

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269L01.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:09 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:35 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:35 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.08	9.03	0.005	96	70	760.366	10.00		0.05	
47	Cl-benzene-d5, I2	12.72	12.66	0.005	82	119	211.140	10.00		0.06	
62	1,4-DCB-d4 150 15	15.23	15.15	0.005	152	150	183.003	10.00		0.07	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.49	7.44	0.003	111	113	518.428	19.24		19.2	%Recovery
29	1,2-di-Cl-ethane-	8.06	8.02	0.003	65	102	218.761	17.41		17.4	96.22%
55	toluene-d8(S2)	11.21	11.15	0.004	100	99	765.373	20.15		20.2	87.07%
70	4-Br-1-F-Bz (S3)	13.96	13.90	0.004	174	95	332.480	20.33		20.3	100.76%
											101.67%

Target Compounds

<<< I1 : ISTD ID = 1 >>>

Qvalue

3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	383.100	17.63		17.6	99	
4	Chloromethane	2.78	2.78	0.000	50	52	292.753	15.82		15.8	96	
9	F114 85 135	2.83	2.83	0.000	85	135	484.444	19.95		20.0	97	
5	vinyl chloride	2.96	2.96	0.000	62	64	318.677	18.16		18.2	97	
6	bromomethane	3.38	3.37	0.001	94	96	289.007	18.74		18.7	99	
7	Chloroethane	3.53	3.52	0.002	64	66	251.941	19.39		19.4	98	
8	tri-Cl-F-methane	4.18	4.14	0.004	101	103	621.777	20.96		21.0	99	
111	isopropyl alcoho	4.29	4.27	0.002	45	43	61.593	179.53		179.5	56	
100	ethyl ether x5	4.47	4.44	0.004	59	74	911.831	84.14		84.1	98	
102	Acrolein x10	4.18	4.15	0.003	56	55	180.850	209.16		209.2	95	
119	methyl acetate	5.10	5.06	0.005	43	74	200.521	21.50		21.5	95	
104	Carbon disulfide	5.22	5.18	0.004	76	78	960.634	15.16		15.2	99	
103	Acrylonitrilex10	4.91	4.89	0.003	53	52	323.435	178.02		178.0	100	
95	Acetone x10	4.34	4.31	0.004	43	58	228.722	174.91		174.9	96	
108	F-113	5.06	5.02	0.004	151	101	512.767	22.50		22.5	99	
13	11-dichloroethene	4.79	4.76	0.004	61	96	558.505	18.74		18.7	98	
101	Acetonitrilex10	4.22	4.20	0.003	41	40	86.265	177.07		177.1	96	
109	Iodomethane	4.83	4.80	0.003	142	127	569.024	23.67		23.7	98	
113	Tert butyl alcoh	4.88	4.86	0.002	59	57	132.404	201.56		201.6	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269L01.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:09 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:35 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:36 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	QI	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.00	4.97	0.004	84	49	575.899	19.19	19.2	96	
112	Allyl chloride	5.10	5.07	0.004	41	76	618.233	18.75	18.7	93	?
200	200 Nitro methane x1	5.85	5.82	0.004	61	46	649.609	195.82	195.8	98	?
10	10 t-Bu-Me-ether	6.03	6.00	0.004	73	57	610.777	18.97	19.0	98	
19	19 t-12-di-Cl-ethene	5.85	5.82	0.004	96	61	434.441	19.01	19.0	96	?
98	98 Vinyl acetate x5	6.45	6.41	0.004	43	86	1703.772	82.85	82.9	99	
21	21 11-dichloroethane	6.19	6.15	0.004	63	83	810.889	20.32	20.3	99	
91	91 2-butanone MEKx10	6.87	6.83	0.004	43	72	1197.008	167.86	167.9	97	?
115	115 Di isoprop ether	6.89	6.85	0.004	45	87	1858.495	18.45	18.5	98	?
22	22 c-12-di-Cl-ethene	7.02	6.97	0.005	96	61	417.889	18.69	18.7	100	
23	23 22-Dichloropropan	7.40	7.35	0.005	77	97	679.257	19.73	19.7	99	
24	24 Br-Cl-methane	7.24	7.18	0.006	128	130	141.841	17.68	17.7	99	
25	25 chloroform	7.31	7.27	0.004	83	85	731.109	18.31	18.3	100	
201	201 Ethyl acetate x2	7.36	7.31	0.005	43	61	357.480	35.16	35.2	97	?
116	116 ETBE	7.45	7.40	0.005	59	87	1080.021	17.60	17.6	99	
117	117 Iso-butyl alcoho	7.36	7.31	0.005	43	42	388.200	190.83	190.8	73	#?
26	26 tetrahydrofuranx5	7.75	7.71	0.005	72	42	67.594	86.75	86.7	97	
34	34 111-tri-Cl-ethane	8.27	8.23	0.004	97	99	637.647	19.17	19.2	100	
30	30 12-dichloroethane	8.18	8.13	0.005	62	64	274.618	17.65	17.7	99	
35	35 11-Di-Cl-propene	8.52	8.48	0.005	75	110	562.608	19.36	19.4	99	
36	36 benzene	8.79	8.75	0.005	78	52	1326.604	20.08	20.1	99	
37	37 CCl4	8.73	8.68	0.005	117	119	523.785	20.58	20.6	100	
97	97 thiophene	8.94	8.89	0.005	84	58	620.752	18.90	18.9	96	
118	118 TAME	9.05	9.00	0.005	73	43	756.627	17.91	17.9	95	
39	39 12-di-Cl-propane	9.54	9.49	0.005	63	76	365.865	18.92	18.9	98	
40	40 trichloroethene	9.59	9.54	0.005	130	132	405.826	19.82	19.8	99	
96	96 Me-methacrylate	9.89	9.85	0.004	69	100	123.368	18.37	18.4	97	
42	42 Br-di-Cl-methane	9.65	9.61	0.004	83	85	474.566	17.60	17.6	98	
41	41 dibromomethane	9.49	9.45	0.004	174	172	183.214	19.64	19.6	99	
45	45 c-13-di-Cl-propen	10.42	10.37	0.006	75	110	432.009	18.17	18.2	100	
92	92 2-ClEt-Vi-ether10	10.18	10.15	0.004	63	43	302.903	74.28	74.3	98	
56	56 toluene	11.28	11.23	0.006	91	92	1222.138	19.68	19.7	100	

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 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:09 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:35 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:36 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Ql	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.41	11.36	0.006	69	99	228.934	17.82	17.8	99	
93	2-Hexanone x5	11.54	11.48	0.007	43	58	378.055	87.53	87.5	99	
48	112-tri-Cl-Et	11.09	11.03	0.007	97	83	170.332	18.73	18.7	95	
58	1,2-di-br-ethane	11.88	11.82	0.007	107	109	180.169	18.52	18.5	96	
51	di-Br-Cl-methane	11.62	11.57	0.006	129	127	236.052	19.28	19.3	99	
46	t-13-di-Cl-propen	10.92	10.87	0.006	75	110	293.929	19.18	19.2	99	
105	1-Chlorohexane	12.69	12.64	0.006	55	93	404.758	21.47	21.5	98	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.57	10.52	0.004	43	58	136.432	20.56	20.6	97	
49	1,3-di-Cl-propane	11.34	11.29	0.004	76	78	290.382	21.04	21.0	99	
59	tetra-Cl-ethene	12.06	12.00	0.004	166	168	429.899	23.14	23.1	100	
60	chlorobenzene	12.76	12.70	0.004	112	77	687.288	22.27	22.3	99	
61	1112-tetra-Cl-Et	12.68	12.62	0.004	131	133	277.514	22.46	22.5	99	
64	ethylbenzene	12.95	12.90	0.004	91	106	1357.792	22.84	22.8	100	
65	m/p-Xylenes x2	13.16	13.10	0.005	91	106	2047.539	45.30	45.3	99	
99	1-4-di-Cl-butane	13.51	13.46	0.004	55	41	265.975	19.53	19.5	98	
52	bromoforn	13.27	13.22	0.004	173	175	133.209	21.27	21.3	96	
66	styrene	13.49	13.43	0.005	104	78	714.123	22.37	22.4	98	
67	o-xylene	13.56	13.50	0.005	91	106	978.072	21.99	22.0	99	
68	1122-Tetra-Cl-Et	13.57	13.51	0.005	83	85	160.767	19.69	19.7	98	
110	t-1,4-dichloro-2	13.72	13.67	0.004	89	53	31.577	24.88	24.9	72	#?
106	Cl-benzyl	15.19	15.12	0.005	91	126	236.942	23.36	23.4	99	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.69	13.64	0.003	110	97	42.651	19.14	19.1	92	
71	isopropylbenzene	13.92	13.86	0.004	105	120	1374.707	22.86	22.9	100	
72	bromobenzene	14.15	14.10	0.003	156	158	279.305	21.16	21.2	97	
73	n-propylbenzene	14.34	14.29	0.004	120	78	368.669	22.74	22.7	98	
74	2-Cl-Tl	14.44	14.37	0.004	126	128	215.744	20.03	20.0	99	
75	4-Cl-Tl	14.51	14.45	0.004	126	128	314.992	21.40	21.4	94	m
76	135-tri-Me-Bz	14.62	14.57	0.004	105	120	1055.627	21.74	21.7	100	
79	tert-butylbenzene	14.89	14.84	0.004	119	91	1114.768	22.36	22.4	98	
78	124-tri-Me-Bz	15.01	14.94	0.004	105	120	906.226	21.95	22.0	100	

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 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:09 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:35 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:36 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.19	15.11	0.005	146	148	438.952	20.87	20.9	90
82	14-di-Cl-Bz	146	15.24	15.19	0.004	146	148	589.866	19.30	19.3	94
81	sec-butylbenzene		15.11	15.04	0.004	105	134	1597.369	21.91	21.9	100
77	4-iso-Pr-toluene		15.28	15.22	0.004	119	134	1169.490	22.24	22.2	100
84	12-di-Cl-benzene		15.59	15.52	0.005	146	148	393.257	19.38	19.4	98
85	n-butylbenzene		15.67	15.60	0.005	91	134	1160.225	21.05	21.1	100
86	12-diBr-3-Cl-Pra		16.06	15.97	0.006	157	155	28.220	18.98	19.0	95
87	124-tri-Cl-Bz		17.32	17.24	0.005	180	182	295.683	18.55	18.5	95
88	naphthalene		17.56	17.49	0.005	128	129	219.618	18.04	18.0	100
90	123-tri-Cl-Bz		17.75	17.68	0.004	180	182	229.553	20.76	20.8	100
89	hx-Cl-butadiene		17.59	17.52	0.005	225	260	292.763	24.31	24.3	96

