88.73		87.30		1.6	
5	Beryllium	0.11	B	0.08	U	100.0	
6	Cadmium	1.06	B	1.85	B	74.3	
7	Calcium	107992.05		108495.02		0.5	
8	Chromium	8.76		8.22	B	6.1	
9	Cobalt	-0.12	U	-1.40	U		
10	Copper	5.46	B	9.66	B	77.1	
11	Iron	115.17		121.96	B	5.9	
12	Lead	1.69	B	-2.86	U	100.0	
13	Magnesium	39087.80		34874.37		10.8	E
14	Manganese	27.67		28.80		4.1	
15	Nickel	4.48	B	17.50	B	290.7	
16	Potassium	3111.40		2753.40		11.5	E
17	Selenium	3.04	B	17.57	B	477.2	
18	Silver	0.72	B	1.00	U	100.0	
19	Sodium	30696.18		29670.45		3.3	
20	Thallium	0.59	U	10.49	B		
21	Vanadium	2.83	B	2.84	B	0.6	
22	Zinc	17.69		47.57	B	168.9	
23	Molybdenum	3.83	B	5.57	B	45.5	

FORM-9 Metal
Applied P & Ch Laboratory
Serial Dilution

Client Name:	GEOFON, Inc.	Project No:	04-4428.10	Lab Code:	APCL
Project Name:	JPL GW Mon-2Q03	Service ID:	033130	Sequence No.:	03M1458E
		Batch No.:	03M1458	Method:	200.9
Dilution Sample No.:	03-3082-04	Matrix:	WATER	Instrument:	GFAA-E
Client Sample No.:	MW-11-3	Analysis Date:	05/13/03		

Concentration Units: **UG/L**

#	Analyte	Initial Sample	11:42	Serial Dilut	11:55	% Diff.	Q
		Results(I)	C	Results(S)	C		
1	Arsenic	0.90	U	-0.50	U		

FORM-13 Metal
Applied P & Ch Laboratory
Preparation Log

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Lab Code: APCL
Project Name: JPL GW Mon-2Q03	Service ID: 033130	Sequence No.: 03M1452L
	Batch No.: 03M1452	Method: 200.9
Preparation Matrix: WATER	Instrument: ICP -L	

#	Client Sample No.	APCL Sample No.	Preparation Date	Weight (gram)	Volume (ml)
1	1990-065-208	03-3113-02	05/12/03		50.0
2	1990-065-213	03-3113-07	05/12/03		50.0
3	MW-22-1	03-3130-02DM	05/12/03		50.0
4	EB-12-5/8/03	03-3130-01	05/12/03		50.0
5	MW-22-2	03-3130-03	05/12/03		50.0
6	MW-22-3	03-3130-04	05/12/03		50.0
7	MW-22-4	03-3130-05	05/12/03		50.0
8	MW-22-5	03-3130-06	05/12/03		50.0
9	Composite	03-3132-01	05/12/03		50.0
10	Composite	03-3135-01	05/12/03		50.0
11	S-5	03-3143-01	05/12/03		50.0
12	030507002	03-3144-01	05/12/03		50.0
13	030507004	03-3144-02	05/12/03		50.0
14	030507006	03-3144-03	05/12/03		50.0
15	030507018	03-3144-04	05/12/03		50.0
16		03M1452MB	05/12/03		50.0
17		03M1452LCS	05/12/03		50.0
18		03M1452LCSD	05/12/03		50.0
19	MW-22-1 Dup.	03M1452MD	05/12/03		50.0
20	MW-22-1 MS	03M1452MS	05/12/03		50.0
21	MW-22-1 MSD	03M1452MSD	05/12/03		50.0

FORM-13 Metal
Applied P & Ch Laboratory
Preparation Log

Client Name:	GEOFON, Inc.	Project No:	04-4428.10	Lab Code:	APCL
Project Name:	JPL GW Mon-2Q03	Service ID:	033130	Sequence No.:	03M1458E
		Batch No.:	03M1458	Method:	200.9
Preparation Matrix:	WATER	Instrument:	GFAA-E		

#	Client Sample No.	APCL Sample No.	Preparation Date	Weight (gram)	Volume (ml)
1	MW-11-3	03-3082-04DM	05/13/03		50.0
2	EB-10-5/6/03	03-3082-01	05/13/03		50.0
3	MW-11-1	03-3082-02	05/13/03		50.0
4	MW-11-2	03-3082-03	05/13/03		50.0
5	MW-11-4	03-3082-05	05/13/03		50.0
6	MW-11-5	03-3082-06	05/13/03		50.0
7	DUPE-6-2Q03	03-3102-01	05/13/03		50.0
8	EB-11-5/7/03	03-3102-02	05/13/03		50.0
9	MW-12-1	03-3102-03	05/13/03		50.0
10	MW-12-2	03-3102-04	05/13/03		50.0
11	MW-12-3	03-3102-05	05/13/03		50.0
12	MW-12-4	03-3102-06	05/13/03		50.0
13	MW-12-5	03-3102-07	05/13/03		50.0
14	EB-12-5/8/03	03-3130-01	05/13/03		50.0
15	MW-22-1	03-3130-02	05/13/03		50.0
16	MW-22-2	03-3130-03	05/13/03		50.0
17	MW-22-3	03-3130-04	05/13/03		50.0
18	MW-22-4	03-3130-05	05/13/03		50.0
19	MW-22-5	03-3130-06	05/13/03		50.0
20		03M1458MB	05/13/03		50.0
21		03M1458LCS	05/13/03		50.0
22		03M1458LCSD	05/13/03		50.0
23	MW-11-3 Dup.	03M1458MD	05/13/03		50.0
24	MW-11-3 MS	03M1458MS	05/13/03		50.0
25	MW-11-3 MSD	03M1458MSD	05/13/03		50.0

FORM-14 Metal
Applied P & Ch Laboratory
Analysis Run Log

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03
Batch No.(s): 03M1452

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L
Start Date: 05/12/03

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9
End Date: 05/12/03

#	APCL Sample No.	D/F	Time	Al	Sb	As	Ba	Be	Cd	Ca	Cr	Co	Cu	Fe	Pb	Mg	Mn	Hg	Ni	K	Se	Ag	Na	Ti	V	Zn	Mo	Sr	Ti	Sn	Li	B	Si
1	Calib Blank	1.00	12:39	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
2	STD1 1423A	1.00	12:41	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
3	STD2 1423B	1.00	12:44	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
4	STD3 1423C	1.00	12:46	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
5	ICV 1447A	1.00	12:49	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
6	ICB	1.00	12:54	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
7	CRI A1432	1.00	12:57	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
8	ICSA 1441	1.00	12:59	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
9	ICSAB 1443	1.00	13:02	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
10	CCV 1447B	1.00	13:05	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
11	CCB	1.00	13:12	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
12	M-BL 03M1452 W	1.00	13:17	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
13	LCS-03M1452	1.00	13:24	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
14	LCSD-03M1452	1.00	13:26	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
15	3130-2 S F=1	1.00	13:30	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
16	3130-2 D F=1	1.00	13:33	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
17	3130-2 1/5 F=5	5.00	13:37	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
18	3130-2 MS F=1	1.00	13:39	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
19	3130-2 MSD F=1	1.00	13:42	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
20	3130-2 PS F=1	1.00	13:45	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
21	CCV 1447B	1.00	13:47	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
22	CCB	1.00	13:56	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
23	3130-1 F=1	1.00	14:00	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
24	3130-3 F=1	1.00	14:03	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
25	3130-4 F=1	1.00	14:06	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
26	3130-5 F=1	1.00	14:10	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
27	3130-6 F=1	1.00	14:13	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
28	3144-1 F=1	1.00	14:17	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
29	3144-2 F=1	1.00	14:20	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
30	3144-3 F=1	1.00	14:24	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
31	3144-4 F=1	1.00	14:28	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
32	CCV 1447B	1.00	14:30	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
33	CCB	1.00	14:43	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
34	3143-1 F=1	1.00	14:47	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
35	3113-2 F=1	1.00	14:50	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
36	3113-7 F=1	1.00	14:54	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
37	3132-1 F=1	1.00	14:57	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
38	3135-1 F=1	1.00	15:01	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
39	CCV 1447B	1.00	15:04	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
40	CCB	1.00	15:12	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓

FORM-14 Metal
Applied P & Ch Laboratory
Analysis Run Log

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: ICP -L
Start Date: 05/12/03

Lab Code: APCL
Sequence No.: 03M1452L
Method: 200.9
End Date: 05/12/03

Batch No.(s): 03M1452

#	APCL Sample No.	D/F	Time	Al	Sb	As	Ba	Be	Cd	Ca	Cr	Co	Cu	Fe	Pb	Mg	Mn	Hg	Ni	K	Se	Ag	Na	Tl	V	Zn	Mo	Sr	Ti	Sn	Li	B	Si
41	M-BL 03M1453 W	1.00	15:15	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
42	LCS-03M1453	1.00	15:18	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
43	LCSD-03M1453	1.00	15:20	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
44	3141-2 S F=1	1.00	15:24	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
45	3141-2 D F=1	1.00	15:27	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
46	3141-2 1/5 F=5	5.00	15:31	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
47	3141-2 MS F=1	1.00	15:33	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
48	3141-2 MSD F=1	1.00	15:36	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
49	3141-2 PS F=1	1.00	15:39	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
50	CCV 1447B	1.00	15:41	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
51	CCB	1.00	15:45	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
52	3141-1 F=1	1.00	16:16	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
53	3141-3 F=1	1.00	16:19	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
54	3141-3 F=2	2.00	16:23	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
55	3141-4 F=1	1.00	16:26	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
56	3141-5 F=1	1.00	16:29	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
57	3141-6 F=1	1.00	16:33	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
58	3141-5 F=2	2.00	16:35	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
59	3141-7 F=1	1.00	16:42	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
60	3141-8 F=1	1.00	16:46	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
61	CCV 1447B	1.00	16:51	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
62	CCB	1.00	16:54	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
63	3141-9 F=1	1.00	16:57	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
64	3141-9 F=2	2.00	17:00	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
65	3141-10 F=1	1.00	17:04	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
66	3141-11 F=1	1.00	17:09	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
67	3141-12 F=1	1.00	17:13	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
68	ICSA 1441	1.00	17:15	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
69	ICSAB 1443	1.00	17:18	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
70	CCV 1447B	1.00	17:21	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
71	CCB	1.00	17:24	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓
72	DLC A1427	1.00	17:28	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓	✓

FORM-14 Metal
Applied P & Ch Laboratory
Analysis Run Log

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03
Batch No.(s): 03M1458

Project No: 04-4428.10
Service ID: 033130
Instrument: GFAA-E
Start Date: 05/13/03

Lab Code: APCL
Sequence No.: 03M1458E
Method: 200.9
End Date: 05/13/03

#	APCL Sample No.	D/F	Time	Al	Sb	As	Ba	Be	Cd	Ca	Cr	Co	Cu	Fe	Pb	Mg	Mn	Hg	Ni	K	Se	Ag	Na	Ti	V	Zn	Mo	Sr	Ti	Sn	Li	B	Si
1	AS Position 002	1.00	10:17			✓																											
2	AS Position 001	1.00	10:25			✓																											
3	Calib. Blank	1.00	10:34			✓																											
4	1/2 STD1 1472A	1.00	10:40			✓																											
5	STD1 1472A	1.00	10:46			✓																											
6	STD2 1472B	1.00	10:53			✓																											
7	STD3 1472C	1.00	10:59			✓																											
8	ICV A1474	1.00	11:10			✓																											
9	ICB	1.00	11:17			✓																											
10	M-BL 03M1458	1.00	11:23			✓																											
11	LCS-03M1458	1.00	11:29			✓																											
12	LCSD-03M1458	1.00	11:36			✓																											
13	3082-4 S F=1	1.00	11:42			✓																											
14	3082-4 D F=1	1.00	11:48			✓																											
15	3082-4 1/5 F=5	5.00	11:55			✓																											
16	3082-4 MS F=1	1.00	12:01			✓																											
17	3082-4 MSD F=1	1.00	12:08			✓																											
18	3082-4 PS F=1	1.00	12:14			✓																											
19	3082-1 F=1	1.00	12:21			✓																											
20	CCV A1474	1.00	12:27			✓																											
21	CCB	1.00	12:34			✓																											
22	3082-2 F=1	1.00	12:40			✓																											
23	3082-3 F=1	1.00	12:46			✓																											
24	3082-5 F=1	1.00	12:52			✓																											
25	3082-6 F=1	1.00	12:59			✓																											
26	3130-1 F=1	1.00	13:05			✓																											
27	3130-2 F=1	1.00	13:17			✓																											
28	CCV A1474	1.00	13:42			✓																											
29	CCB	1.00	13:49			✓																											
30	3130-3 F=1	1.00	14:08			✓																											
31	3130-4 F=1	1.00	14:14			✓																											
32	3130-5 F=1	1.00	14:21			✓																											
33	3130-6 F=1	1.00	14:27			✓																											
34	3102-1 F=1	1.00	14:34			✓																											
35	3102-2 F=1	1.00	14:40			✓																											
36	3102-3 F=1	1.00	14:47			✓																											
37	3102-4 F=1	1.00	14:53			✓																											
38	CCV A1474	1.00	14:59			✓																											
39	CCB	1.00	15:06			✓																											
40	CCV A1474	1.00	15:15			✓																											

FORM-14 Metal
Applied P & Ch Laboratory
Analysis Run Log

Client Name: GEOFON, Inc.
Project Name: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 033130
Instrument: GFAA-E
Start Date: 05/13/03

Lab Code: APCL
Sequence No.: 03M1458E
Method: 200.9
End Date: 05/13/03

#	APCL Sample No.	D/F	Time	Al	Sb	As	Ba	Be	Cd	Ca	Cr	Co	Cu	Fe	Pb	Mg	Mn	Hg	Ni	K	Se	Ag	Na	Tl	V	Zn	Mo	Sr	Ti	Sn	Li	B	Si
41	CCB	1.00	15:21			✓																											
42	3102-5 F=1	1.00	15:27			✓																											
43	3102-6 F=1	1.00	15:33			✓																											
44	3102-7 F=1	1.00	15:40			✓																											
45	CCV A1474	1.00	16:06			✓																											
46	CCB	1.00	16:12			✓																											

13760 Magnolia Ave. Chino CA 91710

Metal Digestion (3010/3050) Worksheet

Tel: (909) 590-1828 Fax: (909) 590-1488

Batch # 03M1452 Matrix: W Method used: 3010A Date: 5/16/03 Digested by: X Diluted by: _____Lot #: ASTM Type I water RW1411 HNO₃ 1102/20 H₂SO₄ _____ HCl 4102050 H₂O₂ _____

OP #	Type	Samp ID /Lot #	X (g or mL)	$V_{\text{digest}}/X = f_1$	$V_1/V_i = f_2$	$V_j/V_i = f_3$	$F = f_1 f_2 f_3$	Note
1651	Method Blank	Bl. Lot <u>RW1411</u>	<u>50</u>	<u>50/X =</u>	<u>/ =</u>	<u>/ =</u>		<u>23me</u>
1652	LCS1	Bl. Lot <u>1</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		<u>T-95c</u>
1653	Sample-1	<u>3130-2</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1654	MS1 on S-1	<u>-2</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1655	MS2 on S-1	<u>-2</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1656	Sample 2	<u>-1</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1657	Sample 3	<u>-3</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1658	Sample 4	<u>-4</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1659	Sample 5	<u>-5</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1660	Sample 6	<u>-6</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1661	Sample 7	<u>3104-1</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1662	Sample 8	<u>-2</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1663	Sample 9	<u>-3</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1664	Sample 10	<u>-4</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1665	LCS2	Bl. Lot <u>RW1411</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1666	Sample 11	<u>3103-1</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1667	Sample 12	<u>3113-2</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1668	Sample 13	<u>-7</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1669	Sample 14	<u>3132-1</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1670	Sample 15	<u>3135-1</u>	<u>↓</u>	<u>↓</u> <u>/X =</u> <u>↓</u>	<u>/ =</u>	<u>/ =</u>		
1671	Sample 16			<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1672	Sample 17			<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1673	Sample 18			<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1674	Sample 19			<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1675	Sample 20	<u>X 5/16/03</u>		<u>/X =</u>	<u>/ =</u>	<u>/ =</u>		
1676	Duplicate	<u>3130-2</u>	<u>50</u>	<u>50/X =</u>	<u>/ =</u>	<u>/ =</u>		

Specification of matrix spike and lab control spike

QC Type	Spiked Element *	Spike Stock Solution Lot #	Spike Stock (Rep.) Conc. C_s , $\mu\text{g/mL}$	Spike Stock Volum Used V_s , mL	Spike Level $T' = C_s V_s / V$ ppm or mg/L	Sample Spike T , ppm
MS1	/As/Se/Sb/M ₂₀	AA- /AA- /AA- /AA- <u>1033</u>	/ / / <u>25</u>	/ / / <u>2</u>	/ / / <u>1</u>	
MS2	/As/Se/Sb/M ₂₀	AA- /AA- /AA- /AA- <u>11</u>	/ / / <u>1</u>	/ / / <u>1</u>	/ / / <u>1</u>	
LCS1	/As/Se/Sb/M ₂₀	AA- /AA- /AA- /AA- <u>1482</u>	/ / / <u>1</u>	/ / / <u>1</u>	/ / / <u>1</u>	
LCS2	/As/Se/Sb/M ₂₀	AA- /AA- /AA- /AA- <u>11</u>	/ / / <u>1</u>	/ / / <u>1</u>	/ / / <u>1</u>	

* Notation: T - rep. sample spike level. T' - digest solution spike level. $T = f T' = C_s V_s / X$. M20 (or Mj) represents 20 (or j) metals, (see STD logbook).
 If digest needs dilution for different metals, use dilution worksheet.

APCL form 6-116 April, 03, 1996. Ver. 4.0

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

Root-File:[CUST.DOC.AA]DIGEST.ROOT.TEX

File:[CUST.DOC.AA]DIGEST.TEX

Supervisor Initial

6589

290 Magnolia Ave. Chino CA 91710

Metal Digestion (3010/3050) Worksheet

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

Batch # 03M1458 Matrix: W Method used: 3020A Date: 5/13/03 Digested by: ML Diluted by: _____# : ASTM Type I water AW 1411 HNO₃ 110220 H₂SO₄ _____ HCl _____ H₂O₂ _____

OP #	Type	Samp ID /Lot #	X (g or mL)	$V_{\text{digest}}/X = f_1$	$V_f/V_i = f_2$	$V_f/V_i = f_3$	$F=f_1f_2f_3$	Note
1755	Method Blank	Bl. Lot: <u>AW 1411</u>	<u>50</u>	<u>50</u> /X=	<u>1</u> =	<u>1</u> =		<u>GEAA/AS</u>
1756	LCS1	Bl. Lot: <u>11</u>		/X=	<u>1</u> =	<u>1</u> =		<u>T=95°C</u>
1757	Sample-1	<u>3082-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1758	MS1 on S-1	<u>-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1759	MS2 on S-1	<u>-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1760	Sample 2	<u>-1</u>		/X=	<u>1</u> =	<u>1</u> =		
1761	Sample 3	<u>-2</u>		/X=	<u>1</u> =	<u>1</u> =		
1762	Sample 4	<u>-3</u>		/X=	<u>1</u> =	<u>1</u> =		
1763	Sample 5	<u>-5</u>		/X=	<u>1</u> =	<u>1</u> =		
1764	Sample 6	<u>-6</u>		/X=	<u>1</u> =	<u>1</u> =		
1765	Sample 7	<u>3130-1</u>		/X=	<u>1</u> =	<u>1</u> =		
1766	Sample 8	<u>-2</u>		/X=	<u>1</u> =	<u>1</u> =		<u>dup/ms</u>
1767	Sample 9	<u>-3</u>		/X=	<u>1</u> =	<u>1</u> =		
1768	Sample 10	<u>-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1769	LCS2	Bl. Lot: <u>AW 1411</u>		/X=	<u>1</u> =	<u>1</u> =		
1770	Sample 11	<u>-5</u>		/X=	<u>1</u> =	<u>1</u> =		
1771	Sample 12	<u>-6</u>		/X=	<u>1</u> =	<u>1</u> =		
1772	Sample 13	<u>3102-1</u>		/X=	<u>1</u> =	<u>1</u> =		
1773	Sample 14	<u>-2</u>		/X=	<u>1</u> =	<u>1</u> =		
1774	Sample 15	<u>-3</u>		/X=	<u>1</u> =	<u>1</u> =		
1775	Sample 16	<u>-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1776	Sample 17	<u>-5</u>		/X=	<u>1</u> =	<u>1</u> =		
1777	Sample 18	<u>-6</u>		/X=	<u>1</u> =	<u>1</u> =		
1778	Sample 19	<u>-7</u>		/X=	<u>1</u> =	<u>1</u> =		
1779	Sample 20	<u>3082-4</u>		/X=	<u>1</u> =	<u>1</u> =		
1780	Duplicate	<u>3082-4</u>	<u>50</u>	<u>50</u> /X=	<u>1</u> =	<u>1</u> =		

Specification of matrix spike and lab control spike

QC Type	Spiked Element *	Spike Stock Solution Lot #	Spike Stock (Rep.) Conc. C_s , $\mu\text{g/mL}$	Spike Stock Volum Used V_s , mL	Spike Level $T'=C_s V_s/V$ ppm or mg/L	Sample Spike T , ppm
MS1	/As/Sb/M ₂₀	AA- /AA-103/AA- /AA-	<u>15</u> / /	<u>10.5</u> /	<u>10.5</u> /	
MS2	/As/Sb/M ₂₀	AA- /AA-11/AA- /AA-	<u>1</u> / /	<u>1</u> / /	<u>1</u> / /	
LCS1	/As/Sb/M ₂₀	AA- /AA-11/AA- /AA-	<u>1</u> / /	<u>1</u> / /	<u>1</u> / /	
LCS2	/As/Sb/M ₂₀	AA- /AA-11/AA- /AA-	<u>1</u> / /	<u>1</u> / /	<u>1</u> / /	

* Notation: T - rep. sample spike level. T' - digest solution spike level. $T=f T'=C_s V_s/X$. M20 (or Mj) represents 20 (or j) metals, (see STD logbook).

If digest needs dilution for different metals, use dilution worksheet.