7/15
 57/15

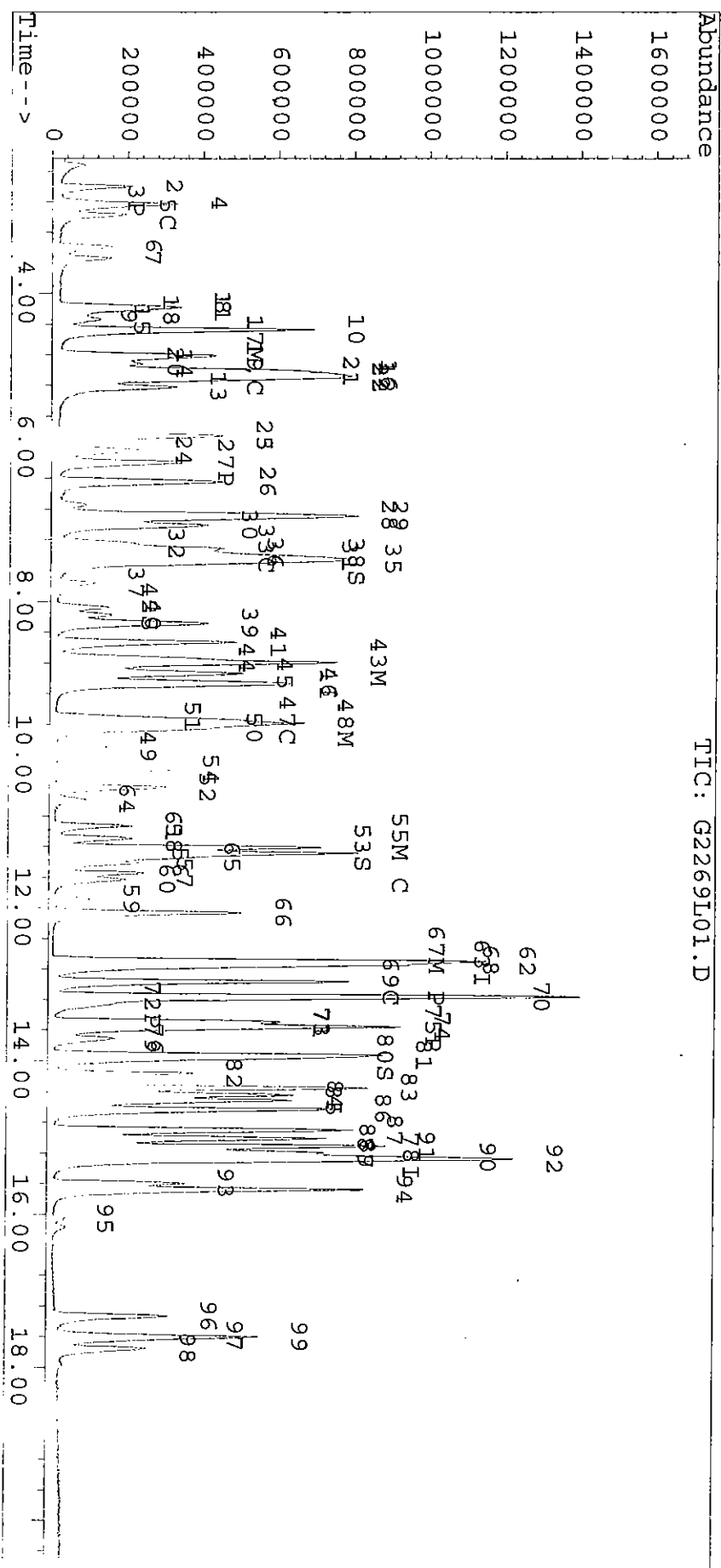
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269L01.D
 Acq On : 30 Apr 03 6:09 pm
 Sample : f=1
 Misc :
 Quant Time: May 1 11:35 2003

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

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



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: 32964

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2269

MS Filename: G2269M01

Date Analyzed: 043003

Time Analyzed: 18:38

MSD Filename: G2269N01

Date Analyzed: 043003

Time Analyzed: 19:08

MS Sample No: MW-17-2

Sample Lab ID: 03-2933-3

Spiked Components	Unit	Spike Added	Concentration		MS Rec% #	QC Limit, % REC
			Unspiked	MS		
BENZENE	µg/L	20	0	20.6	103	65-121
CHLOROBENZENE	µg/L	20	0	22.7	114	65-134
1,1-DICHLOROETHENE	µg/L	20	0	20.0	100	65-127
TOLUENE	µg/L	20	0	20.2	101	65-134
TRICHLOROETHENE	µg/L	20	0.9	20.5	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	20.6	103	0	28	65-121
CHLOROBENZENE	µg/L	20	22.4	112	2	35	65-134
1,1-DICHLOROETHENE	µg/L	20	19.4	97	3	31	65-127
TOLUENE	µg/L	20	20.0	100	1	35	65-134
TRICHLOROETHENE	µg/L	20	20.9	100	2	30	65-125
# of Out-of-control				0	0		

Column to be used to flag recovery and RPD values:

* - Values outside of contract required QC Limits

D - Spiked components diluted out

Comments: _____

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269M01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 30 18:38 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : May 01 11:44 2003
 Print Time : Thu May 01 11:44 2003
 Miscellaneous :

Sample : f=1 \$2933-03
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

6949

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.07	9.03	0.004	96	70	774.675	10.00		0.04	
47	Cl-benzene-d5, I2	12.72	12.66	0.005	82	119	217.575	10.00		0.06	
62	1,4-DCB-d4 150 15	15.24	15.15	0.005	152	150	187.626	10.00		0.08	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.48	7.44	0.003	111	113	528.190	19.24		19.2	%Recovery
29	1,2-di-Cl-ethane-	8.06	8.02	0.003	65	102	215.931	16.87		16.9	96.22%
55	toluene-d8(S2)	11.21	11.15	0.004	100	99	808.687	20.90		20.9	84.35%
70	4-Br-1-F-Bz (S3)	13.96	13.90	0.004	174	95	344.365	20.54		20.5	104.49%
										102.71%	

Target Compounds

Qvalue

<<< I1 : ISTD ID = 1 >>>											
3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	415.115	18.79		18.8	98
4	Chloromethane	2.78	2.78	0.000	50	52	305.857	16.23		16.2	97
9	F114 85 135	2.83	2.83	0.000	85	135	534.423	21.60		21.6	94
5	vinyl chloride	2.97	2.96	0.001	62	64	340.242	19.03		19.0	98
6	bromomethane	3.39	3.37	0.002	94	96	314.373	20.05		20.1	96
7	Chloroethane	3.54	3.52	0.003	64	66	264.625	19.99		20.0	99
8	tri-Cl-F-methane	4.18	4.14	0.004	101	103	680.412	22.51		22.5	99
111	isopropyl alcoh	4.29	4.27	0.002	45	43	47.972	137.24		137.2	1
100	ethyl ether x5	4.47	4.44	0.004	59	74	887.068	80.34		80.3	98
102	Acrolein x10	4.19	4.15	0.005	56	55	222.501	252.58		252.6	89
119	methyl acetate	5.10	5.06	0.005	43	74	253.506	26.61		26.6	100
104	Carbon disulfide	5.22	5.18	0.005	76	78	1039.627	16.10		16.1	100
103	Acrylonitrilex10	4.93	4.89	0.005	53	52	261.166	141.09		141.1	96
95	Acetone x10	4.34	4.31	0.004	43	58	290.029	221.19		221.2	98
108	F-113	5.06	5.02	0.005	151	101	577.916	24.89		24.9	100
13	11-dichloroethene	4.79	4.76	0.004	61	96	606.139	19.96		20.0	99
101	Acetonitrilex10	4.23	4.20	0.004	41	40	73.046	146.82		146.8	92
109	Iodomethane	4.83	4.80	0.004	142	127	564.643	23.01		23.0	99
113	Tert butyl alcoh	4.89	4.86	0.003	59	57	119.163	177.84		177.8	94

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Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269M01.D Sample : f=1 \$2933-03
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:38 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:44 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:45 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.00	4.97	0.004	84	49	583.386	19.07	19.1	95	
112	Allyl chloride	5.11	5.07	0.005	41	76	656.151	19.53	19.5	95	?
200	200 Nitro methane x1	5.85	5.82	0.004	61	46	667.219	197.41	197.4	100	?
10	10 t-Bu-Me-ether	6.04	6.00	0.005	73	57	600.945	18.32	18.3	98	
19	19 t-12-di-Cl-ethene	5.85	5.82	0.004	96	61	459.535	19.74	19.7	100	?
98	98 Vinyl acetate x5	6.45	6.41	0.004	43	86	802.799	38.32	38.3	97	
21	21 11-dichloroethane	6.20	6.15	0.005	63	83	836.385	20.57	20.6	98	
91	91 2-butanone MEKx10	6.88	6.83	0.005	43	72	1192.427	164.13	164.1	98	?
115	115 Di isoprop ether	6.90	6.85	0.005	45	87	1881.344	18.33	18.3	98	
22	22 c-12-di-Cl-ethene	7.01	6.97	0.004	96	61	436.960	19.18	19.2	98	
23	23 22-Dichloropropan	7.39	7.35	0.004	77	97	711.931	20.29	20.3	98	
24	24 Br-Cl-methane	7.24	7.18	0.006	128	130	146.177	17.88	17.9	97	
25	25 chloroform	7.31	7.27	0.004	83	85	752.402	18.50	18.5	100	
201	201 Ethyl acetate x2	7.35	7.31	0.004	43	61	216.135	20.87	20.9	75	#?
116	116 ETBE	7.45	7.40	0.005	59	87	1079.348	17.27	17.3	100	
117	117 Iso-butyl alcoho	7.35	7.31	0.004	43	42	226.183	109.13	109.1	1	#?
26	26 tetrahydrofuranx5	7.76	7.71	0.005	72	42	64.364	81.07	81.1	95	
34	34 111-tri-Cl-ethane	8.27	8.23	0.004	97	99	670.454	19.79	19.8	99	
30	30 12-dichloroethane	8.18	8.13	0.005	62	64	282.037	17.79	17.8	99	
35	35 11-Di-Cl-propene	8.53	8.48	0.006	75	110	601.335	20.31	20.3	98	
36	36 benzene	8.80	8.75	0.005	78	52	1383.046	20.55	20.5	99	
37	37 CCl4	8.74	8.68	0.006	117	119	563.351	21.72	21.7	99	
97	97 thiophene	8.95	8.89	0.006	84	58	633.601	18.93	18.9	96	
118	118 TAME	9.05	9.00	0.005	73	43	732.507	17.02	17.0	96	
39	39 12-di-Cl-propane	9.54	9.49	0.005	63	76	383.147	19.45	19.4	97	
40	40 trichloroethene	9.59	9.54	0.005	130	132	428.177	20.53	20.5	100	
96	96 Me-methacrylate	9.89	9.85	0.004	69	100	119.429	17.49	17.5	100	
42	42 Br-di-Cl-methane	9.65	9.61	0.004	83	85	477.062	17.37	17.4	96	
41	41 dibromomethane	9.50	9.45	0.005	174	172	177.451	18.68	18.7	99	
45	45 c-13-di-Cl-propen	10.41	10.37	0.005	75	110	438.612	18.11	18.1	98	
92	92 2-ClEt-Vi-ether10	10.19	10.15	0.005	63	43	290.585	69.94	69.9	98	
56	56 toluene	11.28	11.23	0.006	91	92	1276.090	20.17	20.2	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269M01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 30 18:38 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : May 01 11:44 2003
 Print Time : Thu May 01 11:45 2003
 Miscellaneous :

Sample : f=1 \$2933-03
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

4951

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.41	11.36	0.006	69	99	130.777	9.99	10.0	98	
93	2-Hexanone x5	11.53	11.48	0.006	43	58	337.629	76.73	76.7	96	
48	112-tri-Cl-Et	11.08	11.03	0.006	97	83	165.941	17.90	17.9	94	
58	1,2-di-br-ethane	11.88	11.82	0.007	107	109	172.190	17.38	17.4	98	
51	di-Br-Cl-methane	11.63	11.57	0.007	129	127	232.721	18.66	18.7	100	
46	t-13-di-Cl-propen	10.91	10.87	0.005	75	110	287.212	18.40	18.4	99	
105	1-Chlorohexane	12.69	12.64	0.006	55	93	439.123	22.94	22.9	99	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.59	10.52	0.006	43	58	126.942	18.51	18.5	98	
49	1,3-di-Cl-propane	11.34	11.29	0.004	76	78	294.443	20.70	20.7	99	
59	tetra-Cl-ethene	12.06	12.00	0.004	166	168	463.358	24.20	24.2	97	
60	chlorobenzene	12.76	12.70	0.004	112	77	722.029	22.71	22.7	98	
61	1112-tetra-Cl-Et	12.68	12.62	0.005	131	133	280.221	22.01	22.0	99	
64	ethylbenzene	12.95	12.90	0.004	91	106	1418.776	23.16	23.2	99	
65	m/p-Xylenes x2	13.16	13.10	0.005	91	106	2156.122	46.29	46.3	98	
99	1-4-di-Cl-butane	13.52	13.46	0.005	55	41	259.306	18.48	18.5	98	
52	bromoforn	13.28	13.22	0.005	173	175	133.503	20.66	20.7	99	
66	styrene	13.49	13.43	0.005	104	78	747.037	22.71	22.7	99	
67	o-xylene	13.56	13.50	0.005	91	106	1015.279	22.15	22.2	99	
68	1122-Tetra-Cl-Et	13.57	13.51	0.005	83	85	164.797	19.58	19.6	96	
110	t-1,4-dichloro-2	13.73	13.67	0.005	89	53	32.323	24.68	24.7	69	
106	Cl-benzyl	15.19	15.12	0.005	91	126	262.898	25.28	25.3	97	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.69	13.64	0.003	110	97	40.952	17.93	17.9	93	
71	isopropylbenzene	13.92	13.86	0.004	105	120	1447.451	23.48	23.5	100	
72	bromobenzene	14.16	14.10	0.004	156	158	293.025	21.65	21.7	98	
73	n-propylbenzene	14.35	14.29	0.004	120	78	392.745	23.63	23.6	98	
74	2-Cl-Tl	14.45	14.37	0.005	126	128	236.513	21.42	21.4	94	
75	4-Cl-Tl	14.52	14.45	0.005	126	128	331.889	21.99	22.0	99	
76	135-tri-Me-Bz	14.63	14.57	0.004	105	120	1108.916	22.27	22.3	98	
79	tert-butylbenzene	14.90	14.84	0.004	119	91	1187.444	23.23	23.2	99	
78	124-tri-Me-Bz	15.01	14.94	0.005	105	120	901.551	21.30	21.3	98	

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

5/10/03

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269M01.D Sample : F=1 \$2933-03
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 18:38 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:44 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:45 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	15.19	15.11	0.005	146	148	425.408	19.73	19.7	90	
82	14-di-Cl-Bz	15.25	15.19	0.004	146	148	658.249	21.01	21.0	94	
81	sec-butylbenzene	15.12	15.04	0.005	105	134	1786.653	23.91	23.9	99	
77	4-iso-Pr-toluene	15.29	15.22	0.004	119	134	1253.502	23.25	23.2	99	
84	12-di-Cl-benzene	15.60	15.52	0.005	146	148	413.326	19.87	19.9	97	
85	n-butylbenzene	15.67	15.60	0.005	91	134	1289.015	22.81	22.8	99	
86	12-diBr-3-Cl-Pra	16.06	15.97	0.005	157	155	29.283	19.21	19.2	99	
87	124-tri-Cl-Bz	17.32	17.24	0.005	180	182	317.616	19.40	19.4	97	
88	naphthalene	17.55	17.49	0.004	128	129	219.459	17.59	17.6	99	
90	123-tri-Cl-Bz	17.75	17.68	0.004	180	182	236.766	20.88	20.9	100	
89	hx-Cl-butadiene	17.59	17.52	0.005	225	260	334.033	27.08	27.1	95	

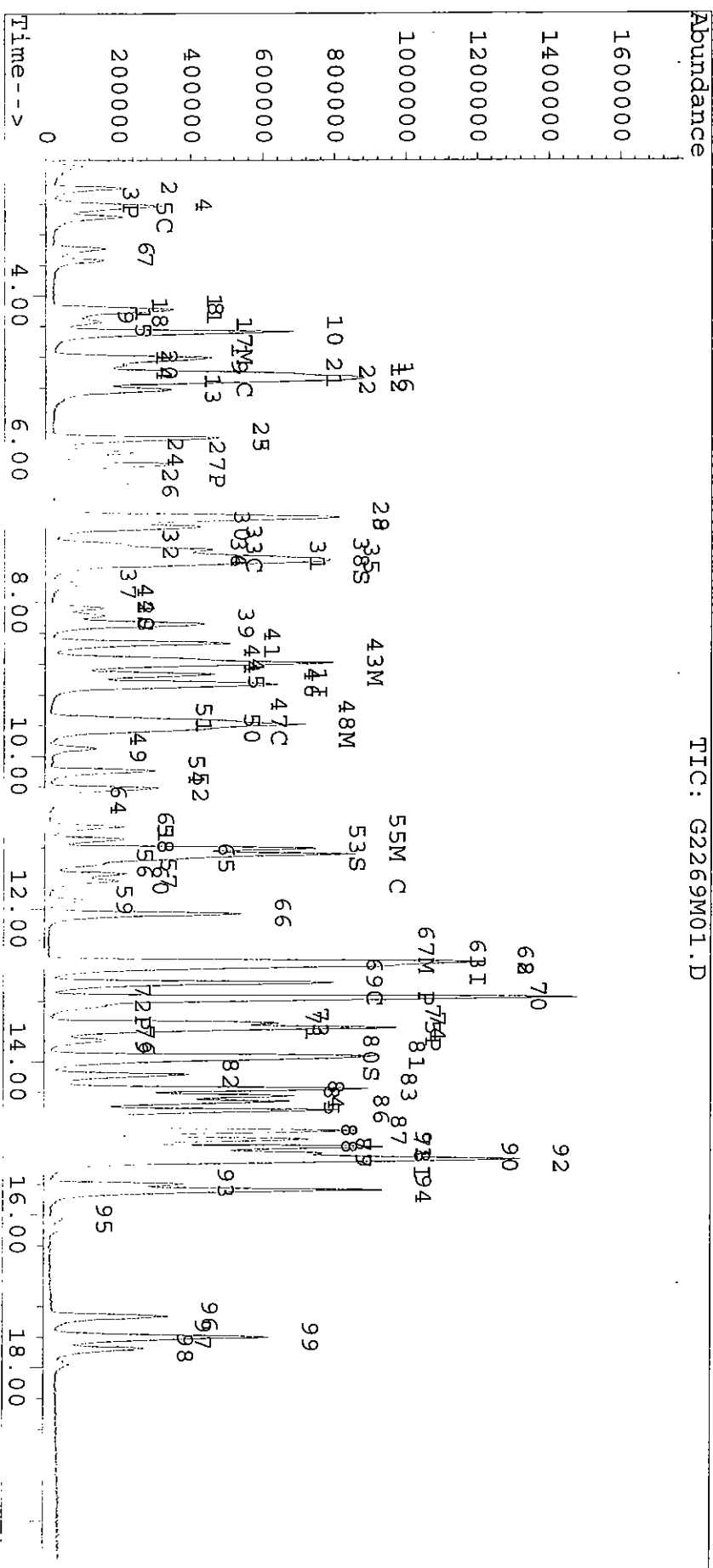
m
 47.163

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

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269M01.D Vial: 12
 Acq On : 30 Apr 03 6:38 pm Operator: Eddie
 Sample : f=1 \$2933-03 Inst : GCMS-G
 Misc : Multiplr: 1.00
 Quant Time: May 1 11:44 2003 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269N01.D Sample : f=1 \$2933-03
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 19:08 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:46 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:47 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.08	9.03	0.005	96	70	799.033	10.00		Dev (Min)	
47	Cl-benzene-d5, I2	12.73	12.66	0.005	82	119	232.426	10.00		0.05	
62	1,4-DCB-d4 150 15	15.24	15.15	0.005	152	150	196.889	10.00		0.06	

System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.49	7.44	0.003	111	113	557.316	19.69		%Recovery	
29	1,2-di-Cl-ethane-	8.07	8.02	0.003	65	102	221.284	16.76		98.46%	
55	toluene-d8(S2)	11.21	11.15	0.003	100	99	830.407	20.81		16.8	83.81%
70	4-Br-1-F-Bz (S3)	13.97	13.90	0.005	174	95	362.036	20.58		20.8	104.03%
										20.6	102.90%

Target Compounds											
<<< I1 : ISTD ID = 1 >>>											
3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	431.107	18.93		18.9	100
4	Chloromethane	2.78	2.78	0.000	50	52	326.254	16.78		16.8	96
9	F114 85 135	2.83	2.83	0.000	85	135	542.850	21.28		21.3	99
5	vinyl chloride	2.97	2.96	0.000	62	64	342.420	18.57		18.6	99
6	bromomethane	3.40	3.37	0.003	94	96	331.116	20.49		20.5	95
7	Chloroethane	3.54	3.52	0.003	64	66	268.905	19.70		19.7	99
8	tri-Cl-F-methane	4.19	4.14	0.005	101	103	693.600	22.25		22.2	99
111	isopropyl alcoho	4.31	4.27	0.004	45	43	55.114	152.87		152.9	77
100	ethyl ether x5	4.47	4.44	0.004	59	74	945.291	83.00		83.0	99
102	Acrolein x10	4.20	4.15	0.005	56	55	189.433	208.49		208.5	94
119	methyl acetate	5.10	5.06	0.005	43	74	239.713	24.42		24.4	93
104	Carbon disulfide	5.22	5.18	0.004	76	78	1053.792	15.82		15.8	100
103	Acrylonitrile x10	4.93	4.89	0.004	53	52	307.021	160.81		160.8	98
95	Acetone x10	4.34	4.31	0.004	43	58	237.913	172.99		173.0	98
108	F-113	5.06	5.02	0.004	151	101	632.383	26.40		26.4	99
13	11-dichloroethene	4.80	4.76	0.005	61	96	608.613	19.43		19.4	98
101	Acetonitrile x10	4.26	4.20	0.007	41	40	92.902	181.52		181.5	85
109	Iodomethane	4.84	4.80	0.004	142	127	587.471	23.23		23.2	98
113	Tert butyl alcoh	4.90	4.86	0.004	59	57	126.386	182.92		182.9	96

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269N01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 30 19:08 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : May 01 11:46 2003
 Print Time : Thu May 01 11:47 2003
 Miscellaneous :

Sample : f=1 \$2933-03
 Inst. : GCMS-G
 RF via : Multiple Level Calibration
 Operator: Eddie
 Multiplr: 1.000000