APCL form 6-116 April. 03, 1996. Ver. 4.0

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

Root File: [CUST.DOC.AA]DIGEST_ROOT.TEX

File: [CUST.DOC.AA]DIGEST.TEX

Supervisor Initial

ML

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 ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130

Anal. Method SM2320B
Collected by:

Component Name: Bicarbonate
CAS No:

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	< 2	U
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	271	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	148	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	130	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	114	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/10/03	03W2817	mg/L	2	49.9	
03W2817-MB-01	03W2817-MB-01	Water	05/10/03	05/10/03	05/10/03	03W2817	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 GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130

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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	<2	U
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	<2	U
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	2.6	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	5.2	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	2.6	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	76.6	
03W2817-MB-01	03W2817-MB-01	Water	05/10/03	05/10/03	05/10/03	03W2817	mg-CaCO ₃ /L	2	<2	U

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

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

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

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

Client Name: GEOFON, Inc.
Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130

Anal. Method 9040B
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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	7.31	
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	7.64	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	8.21	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	8.18	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	8.07	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	9.23	
03W2790-MB-01	03W2790-MB-01	Water	05/08/03	05/08/03	05/08/03	03W2790	pH unit	0.01	6.89	

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 GW Mon-2Q03 Service ID: 33130 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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	<10	U
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	646	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	287	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	287	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	213	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/09/03	03W2800	mg/L	10	208	
03W2800-MB-01	03W2800-MB-01	Water	05/09/03	05/09/03	05/09/03	03W2800	mg/L	10	<10	U

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

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

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

Applied P & Ch Laboratory
Wet Analysis Results for Method 7196

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 7196
Project ID: JPL GW Mon-2Q03 Service ID: 33130 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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/08/03	03W2788	mg/L	0.01	<0.01	U
03W2788-MB-01	03W2788-MB-01	Water	05/08/03	05/08/03	05/08/03	03W2788	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 GW Mon-2Q03

Project No: 04-4428.10
Service ID: 33130

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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	<4	U
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	3.2	B
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	1.6	B
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	1.8	B
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	<4	U
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/14/03	03W2866	µg/L	4	<4	U
03W2866-MB-01	03W2866-MB-01	Water	05/14/03	05/14/03	05/14/03	03W2866	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 No: 04-4428.10 Anal. Method 300.0
 Project ID: JPL GW Mon-2Q03 Service ID: 33130 Collected by:

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

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.25	0.27	
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	2.5	107	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	1	34.7	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	1	33.9	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.4	11.8	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.4	8.1	
03W2777-MB-01	03W2777-MB-01	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.2	<0.2	U

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

Note: Q - Qualifier.

Qualifier: U - Not Detected or less than MDL

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

Applied P & Ch Laboratory
Wet Analysis Results for Method 300.0

Client Name: GEOFON, Inc. Project No: 04-4428.10 Anal. Method 300.0
 Project ID: JPL GW Mon-2Q03 Service ID: 33130 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
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.05	0.087	
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.5	9.1	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.2	6.8	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.2	6.9	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.08	4.1	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.08	0.11	
03W2777-MB-01	03W2777-MB-01	Water	05/08/03	05/08/03	05/08/03	03W2777	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 GW Mon-2Q03

Project No: 04-4428.10
 Service ID: 33130

Anal. Method 300.0
 Collected by:

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

Lab ID	Sample ID	Matrix	Coll. Date	Rcv Date	Anal. Date	Batch	Unit	RL	Result	Q
03-3130-1	EB-12-5/8/03	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	0.63	0.69	
03-3130-2	MW-22-1	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	6.3	138	
03-3130-3	MW-22-2	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	2.5	29.7	
03-3130-4	MW-22-3	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	2.5	34.6	
03-3130-5	MW-22-4	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	1	7.6	
03-3130-6	MW-22-5	Water	05/08/03	05/08/03	05/08/03	03W2777	mg/L	1	22.0	
03W2777-MB-01	03W2777-MB-01	Water	05/08/03	05/08/03	05/08/03	03W2777	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.
 Case No:
 Project ID: JPL GW Mon-2Q03

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03W2777

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

LCS Filename: -
 LCSD Filename: -

Date Analyzed: 050803
 Date Analyzed: 050803
 Time Analyzed: 11:25
 Time Analyzed: 11:38

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHLORIDE CL ⁻	mg/L	4.0	0	4.25	106	80-120
NITRATE AS N	mg/L	1.5	0	1.58	105	80-120
SULFATE SO ₄ ⁻	mg/L	15	0	15.7	105	80-120
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE CL ⁻	mg/L	4.0	4.04	101	5	20	80-120
NITRATE AS N	mg/L	1.5	1.58	105	0	20	80-120
SULFATE SO ₄ ⁻	mg/L	15	17.1	114	8	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.
Case No:
Project ID: JPL GW Mon-2Q03

Contract No:
SAS No:
Project No: 04-4428.10
Batch No: 03W2777

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

MS Filename: -

Date Analyzed: 050803

Time Analyzed: 15:06

MSD Filename: -

Date Analyzed: 050803

Time Analyzed: 15:18

MS Sample No: MW-22-1

Sample Lab ID: 03-3130-2

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHLORIDE CL ⁻	mg/L	100	107	209	102	75-125
NITRATE AS N	mg/L	37.5	9.1	47.7	103	75-125
SULFATE SO ₄ ⁻	mg/L	375	138	540	107	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
CHLORIDE CL ⁻	mg/L	100	211	104	2	20	75-125
NITRATE AS N	mg/L	37.5	47.7	103	0	20	75-125
SULFATE SO ₄ ⁻	mg/L	375	532	105	2	25	75-125
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

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

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

Sample Matrix: Water

Batch No: 03W2866

LCS Filename: -

Date Analyzed: 051403

Time Analyzed: 24:44

LCSD Filename: -

Date Analyzed: -

Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
PERCHLORATE	µg/L	25	0	24.4	98	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.
Case No:
Project ID: JPL GW Mon-2Q03

Contract No:
SAS No:
Project No: 04-4428.10
Batch No: 03W2866

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

MS Filename: -

Date Analyzed: 051403

Time Analyzed: 15:25

MSD Filename: -

Date Analyzed: 051403

Time Analyzed: 15:44

MS Sample No: MW-22-1

Sample Lab ID: 03-3130-2

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
PERCHLORATE	µg/L	50	3.2	57.3	108	75-125
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, % RPD REC	
						RPD	REC
PERCHLORATE	µg/L	50	54.1	102	6	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.
 Case No:
 Project ID: JPL GW Mon-2Q03

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03W2800

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

LCS Filename: -
 LCSD Filename: -

Date Analyzed: 050903
 Date Analyzed: 050903
 Time Analyzed: 10:38
 Time Analyzed: 10:38

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	0	390	98	88-108
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, % RPD REC	
						RPD	REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	386	97	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:	33130
Project ID:	JPL GW Mon-2Q03	Project No:	04-4428.10	Sample Matrix:	Water
		Batch No:	03W2800		
MS Filename:	-	Date Analyzed:	050903	Time Analyzed:	10:38
MSD Filename:	-	Date Analyzed:	050903	Time Analyzed:	10:38
MS Sample No:	MW-22-1	Sample Lab ID:	03-3130-2		

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	646	1030	96	80-119
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	MSD Concentration	MSD Rec% #	RPD% #	QC Limit, % RPD REC
SOLIDS, TOTAL DISSOLVED (TDS)	mg/L	400	1010	91	5	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: 33130

Project ID: JPL GW Mon-2Q03

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03W2788

LCS Filename: -

Date Analyzed: 050803

Time Analyzed: 14:43

LCSD Filename: -

Date Analyzed: 050803

Time Analyzed: 14:43

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
CHROMIUM (VI)	mg/L	0.25	0	0.222	89	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	0.239	96	8	19	80-115
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

FORM-3

Applied P & Ch Laboratory

Matrix Spike/Matrix Spike Duplicate Recovery for Method 7196

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

Contract No:
 SAS No:
 Project No: 04-4428.10
 Batch No: 03W2788

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

MS Filename: -

Date Analyzed: 050803

Time Analyzed: 14:43

MSD Filename: -

Date Analyzed: 050803

Time Analyzed: 14:43

MS Sample No: MW-22-1

Sample Lab ID: 03-3130-2

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
CHROMIUM (VI)	mg/L	0.25	0	0.227	91	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	0.219	88	3	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: _____

Wet Chemistry QC Report B
Duplicate Results

Matrix: Water

APCL Service ID: 03-3130

Analysis	Batch ID	Analysis Date	Sample Name	Unit	Result	Duplicate Result	RPD %	RPD Control limit
Bicarbonate	03W2817	5/10/03	03-3082-2	mg/L	208	210	1	20
Carbonate	03W2817	5/10/03	03-3082-2	mg-CaCO ₃ /L	ND	ND	NC	20
pH	03W2790	5/8/03	EB-12-5/8/03	pH unit	7.31	7.36	1	20

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

6A

INITIAL CALIBRATION DATA

Lab Name: Applied P & Ch Lab Contract: _____

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)

$$r = 0.9997$$

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

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 Table -- Last Modified: 17:42 on Fri, 21 Mar 2003

Component # 1 Fluoride Retention Time 1.32
Reference Comp. Fluoride Window Size 0.15 min.
Amount = K0 + K1*Area
K0 = -9.62851E-004
K1 = 1.37614E-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*Area
K0 = 1.28188E-001
K1 = 1.95287E-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*Area
K0 = 2.38085E-002
K1 = 9.76240E-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 = $K0 + K1 \cdot \text{Area}$
 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

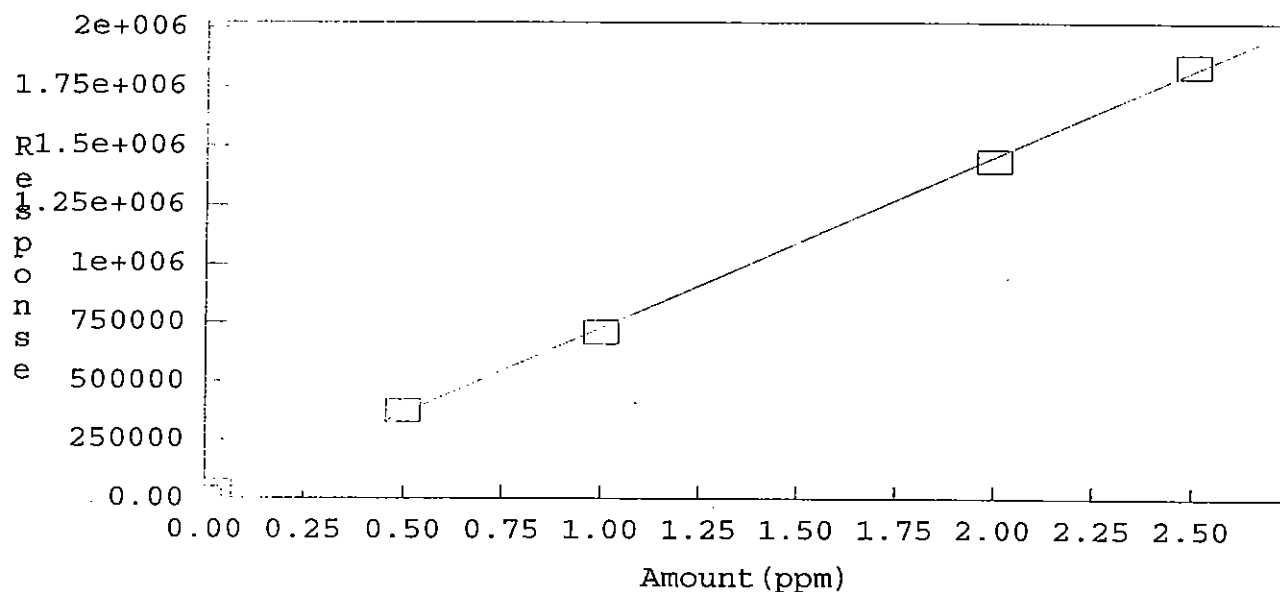
$r^2 = 0.999552$

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

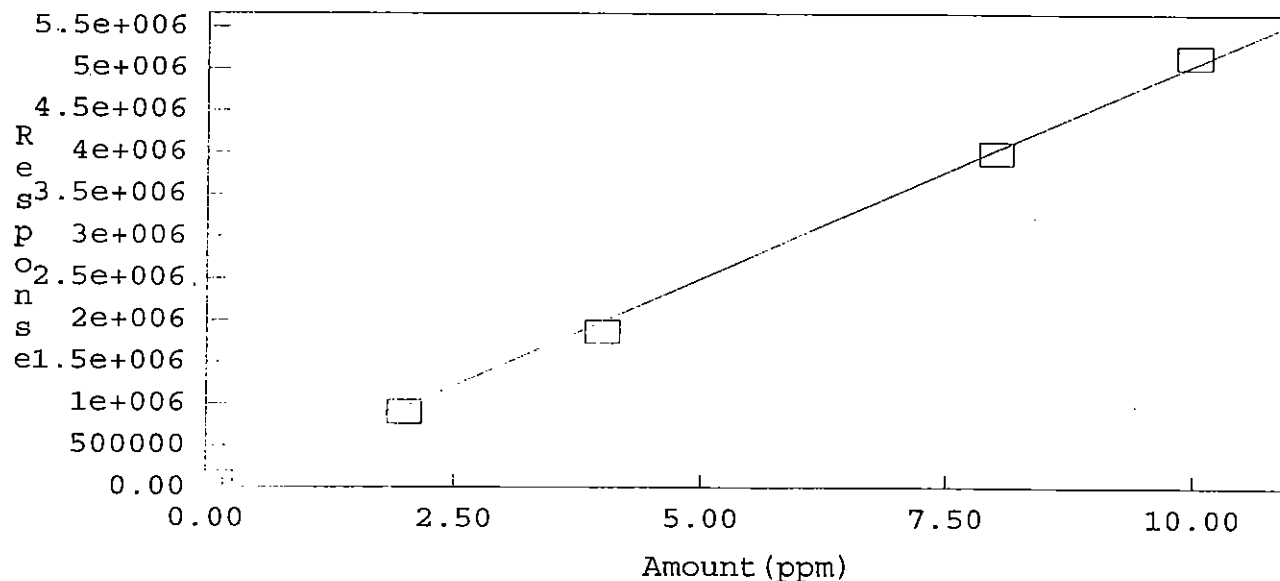
$r^2 = 0.998409$

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

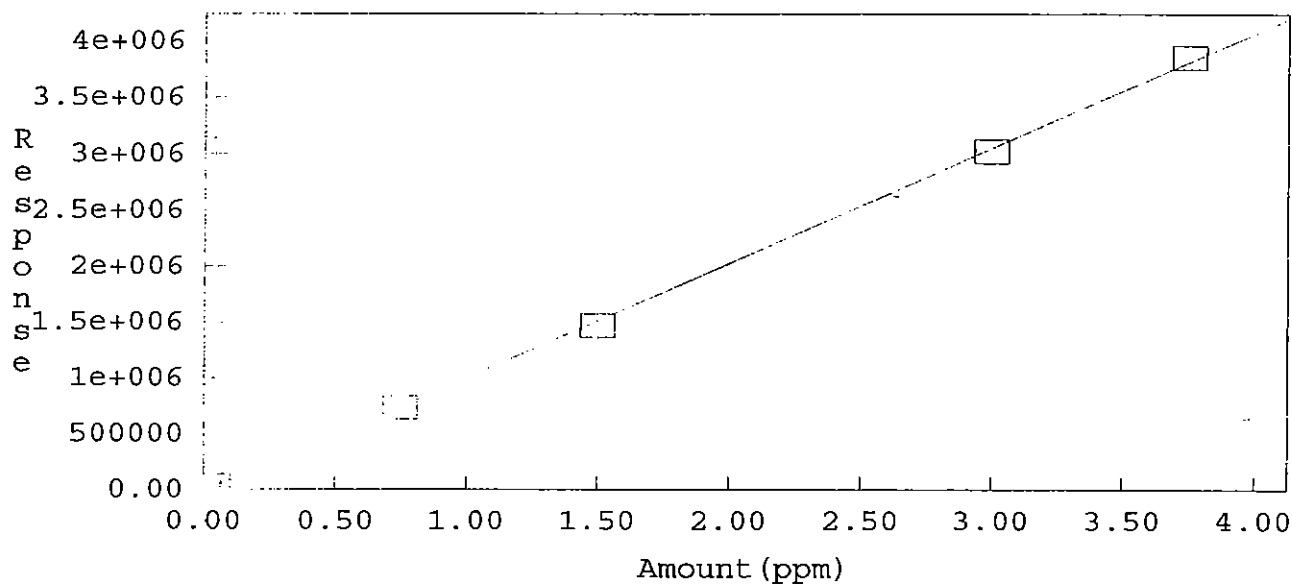
$r^2 = 0.999594$

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

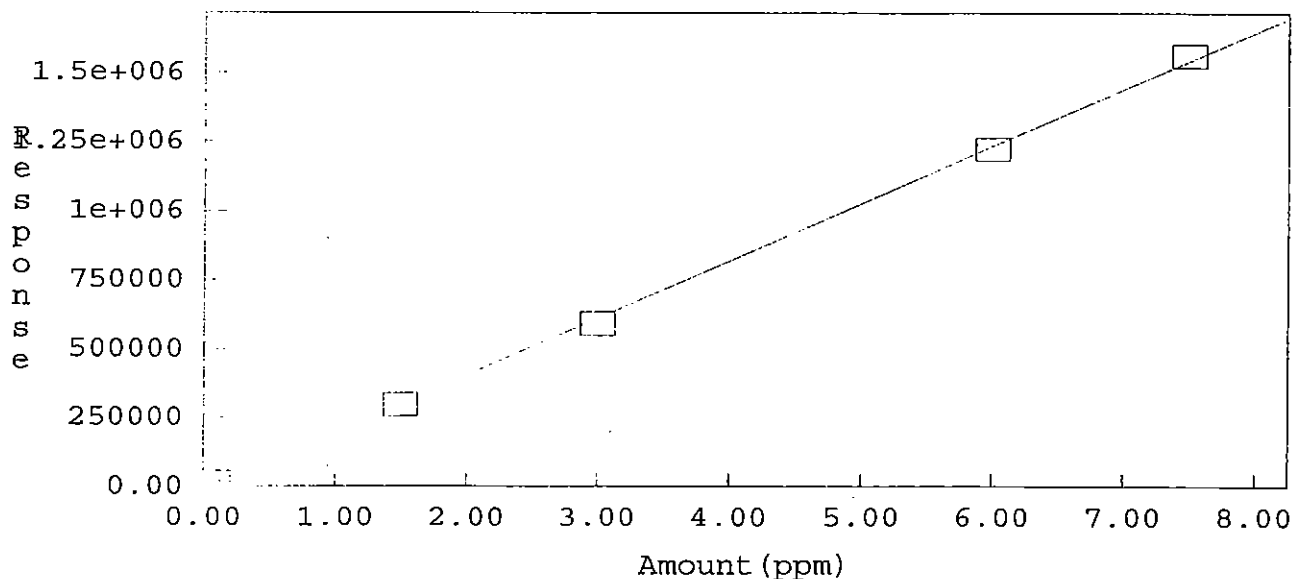
$r^2 = 0.999518$

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

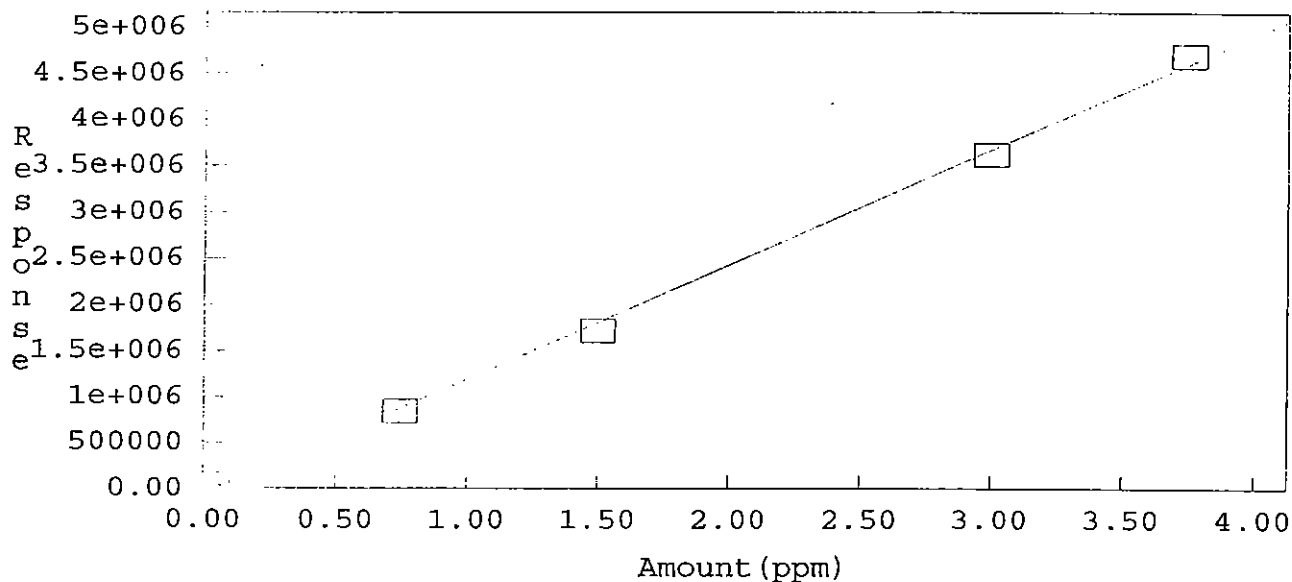
$r^2 = 0.998618$

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

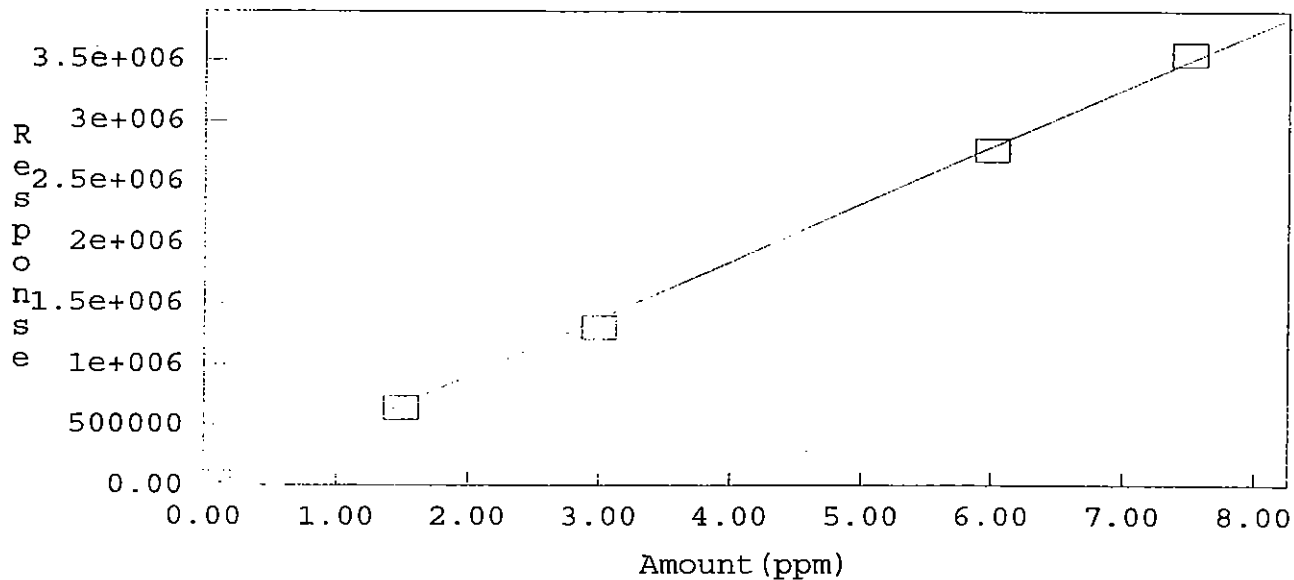
$r^2 = 0.998898$

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

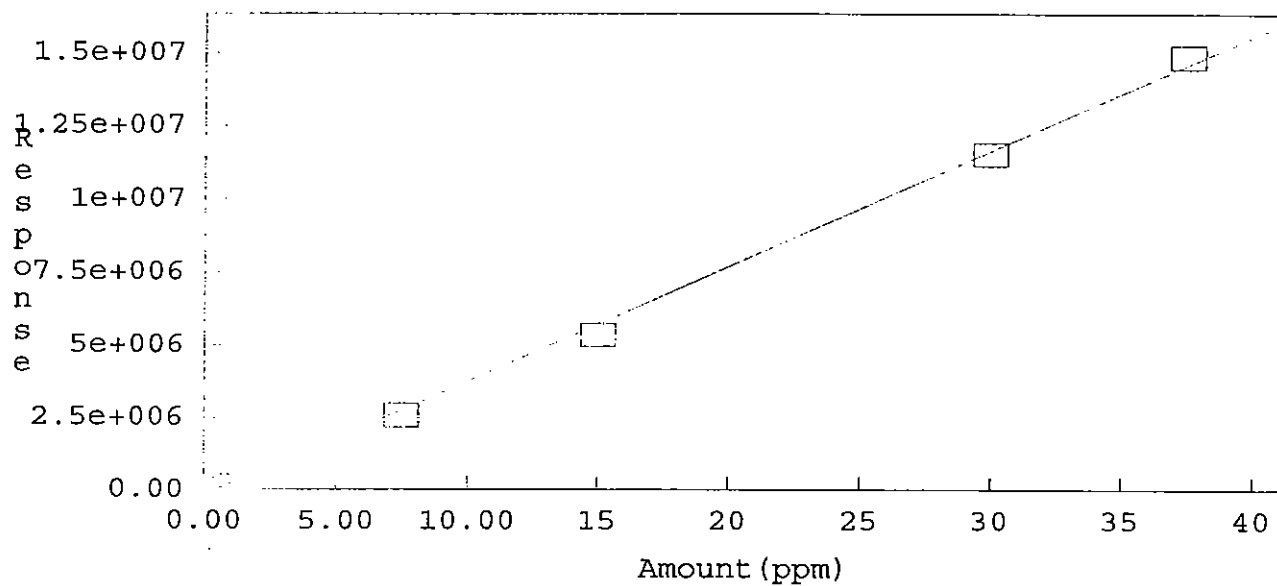
$r^2 = 0.998245$

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

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

Standardization: External

Calibration: Area



```

=====
Sample Name: ICV-W7768-100X                      Date: 03/21/2003 17:56:33
Data File  : C:\DX\DATA\e300-063\W7768Q01.D07
Method     : C:\DX\METHOD\E300-063.MET
ACI Address: 1 System: 1 Inject#: 7                Detector: COND
Analyst    : David                               Column: Dionex AS4A-SC
=====

```

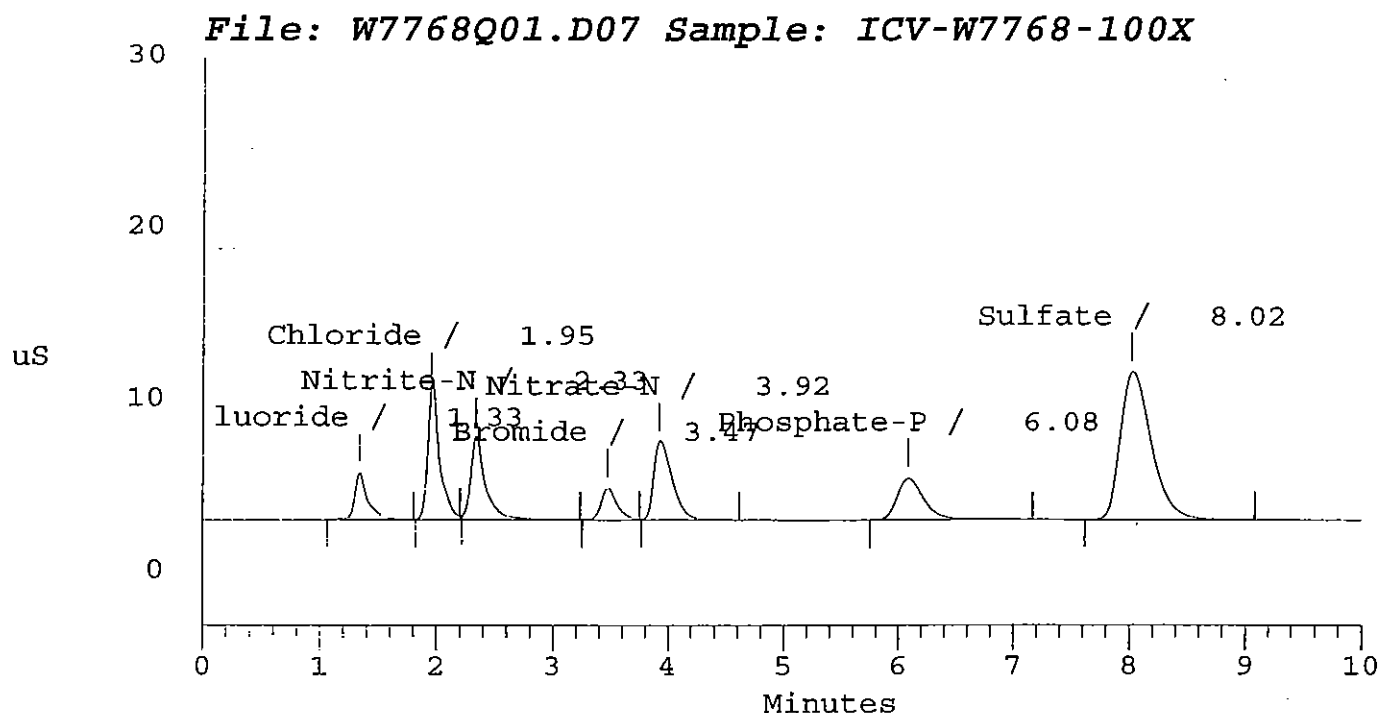
```

-----
Calibration Volume Dilution Points Rate Start Stop Area Reject
-----
External          1          1    3000   5Hz   0.00 10.00      1000
-----

```

***** Component Report: All Components *****

Pk. Num	Ret Time	Component Name	Concentration ppm	Height	Area	Bl. Code	%Delta
1	1.33	Fluoride	0.999	89896	726672	2	0.00
2	1.95	Chloride	3.767	248158	1863483	2	0.00
3	2.33	Nitrite-N	1.445	160603	1455681	2	1.17
4	3.47	Bromide	2.881	61007	586729	2	-0.71
5	3.92	Nitrate-N	1.410	149958	1697030	2	0.00
6	6.08	Phosphate-P	2.852	79916	1302965	1	0.00
7	8.02	Sulfate	13.993	285121	5314983	1	0.00
Totals			27.347	1074659	12947545		



```

=====
Sample Name: ICB                      Date: 03/21/2003 18:21:00
Data File  : C:\DX\DATA\E300-063\W7767Q01.D08
Method     : C:\DX\METHOD\E300-063.MET
ACI Address: 1 System: 1 Inject#: 8    Detector: COND
Analyst    : David                    Column: Dionex AS4A-SC
=====

```

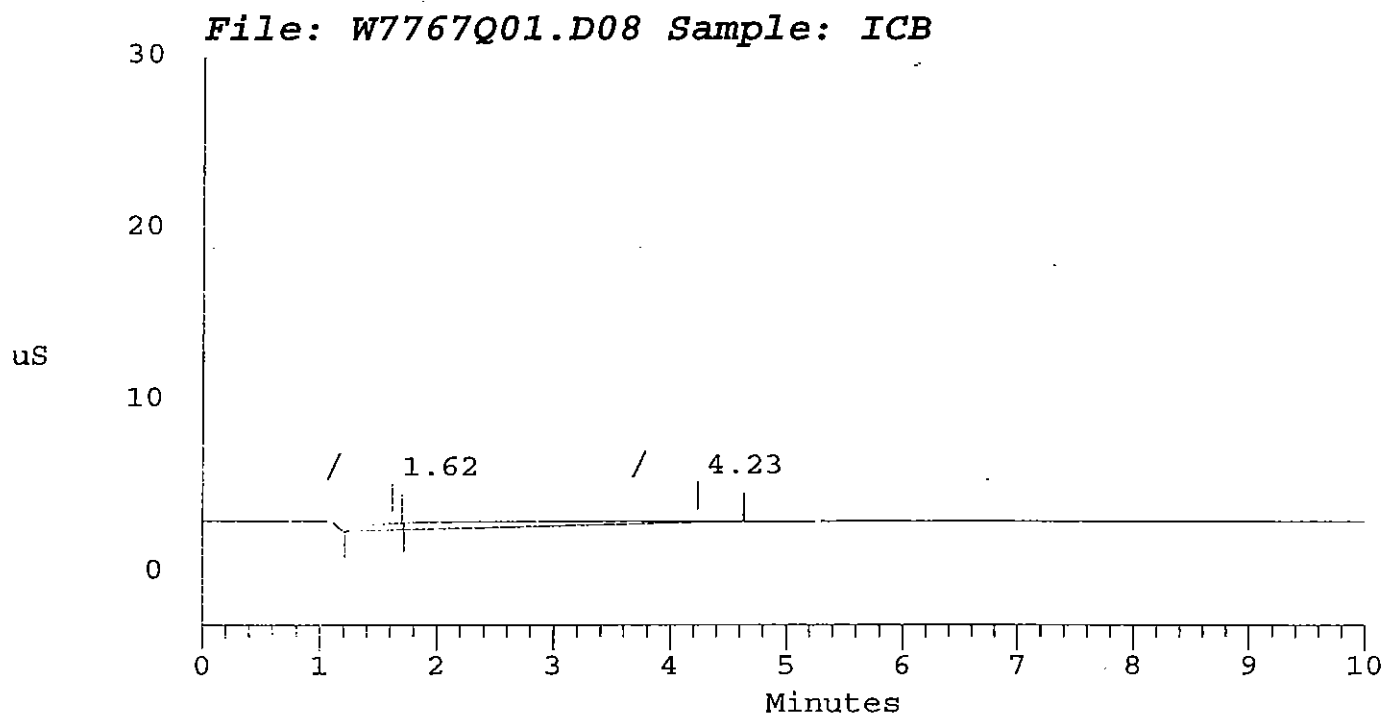
```

Calibration Volume Dilution Points Rate Start Stop Area Reject
-----
External          1          1    3000   5Hz   0.00 10.00      1000

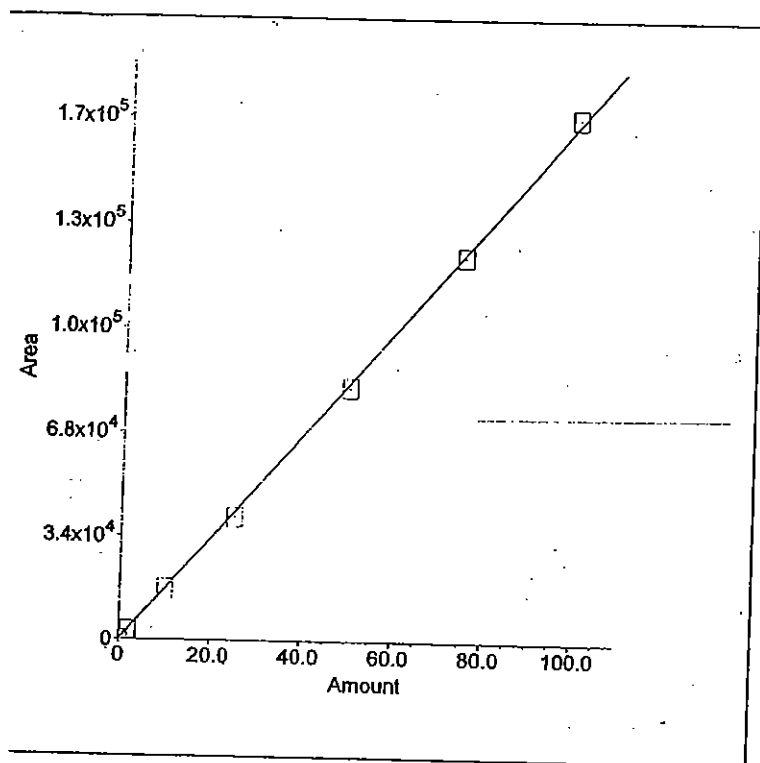
```

***** Component Report: All Components *****

Pk. Num	Ret Time	Component Name	Concentration ppm	Height	Area	Bl. Code	%Delta
0	0.00	Fluoride	0.000	0	0	0	0.00
0	0.00	Chloride	0.000	0	0	0	0.00
0	0.00	Nitrite-N	0.000	0	0	0	0.00
0	0.00	Bromide	0.000	0	0	0	0.00
0	0.00	Nitrate-N	0.000	0	0	0	0.00
0	0.00	Phosphate-P	0.000	0	0	0	0.00
0	0.00	Sulfate	0.000	0	0	0	0.00
Totals			0.000	0	0		



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 IC-C

APCL Perchlorate Analysis Report

Sample Name : ICV 50 ppb w7828a

Data File Name : C:\DATA\E314-011\icv-50pb_009.DXD

Method File Name : C:\PEAKNET\METHOD\E314-011.met

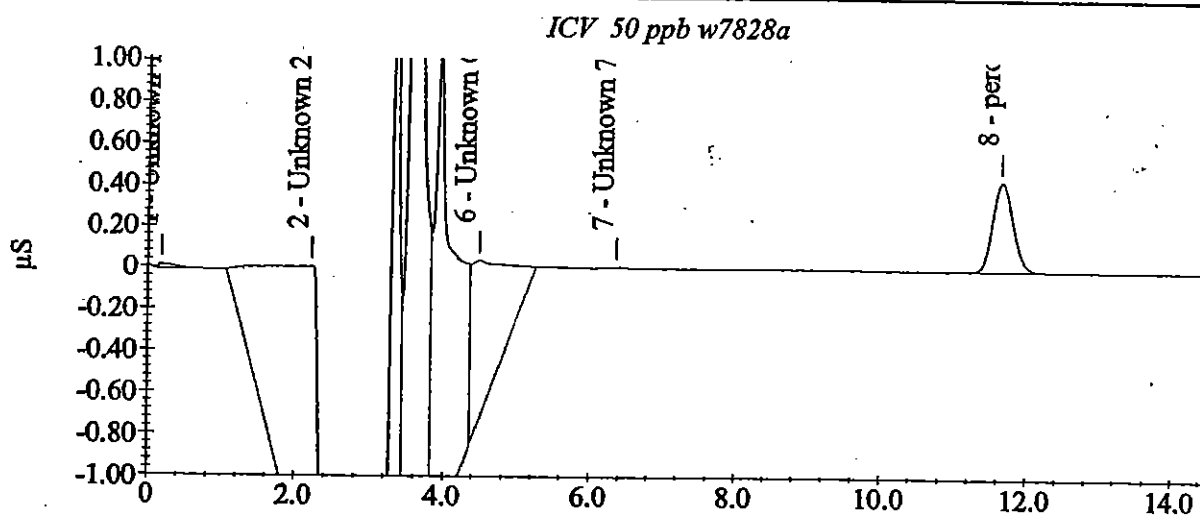
Date Time Collected : 03/12/2003 8:16:15 PM

System Operator : wei wang

Dilution Factor : 1.00

Peak Information : All Components

Peak #	Component Name	Retention Time	Amount (ppb)	Peak Area	Peak Height
8	perchlorate	11.65	49.49	83990	4321



APCL Perchlorate Analysis Report

Sample Name : icb

Data File Name : C:\DATA\E314-011\ICB_010.DXD

Method File Name : c:\PeakNet\method\E314-011.met

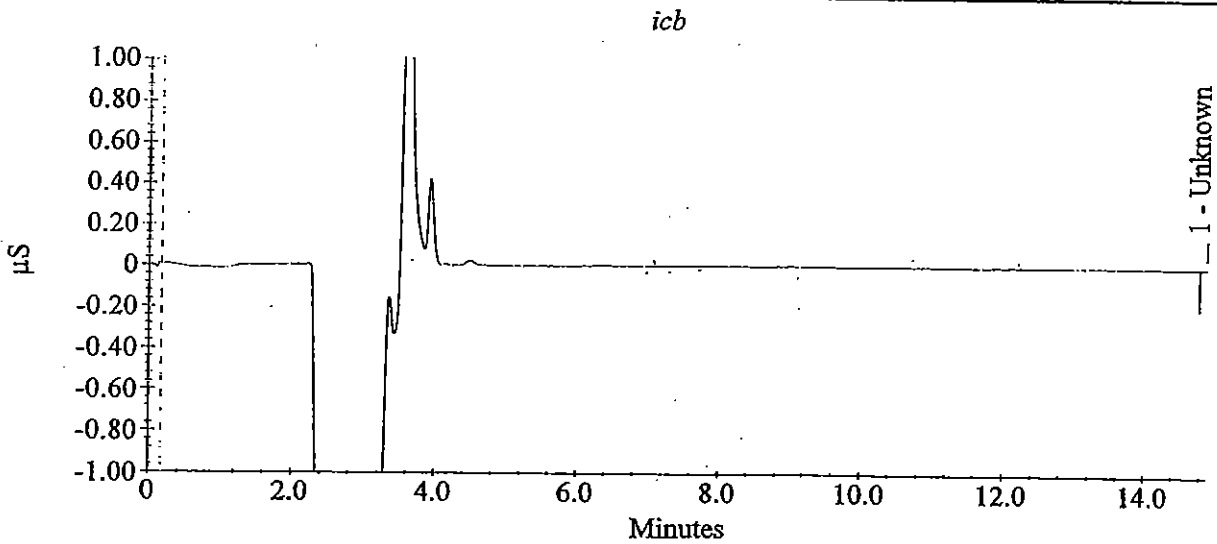
Date Time Collected : 03/12/2003 8:33:51 PM

System Operator : wei wang

Dilution Factor : 1.00

Peak Information : All Components

Peak #	Component Name	Retention Time	Amount (ppb)	Peak Area	Peak Height
--------	----------------	----------------	--------------	-----------	-------------



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

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

Contract No.:
SAS No.:
Project No.: 04-4428.10

Lab Code: APCL
Service ID: 33130

#	Component Name	Method	Batch No.	Unit	Expected	Test Result	Rec. %	Dev. %	Flag	Control Limit, %	Test Date
1	Chloride Cl^-	300.0	03W2777	mg/L	4.0	4.04	101	1	✓	90-110	05/08/2003
	NITRATE as N-NO_3^- , BY	300.0	03W2777	mg/L	1.5	1.57	104	4	✓	90-110	05/08/2003
	SULFATE SO_4^{--} , BY I	300.0	03W2777	mg/L	15	15.6	104	4	✓	90-110	05/08/2003
	Chloride Cl^-	300.0	03W2777	mg/L	4.0	4.05	101	1	✓	90-110	05/08/2003
	NITRATE as N-NO_3^- , BY	300.0	03W2777	mg/L	1.5	1.56	104	4	✓	90-110	05/08/2003
	SULFATE SO_4^{--} , BY I	300.0	03W2777	mg/L	15	15.4	103	3	✓	90-110	05/08/2003
	Chloride Cl^-	300.0	03W2777	mg/L	4.0	4.10	103	3	✓	90-110	05/08/2003
	NITRATE as N-NO_3^- , BY	300.0	03W2777	mg/L	1.5	1.57	104	4	✓	90-110	05/08/2003
	SULFATE SO_4^{--} , BY I	300.0	03W2777	mg/L	15	15.5	103	3	✓	90-110	05/08/2003
2	Perchlorate	314.0	03W2866	$\mu\text{g/L}$	50	49.5	99	-1	✓	90-110	05/14/2003
	Perchlorate	314.0	03W2866	$\mu\text{g/L}$	50	50.1	100	0	✓	90-110	05/14/2003
	Perchlorate	314.0	03W2866	$\mu\text{g/L}$	50	51.2	102	2	✓	90-110	05/14/2003
	Perchlorate	314.0	03W2866	$\mu\text{g/L}$	50	48.7	97	-3	✓	90-110	05/14/2003
3	Chromium (VI)	7196	03W2788	mg/L	0.25	0.242	97	-3	✓	90-110	05/08/2003
	Chromium (VI)	7196	03W2788	mg/L	0.25	0.255	102	2	✓	90-110	05/08/2003

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)	
5/7/03	A-01	Not in Use			✓	B-01	1.00	10.01	200.00	✓✓✓	C-01	1.0000	10.0001	200.0000	✓✓✓	1/2
	A-02					B-05	1.00	10.00	200.01	✓✓✓	C-05	1.0001	10.0000	200.0001	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	9.99	200.00	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/8/03	A-01	Not in Use			✓	B-01	1.00	10.01	200.00	✓✓✓	C-01	1.0000	10.0001	200.0000	✓✓✓	1/2
	A-02					B-05	1.00	10.00	200.00	✓✓✓	C-02	1.0001	10.0000	200.0000	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	9.99	199.99	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/9/03	A-01	Not in Use			✓	B-01	1.00	10.01	200.00	✓✓✓	C-01	1.0000	10.0000	200.0001	✓✓✓	1/2
	A-02					B-05	1.00	10.00	200.00	✓✓✓	C-02	1.0001	10.0001	199.9989	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	9.99	199.99	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					
5/12/03	A-01	Not in Use			✓	B-01	1.00	10.01	200.00	✓✓✓	C-01	1.0000	10.0000	200.0000	✓✓✓	1/2
	A-02					B-05	1.00	10.00	200.00	✓✓✓	C-05	1.0001	10.0001	199.9989	✓✓✓	
	A-03	1.00	9.99	200.00	✓	B-06	1.00	9.99	199.99	✓✓✓	C-					
	A-04					B-07	1.00	10.00	200.00	✓✓✓	C-					
	A-					B-					C-					

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

Alkalinity/OH/CO₃/HCO₃ (310.1/SM12320B) Worksheet

Batch # 0301287 Matrix: W Titrant H₂SO₄ Lot # 207900 Concentration (C) 0.02555 N₁ Test Date: 1/6/03 Analysis: W SOP: G-51

#	Sample ID	Dilution V ₁ /V ₂ =f ₁	Smpl Amt V ₁ , mL	H ₂ SO ₄ (mL) by Phln S _A	E _A	A	H ₂ SO ₄ (mL) by MR-BOG S _B	E _B	B	Phln-Alk., P	Tot. Alk., T (in unit of mgCaCO ₃ /L)	OH ⁻	CO ₃ ²⁻	HCO ₃ ⁻	Note & Anomaly
1	MB: T116	1 =	100			0			0		0	0	0	0	
2	445	1 =	100						7.90		100.9				
3	445	1 =	100						29.0		100.9				
4	3082-1	1 =	100			0			0		0	0	0	0	
5	-2	1 =	100			0			8.15		208.2	0	0	208.2	
6	-3	1 =	100			0.05			6.75	1.3	173.7	0	2.6	171.1	
7	-4	1 =	100			0			6.90		176.3	0	0	176.3	
8	44	1 =	100			0.40			3.15	10.2	100.9	0	20.4	80.5	
9	16	1 =	100			0.10			4.51	2.6	118.8	0	5.2	113.6	
10	3130-1	1 =	100			0			0		0	0	0	0	
11	-2	1 =	100			0			10.60		270.8	0	0	270.8	
12	-3	1 =	100			0.01			5.81	1.3	150.7	0	2.6	148.1	
13	-4	1 =	100			0.10			5.20	2.6	131.4	0	5.2	130.2	
14	-5	1 =	100			0.05			5.70	1.3	146.9	0	2.6	144.3	
15	-6	1 =	100			1.50			3.45	38.3	126.5	0	76.6	49.9	
16		1 =													
17		1 =													
18		1 =													
19		1 =													
20		1 =													
Dup.	3082-2	1 =	100			0			8.20		209.5	0	0	209.5	

Calculation:

$$A = S_A - E_A$$

$$B = S_B - E_B$$

$$P = 50,000 f_1 A C / V$$

$$T = 50,000 f_1 (A+B) C / V$$

APCL Form 5-101, Rev. 28, 1998 Ver 3.2
Using blue pen. Correcting by red pen.