4955

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.01	4.97	0.004	84	49	602.598	19.10	19.1	95	
112	Allyl chloride	5.11	5.07	0.004	41	76	686.580	19.81	19.8	96	?
200	200 Nitro methane x1	5.86	5.82	0.005	61	46	712.079	204.26	204.3	99	?
10	10 t-Bu-Me-ether	6.04	6.00	0.005	73	57	616.032	18.21	18.2	98	
19	19 t-12-di-Cl-ethene	5.86	5.82	0.005	96	61	475.495	19.80	19.8	96	?
98	98 Vinyl acetate x5	6.45	6.41	0.005	43	86	1383.630	64.03	64.0	96	
21	21 11-dichloroethane	6.21	6.15	0.006	63	83	887.654	21.17	21.2	99	
91	91 2-butanone MEKx10	6.88	6.83	0.005	43	72	1239.483	165.40	165.4	98	?
115	115 Di isoprop ether	6.89	6.85	0.005	45	87	1959.678	18.51	18.5	97	?
22	22 c-12-di-Cl-ethene	7.01	6.97	0.005	96	61	444.345	18.91	18.9	99	
23	23 22-Dichloropropan	7.40	7.35	0.006	77	97	731.997	20.23	20.2	98	
24	24 Br-Cl-methane	7.25	7.18	0.007	128	130	147.335	17.48	17.5	98	
25	25 chloroform	7.32	7.27	0.005	83	85	772.781	18.42	18.4	100	
201	201 Ethyl acetate x2	7.36	7.31	0.006	43	61	327.044	30.61	30.6	98	?
116	116 ETBE	7.45	7.40	0.006	59	87	1143.313	17.73	17.7	99	
117	117 Iso-butyl alcoho	7.36	7.31	0.006	43	42	358.178	167.55	167.5	90	#?
26	26 tetrahydrofuranx5	7.77	7.71	0.007	72	42	64.183	78.38	78.4	90	
34	34 111-tri-Cl-ethane	8.29	8.23	0.006	97	99	707.538	20.25	20.2	99	
30	30 12-dichloroethane	8.18	8.13	0.006	62	64	291.468	17.83	17.8	98	
35	35 11-Di-Cl-propene	8.53	8.48	0.006	75	110	630.446	20.65	20.6	99	
36	36 benzene	8.80	8.75	0.006	78	52	1433.228	20.64	20.6	99	
37	37 CCl4	8.74	8.68	0.006	117	119	580.631	21.71	21.7	100	
97	97 thiophene	8.94	8.89	0.006	84	58	673.421	19.51	19.5	96	
118	118 TAME	9.05	9.00	0.005	73	43	764.544	17.22	17.2	97	
39	39 12-di-Cl-propane	9.54	9.49	0.006	63	76	402.987	19.83	19.8	97	
40	40 trichloroethene	9.59	9.54	0.006	130	132	449.209	20.88	20.9	100	
96	96 Me-methacrylate	9.89	9.85	0.005	69	100	132.657	18.78	18.8	98	
42	42 Br-di-Cl-methane	9.66	9.61	0.005	83	85	509.430	17.98	18.0	98	
41	41 dibromomethane	9.50	9.45	0.005	174	172	182.470	18.62	18.6	100	
45	45 c-13-di-Cl-propen	10.42	10.37	0.006	75	110	458.034	18.33	18.3	98	
92	92 2-ClEt-Vl-etherx10	10.20	10.15	0.006	63	43	299.817	69.96	70.0	97	
56	56 toluene	11.29	11.23	0.007	91	92	1304.268	19.99	20.0	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269N01.D Sample : f=1 \$2933-03
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 19:08 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:46 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:47 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.42	11.36	0.007	69	99	199.361	14.76	14.8	97	
93	2-Hexanone x5	11.55	11.48	0.007	43	58	370.568	81.64	81.6	93	
48	1,2-di-tri-Cl-Et	11.08	11.03	0.006	97	83	172.022	17.99	18.0	95	
58	1,2-di-br-ethane	11.89	11.82	0.008	107	109	178.705	17.49	17.5	100	
51	di-Br-Cl-methane	11.63	11.57	0.007	129	127	236.128	18.36	18.4	100	
46	t-13-di-Cl-propen	10.93	10.87	0.007	75	110	302.557	18.79	18.8	100	
105	1-Chlorohexane	12.70	12.64	0.007	55	93	470.284	23.85	23.9	99	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.58	10.52	0.005	43	58	136.321	18.61	18.6	96	
49	1,3-di-Cl-propane	11.35	11.29	0.005	76	78	295.029	19.42	19.4	99	
59	tetra-Cl-ethene	12.07	12.00	0.005	166	168	481.078	23.52	23.5	99	
60	chlorobenzene	12.76	12.70	0.004	112	77	760.654	22.39	22.4	98	
61	1,1,2-tetra-Cl-Et	12.69	12.62	0.006	131	133	296.618	21.81	21.8	99	
64	ethylbenzene	12.97	12.90	0.006	91	106	1456.436	22.25	22.3	99	
65	m/p-Xylenes x2	13.17	13.10	0.005	91	106	2218.648	44.59	44.6	98	
99	1-4-di-Cl-butane	13.52	13.46	0.005	55	41	268.410	17.90	17.9	98	
52	bromoforn	13.28	13.22	0.004	173	175	133.514	19.27	19.3	96	
66	styrene	13.50	13.43	0.005	104	78	757.360	21.56	21.6	99	
67	o-xylene	13.56	13.50	0.005	91	106	1055.350	21.56	21.6	99	
68	1,1,2,2-Tetra-Cl-Et	13.57	13.51	0.005	83	85	167.760	18.66	18.7	99	
110	t-1,4-dichloro-2	13.73	13.67	0.005	89	53	31.791	22.36	22.4	72	
106	Cl-benzyl	15.20	15.12	0.006	91	126	245.579	21.89	21.9	98	
<<< I3 : ISTD ID = 62 >>>											
69	1,2,3-tri-Cl-Pr	13.71	13.64	0.005	110	97	41.377	17.26	17.3	90	
71	isopropylbenzene	13.93	13.86	0.005	105	120	1556.345	24.06	24.1	100	
72	bromobenzene	14.17	14.10	0.004	156	158	288.810	20.33	20.3	98	
73	n-propylbenzene	14.35	14.29	0.004	120	78	404.898	23.22	23.2	100	
74	2-Cl-Tl	14.45	14.37	0.005	126	128	236.570	20.42	20.4	93	
75	4-Cl-Tl	14.52	14.45	0.005	126	128	329.313	20.79	20.8	100	
76	1,3,5-tri-Me-Bz	14.63	14.57	0.004	105	120	1197.414	22.92	22.9	99	
79	tert-butylbenzene	14.91	14.84	0.005	119	91	1246.176	23.24	23.2	100	
78	1,2,4-tri-Me-Bz	15.02	14.94	0.005	105	120	954.190	21.48	21.5	100	

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269N01.D Sample : f=1 \$2933-03
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 19:08 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:46 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:47 2003
 Miscellaneous :

4957

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.20	15.11	0.005	146	148	443.615	19.61	19.6	91
82	14-di-Cl-Bz	146	15.26	15.19	0.005	146	148	689.114	20.96	21.0	93
81	sec-butylbenzene		15.12	15.04	0.005	105	134	1798.695	22.94	22.9	100
77	4-iso-Pr-toluene		15.29	15.22	0.005	119	134	1311.559	23.18	23.2	99
84	12-di-Cl-benzene		15.61	15.52	0.006	146	148	425.827	19.51	19.5	97
85	n-butylbenzene		15.68	15.60	0.005	91	134	1365.645	23.03	23.0	100
86	12-diBr-3-Cl-Pra		16.05	15.97	0.005	157	155	29.519	18.45	18.5	97
87	124-tri-Cl-Bz		17.33	17.24	0.006	180	182	330.279	19.23	19.2	97
88	naphthalene		17.57	17.49	0.005	128	129	223.073	17.03	17.0	99
90	123-tri-Cl-Bz		17.76	17.68	0.005	180	182	240.502	20.22	20.2	97
89	hx-Cl-butadiene		17.60	17.52	0.005	225	260	345.375	26.68	26.7	97

2/2
5/1/03

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

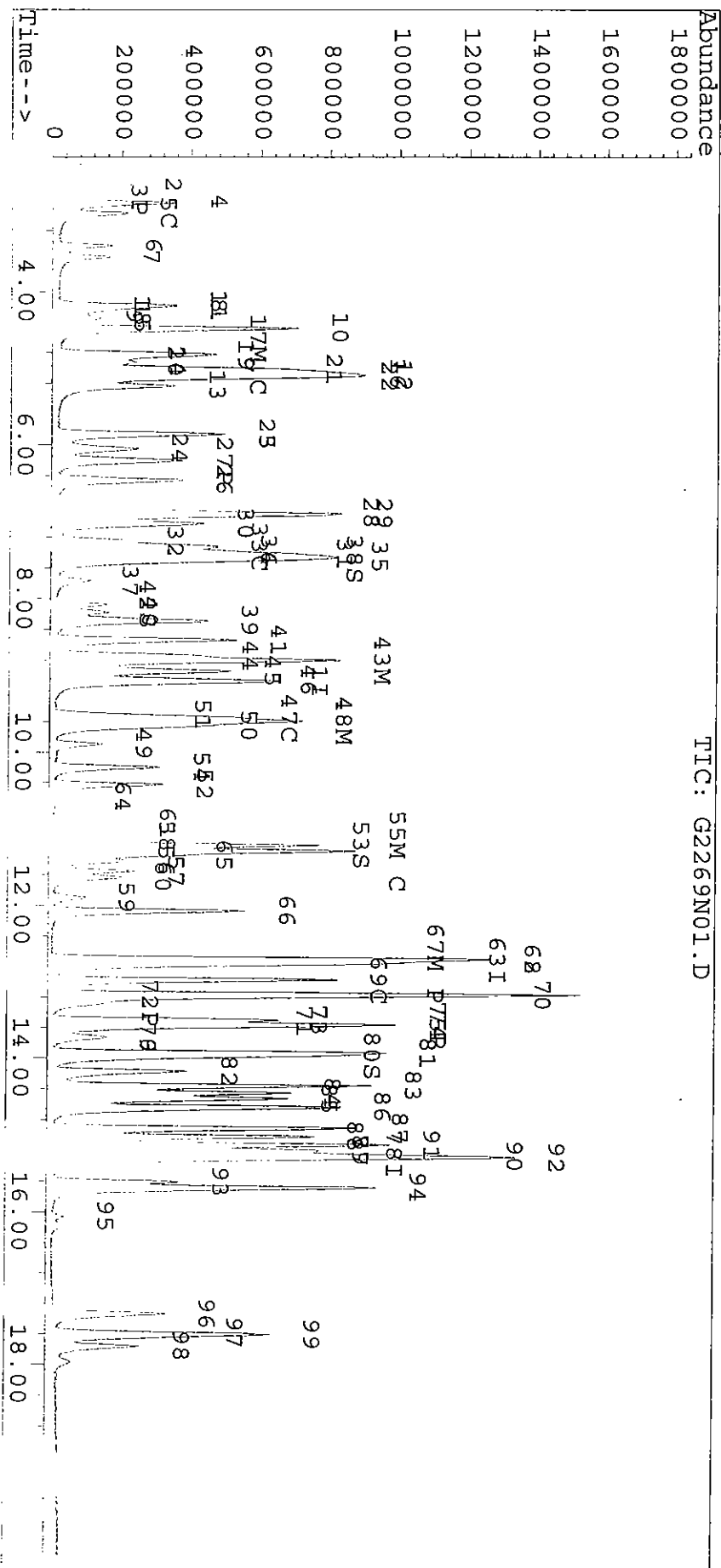
Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269N01.D
 Acq On : 30 Apr 03 7:08 pm
 Sample : f=1 \$2933-03
 Misc :
 Quant Time: May 1 11:46 2003

Quant Results File: quant.res

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

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



Data Filename: C:\HPCHEM\1\DATA\03G2269\2933-03.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 22:28 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Apr 30 22:49 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:49 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
----	----------------	------	-----	------	------	----	---------	--------	-------	---------	------

Internal Standards											
1	Fluorobenzene I1	9.12	9.03	0.010	96	70	791.241	10.00		0.09	
47	Cl-benzene-d5, I2	12.77	12.66	0.008	82	119	218.903	10.00		0.10	
62	1,4-DCB-d4 150 15	15.26	15.15	0.007	152	150	178.189	10.00		0.11	

System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.53	7.44	0.006	111	113	493.993	17.60		17.6	87.99%
29	1,2-di-Cl-ethane-	8.12	8.02	0.006	65	102	228.093	17.45		17.4	87.24%
55	toluene-d8(S2)	11.25	11.15	0.007	100	99	795.479	20.13		20.1	100.63%
70	4-Br-1-F-Bz (S3)	13.99	13.90	0.006	174	95	322.931	20.28		20.3	101.42%

Target Compounds	<<< I1 : ISTD ID = 1 >>>	Qvalue
111 111 isopropyl alcoho	4.26 4.27 0.000 45 43 2.725 7.63 7.6	1
119 119 methyl acetate	5.05 5.06 0.000 43 74 1.787 0.45 0.4	56
104 104 Carbon disulfide	5.26 5.18 0.008 76 78 23.344 0.35 0.4	84
91 91 2-butanone MEKx10	6.83 6.83 0.000 43 72 4.291 0.58 0.6	74
117 117 Iso-butyl alcoho	7.31 7.31 0.000 43 42 1.598 0.75 0.8	1
40 40 trichloroethene	9.65 9.54 0.012 130 132 18.520 0.87 0.9	89
48 48 112-tri-Cl-Et	11.27 11.03 0.027 97 83 20.768 1.97 2.0	4
<<< I2 : ISTD ID = 47 >>>		
54 54 MIBK	10.63 10.52 0.009 43 58 39.323 5.28 5.3	98
49 49 1,3-di-Cl-propane	11.27 11.29 -0.002 76 78 9.228 0.64 0.6	91