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

Root-File: [CUST.DOC.WET]ALKROOT.TEX

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

Titration Results	OH ⁻ (CaCO ₃ mg/L)	CO ₃ ²⁻ (CaCO ₃ mg/L)	HCO ₃ ⁻ (CaCO ₃ mg/L)
P=0	0	0	T
P<T/2	0	2P	T-2P
P=T/2	0	2P	0
P>T/2	2P-T	2(T-P)	0
P=T	T	0	0

Applied P & Ch Laboratory

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

pH (150.1/9045B) Worksheet

043

Temperature compensation must be performed by the instrument automatically.

Analyst DC

SOP: G-44

Batch # 03W2790 Analysis Date: 5, 8, 03

Starting Time: 16:27 Ending Time: _____

Matrix ☒ Aqueous ☐ Soil

Batch # 03W2793 Analysis Date: 5, 8, 03

Starting Time: 18:05 Ending Time: _____

Matrix ☒ Aqueous ☐ Soil

Standard	4.00	7.00	10.00	Standard	4.00	7.00	10.00
Lot #		<u>030660-24</u>	<u>030659-24</u>	Lot #		<u>030660-24</u>	<u>030659-24</u>
Temperature °C		<u>23.8</u>	<u>23.8</u>	Temperature °C		<u>24.0</u>	<u>24.0</u>
pH Reading		<u>7.01</u>	<u>10.02</u>	pH Reading		<u>6.99</u>	<u>10.00</u>
T-corrected pH		<u>7.00</u>	<u>10.01</u>	T-corrected pH		<u>7.00</u>	<u>10.01</u>
Control Limit	±0.05 pH unit			Control Limit	±0.05 pH unit		

#	Sample ID	Pre-treat	pH	Note	#	Sample ID	Pre-treat	pH	Note
MB	<u>T1116</u>		<u>6.89</u>		MB	<u>T1116</u>		<u>6.92</u>	
1	<u>3130-1</u>		<u>7.31</u>		1	<u>3143-1</u>		<u>8.29</u>	
2	<u>-2</u>		<u>7.64</u>		2	<u>3141-1</u>		<u>6.89</u>	
3	<u>-3</u>		<u>8.21</u>		3	<u>-2</u>		<u>7.12</u>	
4	<u>-4</u>		<u>8.18</u>		4	<u>-3</u>		<u>6.79</u>	
5	<u>-5</u>		<u>8.07</u>		5	<u>-4</u>		<u>6.85</u>	
6	<u>-6</u>		<u>9.23</u>		6	<u>-5</u>		<u>6.80</u>	
7	<u>3132-2</u>		<u>9.08</u>		7	<u>-6</u>		<u>6.80</u>	
8	<u>3135-2</u>		<u>9.76</u>		8	<u>-7</u>		<u>6.97</u>	
9					9	<u>-8</u>		<u>7.00</u>	
10					10	<u>-9</u>		<u>6.78</u>	
11					11	<u>-10</u>		<u>6.97</u>	
12					12	<u>-11</u>		<u>7.87</u>	
13					13	<u>-12</u>		<u>7.54</u>	
14					14				
15					15				
16					16				
17					17				
18					18				
19					19				
20					20				
Dup.	<u>3130-1</u>		<u>7.36</u>		Dup.	<u>3141-12</u>		<u>7.57</u>	

Applied P & Ch Laboratory

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Chromium (VI) (7196) Worksheet

Batch # 03W2788 Matrix: W

[Holding Time: 24 hours!!]

Test Date: 5/8/02Analyst: h

Lot #: Reagent Water

Diphenylcazide solution

Test Time: 14:43

SOP: G-

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 = 2.000 + 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) $C = f_1 f_2 C'$	Anoma Note
CCV	Lot: W- <u>7853</u>	Expected Conc.: x	/ = <u>0.75</u> mg/L		<u>0.202</u>	<u>0.242</u> mg/L	REC. %	90-110
Method Blank	Bl. Lot: <u>T1116</u>		$/X_0 = 1$	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</u> ppm	
LCS1	Bl. Lot: <u>1</u>		$/X_0 =$	95.0/ =	<u>0.186</u>	mg/L	<u>0.222</u> ppm	
Sample-1	<u>3130-1</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>0.190</u>	mg/L	<u>0.227</u> ppm	
MSD on S-1	<u>2</u>		$/X_0 =$	95.0/ =	<u>0.183</u>	mg/L	<u>0.219</u> ppm	
Sample 2	<u>2</u>		$/X_0 =$	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</u> ppm	
Sample 3	<u>3</u>		$/X_0 =$	95.0/ =	<u>0.002</u>	mg/L	<u>0.002</u> ppm	
Sample 4	<u>4</u>		$/X_0 =$	95.0/ =	<u>0.001</u>	mg/L	<u>0.001</u> ppm	
Sample 5	<u>5</u>		$/X_0 =$	95.0/ =	<u>0.002</u>	mg/L	<u>0.002</u> ppm	
Sample 6	<u>6</u>		$/X_0 =$	95.0/ =	<u>0.000</u>	mg/L	<u>0.000</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>T1116</u>		$/X_0 = 1$	95.0/ =	<u>0.200</u>	mg/L	<u>0.239</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>0.213</u>		$/X_0 =$	95.0/ =	<u>0.213</u>	<u>0.255</u> mg/L	ppm	

Type	STD Lot #	$C_{STD}(\mu\text{g/mL}) \times V_{STD}(\text{mL}) / X(\text{g or mL}) = T$	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
MS	W- <u>7853</u>	x / = <u>0.75</u> ppm	%	80-120 %/80-120 %	PQL(w) 0.01
MSD	W- <u>1</u>	x / = ppm	%		PQL(s) 0.05
LCS	W- <u>7759</u>	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W- <u>1</u>	x / = ppm	%		MDL(s) 0.025

Chromium (VI) (7196) Worksheet

Batch # 02W1295 Matrix: W

[Holding Time: 24 hours!!]

Test Date: 1/24/03Analyst: BV

SOP: G-22

Lot #: Reagent Water

Diphenylcazide solution

Test Time:

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 / = 0.050 mg/L	0.001		C.C.=0.997 (> 0.995)	
STD-4	W-	x / = 0.15 mg/L	0.009		RSD= % (< 15%)	
STD-5	W-	x / = 0.250 mg/L	0.014		Ref. page	
STD-6	W-✓	x / = 0.50 mg/L	0.015		A=0.003+0.846C	

Analysis Type	Sample ID or Lot #	Samp. Amnt X ₀ (g or mL)	Dilu./Ext X/X ₀ =f ₁	Treat. Ratio V/X=f ₂	540 nm A	Concentration C'=A/RF	C (Sample) C=f ₁ f ₂ C'	Anomaly Note
CCV	Lot: W-7076	Expected Conc.: x	/ = 0.25 mg/L		0.246	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: 1		/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.	closing 0.25 mg/L		/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-✓	x / = ppm	%		PQL(s) 0.05
LCS	W-7191	x / = ppm	%	80-120 %/80-120 %	MDL(w) 0.005
LCSD	W-✓	x / = ppm	%		MDL(s) 0.025

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

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

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

Applied P & Ch Laboratory

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

Conductance (120.1) Worksheet

Test Date: 5/8/03

Analyst: w

Matrix: w

Calibration STD: 0.0100M KCl

uS/cm

SOP: G-45

#	Sample ID	Treatment $V/X=f_0$	Dilution $V_f/V_i=f_1$	Temperature $T, ^\circ\text{C}$	$C_{25}=C_T f_1/[1-0.0191(25-T)]$ $C_T, \mu\text{mhos/cm}$	$C_{25}, \mu\text{mhos/cm}$	$\rho=1/C_{25}$ $\text{M}\Omega/\text{cm}$	Note & Anomaly
Cal.	Lot #:	-	-	25°C				$C_{25}=1,413 \pm 20$
MB		/ =	/ =					for CM
1	3152-1	/ =	/ =					
2	-2	/ =	/ =					
3	-3	/ =	/ =					
4	-4	/ =	/ =					
5	-5	/ =	/ =					
6	-6	/ =	/ =					
7	3082-1	/ =	/ =					
8	-2	/ =	/ =					
9	-3	/ =	/ =					
10	-4	/ =	/ =					
11	-5	/ =	/ =					
12	✓ -6	/ =	/ =					
13	3130-1	/ =	/ =					
14	-2	/ =	/ =					
15	-3	/ =	/ =					
16	-4	/ =	/ =					
17	-5	/ =	/ =					
18	✓ -6	/ =	/ =					
19		/ =	/ =					
20		/ =	/ =					
Dup.		/ =	/ =					

13760 Magnolia Ave. Chino CA 91710

Solid Analysis (160.1, 160.2, 160.3) Worksheet

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

Batch # 03W2800 Matrix W Method: 160.1 Balance No. _____Date: 5/9/03 Analyst: DL

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	T1116	1 =	100	103.1172	103.1174	103.1175	0.0003	3	GL
2	LCS	T1116	1 =	100	112.2993	112.3385	112.3383	0.0390	390	K
3	Sample-1	3090-1	1 =	100	115.9188	115.9476	115.9478	0.0290	290	Y9
4	MS on S-1	3130-2	1 =	100	103.3079	103.4205	103.4107	0.1028	1028	L
5	MSD on S-1	-2	1 =	100	113.8619	113.9633	113.9631	0.1012	1012	6
6	Sample-2	3090-2	1 =	100	114.1383	114.1677	114.1679	0.0296	296	G3
7	Sample-3	-3	1 =	100	113.0683	113.1017	113.1019	0.0336	336	PR
8	Sample-4	-4	1 =	100	103.9417	103.9948	103.9949	0.0492	492	14
9	Sample-5	3090-1	1 =							
10	Sample-6	3130-1	1 =	100	113.9026	113.9028	113.9030	0.0004	4	P
11	Sample-7	-2	1 =	100	113.716	113.2364	113.2362	0.0646	646	25
12	Sample-8	-3	1 =	100	114.3837	114.4127	114.4124	0.0287	287	A
13	Sample-9	-4	1 =	100	115.8624	115.8910	115.8911	0.0287	287	30
14	Sample-10	-5	1 =	100	106.3250	106.3467	106.3469	0.0213	213	8
15	LCSD	T1116	1 =	100	115.3109	115.3493	115.3495	0.0386	386	38
16	Sample-11	3130-6	1 =	100	113.8234	113.8440	113.8442	0.0208	208	33
17	Sample-12	3086-1	1 =	100	104.3118	104.4449	104.4446	0.1028	1028	3
18	Sample-13		/ =							
19	Sample-14		/ =							
20	Sample-15		/ =							
21	Sample-16		/ =							
22	Sample-17		/ =							
23	Sample-18		/ =							
24	Sample-19		/ =							
25	Sample-20		/ =							
26	Mix Dup.		/ =							

Type	STD Lot #	$C_{STD}(\mu\text{g/mL}) \times V_{STD}(\text{mL}) / X(\text{g or mL}) = T$	Spike Rec.	Ctl Limit (W/S)	PQL/MDL (in ppm)
MS	W- 7618	x / = 400 ppm	%	85-115 %/80-120 %	PQL(w) 10
MSD	W- ↓	x / = ↓ ppm	%		PQL(s) 50
LCS	W- 7619	x / = ↓ ppm	%	90-110 %/85-115 %	MDL(w) 4
LCSD	W- ↓	x / = ↓ ppm	%		MDL(s) 20

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

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

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

Sample	Sample Type	Level	Method	Data File	Volume	Dilution
##03w2866kw ipc 25ppb w7759	Sample		e314-011.met	c:\data\03w2866kw2866k ipc 25ppb	1	1
ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2866kw2866k q01	1	1
ccb	Sample		e314-011.met	c:\data\03w2866kw2866k ccb01	1	1
lcs 25ppb w7827d	Sample		e314-011.met	c:\data\03w2866kw2866k l01	1	1
LCS 18PPB W7685D	Sample		e314-011.met	c:\data\03w2866kw2866k j01	1	1
ICCS 4ppb w7827b	Sample		e314-011.met	c:\data\03w2866kw2866k iccs 4ppb	1	1
mb	Sample		e314-011.met	c:\data\03w2866kw2866k k01	1	1
3177-01 F=1	Sample		e314-011.met	c:\data\03w2866k\3177-01	1	1
3178-01 f=1	Sample		e314-011.met	c:\data\03w2866k\3178-01	1	1
3130-01 F=1	Sample		e314-011.met	c:\data\03w2866k\3130-01	1	1
3130-02 F=1	Sample		e314-011.met	c:\data\03w2866k\3130-02	1	1
3130-03 f=1	Sample		e314-011.met	c:\data\03w2866k\3130-03	1	1
ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2866kw2866k q02	1	1
ccb	Sample		e314-011.met	c:\data\03w2866kw2866k k02	1	1
3130-04 f=1	Sample		e314-011.met	c:\data\03w2866k\3130-04	1	1
3130-05 F=1	Sample		e314-011.met	c:\data\03w2866k\3130-05	1	1
3130-06 F=1	Sample		e314-011.met	c:\data\03w2866k\3130-06	1	1
3130-02 MS 50PPB F=1	Sample		e314-011.met	c:\data\03w2866kw2866k m01	1	1
3130-02 MSD 50PPB F=1	Sample		e314-011.met	c:\data\03w2866kw2866k n01	1	1
3205-01 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-01	1	1
3205-02 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-02	1	1
3205-03 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-03	1	1
3205-04 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-04	1	1
ccv 50ppb w7827e	Sample		e314-011.met	c:\data\03w2866kw2866k q03	1	1
CCB	Sample		e314-011.met	c:\data\03w2866kw2866k k03	1	1
3205-05 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-05	1	1
3205-06 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-06	1	1
3205-07 F=1	Sample		e314-011.met	c:\data\03w2866k\3205-07	1	1
3178-01 f=1	Sample		e314-011.met	c:\data\03w2866k\3178-01	1	1
CCV 50PPB W7827E	Sample		e314-011.met	c:\data\03w2866kw2866k q04	1	1
	Sample		aastopcl.met		1	1

Analyst alex cy
Date 5/14/03
Instrument ZC-K

Line	Sample	Sample Type	Level	Method	Data File	Volume	Dilution
1	Cal blank	Sample		e314-011.met	c:\data\314-011\mb_001.dxd	1	1
2	cal standard 2ppb W7827a	Sample		e314-011.met	c:\data\314-011\std-2pb_002.dxd	1	1
3	cal standard 4ppb W7827b	Sample		e314-011.met	c:\data\314-011\std-4pb_003.dxd	1	1
4	cal standard 10ppb W7827c	Sample		e314-011.met	c:\data\314-011\std-10pb_004.dxd	1	1
5	cal standard 25ppb W7827d	Sample		e314-011.met	c:\data\314-011\std-25pb_005.dxd	1	1
6	cal standard 50ppb W7827e	Sample		e314-011.met	c:\data\314-011\std-50pb_006.dxd	1	1
7	cal standard 75ppb W7827f	Sample		e314-011.met	c:\data\314-011\std-75pb_007.dxd	1	1
8	cal standard 100ppb W7827g	Sample		e314-011.met	c:\data\314-011\std-100pb_008.dxd	1	1
9	ICV 50 ppb w7828a	Sample		e314-011.met	c:\data\314-011\vcv-50pb_009.dxd	1	1
0	icb	Sample		e314-011.met	c:\data\314-011\icb_010.dxd	1	1
1	anion 100pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-100_011.dxd	1	1
2	anion 200pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-200_012.dxd	1	1
3	anion 300pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-300_013.dxd	1	1
4	anion 400pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-400_014.dxd	1	1
5	anion 500pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-500_015.dxd	1	1
6	anion 600pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-600_016.dxd	1	1
7	anion 800pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-800_017.dxd	1	1
8	anion 1000pm each ,25pb CLO4	Sample		e314-011.met	c:\data\314-011\mct-1000_018.dxd	1	1
9	anion 400pm each 2pb	Sample		e314-011.met	c:\data\314-011\ipc-2pb_019.dxd	1	1
0	anion 400pm each 4pb	Sample		e314-011.met	c:\data\314-011\ipc-4pb_020.dxd	1	1
1	anion 400pm each 25pb	Sample		e314-011.met	c:\data\314-011\ipc-25pb_021.dxd	1	1
2	ICV 50 ppb	Sample		e314-011.met	c:\data\314-011\ccv-50pb	1	1
3	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-02_023.dxd	1	1
4	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-03_024.dxd	1	1
5	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-04	1	1
6	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-05	1	1
7	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-06	1	1
8	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-07	1	1
9	MDL 4pb	Sample		e314-011.met	c:\data\314-011\mdl-08	1	1
0	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
1	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
2	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
3	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
4	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
5	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
6	IDP and IDA 25pb	Sample		e314-011.met	c:\data\314-011\idap-25pb	1	1
7	MCT anion 800pm each, 25pbCLO4	Sample		e314-011.met	c:\data\314-011\lpo-25pb	1	1
8	MCT anion 800pm each, 25pbCLO4	Sample		e314-011.met	c:\data\314-011\lpo-25pb	1	1
9	MCT anion 800pm each, 4pbCLO4	Sample		e314-011.met	c:\data\314-011\lpc-4pb	1	1
0	MCT anion 800pm each, 4pbCLO4	Sample		e314-011.met	c:\data\314-011\lpc-4pb	1	1
1	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s01	1	5
2	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s02	1	5
3	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s03	1	5
4	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s04	1	5
5	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s05	1	5
6	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s06	1	5
7	MDL 20pb soil	Sample		e314-011.met	c:\data\314-011\mdl-s07	1	5
8	standard 25ppb W7827d	Sample		e314-011.met	c:\data\314-011\std-25pb	1	1
9	anion 100pm each,4pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-100-4pb	1	1
0	anion 200pm each ,4pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-200-4pb	1	1
1	anion 300pm each ,4pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-300-4pb	1	1
2	anion 100pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-100-2pb	1	1
3	anion 200pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-200-2pb	1	1
4	anion 300pm each,2pb CLO4	Sample		e314-011.met	c:\data\314-011\lam-300-2pb	1	1
5	1982-01 B S.C 4450us/cm	Sample		e314-011.met	c:\data\314-011\1982-01	1	1
6	1982-01 B S.C 4450us/cm	Sample		e314-011.met	c:\data\314-011\1982-01	1	2
7	1982-02 f=10	Sample		e314-011.met	c:\data\314-011\1982-02_057.dxd	1	10
8		Sample		aastopcl.met		1	1

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

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

Comment:

Analyst DRDate 3/21/03Instrument J

DIONEX SCHEDULE - C:\DX\SCHEDULE\03W2777.SCH

Inj#	Sample Name	Method	Data File	Vol.	Dil.	Int.Std.
1	##03W2777, W CCVW77	..\E300-063	..\W2777Q01.D01	1	1	1
2	MB RW1410	..\E300-063	..\W2777K01.D02	1	1	1
3	LCS W7768-100X	..\E300-063	..\W2777L01.D03	1	1	1
4	LCSD W7768-100X	..\E300-063	..\W2777J01.D04	1	1	1
5	3102-1 F=2.5	..\E300-063	..\3102-101.D05	1	2.5	1
6	3102-3 F=4	..\E300-063	..\3102-301.D06	1	4	1
7	3102-4 F=2.5	..\E300-063	..\3102-401.D07	1	2.5	1
8	3102-5 F=2.5	..\E300-063	..\3102-501.D08	1	2.5	1
9	3102-6 F=2.5	..\E300-063	..\3102-601.D09	1	2.5	1
10	3102-7 F=2.5	..\E300-063	..\3102-701.D10	1	2.5	1
11	3102-2 F=1.25	..\E300-063	..\3102-201.D11	1	1.25	1
12	CCV2W7767-100X	..\E300-063	..\W2777Q01.D12	1	1	1
13	MB RW1410	..\E300-063	..\W2777K11.D13	1	1	1
14	\$3130-2 MS F=25	..\E300-063	..\W2777M01.D14	1	25	1
15	\$3130-2 MSD F=25	..\E300-063	..\W2777N01.D15	1	25	1
16	3130-2 F=12.5	..\E300-063	..\3130-201.D16	1	12.5	1
17	3130-3 F=5	..\E300-063	..\3130-301.D17	1	5	1
18	3130-4 F=5	..\E300-063	..\3130-401.D18	1	5	1
19	3130-5 F=2	..\E300-063	..\3130-501.D19	1	2	1
20	3130-6 F=2	..\E300-063	..\3130-601.D20	1	2	1
21	3130-1 F=1.25	..\E300-063	..\3130-101.D21	1	1.25	1
22	CCV3W7767-100X	..\E300-063	..\W2777Q01.D22	1	1	1
23		..\STOP.MET		1	1	1

Comment:

LCS/LCSD LOT # W7768

MS/MSD LOT # W7767

ELUENT LOT # W7868

ANALYTICAL METHOD 9056/E300 MATRIX W

Analyst TM
Date 5/8/03
Instrument J



Applied Physics & Chemistry Laboratory

13760 Magnolia Ave. Chino CA 91710

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

June 6, 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 03-2987 and your project : 04-4428.10 JPL.
Enclosed please find:

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

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

Respectfully submitted,

Regina Kirakozova

Associate QA/QC Director

Applied P & Ch Laboratory

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Submitted to:

GEOFON, Inc.

Attention: Leo Williamson

22632 Golden Spring Dr Ste 270

Diamond Bar CA 91765

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

APCL Analytical Report

Service ID #: 801-032987

Collected by: Leo Williamson

Collected on: 04/30/03

Received: 04/30/03

Extracted: N/A

Tested: 04/30-05/13/03

Reported: 05/15/03

Sample Description: Water

Project Description: 04-4428.10 JPL

Analysis of Water Samples

Component Analyzed	Method	Unit	PQL	Analysis Result		
				EB-8-4/30/03 03-02987-1	MW-23-1 03-02987-2	MW-23-2 03-02987-3
BICARBONATE	SM2320B	mg/L	2	< 2	243	215
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	< 2	< 2	< 2
PH	9040B	pH unit	0.01	6.48	7.00	7.70
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	6.0J	765	606
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	2.9J	3.8J
Dilution Factor				1.25	20	10
CHLORIDE CL ⁻	300.0	mg/L	0.2	0.20J	118	96.5
NITRATE AS N	300.0	mg/L	0.04	0.067	13.8	12.5
SULFATE SO ₄ ²⁻	300.0	mg/L	0.5	< 0.63	183	124
Dilution Factor				1	1	1
ARSENIC	200.9	µg/L	5	< 5	< 5	< 5
CALCIUM	200.7	µg/L	200	< 200	147,000	106,000
IRON	200.7	µg/L	50	23.3J	447	49.2J
MAGNESIUM	200.7	µg/L	100	39.0J	50,300	39,900
POTASSIUM	200.7	µg/L	400	146J	3,030	2,950
SODIUM	200.7	µg/L	2000	< 2000	35,900	36,700
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
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
2-BUTANONE	524.2	µg/L	10	< 10	< 10	< 10
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-8-4/30/03 03-02987-1	MW-23-1 03-02987-2	MW-23-2 03-02987-3
4-CHLOROTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
1,2-DIBROMO-3-CHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5
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
METHYLENE CHLORIDE	524.2	µg/L	0.5	0.5J	0.5J	0.4J
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	4J	4J	3J
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	0.8	0.4J
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	1	0.6
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
1,1,2,2-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

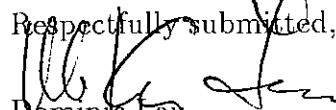
APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				MW-23-3 03-02987-4	MW-23-4 03-02987-5	MW-23-5 03-02987-6	TB-8-4/30/03 03-02987-7
BICARBONATE	SM2320B	mg/L	2	146	125	95.8	-
CARBONATE	SM2320B	mg-CaCO ₃ /L	2	< 2	5.2	92.0	-
PH	9040B	pH unit	0.01	7.88	8.28	9.55	-
SOLIDS, TOTAL DISSOLVED (TDS)	160.1	mg/L	10	284	220	263	-
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	< 4	< 4	< 4	-
Dilution Factor				4	2	2.5	1
CHLORIDE CL ⁻	300.0	mg/L	0.2	20.0	17.7	14.5	-
NITRATE N-NO ₃ ⁻ AS N	300.0	mg/L	0.04	7.2	6.5	0.14	-
SULFATE SO ₄ ⁻	300.0	mg/L	0.5	14.4	8.5	2.2	-
Dilution Factor				1	1	1	1
ARSENIC	200.9	µg/L	5	< 5	< 5	3.2J	-
CALCIUM	200.7	µg/L	200	40,700	27,400	5,270	-
IRON	200.7	µg/L	50	677	59.3	189	-
MAGNESIUM	200.7	µg/L	100	13,800	12,500	450	-
POTASSIUM	200.7	µg/L	400	1,760	1,990	2,530	-
SODIUM	200.7	µg/L	2000	27,000	30,500	97,700	-
VOLATILE ORGANIC COMPOUNDS							
Dilution Factor				1	1	1	1
BENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
BROMOBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
BROMOCHLOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
BROMODICHLOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
BROMOFORM	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
BROMOMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
N-BUTYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
SEC-BUTYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TERT-BUTYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
2-BUTANONE	524.2	µg/L	10	<10	<10	<10	<10
CARBON TETRACHLORIDE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CHLORODIBROMOMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CHLOROFORM	524.2	µg/L	0.5	<0.5	<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	0.5	<0.5	<0.5	<0.5	<0.5
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

APCL Analytical Report

Component Analyzed	Method	Unit	PQL	Analysis Result			
				MW-23-3 03-02987-4	MW-23-4 03-02987-5	MW-23-5 03-02987-6	TB-8-4/30/03 03-02987-7
DICHLORODIFLUOROMETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,2-DICHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
2,2-DICHLOROPROPANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
CIS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRANS-1,3-DICHLOROPROPENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
HEXACHLOROBUTADIENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
ISOPROPYLBENZENE (CUMENE)	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
P-ISOPROPYLTOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
METHYLENE CHLORIDE	524.2	µg/L	0.5	0.5J	0.5J	0.6	0.7
METHYL-T-BUTYL ETHER (MTBE)	524.2	µg/L	1	<1	<1	<1	<1
4-METHYL-2-PENTANONE (MIBK)	524.2	µg/L	10	3J	5J	3J	<10
NAPHTHALENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
N-PROPYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
STYRENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2,2-TETRACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TETRACHLOROETHENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TOLUENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,3-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRICHLOROBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,1-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,1,2-TRICHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
TRICHLOROETHENE	524.2	µg/L	0.5	<0.5	<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
1,1,2,3,4-PENTACHLORO-1,2,3,4,5-PENTACHLOROETHANE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,2,4-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
1,3,5-TRIMETHYLBENZENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
VINYL CHLORIDE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
O-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5
M/P-XYLENE	524.2	µg/L	0.5	<0.5	<0.5	<0.5	<0.5

PQL: Practical Quantitation Limit. MDL: Method Detection Limit. CRDL: Contract Required Detection Limit
 N.D.: Not Detected or less than the practical quantitation limit. "-": Analysis is not required.
 J: Reported between PQL and MDL.
 Listed Dilution Factors (DF) are relative to the method default DF. All unlisted DFs are 1.0

Respectfully submitted,

 Dominic Lau
 Laboratory Director
 Applied P & Ch Laboratory

Level C Data Package Deliverables

General Information

Project: 04-4428.10 JPL

APCL Service ID: 03-2987



Applied P & Ch Laboratory

13760 Magnolia Ave. Chino, CA 91710

Telephone (909)590-1828

Fax (909)590-1498

Case Narrative

Project: JPL/04-4428.10

For GEOFON, Inc.

APCL Service No: 03-2987

1. Sample Identification

The sample identifications are listed in the following table:

GEOFON, Inc. Sample ID	APCL Sample ID
MW-23-5	03-02987-6
MW-23-4	03-02987-5
MW-23-3	03-02987-4
MW-23-2	03-02987-3
MW-23-1	03-02987-2
TB-8-4/30/03	03-02987-7
EB-8-4/30/03	03-02987-1

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 (Bicarbonate, Carbonate),
9040B (pH),
160.1 (Solids, Total Dissolved (TDS)),
200.7/200.9 (Metals),

3. Holding Time

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

4. Preservation

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

5. Tele-log

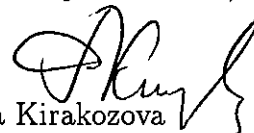
None

6. Anomaly

None

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

Respectfully submitted,


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



LABORATORY COPY

MMV-23 0027

MAIL REPORT (COMPANY NAME)

MAIL REPORT (COMPANY NAME)
WEOFFAY, INC.

RECIPIENT NAME
Leo W. Williams

ADDRESS
7632 Golden Springs Dr. #210

CITY, STATE AND ZIP CODE
Diamond Bar CA 91765

25

2

Comments	Na/K/Ca/As/Mg/Fe

SLMSPD

2002

TEMPERATURE UPON RECEIPT

Project Data Manager

Sample Receiving Checklist

APCL Service ID: **2987**Client Name/Project: Geolon JPL

1. Sample Arrival

Date/Time Received 4/30/03 1220Date/Time Opened 4/30/03 1220By (name): Kenneth Chan

Custody Transfer:

☐ Client☐ Golden State☐ UPS☐ US Mail☐ FedEx☒ APCL Empl: Adm

2. Chain-of-Custody (CoC)

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

on Hold _____

Received 7☒ 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☐ NoneTemp °C 4.5

(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: 30 Apr 2003

Time: 7:42 a.m.

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

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Sample Login: Check List

03-02987 (0470_ 138) (2202777_ 138)

04/30/03

Part 1: General Information

☐ Company Information	Name:	<i>GEOFON, Inc.</i>
	Address:	<i>22632 Golden Spring Dr Ste 270 ,Diamond Bar ,CA 91765</i>
☐ Project Information	Project Description:	<i>JPL</i>
	Project #:	<i>04-4428.10</i>
☐ Billing Information	P.O. #:	
	Bill Address:	<i>22632 Golden Spring Dr Ste 270 ,Diamond Bar ,CA 91765</i>
	Lab Project ID:	
	Client Database #:	<i>3</i>
☐ Receiving Information	Who Received Sample?	<i>Kenny Chan</i>
	Receiving Date/Time:	<i>04/30/03 1220</i>
	COC No.	
☐ Shipping Information	Shipping Company	<i>APCL pick up</i>
	Packing Information:	<i>Cooler/Ice Chester</i>
	Cooler Temperature:	<i>4.5 °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-23-5	VOC	03-02987-6- α	W	V	C	40	6	G	043003	N	0	7 ☐
	MW-23-5	Metal	03-02987-6- β	W	P	N	500	2	G	043003	N	0	7 ☐
	MW-23-5	300	03-02987-6- γ	W	P		1000	2	G	043003	N	0	7 ☐
2	MW-23-4	VOC	03-02987-5- α	W	V	C	40	3	G	043003	N	0	7 ☐
	MW-23-4	Metal	03-02987-5- β	W	P	N	500	1	G	043003	N	0	7 ☐
	MW-23-4	300	03-02987-5- γ	W	P		1000	1	G	043003	N	0	7 ☐
3	MW-23-3	VOC	03-02987-4- α	W	V	C	40	3	G	043003	N	0	7 ☐
	MW-23-3	Metal	03-02987-4- β	W	P	N	500	1	G	043003	N	0	7 ☐
	MW-23-3	300	03-02987-4- γ	W	P		1000	1	G	043003	N	0	7 ☐
4	MW-23-2	VOC	03-02987-3- α	W	V	C	40	3	G	043003	N	0	7 ☐
	MW-23-2	Metal	03-02987-3- β	W	P	N	500	1	G	043003	N	0	7 ☐
	MW-23-2	300	03-02987-3- γ	W	P		1000	1	G	043003	N	0	7 ☐
5	MW-23-1	VOC	03-02987-2- α	W	V	C	40	3	G	043003	N	0	7 ☐
	MW-23-1	Metal	03-02987-2- β	W	P	N	500	1	G	043003	N	0	7 ☐
	MW-23-1	300	03-02987-2- γ	W	P		1000	1	G	043003	N	0	7 ☐
6	TB-8-4/30/03	VOC	03-02987-7	W	V	C	40	2	G	043003	N	0	7 ☐
7	EB-8-4/30/03	VOC	03-02987-1- α	W	V	C	40	3	G	043003	N	0	7 ☐
	EB-8-4/30/03	Metal	03-02987-1- β	W	P	N	500	1	G	043003	N	0	7 ☐
	EB-8-4/30/03	300	03-02987-1- γ	W	P		1000	1	G	043003	N	0	7 ☐

Part 3: Analysis Information

Test Items:

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

☐ 206.2/7060A Arsenic, As, by GFAA

☐ 8270-SIM 1,4-Dioxane

Seq. #	Client's Sample ID (as given on COC)	Sample Sub-ID	APCL Sample ID	Matrix	524.2	CHROMIUM	PERCH	CL	SO4	NO3	CARBON	BICARB
1	MW-23-5	VOC	03-02987-6- α	W	X							☐
	MW-23-5	Metal	03-02987-6- β	W								☐
	MW-23-5	300	03-02987-6- γ	W		X	X	X	X	X	X	☐
2	MW-23-4	VOC	03-02987-5- α	W	X							☐
	MW-23-4	Metal	03-02987-5- β	W								☐
	MW-23-4	300	03-02987-5- γ	W		X	X	X	X	X	X	☐
3	MW-23-3	VOC	03-02987-4- α	W	X							☐
	MW-23-3	Metal	03-02987-4- β	W								☐
	MW-23-3	300	03-02987-4- γ	W		X	X	X	X	X	X	☐
4	MW-23-2	VOC	03-02987-3- α	W	X							☐
	MW-23-2	Metal	03-02987-3- β	W								☐
	MW-23-2	300	03-02987-3- γ	W		X	X	X	X	X	X	☐
5	MW-23-1	VOC	03-02987-2- α	W	X							☐
	MW-23-1	Metal	03-02987-2- β	W								☐
	MW-23-1	300	03-02987-2- γ	W		X	X	X	X	X	X	☐
6	TB-8-4/30/03	VOC	03-02987-7	W	X							☐
7	EB-8-4/30/03	VOC	03-02987-1- α	W	X							☐
	EB-8-4/30/03	Metal	03-02987-1- β	W								☐
	EB-8-4/30/03	300	03-02987-1- γ	W		X	X	X	X	X	X	☐

Seq. #	Client's Sample ID (as given on COC)	Sample Sub-ID	APCL Sample ID	Matrix	PH	TDS	NA	CA	K	MG	FE	AS
1	MW-23-5	VOC	03-02987-6- α	W								☐
	MW-23-5	Metal	03-02987-6- β	W			X	X	X	X	X	☐
	MW-23-5	300	03-02987-6- γ	W	X	X						☐
2	MW-23-4	VOC	03-02987-5- α	W								☐
	MW-23-4	Metal	03-02987-5- β	W			X	X	X	X	X	☐
	MW-23-4	300	03-02987-5- γ	W	X	X						☐
3	MW-23-3	VOC	03-02987-4- α	W								☐
	MW-23-3	Metal	03-02987-4- β	W			X	X	X	X	X	☐
	MW-23-3	300	03-02987-4- γ	W	X	X						☐
4	MW-23-2	VOC	03-02987-3- α	W								☐
	MW-23-2	Metal	03-02987-3- β	W			X	X	X	X	X	☐
	MW-23-2	300	03-02987-3- γ	W	X	X						☐
5	MW-23-1	VOC	03-02987-2- α	W								☐
	MW-23-1	Metal	03-02987-2- β	W			X	X	X	X	X	☐
	MW-23-1	300	03-02987-2- γ	W	X	X						☐
6	TB-8-4/30/03	VOC	03-02987-7	W								☐
7	EB-8-4/30/03	VOC	03-02987-1- α	W								☐
	EB-8-4/30/03	Metal	03-02987-1- β	W			X	X	X	X	X	☐

EB-8-4/30/03

300

03-02987-1- γ .

W

X

X

☐

Seq. #	Client's Sample ID (as given on COC)	Sample Sub-ID	APCL Sample ID	Matrix	DIOXANE
1	MW-23-5	VOC	03-02987-6- α	W	☐
	MW-23-5	Metal	03-02987-6- β	W	☐
	MW-23-5	300	03-02987-6- γ	W	☐
2	MW-23-4	VOC	03-02987-5- α	W	☐
	MW-23-4	Metal	03-02987-5- β	W	☐
	MW-23-4	300	03-02987-5- γ	W	☐
3	MW-23-3	VOC	03-02987-4- α	W	☐
	MW-23-3	Metal	03-02987-4- β	W	☐
	MW-23-3	300	03-02987-4- γ	W	☐
4	MW-23-2	VOC	03-02987-3- α	W	☐
	MW-23-2	Metal	03-02987-3- β	W	☐
	MW-23-2	300	03-02987-3- γ	W	☐
5	MW-23-1	VOC	03-02987-2- α	W	☐
	MW-23-1	Metal	03-02987-2- β	W	☐
	MW-23-1	300	03-02987-2- γ	W	☐
6	TB-8-4/30/03	VOC	03-02987-7	W	☐
7	EB-8-4/30/03	VOC	03-02987-1- α	W	☐
	EB-8-4/30/03	Metal	03-02987-1- β	W	☐
	EB-8-4/30/03	300	03-02987-1- γ	W	☐

☐ Client's Requirement:

RUN MS/MSD ON SAMPLE #6 ~

Login By En-Yu Paul KouCheck By 16X

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: 05/13/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: 03G2456-MB-01	Lab Sample ID: 03G2456-MB-01	Received Date: 05/13/2003
Sample Type: Method Blank	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: G2456K01	Prep. No: -	Anal. Time: 15:55
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	BENZENE	71-43-2	µg/L	0.5	<0.5	U
2	BROMOBENZENE	108-86-1	µg/L	0.5	<0.5	U
3	BROMOCHLOROMETHANE	74-97-5	µg/L	0.5	<0.5	U
4	BROMODICHLOROMETHANE	75-27-4	µg/L	0.5	<0.5	U
5	BROMOFORM	75-25-2	µg/L	0.5	<0.5	U
6	BROMOMETHANE	74-83-9	µg/L	0.5	<0.5	U
7	N-BUTYLBENZENE	104-51-8	µg/L	0.5	<0.5	U
8	SEC-BUTYLBENZENE	135-98-8	µg/L	0.5	<0.5	U
9	TERT-BUTYLBENZENE	98-06-6	µg/L	0.5	<0.5	U
10	2-BUTANONE	78-93-3	µg/L	10	3	J
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	CHLROMETHANE	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	0.5	<0.5	U
20	1,2-DIBROMOETHANE (EDB)	106-93-4	µg/L	0.5	<0.5	U
21	DIBROMOMETHANE	74-95-3	µg/L	0.5	<0.5	U
22	1,2-DICHLOROBENZENE	95-50-1	µg/L	0.5	<0.5	U
23	1,3-DICHLOROBENZENE	541-73-1	µg/L	0.5	<0.5	U
24	1,4-DICHLOROBENZENE	106-46-7	µg/L	0.5	<0.5	U
25	DICHLORODIFLUOROMETHANE	75-71-8	µg/L	0.5	<0.5	U
26	1,1-DICHLOROETHANE	75-34-3	µg/L	0.5	<0.5	U
27	1,2-DICHLOROETHANE	107-06-2	µg/L	0.5	<0.5	U
28	1,1-DICHLOROETHENE	75-35-4	µg/L	0.5	<0.5	U
29	CIS-1,2-DICHLOROETHENE	156-59-2	µg/L	0.5	<0.5	U
30	TRANS-1,2-DICHLOROETHENE	156-60-5	µg/L	0.5	<0.5	U
31	1,2-DICHLOROPROPANE	78-87-5	µg/L	0.5	<0.5	U
32	1,3-DICHLOROPROPANE	142-28-9	µg/L	0.5	<0.5	U
33	2,2-DICHLOROPROPANE	594-20-7	µg/L	0.5	<0.5	U
34	1,1-DICHLOROPROPENE	563-58-6	µg/L	0.5	<0.5	U
35	CIS-1,3-DICHLOROPROPENE	10061-01-5	µg/L	0.5	<0.5	U
36	TRANS-1,3-DICHLOROPROPENE	10061-02-6	µg/L	0.5	<0.5	U
37	ETHYLBENZENE	100-41-4	µg/L	0.5	<0.5	U
38	HEXACHLOROBUTADIENE	87-68-3	µg/L	0.5	<0.5	U
39	ISOPROPYLBENZENE (CUMENE)	98-82-8	µg/L	0.5	<0.5	U

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	4.5	
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U

Surrogates

Control Limit, %

Surro. Rec.%

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

70-129

102

70-129

97

70-122

101

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

94

50-200

88

50-200

90

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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: EB-8-4/30/03	Lab Sample ID: 03-2987-1	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-01	Prep. No: -	Anal. Time: 16:49
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.5	J
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	4	J
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

101

70-129

103

70-122

105

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

96

50-200

92

50-200

90

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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: MW-23-1	Lab Sample ID: 03-2987-2	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-02	Prep. No: -	Anal. Time: 17:43
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.5	J
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	4	J
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	0.8	
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	1	
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

102

70-129

102

70-122

104

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

94

50-200

89

50-200

88

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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: MW-23-2	Lab Sample ID: 03-2987-3	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-03	Prep. No: -	Anal. Time: 18:10
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.4	J
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	3	J
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	0.4	J
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	0.6	
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U
Surrogates				Control Limit, %	Surro. Rec.%	
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	460-00-4		70-129	101	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	101	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	104	
4	TOLUENE-D8	2037-26-5		73-129	103	
# 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	87	
3	FLUOROBENZENE	462-06-6		50-200	87	
# 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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: MW-23-3	Lab Sample ID: 03-2987-4	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-04	Prep. No: -	Anal. Time: 18:37
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

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

Surrogates

		Control Limit, %	Surro. Rec. %
1	1-BROMO-4-FLUOROBENZENE (4-BROMOFL)	70-129	101
2	1,2-DICHLOROETHANE-D4	70-129	100
3	DIBROMOFLUOROMETHANE	70-122	103
4	TOLUENE-D8	73-129	104
# of out-of-control			0