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

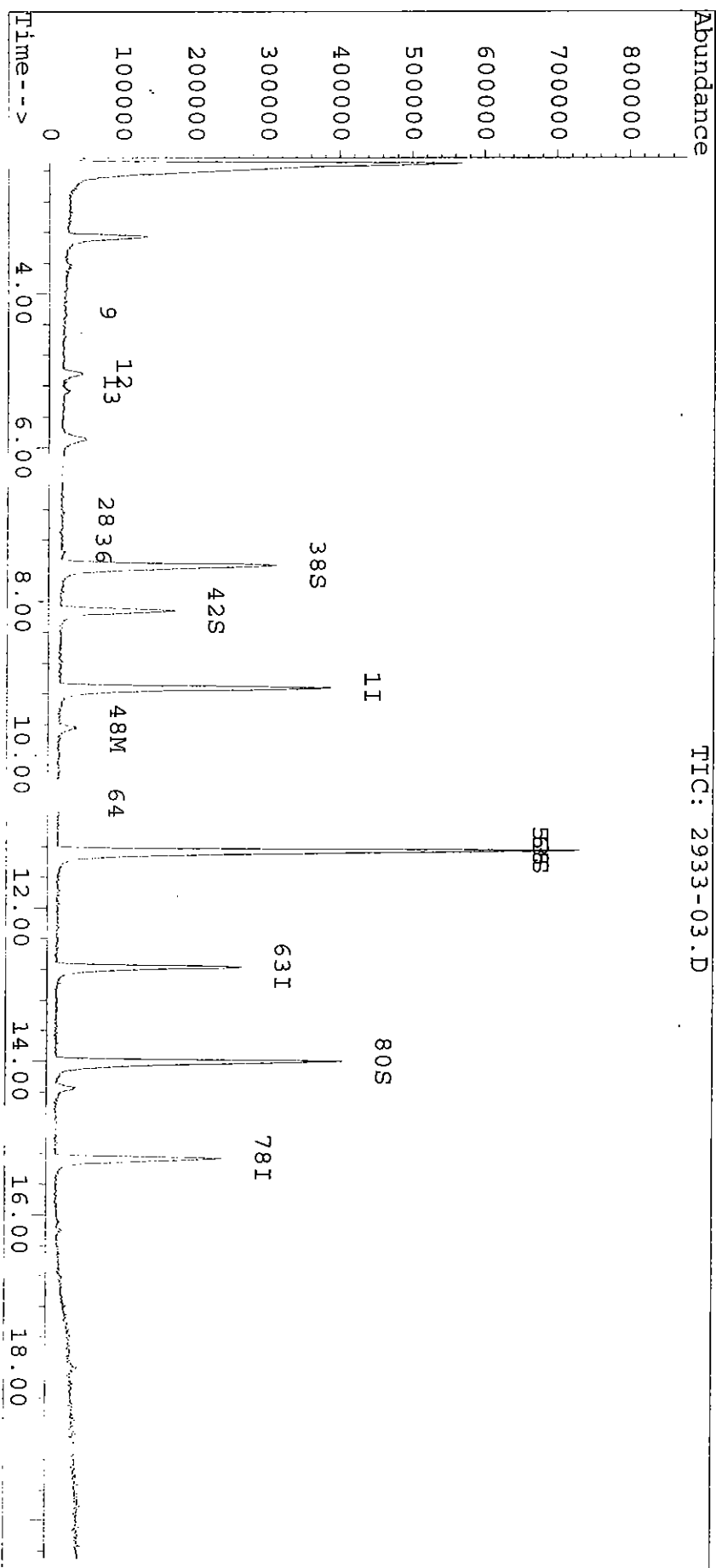
Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2269\2933-03.D
 Acq On : 30 Apr 03 10:28 pm
 Sample : f=1
 Misc :
 Quant Time: Apr 30 22:49 2003

Quant Results File: quant.res

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

Vial: 29
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00



FORM-4A

Applied P & Ch Laboratory

Method Blank Summary for Method 524.2

Client Name: GEOFON, Inc.

Case No:

Project ID: JPL

Contract No:

SAS No:

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2269

Data File Name: G2269K02

Heated Purge: (Y/N) N

Lab Code: APCL

Service ID: 32964

Analysis Date: 04/30/03

Analysis Time: 21:03

Instrument ID: GC/MS: G

GC Column: DB-VEX

Column ID: 0.45 mm

Sample ID: 03G2269-MB-01

Lab Sample ID: 03G2269-MB-01

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

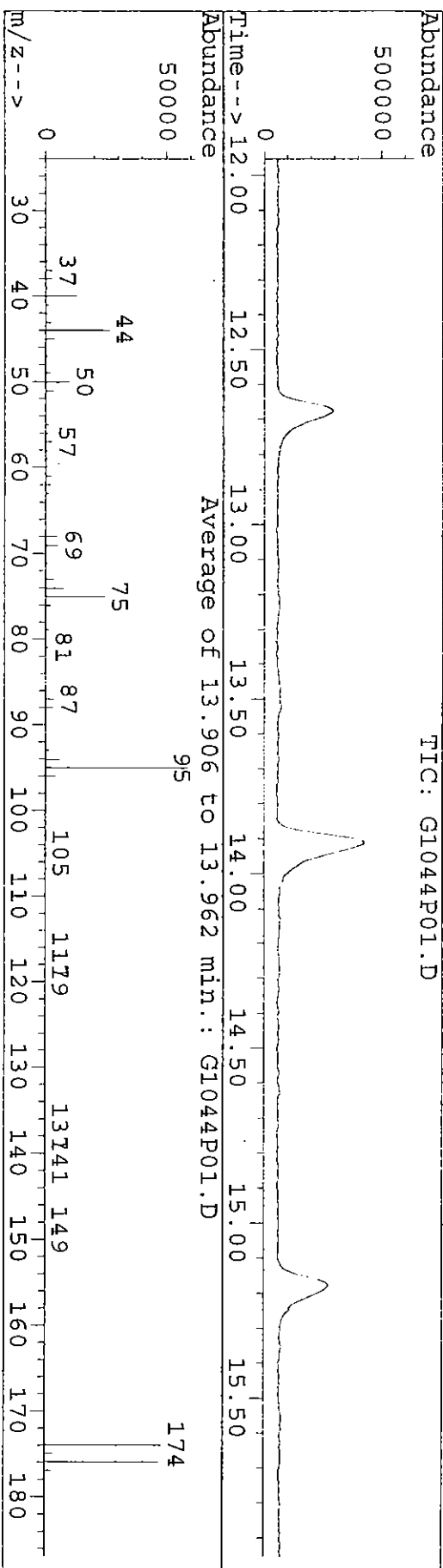
#	Client Sample No	Lab Sample ID	Sample Type	Data Filename	Analysis Date	Analysis Time
1	03G2269-LCS-01	03G2269-LCS-01	Lab Control Spike	G2269L01	04/30/03	18:09
2	MW-17-2MS	03-2933-3MS	Matrix Spike	G2269M01	04/30/03	18:38
3	MW-17-2MSD	03-2933-3MSD	Matrix Spike Duplicate	G2269N01	04/30/03	19:08
4	DUPE-4-2Q03	03-2964-1	Field Sample	2964-01	05/01/03	00:53
5	EB-7-4/29/03	03-2964-2	Field Sample	2964-02	05/01/03	01:21
6	MW-24-1	03-2964-3	Field Sample	2964-03	05/01/03	01:50
7	MW-24-2	03-2964-4	Field Sample	2964-04	05/01/03	02:19
8	MW-24-3	03-2964-5	Field Sample	2964-05	05/01/03	02:48
9	MW-24-4	03-2964-6	Field Sample	2964-06	05/01/03	03:17
10	MW-24-5	03-2964-7	Field Sample	2964-07	05/01/03	03:46
11	TB-7-4/29/03	03-2964-8	Field Sample	2964-08	05/01/03	04:15
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						
23						
24						
25						

BFB

Data File : C:\HPCHEM\1\DATA\03G1044\G1044P01.D
 Acq On : 10 Jan 03 1:51 pm
 Sample : ##03g1044, w
 Misc :

Vial: 16
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2



Peak Apex is scan: 1479

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
50	95	15	40	16.9	9968	PASS
75	95	30	60	41.9	24658	PASS
95	95	100	100	100.0	58860	PASS
96	95	5	9	7.2	4263	PASS
173	174	0	2	0.0	0	PASS
174	95	50	100	82.4	48487	PASS
175	174	5	9	7.2	3468	PASS
176	174	95	101	96.8	46953	PASS
177	176	5	9	6.3	2949	PASS

G1044 L.D E524G003.M

Mon Jan 13 09:24:09 '93

5A
VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name : APPLIED P & CH LAB

Contract: _____

Lab Code: _____ Case No.: _____

SAS No.: _____ SDG No.: 03-2964

Lab File ID: G1044 P01

BFB Injection Date: 01/10/2003

Instrument ID: GCMS-G

BFB Injection Time: 1351

GC Column: VOCOL ID: 0.32 (mm)

Heated Purge: (Y/N) N

m/e	ION ABUNDANCE CRITERIA	%RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	16.9
75	30.0 - 60.0% of mass 95	41.9
95	Base peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	7.2
173	Less than 2.0% of mass 174	0.0 (0.0)1
174	50.0 - 100.0% of mass 95	82.4
175	5.0 - 9.0% of mass 174	5.9 (7.2)1
176	95.0 - 101.0% of mass 174	79.8 (96.8)1
177	5.0 - 9.0% of mass 176	5.0 (6.3)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	003-A0003	003-A0003.D	01/10/03	1427
02	VSTD002	003-002	003-0002.D	01/10/03	1526
03	VSTD010	003-0010	003-0010.D	01/10/03	1555
04	VSTD020	003-0020	003-0020.D	01/10/03	1624
05	VSTD040	003-0040	003-0040.D	01/10/03	1654
06	VSTD080	003-0080	003-0080.D	01/10/03	1723
07					
08					
09					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Mon Jan 13 09:57:17 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =3-0003.D 2 =3-002.D 10 =3-010.D
 20 =3-020.D 40 =3-040.D 80 =3-080.D

Compound		0.3	2	10	20	40	80	Avg	%RSD	
1) I	1 Fluorobenzene I1	---	---	---	ISTD	---	---	---		
2) 3	di-Cl-di-F-metha	0.401	0.288	0.312	0.289	0.285	0.276	0.308	15.17	1.000
3) P	4 Chloromethane		0.201	0.216	0.253	0.278	0.269	0.243	13.88	
4) 9	Fl14 85 135	0.350	0.300	0.334	0.313	0.315	0.304	0.319	5.93	
5) C	5 vinyl chloride	0.217	0.232	0.243	0.233	0.237	0.223	0.231	3.96	
6) 6	bromomethane	0.255	0.283	0.240	0.190	0.198	0.198	0.227	16.60	0.999
7) 7	Chloroethane	0.158	0.177	0.177	0.169	0.175	0.169	0.171	4.25	
8) 8	tri-Cl-F-methane	0.318	0.367	0.432	0.414	0.413	0.398	0.390	10.70	
9) 9	111 isopropyl alcoh		0.005	0.005	0.004	0.005	0.004	0.005	12.57	
10) 10	100 ethyl ether x5	0.177	0.141	0.135	0.133	0.137	0.133	0.143	12.11	
11) 11	102 Acrolein x10		0.010	0.010	0.015	0.012	0.011	0.011	17.19	1.000
12) 12	119 methyl acetate	0.177	0.103	0.110	0.126		0.124	0.128	22.73	
13) 13	104 Carbon disulfid	0.919	0.866	0.830	0.793	0.817	0.777	0.834	6.25	
14) 14	103 Acrylonitrilex1		0.024	0.024	0.024	0.024	0.024	0.024	1.28	
15) 15	95 Acetone x10		0.027	0.017	0.018	0.016	0.016	0.019	25.00	0.999
16) 16	108 F-113	0.232	0.267	0.343	0.325	0.321	0.311	0.300	13.95	
17) 17	M, C 13 11-dichloroethen	0.376	0.399	0.409	0.389	0.398	0.381	0.392	3.13	0.998
18) 18	101 Acetonitrilex1	0.009	0.001	0.007	0.007	0.007	0.006	0.006	44.01	0.996
19) 19	109 Iodomethane	0.170	0.363	0.385	0.320	0.331	0.296	0.311	24.45	0.999
20) 20	113 Tert butyl alco		0.012	0.008	0.009	0.008	0.009	0.009	17.38	
21) 21	18 methylene chlori	1.220	0.724	0.498	0.407	0.365		0.643	54.62	0.993
22) 22	112 Allyl chloride	0.514	0.479	0.422	0.416	0.405	0.365	0.434	12.39	
23) 23	200 Nitro methane x	0.044	0.044	0.044	0.043	0.044	0.042	0.044	1.92	
24) 24	10 t-Bu-Me-ether	0.842	0.479	0.398	0.418	0.420	0.429	0.498	34.33	1.000
25) 25	19 t-12-di-Cl-ethen	0.324	0.294	0.299	0.296	0.301	0.289	0.301	4.05	
26) 26	98 Vinyl acetate x5	0.312	0.304	0.286	0.157	0.315	0.248	0.270	22.48	
27) 27	P 21 11-dichloroethan	0.500	0.546	0.535	0.527	0.534	0.507	0.525	3.36	
28) 28	91 2-butanone MEKx1	0.104	0.095	0.090	0.088	0.094	0.092	0.094	6.20	