Internal Standard

		Control Limit, %	IS Rec. %
1	CHLOROBENZENE-D5	50-200	91
2	1,4-DICHLOROBENZENE-D4	50-200	86
3	FLUOROBENZENE	50-200	86
# 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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: MW-23-4	Lab Sample ID: 03-2987-5	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-05	Prep. No: -	Anal. Time: 19:04
Methanol Vol. -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.5	J
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	5	J
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	102	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	102	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	104	
4	TOLUENE-D8	2037-26-5		73-129	105	
# of out-of-control					0	
Internal Standard				Control Limit, %	IS Rec.%	
1	CHLOROBENZENE-D5	3114-55-4		50-200	88	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	83	
3	FLUOROBENZENE	462-06-6		50-200	84	
# 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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: MW-23-5	Lab Sample ID: 03-2987-6	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-06	Prep. No: -	Anal. Time: 17:16
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	< 0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.6	B
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	< 1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	3	J
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	101	
2	1,2-DICHLOROETHANE-D4	17060-07-0		70-129	99	
3	DIBROMOFLUOROMETHANE	1868-53-7		70-122	104	
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	91	
2	1,4-DICHLOROBENZENE-D4	3855-82-1		50-200	86	
3	FLUOROBENZENE	462-06-6		50-200	87	
# 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: 04/30/2003
Project ID: JPL	Service ID: 32987	Collected by:
Sample ID: TB-8-4/30/03	Lab Sample ID: 03-2987-7	Received Date: 04/30/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 524.2	Prep. Method: 5030	Instrument ID: GC/MS: A
Batch No: 03G2456	Prep. Date: 05/13/03	Anal. Date: 05/13/03
Data File Name: 2987-07	Prep. No: -	Anal. Time: 19:31
Methanol Vol: -	Sample Amount: 25.0 mL	Dilution Factor: 1
Test Level: Low	Sparge Size: 25 mL	Heated Purge: (Y/N) N

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

#	Component Name	CAS No	Unit	RL	Result	Qualifier
40	P-ISOPROPYLTOLUENE	99-87-6	µg/L	0.5	<0.5	U
41	METHYLENE CHLORIDE	75-09-2	µg/L	0.5	0.7	B
42	METHYL-T-BUTYL ETHER (MTBE)	1634-04-4	µg/L	1	<1	U
43	4-METHYL-2-PENTANONE (MIBK)	108-10-1	µg/L	10	<10	U
44	NAPHTHALENE	91-20-3	µg/L	0.5	<0.5	U
45	N-PROPYLBENZENE	103-65-1	µg/L	0.5	<0.5	U
46	STYRENE	100-42-5	µg/L	0.5	<0.5	U
47	1,1,1,2-TETRACHLOROETHANE	630-20-6	µg/L	0.5	<0.5	U
48	1,1,2,2-TETRACHLOROETHANE	79-34-5	µg/L	0.5	<0.5	U
49	TETRACHLOROETHENE	127-18-4	µg/L	0.5	<0.5	U
50	TOLUENE	108-88-3	µg/L	0.5	<0.5	U
51	1,2,3-TRICHLOROBENZENE	87-61-6	µg/L	0.5	<0.5	U
52	1,2,4-TRICHLOROBENZENE	120-82-1	µg/L	0.5	<0.5	U
53	1,1,1-TRICHLOROETHANE	71-55-6	µg/L	0.5	<0.5	U
54	1,1,2-TRICHLOROETHANE	79-00-5	µg/L	0.5	<0.5	U
55	TRICHLOROETHENE	79-01-6	µg/L	0.5	<0.5	U
56	TRICHLOROFLUOROMETHANE	75-69-4	µg/L	0.5	<0.5	U
57	1,2,3-TRICHLOROPROPANE	96-18-4	µg/L	0.5	<0.5	U
58	1,1,2,2-TETRACHLOROETHANE	76-13-1	µg/L	0.5	<0.5	U
59	1,2,4-TRIMETHYLBENZENE	95-63-6	µg/L	0.5	<0.5	U
60	1,3,5-TRIMETHYLBENZENE	108-67-8	µg/L	0.5	<0.5	U
61	VINYL CHLORIDE	75-01-4	µg/L	0.5	<0.5	U
62	O-XYLENE	95-47-6	µg/L	0.5	<0.5	U
63	M/P-XYLENE	108-38-3	µg/L	0.5	<0.5	U

Surrogates

Control Limit, %

Surro. Rec. %

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

70-129

101

70-129

103

70-122

105

73-129

102

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

90

50-200

85

50-200

84

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

FORM-2A

Applied P & Ch Laboratory

Surrogate Recovery Summary for Method 524.2

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

SDG Number: 032987

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2456

#	Client Sample No	Lab Sample ID	S1 % #	S2 % #	S3 % #	S4 % #	TOT OUT
1	03G2456-LCS-01	03G2456-LCS-01	95	89	97	100	0
2	MW-23-5MS	03-2987-6MS	92	84	91	96	0
3	MW-23-5MSD	03-2987-6MSD	95	92	98	99	0
4	03G2456-MB-01	03G2456-MB-01	102	97	101	103	0
5	EB-8-4/30/03	03-2987-1	101	103	105	101	0
6	MW-23-5	03-2987-6	101	99	104	104	0
7	MW-23-1	03-2987-2	102	102	104	103	0
8	MW-23-2	03-2987-3	101	101	104	103	0
9	MW-23-3	03-2987-4	101	100	103	104	0
10	MW-23-4	03-2987-5	102	102	104	105	0
11	TB-8-4/30/03	03-2987-7	101	103	105	102	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.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32987

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2456

LCS Filename: G2456L01

Date Analyzed: 051303

Time Analyzed: 13:41

LCSD Filename: -

Date Analyzed: -

Time Analyzed: -

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
BENZENE	µg/L	20	0	19.9	100	65-120
CHLOROBENZENE	µg/L	20	0	19.9	100	65-134
1,1-DICHLOROETHENE	µg/L	20	0	17.9	90	65-127
TOLUENE	µg/L	20	0	20.1	101	65-134
TRICHLOROETHENE	µg/L	20	0	20.4	102	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.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32987

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2456

MS Filename: G2456M01

Date Analyzed: 051303

Time Analyzed: 14:08

MSD Filename: G2456N01

Date Analyzed: 051303

Time Analyzed: 14:35

MS Sample No: MW-23-5

Sample Lab ID: 03-2987-6

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
BENZENE	µg/L	20	0	18.8	94	65-121
CHLOROBENZENE	µg/L	20	0	19.0	95	65-134
1,1-DICHLOROETHENE	µg/L	20	0	17.3	87	65-127
TOLUENE	µg/L	20	0	19.3	97	65-134
TRICHLOROETHENE	µg/L	20	0	19.5	98	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	19.6	98	4	28	65-121
CHLOROBENZENE	µg/L	20	19.8	99	4	35	65-134
1,1-DICHLOROETHENE	µg/L	20	17.9	90	3	31	65-127
TOLUENE	µg/L	20	19.8	99	2	35	65-134
TRICHLOROETHENE	µg/L	20	20.4	102	4	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: 32987

Project ID: JPL

Project No: 04-4428.10

Analysis Date: 05/13/03

Sample Matrix: Water

Analysis Time: 15:55

Sample ID: 03G2456-MB-01

Batch No: 03G2456

Instrument ID: GC/MS: A

Lab Sample ID: 03G2456-MB-01

Data File Name: G2456K01

GC Column: HP-VOC

Heated Purge: (Y/N) N

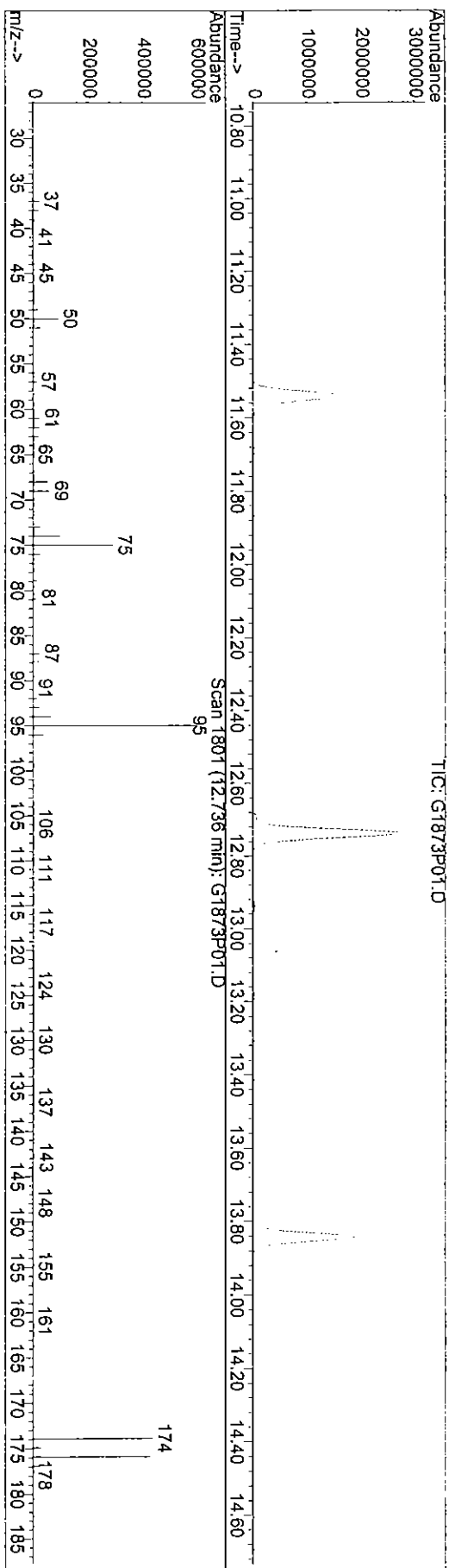
Column ID: 0.20 mm

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	03G2456-LCS-01	03G2456-LCS-01	Lab Control Spike	G2456L01	05/13/03	13:41
2	MW-23-5MS	03-2987-6MS	Matrix Spike	G2456M01	05/13/03	14:08
3	MW-23-5MSD	03-2987-6MSD	Matrix Spike Duplicate	G2456N01	05/13/03	14:35
4	EB-8-4/30/03	03-2987-1	Field Sample	2987-01	05/13/03	16:49
5	MW-23-5	03-2987-6	Field Sample	2987-06	05/13/03	17:16
6	MW-23-1	03-2987-2	Field Sample	2987-02	05/13/03	17:43
7	MW-23-2	03-2987-3	Field Sample	2987-03	05/13/03	18:10
8	MW-23-3	03-2987-4	Field Sample	2987-04	05/13/03	18:37
9	MW-23-4	03-2987-5	Field Sample	2987-05	05/13/03	19:04
10	TB-8-4/30/03	03-2987-7	Field Sample	2987-07	05/13/03	19:31
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						
23						
24						
25						

Data File : C:\MSDCHEM\1\DATA\03G1873\G1873P01.D
 Acq On : 30 Mar 2003 2:29 pm
 Sample : ##03g1873,w
 Misc :
 MS Integration Params: Lscint.p
 Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P &ch Lab** EPA 8260

Vial: 1
 Operator: zou
 Inst : GCMS-A
 Multiplr: 1.00



Spectrum Information: Scan 1801

Target Mass	Rel. Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
50	95	15	40	15.2	92424	PASS
75	95	30	60	48.4	293504	PASS
95	95	100	100	100.0	606592	PASS
96	95	5	9	6.6	39960	PASS
173	174	0.00	2	0.0	0	PASS
174	95	50	100	72.6	440576	PASS
175	174	5	9	7.4	32632	PASS
176	174	95	101	98.1	432384	PASS
177	176	5	9	6.7	29056	PASS

5A
VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name : APPLIED P & CH LAB Contract: _____
 Lab Code: _____ Case No.: _____ SAS No.: _____ SDG No.: 03-2987
 Lab File ID: G1873P01 BFB Injection Date: 3/30/03
 Instrument ID: GCMS-A BFB Injection Time: 1429
 GC Column: HP-VOC ID: 0.20 (mm) Heated Purge: (Y/N) N

m/e	ION ABUNDANCE CRITERIA	%RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	15.2
75	30.0 - 60.0% of mass 95	48.4
95	Base peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.6
173	Less than 2.0% of mass 174	0.0 (0.0)1
174	50.0 - 100.0% of mass 95	72.6
175	5.0 - 9.0% of mass 174	5.4 (7.4)1
176	95.0 - 101.0% of mass 174	71.3 (98.1)1
177	5.0 - 9.0% of mass 176	4.8 (6.7)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	VSTD0003	10-0003	10-00003.D	3/30/03	1550
02	VSTD002	10-0002	10-0002.D	3/30/03	1643
03	VSTD010	10-0010	10-0010.D	3/30/03	1737
04	VSTD020	10-0020	10-0020.D	3/30/03	1830
05	VSTD040	10-0040	10-0040.D	3/30/03	1924
06	VSTD060	10-0060	10-0060.D	3/30/03	2017
07					
08					
09					
10					
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14					
15					
16					
17					
18					
19					
20					
21					
22					

Response Factor Report GCMS-A

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
Title : **Applied P & Ch Lab** EPA 8260
Last Update : Mon Mar 31 10:25:55 2003
Response via : Initial Calibration

Calibration Files
.3 =10-0003A.D 2 =10-0002.D 10 =10-0010.D
20 =10-0020.D 40 =10-0040.D 60 =10-0060.D

Compound	.3	2	10	20	40	60	Avg	%RSD
-----ISTD-----								
1) I 1 Fluorobenzene	0.258	0.233	0.242	0.252	0.272	0.235	0.249	6.08
2) 3 di-Cl-di-F-m	0.339	0.254	0.236	0.247	0.236	0.224	0.256	16.32 0.997
3) P 4 Chloromethan	0.122	0.119	0.120	0.134	0.130	0.116	0.124	5.77
4) 2 F114	0.249	0.237	0.241	0.256	0.266	0.244	0.249	4.29
5) C 5 vinyl chlori	0.185	0.163	0.141	0.134	0.139	0.137	0.150	13.41
6) 6 bromomethane	0.238	0.150	0.151	0.162	0.163	0.154	0.170	20.13 0.998
7) 7 chloroethane	0.387	0.355	0.362	0.395	0.397	0.347	0.374	5.80
8) 8 tri-Cl-F-met	0.030	0.029	0.035	0.035	0.035	0.030	0.032	9.40
9) 91 Acetonitrile	0.025	0.023	0.024	0.024	0.024	0.023	0.024	3.26
10) 9 acrolein	0.037	0.023	0.023	0.023	0.021	0.021	0.025	27.74 0.999
11) 11 acetone X	0.130	0.126	0.124	0.133	0.128	0.123	0.127	3.13
12) 12 ethyl ether	0.318	0.295	0.295	0.321	0.317	0.288	0.306	4.78#
13) M, C13 11-dichloroe	0.079	0.078	0.124	0.222	0.252	0.244	0.166	49.28 0.994
14) 14 Iodomethane	0.230	0.214	0.214	0.239	0.238	0.204	0.223	6.41
15) 15 F-113	0.034	0.033	0.033	0.030	0.034	0.033	0.033	4.16
16) 16 acrylonitril	1.009	0.823	0.821	0.906	0.913	0.826	0.883	8.47
17) 17 carbon disul	0.009	0.003	0.001	0.001	0.001	0.003	0.003	113.71 0.995
18) 94 Isopropyl Al	0.432	0.236	0.219	0.225	0.229	0.223	0.260	32.37 0.999
19) 18 methylene ch	0.264	0.237	0.244	0.255	0.272	0.257	0.255	4.99
20) 19 t-12-di-Cl-e	0.448	0.431	0.450	0.472	0.497	0.487	0.464	5.52
21) 20 t-Bu-Me-ethe	0.005	0.002	0.004	0.004	0.005	0.003	0.004	33.63
22) 95 Tert butyl a	0.299	0.287	0.348	0.348	0.351	0.303	0.317	9.40
23) 94 allyl chlori	0.443	0.403	0.408	0.418	0.443	0.427	0.424	4.06
24) P 21 11-dichloroe	0.012	0.012	0.010	0.010	0.013	0.012	0.012	10.29
25) 97 propionitril	0.260	0.233	0.243	0.255	0.269	0.263	0.254	5.32
26) 22 c-12-di-Cl-e	0.312	0.265	0.290	0.322	0.339	0.320	0.308	8.54
27) 23 22-Dichlorop	0.096	0.098	0.098	0.101	0.105	0.103	0.100	3.22
28) 24 Br-Cl-methan								

(#) = Out of Range
826WA010.M

Mon Mar 31 10:32:53 2003

Response Factor Report GCMS-A

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P & Ch Lab** EPA 8260
 Last Update : Mon Mar 31 10:25:55 2003
 Response via : Initial Calibration

Calibration Files
 .3 =10-0003A.D 2 =10-0002.D 10 =10-0010.D
 20 =10-0020.D 40 =10-0040.D 60 =10-0060.D

Compound	.3	2	10	20	40	60	Avg	%RSD
29) C 25 chloroform		0.417	0.419	0.428	0.442	0.429	0.427	2.30
30) 26 tetrahydrofu	0.030	0.027	0.028	0.024	0.028	0.028	0.027	7.27
31) 98 Diisopropyl	0.675	0.662	0.692	0.667	0.728	0.707	0.688	3.70
32) S 27 Di-Br-F-Me (	0.263	0.216	0.216	0.227	0.232	0.226	0.230	7.61
33) 99 ETBE	0.484	0.471	0.516	0.535	0.583	0.584	0.529	9.11
34) S 29 1,2-Di-Cl-Et	0.258	0.203	0.200	0.209	0.208	0.203	0.214	10.24
35) 30 12-dichloroe	0.084	0.078	0.076	0.080	0.078	0.075	0.079	4.09
36) 32 vinyl acetat	0.304	0.296	0.313	0.295	0.332	0.315	0.309	4.47
37) 92 Nitro Methan		0.035	0.035	0.036	0.039	0.038	0.037	4.34
38) 33 2-butanoneME		0.039	0.027	0.028	0.032	0.031	0.032	14.75
39) 93 Ethyl Acetat		0.123	0.141	0.105	0.150	0.133	0.130	13.15
40) 34 111-trichlor	0.392	0.351	0.363	0.384	0.413	0.389	0.382	5.76
41) 35 11-Di-Cl-pro	0.330	0.289	0.312	0.337	0.365	0.336	0.328	7.84
42) M 36 benzene	1.005	0.924	0.978	1.055	1.057	1.017	1.006	5.00
43) 37 CCl4	0.324	0.282	0.308	0.324	0.363	0.335	0.323	8.40
44) 100 Isobutyl al		0.002	0.001	0.002	0.002	0.002	0.002	30.05
45) 38 thiophene	0.482	0.458	0.496	0.537	0.546	0.535	0.509	7.01
46) C 39 12-di-Cl-pro	0.215	0.196	0.209	0.227	0.226	0.221	0.216	5.42#
47) M 40 trichloroeth	0.281	0.249	0.258	0.285	0.295	0.280	0.275	6.33
48) 41 dibromometha	0.102	0.102	0.102	0.107	0.108	0.105	0.104	2.67
49) 101 TAME	0.410	0.403	0.454	0.511	0.530	0.533	0.474	12.47
50) 42 Br-di-Cl-met		0.278	0.288	0.308	0.309	0.304	0.297	4.56
51) 43 Me-methacryl	0.085	0.087	0.099	0.107	0.116	0.113	0.101	12.85
52) 44 2-ClEt-Vi-et		0.015	0.019	0.024	0.027	0.027	0.023	24.02
53) 45 C-13-di-Cl-p	0.286	0.278	0.309	0.342	0.351	0.346	0.319	10.09
54) 46 t-1,3-dichlo	0.200	0.205	0.239	0.267	0.287	0.280	0.246	15.20
55) I 47 Chlorobezene-d5								
56) 48 112-tri-Cl-E	0.222	0.199	0.200	0.208	0.214	0.206	0.208	4.27

0.998
0.999

(#) = Out of Range
 826WA010.M

Mon Mar 31 10:32:54 2003

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 Title : **Applied P & Ch Lab** EPA 8260
 Last Update : Mon Mar 31 10:25:55 2003
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Calibration Files

.3 =10-0003A.D 2 =10-0002.D 10 =10-0010.D
 20 =10-0020.D 40 =10-0040.D 60 =10-0060.D

Compound	.3	2	10	20	40	60	Avg	%RSD
57) 49 13-di-Cl-pro	0.374	0.361	0.358	0.372	0.382	0.362	0.368	2.58
58) 50 Et methacryl	0.226	0.251	0.288	0.304	0.346	0.334	0.291	16.07
59) 51 di-Br-Cl-met	0.234	0.247	0.256	0.276	0.267	0.256	0.256	6.45
60) P 52 bromoform	0.146	0.119	0.128	0.134	0.148	0.143	0.136	8.58
61) 53 1,4-dichloro	0.320	0.326	0.335	0.351	0.331	0.333	0.333	3.55
62) 54 MIBK	0.132	0.133	0.132	0.158	0.151	0.141	0.141	8.61
63) S 55 toluene-d8	1.172	1.220	1.322	1.363	1.290	1.273	1.273	6.06
64) M,C 56 toluene	1.483	1.381	1.458	1.571	1.597	1.490	1.497	5.23
65) 57 2-hexanone X	0.089	0.091	0.094	0.101	0.095	0.094	0.094	4.99
66) 58 12-dibromoet	0.185	0.188	0.190	0.198	0.209	0.201	0.195	4.69
67) 59 tetra-Cl-eth	0.404	0.360	0.360	0.395	0.415	0.376	0.385	6.08
68) M,P 60 chlorobenzen	1.025	0.902	0.921	0.980	0.998	0.958	0.964	4.84
69) 61 1112-tetra-C	0.282	0.281	0.298	0.322	0.337	0.329	0.308	7.99
70) I 62 1,4-Dichlorobenzen	0.403	0.341	0.366	0.424	0.462	0.419	0.402	10.81
71) 63 1-chlorohexa	3.443	3.192	3.357	3.680	3.841	3.609	3.520	6.68#
72) C 64 Et-Bz	2.731	2.487	2.612	2.843	2.884	2.654	2.702	5.50
73) 65 m/p-Xylenes	1.866	1.872	2.002	2.189	2.278	2.176	2.064	8.50
74) 66 styrene	2.643	2.489	2.693	2.927	3.014	2.852	2.770	7.08
75) 67 o-xylene	0.507	0.479	0.460	0.478	0.510	0.482	0.486	3.89
76) P 68 1122-Tetra-C	0.160	0.134	0.134	0.137	0.145	0.138	0.141	7.08
77) 69 123-tri-Cl-P	0.718	0.718	0.718	0.774	0.810	0.785	0.761	5.45
78) S 70 4-Br-1-F-Bz	3.250	2.950	3.272	3.652	3.847	3.577	3.425	9.53
79) 71 isopropylben	0.751	0.698	0.726	0.785	0.825	0.793	0.763	6.13
80) 72 bromobenzene	0.050	0.053	0.065	0.075	0.092	0.092	0.071	25.62
81) 92 t-1,4-dichlo	0.993	0.867	0.933	1.055	1.134	1.069	1.009	9.66
82) 73 n-propylbenz	0.851	0.749	0.802	0.880	0.934	0.897	0.852	7.89
83) 74 2-Cl-Toluene	0.910	0.783	0.817	0.890	0.944	0.899	0.874	6.98
84) 75 4-Cl-Toluene								

0.994

(#) = Out of Range
 826WA010.M

Mon Mar 31 10:32:54 2003

Response Factor Report GCMS-A

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P & Ch Lab** EPA 8260
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Calibration Files
 .3 =10-0003A.D 2 =10-0002.D 10 =10-0010.D
 20 =10-0020.D 40 =10-0040.D 60 =10-0060.D

Compound	.3	2	10	20	40	60	Avg	%RSD
85) 76 135-tri-Me-B	2.886	2.570	2.825	3.101	3.252	3.022	2.943	8.09
86) 77 4-iso-Pr-tol	3.039	2.843	3.056	3.311	3.633	3.321	3.201	8.70
87) 78 124-tri-Me-B	3.100	2.689	2.842	3.139	3.264	3.085	3.020	7.03
88) 79 tert-butylbe	2.468	2.285	2.505	2.811	2.989	2.782	2.640	9.95
89) 80 13-DCB	1.801	1.498	1.518	1.628	1.720	1.640	1.634	7.10
90) 81 sec-butylben	3.686	3.443	3.710	4.114	4.369	3.965	3.881	8.61
91) 82 14-DCB	1.902	1.503	1.501	1.597	1.707	1.630	1.640	9.17
92) 83 Cl-benzyl	0.068	0.087	0.115	0.123	0.165	0.162	0.120	32.61
93) 84 12-DCB	1.551	1.321	1.324	1.365	1.479	1.417	1.409	6.51
94) 85 n-butylbenze	0.751	0.714	0.798	0.847	0.984	0.919	0.835	12.24
95) 86 12-diBr-2-Cl	0.058	0.072	0.079	0.078	0.098	0.095	0.080	18.35
96) 87 124-tri-Cl-B	0.887	0.929	1.087	1.119	1.063	1.017	1.017	10.07
97) 88 naphthalene	1.196	1.313	1.647	1.687	1.607	1.490	1.490	14.80
98) 89 hx-Cl-butadi	0.613	0.570	0.556	0.614	0.669	0.618	0.607	6.59
99) 90 123-Tri-Cl-B	0.743	0.767	0.924	0.919	0.874	0.845	0.845	10.10

0.990
0.994

(#) = Out of Range
 826WA010.M

Mon Mar 31 10:32:55 2003

INITIAL CALIBRATION SUMMARY

<u>Method File</u>	826WA010
<u>Last Calibration Update</u>	Mon Mar 31 10:32:37 2003
<u>Level 1 File Name</u>	10-0003A.D
<u>Level 2 File Name</u>	10-0002.D
<u>Level 3 File Name</u>	10-0010.D
<u>Level 4 File Name</u>	10-0020.D
<u>Level 5 File Name</u>	10-0040.D
<u>Level 6 File Name</u>	10-0060.D
<u>Level 7 File Name</u>	10-0020.D
<u>Level 1 ID</u>	3
<u>Level 2 ID</u>	2
<u>Level 3 ID</u>	10
<u>Level 4 ID</u>	20
<u>Level 5 ID</u>	40
<u>Level 6 ID</u>	60
<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	1402185	1401700	1425417	1419632	1412433	1400715	-1	-----	-----	-----	-----
3 di-Cl-di-F-methane	10847	65256	344729	715366	1537518	1972057	-1	0.0120	0.2437	0.0000	0.9908
4 Chloromethane	14243	71226	336320	702444	1333124	1882279	-1	0.0000	0.2560	0.0000	0.1632
2 F 114	5136	33302	171012	381760	733605	974692	-1	0.0000	0.1235	0.0000	0.0577
5 vinyl chloride	10455	66474	343348	725558	1503869	2051199	-1	0.0000	0.2487	0.0000	0.0429
6 bromomethane	7777	45613	200938	379493	785517	1151335	-1	0.0000	0.1497	0.0000	0.1341
7 chloroethane	10022	41956	214551	459359	920537	1294623	-1	0.0034	0.1562	0.0000	0.9986
8 tri-Cl-F-methane	16293	99495	515840	1121757	2241613	2920268	-1	0.0000	0.3739	0.0000	0.0580
91 Acetonitrile X10	20424	83795	408495	988740	1981444	2545093	-1	0.0000	0.0317	0.0000	0.0940
9 acrolein X10	11130	69228	329654	676073	1354015	1908483	-1	0.0000	0.0237	0.0000	0.0326
11 acetone X10	28459	104684	332887	639556	1203415	1741743	-1	0.0375	0.0202	0.0000	0.9998
12 ethyl ether X5	27298	176671	881117	1890393	3629304	5150102	-1	0.0000	0.1273	0.0000	0.0313
13 11-dichloroethene	13394	82732	420004	911032	1792018	2421004	-1	0.0000	0.3057	0.0000	0.0478
14 Iodomethane	3342	21843	176232	629497	1424862	2050698	-1	-0.0514	0.2544	0.0000	0.9940
15 F-113	9667	60030	305521	677448	1343471	1714509	-1	0.0000	0.2231	0.0000	0.0641
16 acrylonitrile X10	14334	91693	475343	863383	1929340	2786466	-1	0.0000	0.0330	0.0000	0.0416
17 carbon disulfide	42449	230738	1170329	2572254	5155522	6945088	-1	0.0000	0.8830	0.0000	0.0847
94 Isopropyl AlcoholX10	3968	8959	17262	31346	53652	-1	-1	0.0037	0.0009	0.0000	0.9956
18 methylene chloride	18179	66074	311459	637490	1291619	1876442	-1	0.0027	0.2242	0.0000	0.9997
19 t-12-d-Cl-ethene	11094	66357	348511	724140	1535637	2159090	-1	0.0000	0.2548	0.0000	0.0499
20 t-Bu-Me-ether	18835	120775	641266	1340185	2809850	4095292	-1	0.0000	0.4642	0.0000	0.0552
95 Tert butyl alcoholX10	-1	14346	35248	105910	309880	244429	-1	0.0000	0.0039	0.0000	0.3363

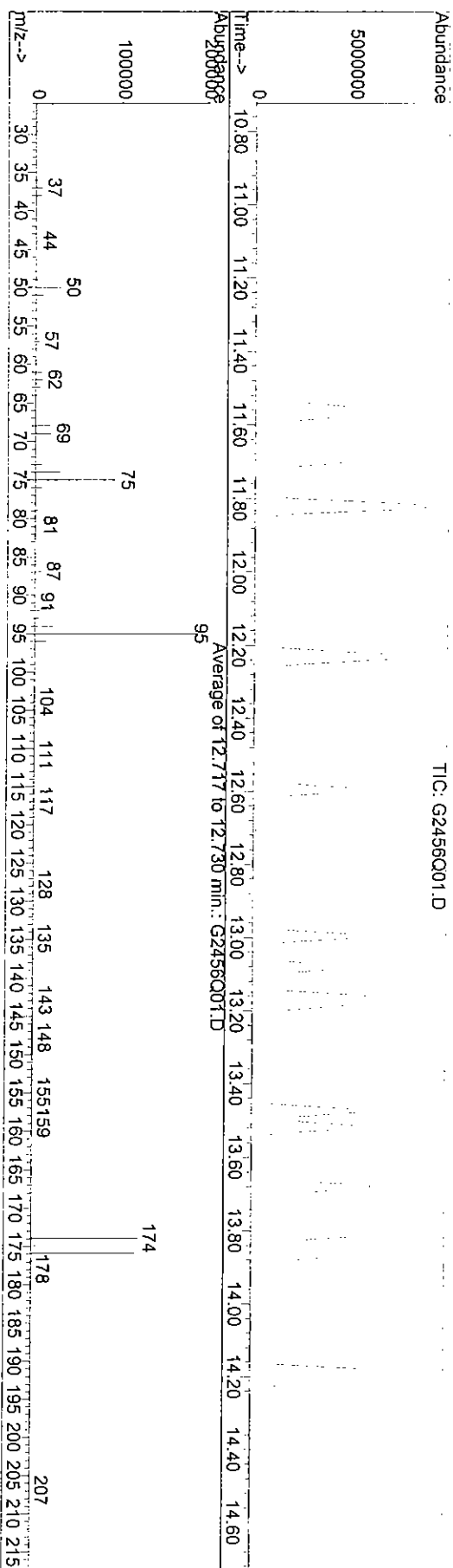
94 allyl chloride	-1	83795	408495	988740	1981444	2545093	-1	0.0000	0.3175	0.0000	0.0940
21 11-dichloroethane	18621	112842	581855	1186542	2504419	3588511	-1	0.0000	0.4236	0.0000	0.0406
97 propionitrile	136	3425	17147	28049	74203	104848	-1	0.0000	0.0119	0.0000	0.1029
22 c-12-di-Cl-ethene	10945	65228	346763	722727	1519062	2211008	-1	0.0000	0.2538	0.0000	0.0532
23 22-Dichloropropane	13116	74347	413858	914214	1915303	2688487	-1	0.0000	0.3080	0.0000	0.0854
24 Br-Cl-methane	4057	27612	140152	286516	593528	865536	-1	0.0000	0.1004	0.0000	0.0322
25 chloroform	22993	116777	597861	1214377	2494537	3609401	-1	0.0000	0.4269	0.0000	0.0230
26 tetrahydrofuranX5	6239	38117	198933	336272	790362	1173186	-1	0.0000	0.0274	0.0000	0.0727
98 Diisopropyl ether	28399	185552	986248	1893560	4112776	5938451	-1	0.0000	0.6884	0.0000	0.0370
27 Di-Br-F-Me (surr)	11084	60638	307963	643560	1311195	1902499	-1	0.0000	0.2302	0.0000	0.0761
99 ETBE	20365	132029	735024	1518648	3294293	4912092	-1	0.0000	0.5289	0.0000	0.0911
29 1,2-Di-Cl-Et-d4 (S1)	10836	56975	284710	593863	1176956	1708276	-1	0.0000	0.2136	0.0000	0.1024
30 12-dichloroethane	3534	21980	108018	227029	439257	631642	-1	0.0000	0.0785	0.0000	0.0409
32 vinyl acetate X5	63860	414860	2230480	4194574	9370203	13254307	-1	0.0000	0.3092	0.0000	0.0447
92 Nitro Methane(x10)	16099	98654	503941	1026662	2197883	3164808	-1	0.0000	0.0367	0.0000	0.0434
33 2-butanoneMEK X10	23496	109206	386182	800835	1835004	2621391	-1	0.0000	0.0316	0.0000	0.1475
93 Ethyl Acetate x2	10913	69080	401801	597072	1689794	2243912	-1	0.0000	0.1305	0.0000	0.1315
34 111-trichloroethane	16490	98389	517930	1091679	2332259	3272931	-1	0.0000	0.3822	0.0000	0.0576
35 11-Di-Cl-propene	13873	81037	444769	957310	2063877	2822864	-1	0.0000	0.3282	0.0000	0.0784
36 benzene	42274	259026	1393453	2994641	5972223	8547309	-1	0.0000	1.0059	0.0000	0.0500
37 CCl4	13612	78957	439090	920143	2050691	2812796	-1	0.0000	0.3225	0.0000	0.0840

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
100 Isobutyl alcoholx10	-1	4726	14196	70592	117355	167235	-1	-----	-----	-----	-----
38 thiophene	20269	128315	706455	1525558	3083581	4494310	-1	0.0000	0.5088	0.0000	0.0701
39 12-di-Cl-propane	9049	55011	298031	645043	1275427	1856699	-1	0.0000	0.2157	0.0000	0.0542
40 trichloroethene	11826	69743	368318	809129	1664792	2353937	-1	0.0000	0.2747	0.0000	0.0633
41 dibromomethane	4302	28599	144754	304462	609801	879445	-1	0.0000	0.1043	0.0000	0.0267
101 TAME	17251	113069	647400	1450103	2994595	4479028	-1	0.0000	0.4736	0.0000	0.1247
42 Br-di-Cl-methane	15590	77950	410468	873215	1744848	2551794	-1	0.0000	0.2972	0.0000	0.0456
43 Me-methacrylate	3581	24393	141762	302947	655401	947564	-1	0.0000	0.1012	0.0000	0.1285
44 2-ClEt-VI-ether10	4864	41895	276079	689334	1551121	2307334	-1	-0.0618	0.0285	0.0000	0.9983
45 c-13-di-Cl-propene	12016	78024	440533	972065	1981413	2911762	-1	0.0000	0.3188	0.0000	0.1009
46 t-1,3-dichloropropene	8432	57535	340215	757085	1619889	2349418	-1	-0.0204	0.2846	0.0000	0.9990
47 Chlorobenzene-d5	98828	690172	1024243	1043945	1007385	1007143	-1	0.0000	1.0000	0.0000	0.0000

48 112-tri-Cl-Et	6594	39408	204447	433455	862039	1245491	-1	0.0000	0.2081	0.0000	0.0427
49 13-di-Cl-propane	11088	71526	366282	776054	1541261	2185204	-1	0.0000	0.3681	0.0000	0.0258
50 Et methacrylate	6698	49628	294482	634508	1395321	2018328	-1	-0.0285	0.3411	0.0000	0.9981
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
51 di-Br-Cl-methane	9920	46263	252673	533500	1110440	1612394	-1	0.0000	0.2556	0.0000	0.0645
52 bromoform	4340	23486	130929	280129	598350	865181	-1	0.0000	0.1364	0.0000	0.0858
53 1,4-dichlorobutane-2	11561	63322	334210	699010	1415056	1998159	-1	0.0000	0.3325	0.0000	0.0355
54 MIBK	4102	26191	136436	276346	635297	913350	-1	0.0000	0.1413	0.0000	0.0861
55 toluene-d8	40331	232082	1249806	2759643	5492872	7796171	-1	0.0000	1.2734	0.0000	0.0606
56 toluene	44000	273555	1493556	3279721	6436941	9003846	-1	0.0000	1.4968	0.0000	0.0523
57 2-hexanone X5	18952	87657	465288	979048	2031915	2868357	-1	0.0000	0.0938	0.0000	0.0499
58 12-dibromoethane	5481	37268	194331	412749	842422	1213791	-1	0.0000	0.1950	0.0000	0.0469
59 tetra-Cl-ethene	11997	71322	368264	825724	1672283	2271115	-1	0.0000	0.3851	0.0000	0.0608
60 chlorobenzene	30395	178605	943078	2046929	4023278	5789041	-1	0.0000	0.9640	0.0000	0.0484
61 1112-tetra-Cl-Et	8360	55612	304924	672900	1358601	1988434	-1	0.0000	0.3081	0.0000	0.0799
62 1,4-Dichlorobenzene-d4	475933	477645	496828	495828	473305	466660	-1	0.0000	1.0000	0.0000	0.0000
63 1-chlorohexane	5749	32590	181618	420352	875550	1173405	-1	0.0000	0.4025	0.0000	0.1081
64 Et-Bz	49158	304901	1668070	3648966	7272072	10104426	-1	0.0000	3.5203	0.0000	0.0668
65 m/p-Xylenes X2	77987	475103	2595435	5637783	10918251	14861766	-1	0.0000	2.7016	0.0000	0.0550
66 styrene	26636	178864	994724	2170590	4313332	6092272	-1	0.0000	2.0638	0.0000	0.0850
67 o-xylene	37736	237765	1338059	2902842	5706977	7984326	-1	0.0000	2.7697	0.0000	0.0708
68 1122-Tetra-Cl-Et	7232	45751	228508	474459	964820	1348818	-1	0.0000	0.4859	0.0000	0.0389
69 123-tri-Cl-Pr	2281	12812	66381	135429	274408	386049	-1	0.0000	0.1411	0.0000	0.0708
70 4-Br-1-F-Bz (S3)	14323	68592	356594	767770	1533349	2198848	-1	0.0000	0.7610	0.0000	0.0545
71 isopropylbenzene	46404	281854	1625443	3621879	7283454	10015298	-1	0.0000	3.4248	0.0000	0.0953
72 bromobenzene	10720	66727	360820	778815	1562791	2219172	-1	0.0000	0.7632	0.0000	0.0613
92 t-1,4-dichloro-2-butene	721	5096	32426	74114	174039	258524	-1	-0.0165	0.0943	0.0000	0.9958
73 n-propylbenzene	14176	82858	463721	1046004	2147440	2993047	-1	0.0000	1.0086	0.0000	0.0966
74 2-Cl-Toluene	12148	71589	398415	872352	1768651	2511389	-1	0.0000	0.8522	0.0000	0.0789
75 4-Cl-Toluene	12996	74825	405718	882193	1787025	2516053	-1	0.0000	0.8737	0.0000	0.0698
76 135-tri-Me-Benzene	41200	245516	1403326	3075105	6157544	8461302	-1	0.0000	2.9426	0.0000	0.0809
77 4-iso-Pr-toluene	43393	271628	1518534	3283328	6877345	9299329	-1	0.0000	3.2006	0.0000	0.0870
78 124-tri-Me-Benzene	44257	256888	1411774	3112878	6178974	8636660	-1	0.0000	3.0196	0.0000	0.0703
79 tert-butylbenzene	35232	218303	1244711	2787927	5658720	7790717	-1	0.0000	2.6401	0.0000	0.0995

80 13-DCB	25714	143111	754344	1614275	3257265	4592008	-1	0.0000	1.6343	0.0000	0.0710
81 sec-butylbenzene	52624	328900	1843396	4079544	8272245	11100655	-1	0.0000	3.8811	0.0000	0.0861
82 14-DCB	27152	143617	745545	1584107	3232580	4565239	-1	0.0000	1.6402	0.0000	0.0917
83 Cl-benzyl	971	8277	57140	121481	312170	454087	-1	-0.0325	0.1668	0.0000	0.9932
84 12-DCB	22144	126151	657679	1353599	2799301	3968647	-1	0.0000	1.4094	0.0000	0.0651
85 n-butylbenzene	10725	68240	396229	839651	1862723	2572119	-1	0.0000	0.8354	0.0000	0.1224
86 12-diBr-2-Cl-Pra	829	6904	39462	77382	184762	265893	-1	-0.0119	0.0970	0.0000	0.9959
87 124-tri-Cl-Bz	17557	84776	461305	1077695	2118549	2975216	-1	0.0000	1.0169	0.0000	0.1007
88 naphthalene	25997	114299	652424	1633734	3193154	4499282	-1	0.0000	1.4901	0.0000	0.1480
89 hx-Cl-butadiene	8755	54493	276180	608951	1266400	1730110	-1	0.0000	0.6067	0.0000	0.0659
90 123-Tri-Cl-Bz	14845	70983	380915	916514	1740600	2447267	-1	0.0000	0.8455	0.0000	0.1010

Data File : C:\MSDCHEM\1\DATA\03G2456\G2456Q01.D Vial: 3
 Acq On : 13 May 2003 1:15 pm Operator: zou
 Sample : f=1 Inst : GCMS-A
 Misc : Multiplr: 1.00
 MS Integration Params: Lscint.p
 Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P &h Lab** EPA 8260



Spectrum Information: Average of 12.717 to 12.730 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
50	95	15	40	15.6	29790	PASS
75	95	30	60	48.9	93562	PASS
95	95	100	100	100.0	191189	PASS
96	95	5	9	7.3	13954	PASS
173	174	0.00	2	0.0	0	PASS
174	95	50	100	64.8	123936	PASS
175	174	5	9	7.8	9688	PASS
176	174	95	101	97.7	121088	PASS
177	176	5	9	7.3	8887	PASS

FORM-5A

Applied P & Ch Laboratory

Volatile Organic Instrument Performance Check for Method 524.2

Bromofluorobenzene (BFB), Part II

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 032987

Project ID: JPL

BFB Inj. Date: 05/13/03

Batch No: 03G2456

BFB Inj. Time: 13:15

Sequence No: 03G2456

Project No: 04-4428.10

Instrument ID: A

GC Column: HP-VOC

Data File Name: G2456Q01

Heated Purge: (Y/N) N

Column ID: 0.20 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	03G2456-CCV-01	03G2456-CCV-01	G2456Q01	05/13/03	13:15
2	03G2456-LCS-01	03G2456-LCS-01	G2456L01	05/13/03	13:41
3	MW-23-5MS	03-2987-6MS	G2456M01	05/13/03	14:08
4	MW-23-5MSD	03-2987-6MSD	G2456N01	05/13/03	14:35
5	03G2456-MB-01	03G2456-MB-01	G2456K01	05/13/03	15:55
6	EB-8-4/30/03	03-2987-1	2987-01	05/13/03	16:49
7	MW-23-5	03-2987-6	2987-06	05/13/03	17:16
8	MW-23-1	03-2987-2	2987-02	05/13/03	17:43
9	MW-23-2	03-2987-3	2987-03	05/13/03	18:10
10	MW-23-3	03-2987-4	2987-04	05/13/03	18:37
11	MW-23-4	03-2987-5	2987-05	05/13/03	19:04
12	TB-8-4/30/03	03-2987-7	2987-07	05/13/03	19:31
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					

Evaluate Continuing Calibration Report

Data File : C:\MSDCHEM\1\DATA\03G2456\G2456Q01.D Vial: 3
 Acq On : 13 May 2003 1:15 pm Operator: zou
 Sample : f=1 Inst : GCMS-A
 Misc : Multiplr: 1.00
 MS Integration Params: Lscint.p

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P & Ch Lab** EPA 8260
 Last Update : Thu Apr 10 14:53:47 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 200%

Compound	AVGRF	CCRF	%Dev	Area	% Dev	(min)
1 I	1.000	1.000	0.0	116	0.01	
2	0.249	0.221	11.2	102	0.02	
3 P	0.256	0.195	23.8	92	0.02	
4	0.124	0.110	11.3	95	0.03	
5 C	0.249	0.224	10.0	102	0.02	
6	0.150	0.118	21.3	102	0.02	
7	0.170	0.139	18.2	99	0.02	
8	0.374	0.346	7.5	101	0.02	
9	0.032	0.039	-21.9	131	0.06	
10	0.024	0.020	16.7	95	0.02	
11	0.025	0.031	-24.0	160	0.03	
12	0.127	0.115	9.4	100	0.02	
13 M, C	0.306	0.272	11.1	98	0.02	
14	0.166	0.241	-45.2	126	0.01	
15	0.223	0.217	2.7	105	0.02	
16	0.033	0.034	-3.0	131	0.02	
17	0.883	0.703	20.4	90	0.02	
18	0.003	0.001	66.7	95	0.02	
19	0.260	0.229	11.9	118	0.02	
20	0.255	0.242	5.1	110	0.01	
21	0.464	0.463	0.2	114	0.02	
22	0.004	0.008	-100.0	242	0.06	
23	0.317	0.393	-24.0	131	0.02	
24 P	0.424	0.392	7.5	109	0.01	
25	0.012	0.013	-8.3	150	0.03	
26	0.254	0.249	2.0	113	0.01	