(#) = Out of Range
 F524G003.M Mon Jan 13 09:57:29 2003

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Mon Jan 13 09:57:17 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =3-0003.D 2 =3-002.D 10 =3-010.D
 20 =3-020.D 40 =3-040.D 80 =3-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD
29) 115 Di isoprop ethe	0.108	1.340	1.315	1.320	1.351	1.317	1.125	44.32
30) 22 c-12-di-Cl-ethen	0.299	0.300	0.291	0.291	0.297	0.287	0.294	1.83
31) 23 22-Dichloropropa	0.573	0.467	0.437	0.419	0.418	0.403	0.453	13.83
32) 24 Br-Cl-methane	0.042	0.099	0.104	0.104	0.108	0.105	0.094	27.43
33) 25 chloroform	0.648	0.531	0.493	0.488	0.504	0.487	0.525	11.86
34) 201 Ethyl acetate x	0.144	0.137	0.100	0.151	0.136	0.134	0.134	14.75
35) 116 ETBE	0.910	0.810	0.787	0.780	0.785	0.769	0.807	6.50
36) 117 Iso-butyl alcoh	0.024	0.029	0.029	0.020	0.031	0.027	0.027#	14.60
37) 26 tetrahydrofuranx	0.011	0.010	0.010	0.010	0.010	0.010	0.010#	4.18
38) S 27 Di-Br-F-Methane	0.604	0.385	0.352	0.350	0.360	0.349	0.400	25.18
39) 34 111-tri-Cl-ethan	0.542	0.428	0.418	0.409	0.414	0.413	0.437	11.85
40) 30 12-dichloroethan	0.024	0.205	0.204	0.201	0.208	0.205	0.175	42.30
41) 35 11-Di-Cl-propene	0.413	0.378	0.379	0.371	0.378	0.373	0.382	4.04
42) S 29 1,2-di-Cl-ethane	0.173	0.165	0.164	0.164	0.165	0.159	0.165	3.14
43) M 36 benzene	0.854	0.901	0.876	0.863	0.883	0.837	0.869	2.63
44) 37 CCl4	0.341	0.311	0.354	0.334	0.340	0.329	0.335	4.32
45) 97 thiophene	0.441	0.456	0.429	0.421	0.430	0.415	0.432	3.39
46) 118 TAME	0.635	0.539	0.536	0.527	0.527	0.541	0.556	8.06
47) C 39 12-di-Cl-propane	0.255	0.253	0.252	0.254	0.257	0.253	0.254	0.66
48) M 40 trichloroethene	0.253	0.267	0.275	0.273	0.279	0.269	0.269	3.39
49) 96 Me-methacrylate	0.128	0.086	0.083	0.087	0.092	0.092	0.095	19.43
50) 42 Br-di-Cl-methane	0.441	0.364	0.340	0.327	0.331	0.324	0.355	12.65
51) 41 dibromomethane	0.100	0.117	0.128	0.132	0.133	0.127	0.123	10.29
52) 45 c-13-di-Cl-prope	0.341	0.316	0.303	0.301	0.309	0.307	0.313	4.71
53) S 55 toluene-d8(S2)	0.509	0.500	0.491	0.504	0.493	0.500	0.500	1.57
54) 92 2-ClEt-Vi-ether1	0.045	0.052	0.054	0.055	0.057	0.057	0.054	8.68
55) M C 56 toluene	0.926	0.820	0.796	0.777	0.794	0.786	0.817	6.80
56) 107 Et methacrylate	0.232	0.176	0.112	0.112	0.178	0.147	0.169	26.25
57) 93 2-Hexanone x5	0.062	0.058	0.050	0.058	0.057	0.057	0.057	7.78

(#) = Out of Range
 F524G003.M Mon Jan 13 09:57:32 2003

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Mon Jan 13 09:57:17 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =3-0003.D 2 =3-002.D 10 =3-010.D
 20 =3-020.D 40 =3-040.D 80 =3-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD
58) 48 112-tri-Cl-Et	0.226	0.143	0.120	0.116	0.121	0.118	0.141	30.67
59) 58 1,2-di-br-ethane	0.050	0.125	0.124	0.129	0.129	0.128	0.114	27.70
60) 51 di-Br-Cl-methane	0.117	0.177	0.166	0.165	0.170	0.172	0.161	13.59
61) 46 t-13-di-Cl-prope	0.200	0.197	0.206	0.204	0.206	0.197	0.202	2.11
62) 105 1-Chlorohexane	0.564	0.285	0.272	0.253	0.249	0.237	0.310	40.49
63) I 47 Cl-benzene-d5, I2	0.207	0.298	0.328	0.319	0.325	0.304	0.297	15.38
64) 54 MIBK	0.624	0.665	0.699	0.690	0.647	0.598	0.654	5.95
65) 49 1,3-di-Cl-propan	0.767	0.892	0.960	0.927	0.884	0.850	0.880	7.60
66) 59 tetra-Cl-ethene	1.465	1.556	1.560	1.485	1.424	1.278	1.461	7.13
67) M P 60 chlorobenzene	0.473	0.619	0.613	0.631	0.602	0.573	0.585	9.96
68) 61 112-tetra-Cl-Et	3.150	3.064	2.822	2.754	2.619	2.485	2.816	9.06
69) C 64 ethylbenzene	2.309	2.321	2.213	2.124	2.010	1.868	2.141	8.30
70) 65 m/p-Xylenes x2	0.809	0.710	0.649	0.615	0.570	0.518	0.645	16.06
71) 99 1-4-di-Cl-butane	0.118	0.305	0.317	0.319	0.302	0.282	0.274	28.23
72) P 52 bromoform	1.595	1.641	1.561	1.507	1.441	1.326	1.512	7.58
73) 66 styrene	2.457	2.410	2.147	2.035	1.889	1.700	2.106	13.98
74) 67 o-xylene	0.391	0.469	0.403	0.385	0.349	0.323	0.387	12.94
75) P 68 112-Tetra-Cl-Et	0.153	0.078	0.060	0.058	0.053	0.081	0.081	51.75
76) 110 t-1,4-dichloro-	0.890	0.534	0.539	0.531	0.489	0.448	0.572	27.91
77) 106 Cl-benzyl								
78) I 62 1,4-DCB-d4 150 152	0.112	0.116	0.123	0.127	0.131	0.122	0.122	6.42
79) 69 123-tri-Cl-Pr	0.935	0.877	0.886	0.890	0.880	0.893	0.893	2.69
80) S 70 4-Br-1-F-Bz (S3)	2.780	3.439	3.399	3.298	3.366	3.435	3.286	7.71
81) 71 isopropylbenzene	0.527	0.700	0.760	0.772	0.787	0.782	0.721	13.87
82) 72 bromobenzene	0.727	0.924	0.921	0.910	0.914	0.920	0.886	8.83
83) 73 n-propylbenzene	0.646	0.602	0.566	0.545	0.606	0.566	0.589	6.23
84) 74 2-Cl-Tl 126								

1.0000

0.449

0.449

0.448

0.449

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P &Ch Lab** EPA 524.2
 Last Update : Mon Jan 13 09:57:17 2003
 Response via : Initial Calibration

Calibration Files
 0.3 =3-0003.D 2 =3-002.D 10 =3-010.D
 20 =3-020.D 40 =3-040.D 80 =3-080.D

Compound	0.3	2	10	20	40	80	Avg	%RSD
85) 75 4-Cl-Tl 126	0.683	0.834	0.866	0.784	0.843	0.816	0.804	8.13
86) 76 135-tri-Me-Bz	2.482	2.722	2.700	2.625	2.675	2.716	2.653	3.43
87) 79 tert-butylbenzen	2.218	2.871	2.824	2.765	2.829	2.837	2.724	9.19
88) 78 124-tri-Me-Bz	1.891	2.455	2.250	2.296	2.326	2.317	2.256	8.48
89) 80 13-di-Cl-Bz 146	0.934	1.158	1.264	1.084	1.198	1.258	1.149	10.87
90) 82 14-di-Cl-Bz 146	1.527	1.764	1.616	1.744	1.694	1.675	1.670	5.24
91) 81 sec-butylbenzene	3.330	4.148	4.224	3.968	4.057	4.172	3.983	8.35
92) 77 4-iso-Pr-toluene	2.419	3.039	3.032	2.939	2.928	2.886	2.874	8.04
93) 84 12-di-Cl-benzene	0.920	1.150	1.163	1.111	1.152	1.156	1.109	8.49
94) 85 n-butylbenzene	3.034	2.990	3.069	2.970	2.978	3.028	3.012	1.28
95) 86 12-diBr-3-Cl-Pra	0.063	0.077	0.085	0.089	0.091	0.091	0.081	13.95
96) 87 124-tri-Cl-Bz	0.534	0.749	0.838	0.873	0.858	0.901	0.792	17.24
97) 88 naphthalene	0.804	0.553	0.616	0.666	0.652	0.700	0.665	12.68
98) 90 123-tri-Cl-Bz	0.497	0.559	0.642	0.649	0.636	0.642	0.604	10.29
99) 89 hx-Cl-butadiene	0.432	0.625	0.690	0.663	0.662	0.652	0.621	15.29

0.449

1.577

(#) = Out of Range
 524G003.M

Mon Jan 13 09:57:36 2003

INITIAL CALIBRATION SUMMARY

Method File

E524G003

Last Calibration Update

Mon Jan 13 09:57:17 2003

Level 1 File Name	3-0003.D	Level 1 ID	0.3
Level 2 File Name	3-002.D	Level 2 ID	2
Level 3 File Name	3-010.D	Level 3 ID	10
Level 4 File Name	3-020.D	Level 4 ID	20
Level 5 File Name	3-040.D	Level 5 ID	40
Level 6 File Name	3-080.D	Level 6 ID	80
Level 7 File Name	3-020.D	Level 7 ID	cc

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
1 Fluorobenzene 11 1	765834	768596	755237	739905	721219	703721	-1	-----	-----	-----	-----
3 di-Cl-di-F-methane 85 87	9202	44272	235359	427426	820884	1551879	-1	0.0191	0.2750	0.0000	0.9996
4 Chloromethane 50 52	4751	30836	163038	374044	803175	1514667	-1	0.0000	0.2433	0.0000	0.1388
9-F114 85 135	8035	46094	252154	462947	909235	1714061	-1	0.0000	0.3193	0.0000	0.0593
5 Vinyl chloride 62 64	4990	35625	183259	344227	682967	1258237	-1	0.0000	0.2307	0.0000	0.0396
6 bromomethane 94 96	5861	43488	181174	281305	570773	1116110	-1	0.0126	0.1961	0.0000	0.9990
7 Chloroethane 64 66	3629	27223	133510	250119	505034	952606	-1	0.0000	0.1709	0.0000	0.0425
8-tri-Cl-F-methane 101 103	7295	56379	326202	612174	1192828	2238212	-1	0.0000	0.3902	0.0000	0.1070
111 isopropyl alcohol x10	1236	8181	34708	55395	131600	244204	-1	0.0000	0.0045	0.0000	0.1257
100 ethyl ether x5	20361	108089	509174	980952	1975186	3742221	-1	0.0000	0.1425	0.0000	0.1211
102 Acrolein x10	3234	15074	75489	215710	340415	601157	-1	0.0000	0.0114	0.0000	0.1719
119 methyl acetate	4062	15788	82815	187169	-1	696519	-1	-0.0033	0.1242	0.0000	0.9997
104 Carbon disulfide	21121	133095	626926	1173322	2357195	4372134	-1	0.0000	0.8336	0.0000	0.0625
103 Acrylonitrile x10	4569	37044	178667	348877	701313	1341637	-1	0.0000	0.0239	0.0000	0.0128
95 Acetone x10	989	41993	128289	268909	465240	912905	-1	0.0227	0.0159	0.0000	0.9991
108 F-113	5327	41068	259145	480691	925581	1749444	-1	0.0000	0.2998	0.0000	0.1395
13 11-dichloroethene 61 96	8641	61263	309126	575822	1147783	2146767	-1	0.0000	0.3920	0.0000	0.0313
101 Acetonitrile x10	2063	1527	53569	97490	189257	353648	-1	0.0013	0.0063	0.0000	0.9985
109 Iodomethane	3901	55824	290992	473953	954242	1666625	-1	0.0467	0.2964	0.0000	0.9961
113 Tert butyl alcohol x10	2556	18436	60720	133851	240444	485913	-1	0.0016	0.0086	0.0000	0.9991
18 methylene chloride 49 84	28023	111315	375753	602942	1052979	-1	-1	0.0818	0.3520	0.0000	0.9932

112 Allyl chloride	11807	73643	318868	616004	1168854	2056270	-1	0.0000	0.4336	0.0000	0.1239
200 Nitro methane x10	10089	68361	334791	633827	1264975	2385881	-1	0.0000	0.0436	0.0000	0.0192
10 t-Bu-Me-ether 73 57	19344	73627	300489	618307	1211982	2415631	-1	-0.0086	0.4281	0.0000	0.9997
19 t-12-di-Cl-ethene 96 61	7446	45222	225516	438020	867973	1629626	-1	0.0000	0.3005	0.0000	0.0405
98 Vinyl acetate x5	35861	233914	1080092	1161950	4543111	6984572	-1	0.0000	0.2704	0.0000	0.2248
21 11-dichloroethane 63 83	11490	83925	403704	780313	1539657	2853804	-1	0.0000	0.5248	0.0000	0.0336
91 2-butanone MEKx10	23960	146366	680153	1296553	2708302	5159574	-1	0.0000	0.0938	0.0000	0.0620
115 Di isoprop ether	2475	206054	993426	1953682	3898884	7414434	-1	0.0032	1.3231	0.0000	0.9998
22 c-12-di-Cl-ethene 96 61	6875	46067	219592	430734	857123	1613152	-1	0.0000	0.2941	0.0000	0.0183
23 22-Dichloropropane 77 97	13155	71811	330092	619878	1207116	2269373	-1	0.0000	0.4529	0.0000	0.1383
24 Br-Cl-methane 128 130	956	15248	78733	153832	312235	593248	-1	-0.0008	0.1060	0.0000	0.9998
25 chloroform 83 85	14881	81604	372529	722194	1453363	2739754	-1	0.0000	0.5251	0.0000	0.1186
201 Ethyl acetate x2	-1	44403	206670	296783	873148	1526995	-1	0.0000	0.1337	0.0000	0.1475
116 ETBE	20917	124488	594002	1154600	2264664	4329036	-1	0.0000	0.8068	0.0000	0.0650
117 Iso-butyl alcohol X10	5582	44403	221032	297076	880890	1546333	-1	0.0000	0.0268	0.0000	0.1460
26 tetrahydrofuranx5	1945	8407	36840	75144	146600	287877	-1	0.0000	0.0102	0.0000	0.0418
27 Di-Bi-F-Methane (S1) 111 1	13867	59189	265788	517816	1038870	1963588	-1	0.0096	0.3493	0.0000	0.9997
34 111-tr-Cl-ethane 97 99	12459	65859	315664	604949	1193974	2324189	-1	0.0000	0.4374	0.0000	0.1185
30 12-dichloroethane 64 62	549	31555	154021	298086	600265	1155412	-1	-0.0025	0.2060	0.0000	0.9999
35 11-Di-Cl-propene 75 110	9488	58117	286388	549558	1091787	2099663	-1	0.0000	0.3822	0.0000	0.0404
29 1,2-di-Cl-ethane-d4 [Surf] 10	-1	26643	124715	242470	475685	894560	-1	0.0000	0.1652	0.0000	0.0314
36 benzene 78 52	19613	138554	661631	1277049	2547937	4710056	-1	0.0000	0.8690	0.0000	0.0263
37 CCl4 117 119	7832	47773	267339	493900	981263	1852391	-1	0.0000	0.3348	0.0000	0.0432
97 thiophene	10129	70133	323713	623624	1239916	2337761	-1	0.0000	0.4320	0.0000	0.0339
118 TAME	4367	97622	406861	792558	1521011	3045504	-1	0.0000	0.5555	0.0000	0.0806
39 12-di-Cl-propane 63 76	5869	38931	190695	376101	741594	1426638	-1	0.0000	0.2543	0.0000	0.0066
40 trichloroethene 130 132	5805	41064	207771	403682	804090	1515012	-1	0.0000	0.2693	0.0000	0.0339
96 Me-methacrylate	1472	19691	65318	123371	250724	519513	-1	-0.0068	0.0920	0.0000	0.9983
42 Br-di-Cl-methane 83 85	10138	55994	256691	484084	955698	1823535	-1	0.0000	0.3546	0.0000	0.1265
41 dibromomethane 174 172	2292	17911	96884	194764	383350	714384	-1	0.0000	0.1227	0.0000	0.1029

Compound	Level 1 Response	Level 2 Response	Level 3 Response	Level 4 Response	Level 5 Response	Level 6 Response	Level 7 Response	Coeff X^0	Coeff X^1 / ave RF	Coeff X^2	R^2 / RSD
45 c-13-di-Cl-propene 75 110	7832	48533	229072	445153	890451	1727952	-1	0.0000	0.4995	0.0000	0.0157
55 toluene-d8(S2) 100 99	-1	78299	377922	725871	1455135	2774904	-1	0.0000	0.0536	0.0000	0.0868
92 2-ClEt-Vi-ether10	10352	80060	410701	820365	1655105	3234891	-1	0.0000	0.8167	0.0000	0.0680
56 toluene 91 92	21278	126101	601438	1149709	2291459	4425814	-1	0.0000			

107 Et methacrylate	-1	35707	132945	165590	512644	828103	-1	0.0000	0.1690	0.0000	0.2625
93 2-Hexanone x5	10744	47646	218029	368204	831419	1601350	-1	0.0000	0.0568	0.0000	0.0778
48 112-tri-Cl-Et	5203	21908	90597	172036	347876	664699	-1	0.0031	0.1179	0.0000	0.9998
58 1,2-di-br-ethane	107 109	1144	19207	94012	191133	371448	-1	-0.0016	0.1288	0.0000	1.0000
51 di-Br-Cl-methane	129 127	2693	27173	125300	243477	489254	-1	0.0000	0.1610	0.0000	0.1359
46 t-13-di-cl-propene	75 110	4584	30276	155530	301394	594704	-1	0.0000	0.2015	0.0000	0.0211
105 1-Chlorohexane		12961	43864	205359	374248	719249	-1	0.0247	0.2364	0.0000	0.9994
47 Cl-benzene-d5, 12	226795	229493	225971	224981	233528	241678	-1	0.0000	1.0000	0.0000	0.0000
54 MIBK	1407	13676	74160	143757	303184	587112	-1	0.0184	0.3053	0.0000	0.9988
49 1,3-di-cl-propane	76 78	4247	30516	157929	310529	604220	-1	0.0000	0.6537	0.0000	0.0595
59 tetra-Cl-ethene	166 168	5221	40927	216892	417298	825820	-1	0.0000	0.8800	0.0000	0.0760

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
60 chlorobenzene	9971	71440	352480	668143	1330525	2470738	-1	0.0000	0.5853	0.0000	0.0996
61 1112-tetra-Cl-Et	131 133	3219	28400	138607	283855	562377	1108773	-1	0.0000	0.0000	0.0906
64 ethylbenzene	91 106	21430	140640	637777	1239073	2446417	4804958	-1	0.0000	0.0000	0.0830
65 m/p-Xylenes x2		31419	213036	1000145	1911139	3754507	7222435	-1	0.0000	0.0000	0.1606
99 1-4-di-Cl-butane		5503	32572	146606	276558	532573	1001628	-1	0.0000	0.6450	0.0980
52 bromoform	173 175	806	13982	71698	143599	281887	544898	-1	0.0303	0.2823	0.0758
66 styrene	104 78	10850	75317	352645	677932	1346269	2563056	-1	0.0000	0.0000	0.1398
67 o-xylene	91 106	16715	110603	485194	915641	1764503	3287299	-1	0.0000	0.3868	0.1294
68 1122-Tetra-Cl-Et	83 85	2658	21540	91096	173152	326262	625224	-1	0.0235	0.0507	0.9991
110 t-1,4-dichloro-2-butene		1095	7032	17679	27009	54200	103110	-1	0.0784	0.4469	0.9969
106 Cl-benzyl		6053	24517	121704	238713	457227	866531	-1	0.0000	0.0000	0.0642
62 1,4-DCB-d4	150 152 13	197143	197148	188831	184633	176207	164785	-1	0.0000	1.0000	0.0771
69 123-tri-Cl-Pr	110 97	253	4401	21971	45443	89331	172722	-1	0.0000	0.8935	0.0269
70 4-Br-1-F-Bz (S3)	174 95	-1	36886	165570	327081	627145	1159540	-1	0.0000	0.0000	0.1387
71 Isopropylbenzene	105 120	16442	135584	641793	1217921	2372165	4528346	-1	0.0000	3.2861	0.0883
72 bromobenzene	156 158	3119	27613	143515	285095	554412	1030644	-1	0.0000	0.7214	0.0623
73 n-propylbenzene	120 78	4297	36443	173970	335858	643879	1212584	-1	0.0000	0.8858	0.0813
74 2-Cl-Tl	126 128	3821	23738	106793	201349	427470	745993	-1	0.0000	0.8045	0.0343
75 4-Cl-Tl	126 128	4042	32882	163598	289530	594245	1075809	-1	0.0000	2.6533	0.0919
76 135-tri-Me-Bz	105 120	14680	107323	509842	969387	1885346	3580223	-1	0.0000	2.7239	0.0848
79 tert-butylbenzene	119 91	13115	113211	533176	1020985	1994231	3739615	-1	0.0000	2.2558	0.1087
78 124-tri-Me-Bz	105 120	11184	96787	424953	847796	1639260	3054447	-1	0.0000	1.1491	
80 13-di-Cl-Bz	146 148	5522	45648	238597	400250	844281	1658291	-1	0.0000		