(#) = Out of Range
 G2456Q01.D 826WA010.M Wed May 14 11:19:47 2003

Evaluate Continuing Calibration Report

Data File : C:\MSDCHEM\1\DATA\03G2456\G2456Q01.D Vial: 3
 Acq On : 13 May 2003 1:15 pm Operator: zou
 Sample : f=1 Inst : GCMS-A
 Misc : Multiplr: 1.00
 MS Integration Params: Iscint.p

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P & Ch Lab** EPA 8260
 Last Update : Thu Apr 10 14:53:47 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 200%

Compound	AVGRF	CCRF	%Dev Area	% Dev(min)
27 23 22-Dichloropropane	0.308	0.339	-10.1	122 0.02
28 24 Br-Cl-methane	0.100	0.099	1.0	114 0.01
29 C 25 chloroform	0.427	0.402	5.9	109 0.02
30 26 tetrahydrofuranX5	0.027	0.027	0.0	133 0.02
31 98 Diisopropyl ether	0.688	0.652	5.2	113 0.02
32 S 27 Di-Br-F-Me (surr)	0.230	0.216	6.1	111 0.02
33 99 ETBE	0.529	0.529	0.0	115 0.01
34 S 29 1,2-Di-Cl-Et-d4 (S1)	0.214	0.187	12.6	104 0.01
35 30 12-dichloroethane	0.079	0.068	13.9	99 0.01
36 32 vinyl acetate X5	0.309	0.297	3.9	116 0.02
37 92 Nitro Methane(x10)	0.037	0.032	13.5	102 0.13
38 33 2-butanoneMEK X10	0.032	0.044	-37.5#	180 0.02
39 93 Ethyl Acetate x2	0.130	0.098	24.6#	108 0.02
40 34 111-trichloroethane	0.382	0.382	0.0	115 0.02
41 35 11-Di-Cl-propene	0.328	0.322	1.8	111 0.01
42 M 36 benzene	1.006	0.969	3.7	107 0.01
43 37 CCl4	0.323	0.344	-6.5	123 0.01
44 100 Isobutyl alcoholx10	0.002	0.003	-50.0#	162 0.04
45 38 thiophene	0.509	0.497	2.4	107 0.01
46 C 39 12-di-Cl-propane	0.216	0.204	5.6	104 0.00
47 M 40 trichloroethene	0.275	0.272	1.1	111 0.01
48 41 dibromomethane	0.104	0.105	-1.0	113 0.00
49 101 TAME	0.474	0.478	-0.8	109 0.01
50 42 Br-di-Cl-methane	0.297	0.289	2.7	109 0.00
51 43 Me-methacrylate	0.101	0.108	-6.9	117 0.01
52 44 2-ClEt-VI-ether10	0.023	0.004	82.6#	17# 0.01

(#) = Out of Range

G2456Q01.D 826WA010.M Wed May 14 11:19:48 2003

Evaluate Continuing Calibration Report

Data File : C:\MSDCHEM\1\DATA\03G2456\G2456Q01.D Vial: 3
 Acq On : 13 May 2003 1:15 pm Operator: zou
 Sample : f=1 Inst : GCMS-A
 Misc : Multiplr: 1.00
 MS Integration Params: Lscint.p

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P & Ch Lab** EPA 8260
 Last Update : Thu Apr 10 14:53:47 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 200%

Compound	AvgRF	CCRF	%Dev	Area	Dev(min)
53 45 c-13-di-Cl-propene	0.319	0.328	-2.8	111	0.00
54 46 t-1,3-dichloropropene	0.246	0.265	-7.7	115	0.00
55 I 47 Chlorobezene-d5	1.000	1.000	0.0	115	0.00
56 48 112-tri-Cl-Et	0.208	0.206	1.0	114	0.00
57 49 13-di-Cl-propane	0.368	0.357	3.0	110	0.00
58 50 Et methacrylate	0.291	0.314	-7.9	118	0.00
59 51 di-Br-Cl-methane	0.256	0.266	-3.9	119	0.00
60 P 52 bromoform	0.136	0.145	-6.6	124	0.00
61 53 1,4-dichlorobutane-2	0.333	0.311	6.6	107	0.00
62 54 MIBK	0.141	0.142	-0.7	123	0.00
63 s 55 toluene-d8	1.273	1.239	2.7	108	0.00
64 M,C 56 toluene	1.497	1.463	2.3	107	0.00
65 57 2-hexanone X5	0.094	0.104	-10.6	127	0.00
66 58 12-dibromoethane	0.195	0.197	-1.0	114	0.00
67 59 tetra-Cl-ethene	0.385	0.387	-0.5	112	0.00
68 M,P 60 chlorobenzene	0.964	0.936	2.9	110	0.00
69 61 1112-tetra-Cl-Et	0.308	0.320	-3.9	114	0.00
70 I 62 1,4-Dichlorobenzene-d4	1.000	1.000	0.0	127	0.00
71 63 1-chlorohexane	0.402	0.377	6.2	113	0.00
72 C 64 Et-Bz	3.520	3.188	9.4	110	0.00
73 65 m/p-Xylenes X2	2.702	2.428	10.1	108	0.00
74 66 styrene	2.064	1.933	6.3	112	0.00
75 67 o-xylene	2.770	2.531	8.6	110	0.00
76 p 68 1122-Tetra-Cl-Et	0.486	0.431	11.3	114	0.00

(#) = Out of Range
 G2456Q01.D 826WA010.M Wed May 14 11:19:48 2003

Evaluate Continuing Calibration Report

Data File : C:\MSDCHEM\1\DATA\03G2456\G2456Q01.D Vial: 3
 Acq On : 13 May 2003 1:15 pm Operator: zou
 Sample : f=1 Inst : GCMS-A
 Misc : Multiplr: 1.00
 MS Integration Params: lscint.p

Method : C:\MSDCHEM\1\METHODS\826WA010.M (RTE Integrator)
 Title : **Applied P &ch Lab** EPA 8260
 Last Update : Thu Apr 10 14:53:47 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 200%

Compound	AvgRF	CCRF	%Dev	Area	%Dev	Area	%Dev
77 69 123-tri-Cl-Pr	0.141	0.127	9.9	118	0.00		
78 S 70 4-Br-1-F-Bz (S3)	0.761	0.702	7.8	115	0.00		
79 71 isopropylbenzene	3.425	3.246	5.2	113	0.00		
80 72 bromobenzene	0.763	0.706	7.5	114	0.00		
81 92 t-1,4-dichloro-2-butene	0.071	0.070	1.4	119	0.00		
82 73 n-propylbenzene	1.009	0.950	5.8	114	0.00		
83 74 2-Cl-Toluene	0.852	0.790	7.3	114	0.00		
84 75 4-Cl-Toluene	0.874	0.793	9.3	113	0.00		
85 76 135-tri-Me-Benzene	2.943	2.738	7.0	112	0.00		
86 77 4-iso-Pr-toluene	3.201	3.026	5.5	116	0.00		
87 78 124-tri-Me-Benzene	3.020	2.766	8.4	112	0.00		
88 79 tert-butylbenzene	2.640	2.508	5.0	113	0.00		
89 80 13-DCB	1.634	1.498	8.3	117	0.00		
90 81 sec-butylbenzene	3.881	3.718	4.2	115	0.00		
91 82 14-DCB	1.640	1.462	10.9	116	0.00		
92 83 Cl-benzyl	0.120	0.165	-37.5#	170	0.00		
93 84 12-DCB	1.409	1.282	9.0	119	0.00		
94 85 n-butylbenzene	0.835	0.834	0.1	125	0.00		
95 86 12-diBr-2-Cl-Pra	0.080	0.084	-5.0	136	0.00		
96 87 124-tri-Cl-Bz	1.017	0.993	2.4	116	0.00		
97 88 naphthalene	1.490	1.602	-7.5	123	0.00		
98 89 hx-Cl-butadiene	0.607	0.562	7.4	116	0.00		
99 90 123-Tri-Cl-Bz	0.845	0.816	3.4	112	0.00		

(#) = Out of Range SPPC's out = 0 CCC's out = 0
 G2456Q01.D 826WA010.M Wed May 14 11:19:49 2003

Continuing Calibration Concentration Summary

Data File G2456Q01
 Method File 826WA010

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
1 Fluorobenzene	10	10.00	ppb	0.00	1646457
3 di-Cl-di-F-methane	20	17.75	ppb	11.27	726202
4 Chloromethane	20	16.62	ppb	16.90	643403
2 F114	20	17.85	ppb	10.76	362968
5 vinyl chloride	20	18.00	ppb	10.01	737069
6 bromomethane	20	15.73	ppb	21.33	387834
7 chloroethane	20	17.52	ppb	12.39	456239
8 tri-Cl-F-methane	20	18.48	ppb	7.59	1137788
91 Acetonitrile X10	200	247.62	ppb	23.81	1294272
9 acrolein X10	200	165.26	ppb	17.37	643826
11 acetone X10	200	288.92	ppb	44.46	1022772
12 ethyl ether X5	100	90.56	ppb	9.44	1897786
13 11-dichloroethene	20	17.82	ppb	10.91	896873
14 Iodomethane	20	20.94	ppb	4.68	792462
15 F-113	20	19.44	ppb	2.78	714265
16 acrylonitrile X10	200	208.87	ppb	4.43	1133945
17 carbon disulfide	20	15.92	ppb	20.39	2314877
94 Isopropyl Alcoholx10	200	165.06	ppb	17.47	29727
18 methylene chloride	20	20.30	ppb	1.50	753632
19 t-12-di-Cl-ethene	20	18.97	ppb	5.15	795724
20 t-Bu-Me-ether	20	19.93	ppb	0.35	1523160
95 Tert butyl alcoholx10	200	394.66	ppb	97.33	256192
94 allyl chloride	20	24.76	ppb	23.81	1294272
21 11-dichloroethane	20	18.52	ppb	7.41	1291524
97 propionitrile	20	21.32	ppb	6.62	41944
22 c-12-di-Cl-ethene	20	19.61	ppb	1.96	819293
23 22-Dichloropropane	20	22.03	ppb	10.17	1117461
24 Br-Cl-methane	20	19.70	ppb	1.50	325556
25 chloroform	20	18.85	ppb	5.75	1325038
26 tetrahydrofuranX5	100	98.97	ppb	1.03	446372
98 Diisopropyl ether	20	18.95	ppb	5.24	2148005
27 Di-Br-F-Me (surr)	20	18.78	ppb	6.09	711767
99 ETBE	20	20.01	ppb	0.05	1742296
29 1,2-Di-Cl-Et-d4 (S1)	20	17.53	ppb	12.35	616351
30 12-dichloroethane	20	17.32	ppb	13.40	223895
32 vinyl acetate X5	100	95.92	ppb	4.08	4882913
92 Nitro Methane(x10)	200	172.91	ppb	13.54	1043467
33 2-butanoneMEK X10	200	277.96	ppb	38.98	1445478
93 Ethyl Acetate x2	40	29.94	ppb	25.14	643184
34 111-trichloroethane	20	19.99	ppb	0.04	1257987
35 11-Di-Cl-propene	20	19.64	ppb	1.81	1061234
36 benzene	20	19.26	ppb	3.69	3190045
37 CCl4	20	21.30	ppb	6.52	1131176

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
100 Isobutyl alcoholx10	200	375.57	ppb	87.79	114213
38 thiophene	20	19.55	ppb	2.27	1637488
39 12-di-Cl-propane	20	18.94	ppb	5.28	672824
40 trichloroethene	20	19.84	ppb	0.82	897033
41 dibromomethane	20	20.07	ppb	0.34	344537
101 TAME	20	20.19	ppb	0.94	1574036
42 Br-di-Cl-methane	20	19.44	ppb	2.78	951451
43 Me-methacrylate	20	21.29	ppb	6.46	354683
44 2-ClEt-Vi-ether10	200	46.95	ppb	76.52	118721
45 c-13-di-Cl-propene	20	20.56	ppb	2.82	1079280
46 t-1,3-dichloropropene	20	19.32	ppb	3.39	871943
47 Chlorobezene-d5	10	10.00	ppb	0.00	1198131
48 112-tri-Cl-Et	20	19.81	ppb	0.93	493987
49 13-di-Cl-propane	20	19.42	ppb	2.89	856503
50 Et methacrylate	20	19.23	ppb	3.83	751839

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
51 di-Br-Cl-methane	20	20.78	ppb	3.89	636409
52 bromoform	20	21.29	ppb	6.43	347919
53 1,4-dichlorobutane-2	20	18.70	ppb	6.48	745202
54 MIBK	20	20.12	ppb	0.60	340676
55 toluene-d8	20	19.47	ppb	2.67	2970011
56 toluene	20	19.55	ppb	2.24	3506528
57 2-hexanone X5	100	110.66	ppb	10.66	1243497
58 12-dibromoethane	20	20.21	ppb	1.03	472189
59 tetra-Cl-ethene	20	20.08	ppb	0.39	926294
60 chlorobenzene	20	19.42	ppb	2.90	2243020
61 1112-tetra-Cl-Et	20	20.74	ppb	3.71	765753
62 1,4-Dichlorobenzene-d4	10	10.00	ppb	0.00	628384
63 1-chlorohexane	20	18.74	ppb	6.30	473929
64 Et-Bz	20	18.11	ppb	9.43	4006756
65 m/p-Xylenes X2	40	35.95	ppb	10.14	6102368
66 styrene	20	18.73	PPB	6.36	2428805
67 o-xylene	20	18.28	ppb	8.62	3180795
68 1122-Tetra-Cl-Et	20	17.76	ppb	11.20	542201
69 123-tri-Cl-Pr	20	17.99	ppb	10.04	159579
70 4-Br-1-F-Bz (S3)	20	18.45	ppb	7.73	882502
71 isopropylbenzene	20	18.96	ppb	5.22	4079382
72 bromobenzene	20	18.50	ppb	7.49	887249
92 t-1,4-dichloro-2-butene	20	16.60	ppb	17.00	87993
73 n-propylbenzene	20	18.83	ppb	5.86	1193315
74 2-Cl-Toluene	20	18.53	ppb	7.33	992418
75 4-Cl-Toluene	20	18.15	ppb	9.26	996343
76 135-tri-Me-Benzene	20	18.61	ppb	6.97	3440427
77 4-iso-Pr-toluene	20	18.91	ppb	5.46	3802809
78 124-tri-Me-Benzene	20	18.32	ppb	8.39	3476640
79 tert-butylbenzene	20	19.00	ppb	5.01	3151803
80 13-DCB	20	18.33	ppb	8.36	1882277
81 sec-butylbenzene	20	19.16	ppb	4.20	4672968

82 14-DCB	20	17.83	ppb	10.87	1837296
83 Cl-benzyl	20	21.70	ppb	8.52	207100
84 12-DCB	20	18.20	ppb	9.02	1611555
85 n-butylbenzene	20	19.96	ppb	0.18	1048008
86 12-diBr-2-Cl-Pra	20	18.56	ppb	7.22	105620
87 124-tri-Cl-Bz	20	19.54	ppb	2.30	1248512
88 naphthalene	20	21.50	ppb	7.48	2012873
89 hx-Cl-butadiene	20	18.54	ppb	7.29	706919
90 123-Tri-Cl-Bz	20	19.31	ppb	3.44	1026015

Average D % 9.3240463

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: 032987

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

CCV Data File: G2456Q01

Instrument ID: A

Batch No: 03G2456

#	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		05/13/03 13:15	1646457 7.64	1198131 11.54	628384 13.84
	CCV Upper Limit			3292914 8.14	2396262 12.04	1256768 14.34
	CCV Lower Limit			823228 7.14	599065 11.04	314192 13.34
1	03G2456-LCS-01	03G2456-LCS-01	05/13/03 13:41	1536499 7.64	1124739 11.54	592039 13.85
2	MW-23-5MS	03-2987-6MS	05/13/03 14:08	1674220 7.64	1209063 11.54	627928 13.84
3	MW-23-5MSD	03-2987-6MSD	05/13/03 14:35	1632156 7.64	1205698 11.54	637369 13.84
4	03G2456-MB-01	03G2456-MB-01	05/13/03 15:55	1477639 7.64	1123135 11.54	551400 13.85
5	EB-8-4/30/03	03-2987-1	05/13/03 16:49	1477542 7.64	1152738 11.55	576265 13.85
6	MW-23-5	03-2987-6	05/13/03 17:16	1430404 7.64	1094707 11.55	539535 13.85
7	MW-23-1	03-2987-2	05/13/03 17:43	1451912 7.64	1121747 11.54	561231 13.85
8	MW-23-2	03-2987-3	05/13/03 18:10	1428231 7.64	1098884 11.55	547562 13.85
9	MW-23-3	03-2987-4	05/13/03 18:37	1410443 7.64	1088510 11.55	538612 13.85
10	MW-23-4	03-2987-5	05/13/03 19:04	1376588 7.64	1051051 11.55	521934 13.85
11	TB-8-4/30/03	03-2987-7	05/13/03 19:31	1385341 7.64	1075974 11.55	534359 13.85
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

VOC Analysis General Logbook

3760 Magnolia Ave. Chino CA 91710
Tel: (909) 590-1828 Fax: (909) 590-1498

Matrix: W

Date: 3-30-03 Analyst: Zou

Sequence # 0351873 Batch # 0351873
IS/Surrogate: GC 14507/14508 Methanol(mark-M): PEG (mark-PEG): Defoaming(mark-DF):

Op. #	Type	Sample ID	Method	V/X=f ₁	V _f /V _i =f ₂	V _{spg} /V _{inj} =f ₃	F	A-#	Datafile	Note	pH
4969	SP	61873P01	626A	2525=1	/ =	/ =	/		61873P01	3-30-03	
4970	MSD	10-0003A	010	/ =	/ =	/ =	/		10-0003A	61873P01	
4971		10-0002		/ =	/ =	/ =	/		10-0002		
4972		10-0010		/ =	/ =	/ =	/		10-0010		
4973		10-0020		/ =	/ =	/ =	/		10-0020		
4974		10-0040		/ =	/ =	/ =	/		10-0040		
4975	✓	10-0060		/ =	/ =	/ =	/		10-0060		
4976	CW	61873P01		/ =	/ =	/ =	/		61873P01	CW/CW	3-30-03
4977	MS	L01		/ =	/ =	/ =	/		L01	61873P01	12:30 PM
4978	MS	M01		/ =	/ =	/ =	/		M01	\$2301-04	12
4979	MSD	N01		/ =	/ =	/ =	/		N01	\$2301-04	
4980	MS	M02		/ =	/ =	/ =	/		M02	\$2303-03	
4981	MSD	N02		/ =	/ =	/ =	/		N02	\$2303-03	
4982	MS	K01		/ =	/ =	/ =	/		K01		
4983	Sample	2469-01		/ =	/ =	/ =	/		2469-01	tb	12
4984		2301-05		/ =	/ =	/ =	/		2301-05	tb	
4985		2303-04		/ =	/ =	/ =	/		2303-04	tb	
4986		2303-03		/ =	/ =	/ =	/		2303-03	ms	
4987		2301-04		/ =	/ =	/ =	/		2301-04	ms	
4988		01		/ =	/ =	/ =	/		01		
4989		02		/ =	/ =	/ =	/		02		
4990		03		/ =	/ =	/ =	/		03		
4991		2303-01		/ =	/ =	/ =	/		2303-01		
4992		02		/ =	/ =	/ =	/		02		
4993		2351-01		/ =	/ =	/ =	/		2351-01		
4994		2469-02		/ =	/ =	/ =	/		2469-02		
4995		03		/ =	/ =	/ =	/		03		
4996		04		/ =	/ =	/ =	/		04		
4997		2449-01		/ =	/ =	/ =	/		2449-01		
4998		02		/ =	/ =	/ =	/		02		
4999	✓	2353-01A	✓	1.1	1.1	1.1	✓		2353-01A	23:15	

Type	Op #	STD Lot #	C _{std} (ng/μL) × V _{std} (μL) / X(g or mL) = T	Op #	STD Lot #	C _{std} (ng/μL) × V _{std} (μL) / X(g or mL) = T
LCS/LCSD	44177	GC	1.1		GC	1.1
MS/MSD	44177/44179	GC	1.1		GC	1.1

Footnote/Anomaly:

2.5 20

AL 4-2-03

Applied P & Ch Laboratory

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

VOC-150

VOC Analysis General Logbook

Sequence # 03G2456 Batch # 0382456 Matrix: W Date: 05/13/03 Analyst: Zou
Lot #: IS/Surrogate: GC-1514/1515 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	$V/X=f_1$	$V_f/V_i=f_2$	$V_{fpg}/V_{inj}=f_3$	F	A-#	Datafile	Note	pH
5769	SP	G2456P01	826WA 010	25/25 = 1	/ =	/ =	1		G2456P01	05/13/03	
5770	CCV	Q01		/ =	/ =	/ =			Q01	GC15148	1:15 pm
5771	LCS	L01		/ =	/ =	/ =			L01		
5772	MS	M01		/ =	/ =	/ =			M01	\$2987-06	<2
5773	MSD	N01		/ =	/ =	/ =			N01	↓	↓
5774	MB	✓ K01		/ =	/ =	/ =			K01		
5775	sample	2987-01		/ =	/ =	/ =			2987-01	524.2	<2
5776		06		/ =	/ =	/ =			06		
5777		02		/ =	/ =	/ =			02		
5778		03		/ =	/ =	/ =			03		
5779		04		/ =	/ =	/ =			04		
5780		05		/ =	/ =	/ =			05		
5781		✓ 07		/ =	/ =	/ =			✓ 07		
5782		3015-01		/ =	/ =	/ =			3015-01		
5783		02		/ =	/ =	/ =			02		
5784		03		/ =	/ =	/ =			03		
5785		04		/ =	/ =	/ =			04		
5786		05		/ =	/ =	/ =			05		
5787		06		/ =	/ =	/ =			06		
5788		07		/ =	/ =	/ =			07		
5789		✓ 08		/ =	/ =	/ =			✓ 08	✓	
5790		3171-03		/ =	/ =	/ =			3171-03		
5791		04		/ =	/ =	/ =			04		
5792		09		/ =	/ =	/ =			09		
5793	✓	✓ 12	✓	✓ / =	✓ / =	✓ / =	✓		✓ 12		✓
5794				/ =	/ =	/ =					
5795				/ =	/ =	/ =					
5796				/ =	/ =	/ =					
5797				/ =	/ =	/ =					
5798				/ =	/ =	/ =					
5799				/ =	/ =	/ =					
5800				/ =	/ =	/ =					

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	5771	GC-1514	200 x 25 / X = ppb		GC-	x / X = ppb
MS/MSD	5772/5773	GC-1514	200 x 25 / X = 20 ppb		GC-	x / X = ppb

Footnote/Anomaly:

6149