82 14-di-Cl-Bz	146 148	9029	69544	305231	644124	1193857	2207723	-1	0.0000	1.6699	0.0000	0.0524
81 sec-butylbenzene	105 134	19695	163551	797663	1465228	2859247	5499576	-1	0.0000	3.9831	0.0000	0.0835
77 4-iso-Pr-toluene	119 134	14306	119820	572630	1085210	2064029	3804782	-1	0.0000	2.8739	0.0000	0.0804
84 12-di-Cl-benzene	146 148	5444	45355	219614	410082	812246	1524040	-1	0.0000	1.1088	0.0000	0.0849
85 n-butylbenzene	91 134	17944	117913	579573	1096665	2099124	3991637	-1	0.0000	3.0116	0.0000	0.0128
86 12-diBr-3-Cl-Pra	157 155	-1	2498	14611	31395	62914	120318	-1	0.0000	0.0813	0.0000	0.1395
87 124-tri-Cl-Bz	180 182	3158	29532	158333	322231	604462	1187497	-1	-0.0539	0.9002	0.0000	0.9995
88 naphthalene	128 129	4753	21798	116373	245906	459400	922945	-1	0.0000	0.6651	0.0000	0.1268
90 123-tri-Cl-Bz	180 182	2940	22060	121207	239739	448256	845940	-1	0.0000	0.6042	0.0000	0.1029
89 hx-Cl-butadiene	225 260	2553	24644	130257	244690	466707	859594	-1	0.0135	0.6525	0.0000	0.9999

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-0003.D
Method : C:\HPCHEM\1\METHODS\E524G003.M
Acq. Time : Jan 10 14:27 2003
Method Update: Mon Jan 13 10:38 2003
Quant. Time : Jan 13 09:28 2003
Print Time : Mon Jan 13 10:41 2003
Miscellaneous :
Sample : f=1
Inst. : GCMS-G
RF via : Multiple Level Calibration
Operator: Eddie
Multiplier: 1.000000

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.04	9.04	0.000	96	70	765.834	10.00			
47	Cl-benzene-d5, I2	12.67	12.66	0.000	82	119	226.795	10.00			
62	1,4-DCB-d4 150 15	15.16	15.16	0.000	152	150	197.143	10.00			
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.45	7.44	0.000	111	113	13.867	0.45		0.5	2.26%
29	1,2-di-Cl-ethane-	8.03	8.03	0.000	65	102	10.557	0.64		0.6	3.22%
55	toluene-d8(S2)	11.16	11.15	0.000	100	99	18.222	0.43		0.4	2.17%
70	4-Br-1-F-Bz (S3)	13.91	13.90	0.000	174	95	9.453	0.47		0.5	2.37%

Target Compounds

<<< I1 : ISTD ID = 1 >>>

Qvalue

3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	9.202	0.39		0.4	63	#
4	Chloromethane	2.79	2.79	0.000	50	52	4.751	0.25		0.3	55	#?
9	F114 85 135	2.82	2.84	-0.003	85	135	8.035	0.33		0.3	69	?
5	vinyl chloride	2.97	2.97	0.000	62	64	4.990	0.28		0.3	54	#
6	bromomethane	3.39	3.38	0.000	94	96	5.861	0.34		0.3	97	#
7	Chloroethane	3.52	3.54	-0.002	64	66	3.629	0.28		0.3	56	#
8	tri-Cl-F-methane	4.16	4.16	0.000	101	103	7.295	0.25		0.2	76	#
111	isopropyl alcoho	4.31	4.28	0.003	45	43	1.236	3.99		4.0	60	#
100	ethyl ether x5	4.45	4.45	0.000	59	74	20.361	1.87		1.9	77	#
102	Acrolein x10	4.19	4.17	0.002	56	55	3.234	3.71		3.7	70	#
119	methyl acetate	5.08	5.07	0.000	43	74	4.062	0.81		0.8	56	#?
104	Carbon disulfide	5.20	5.19	0.000	76	78	21.121	0.33		0.3	78	#
103	Acrylonitrilex10	4.94	4.91	0.004	53	52	4.569	2.49		2.5	66	m
95	Acetone x10	4.24	4.33	-0.009	43	58	0.989	0.67		0.7	98	#
108	F-113	5.04	5.05	-0.001	151	101	5.327	0.26		0.3	67	#
13	1,1-dichloroethene	4.78	4.78	0.000	61	96	8.641	0.29		0.3	80	?
101	Acetonitrilex10	4.23	4.22	0.001	41	40	2.063	5.25		5.2	1	#?
109	Iodomethane	4.79	4.82	-0.003	142	127	3.901	0.16		0.2	53	#?
113	Tert butyl alcoh	4.83	4.87	-0.004	59	57	2.556	3.73		3.7	1	#

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-0003.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 14:27 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:28 2003
 Print Time : Mon Jan 13 10:41 2003
 Miscellaneous :

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

4973

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	28.023	0.81	0.8	89	
112	Allyl chloride	5.08	5.08	0.000	41	76	11.807	0.44	0.4	72	#?
200	200 Nitro methane x1	5.83	5.83	0.000	61	46	10.089	3.02	3.0	92	#?
10	10 t-Bu-Me-ether	6.02	6.01	0.001	73	57	19.344	0.51	0.5	98	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	7.446	0.32	0.3	93	?
98	98 Vinyl acetate x5	6.41	6.41	0.000	43	86	35.861	1.71	1.7	93	#
21	21 11-dichloroethane	6.16	6.16	0.000	63	83	11.490	0.29	0.3	98	
91	91 2-butanone MEKx10	6.86	6.84	0.002	43	72	23.960	4.00	4.0	84	#
115	115 Di isoprop ether	6.65	6.85	-0.023	45	87	2.475	0.03	0.0	99	
22	22 c-12-di-Cl-ethene	6.98	6.97	0.000	96	61	6.875	0.30	0.3	96	
23	23 22-Dichloropropan	7.36	7.35	0.000	77	97	13.155	0.38	0.4	98	
24	24 Br-Cl-methane	7.21	7.19	0.002	128	130	0.956	0.13	0.1	0	
25	25 chloroform	7.28	7.27	0.000	83	85	14.881	0.46	0.5	96	m
201	201 Ethyl acetate x2	7.34	7.31	0.003	43	61	5.585	0.53	0.5	43	#?
116	116 ETBE	7.41	7.40	0.000	59	87	20.917	0.34	0.3	99	#?
117	117 Iso-butyl alcoho	7.34	7.31	0.003	43	42	5.582	2.89	2.9	1	#?
26	26 tetrahydrofuranx5	7.73	7.71	0.002	72	42	1.945	2.48	2.5	22	
34	34 111-tri-Cl-ethane	8.24	8.23	0.000	97	99	12.459	0.37	0.4	95	
30	30 12-dichloroethane	8.16	8.13	0.003	62	64	0.549	0.04	0.0	77	
35	35 11-Di-Cl-propene	8.48	8.48	0.000	75	110	9.488	0.32	0.3	88	
36	36 benzene	8.76	8.76	0.000	78	52	19.613	0.29	0.3	92	
37	37 CCl4	8.68	8.69	0.000	117	119	7.832	0.31	0.3	92	
97	97 thiophene	8.90	8.89	0.000	84	58	10.129	0.31	0.3	94	
118	118 TAME	8.93	9.01	-0.008	73	43	4.367	0.10	0.1	99	
39	39 12-di-Cl-propane	9.51	9.50	0.001	63	76	5.869	0.30	0.3	63	#
40	40 trichloroethene	9.56	9.55	0.001	130	132	5.805	0.28	0.3	94	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	1.472	0.20	0.2	33	#
42	42 Br-di-Cl-methane	9.60	9.62	-0.002	83	85	10.138	0.46	0.5	98	
41	41 dibromomethane	9.46	9.45	0.001	174	172	2.292	0.24	0.2	28	#
45	45 c-13-di-Cl-propen	10.37	10.37	0.000	75	110	7.832	0.33	0.3	60	#
92	92 2-ClEt-Vi-ether10	10.15	10.15	0.000	63	43	10.352	2.52	2.5	75	#
56	56 toluene	11.24	11.23	0.000	91	92	21.278	0.34	0.3	87	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-0003.D
Method : C:\HPCHEM\1\METHODS\E524G003.M
Acq. Time : Jan 10 14:27 2003
Method Update: Mon Jan 13 10:38 2003
Quant. Time : Jan 13 09:28 2003
Print Time : Mon Jan 13 10:41 2003
Miscellaneous :
Sample : f=1
Inst. : GCMS-G
RF via : Multiple Level Calibration
Operator: Eddie
Multiplier: 1.000000

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.51	11.37	0.016	69	99	0.955	0.08	0.1	83	?
93	2-Hexanone x5	11.49	11.49	0.000	43	58	10.744	3.07	3.1	92	?
48	1,2-tri-Cl-Et	11.03	11.03	0.000	97	83	5.203	0.62	0.6	70	#
58	1,2-di-br-ethane	11.84	11.83	0.001	107	109	1.144	0.13	0.1	16	#
51	di-Br-Cl-methane	11.59	11.57	0.002	129	127	2.693	0.22	0.2	97	#
46	t-13-di-Cl-propen	10.89	10.86	0.003	75	110	4.584	0.30	0.3	66	#
105	1-Chlorohexane	12.65	12.64	0.001	55	93	12.961	0.77	0.8	41	?
<<< 12 : ISTD ID = 47 >>>											
54	MIBK	10.47	10.52	-0.004	43	58	1.407	0.28	0.3	94	#
49	1,3-di-Cl-propane	11.31	11.29	0.002	76	78	4.247	0.29	0.3	42	#
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	5.221	0.26	0.3	99	#
60	chlorobenzene	12.70	12.70	0.000	112	77	9.971	0.36	0.4	48	#
61	1,1,2-tetra-Cl-Et	12.63	12.63	0.000	131	133	3.219	0.24	0.2	98	?
64	ethylbenzene	12.91	12.91	0.000	91	106	21.430	0.25	0.2	84	#
65	m/p-Xylenes x2	13.11	13.11	0.000	91	106	31.419	0.65	0.6	89	#
99	1-4-di-Cl-butane	13.48	13.46	0.002	55	41	5.503	0.46	0.5	59	#
52	bromoforn	13.26	13.23	0.003	173	175	0.806	0.13	0.1	30	m
66	styrene	13.43	13.43	0.000	104	78	10.850	0.32	0.3	85	#
67	o-xylene	13.51	13.50	0.000	91	106	16.715	0.35	0.3	76	#
68	1,1,2,2-Tetra-Cl-Et	13.53	13.50	0.002	83	85	2.658	0.44	0.4	87	?
110	t-1,4-dichloro-2	13.69	13.68	0.000	89	53	1.095	1.21	1.2	1	#
106	Cl-benzyl	15.13	15.13	0.000	91	126	6.053	0.62	0.6	61	#?
<<< 13 : ISTD ID = 62 >>>											
71	isopropylbenzene	13.87	13.85	0.001	105	120	16.442	0.25	0.3	87	#
72	bromobenzene	14.11	14.10	0.000	156	158	3.119	0.22	0.2	98	#
73	n-propylbenzene	14.29	14.28	0.000	120	78	4.297	0.23	0.2	56	#
74	2-Cl-Tl	14.40	14.38	0.001	126	128	3.821	0.33	0.3	39	m
75	4-Cl-Tl	14.46	14.45	0.000	126	128	4.042	0.31	0.3	43	m
76	1,3,5-tri-Me-Bz	14.57	14.56	0.000	105	120	14.680	0.28	0.3	84	#
79	tert-butylbenzene	14.85	14.84	0.000	119	91	13.115	0.24	0.2	81	#
78	1,2,4-tri-Me-Bz	14.94	14.94	0.000	105	120	11.184	0.25	0.3	97	#
80	1,3-di-Cl-Bz	15.12	15.12	0.000	146	148	5.522	0.20	0.2	80	?

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


```

Data Filename: C:\HPCHEM\1\DATA\03GI
Method        : C:\HPCHEM\1\METHODS\E
Acq. Time     : Jan 10 14:27 2003
Method Update: Mon Jan 13 10:38 2003
Quant. Time   : Jan 13 09:28 2003
Print Time    : Mon Jan 13 10:41 2003
Miscellaneous :

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

4975

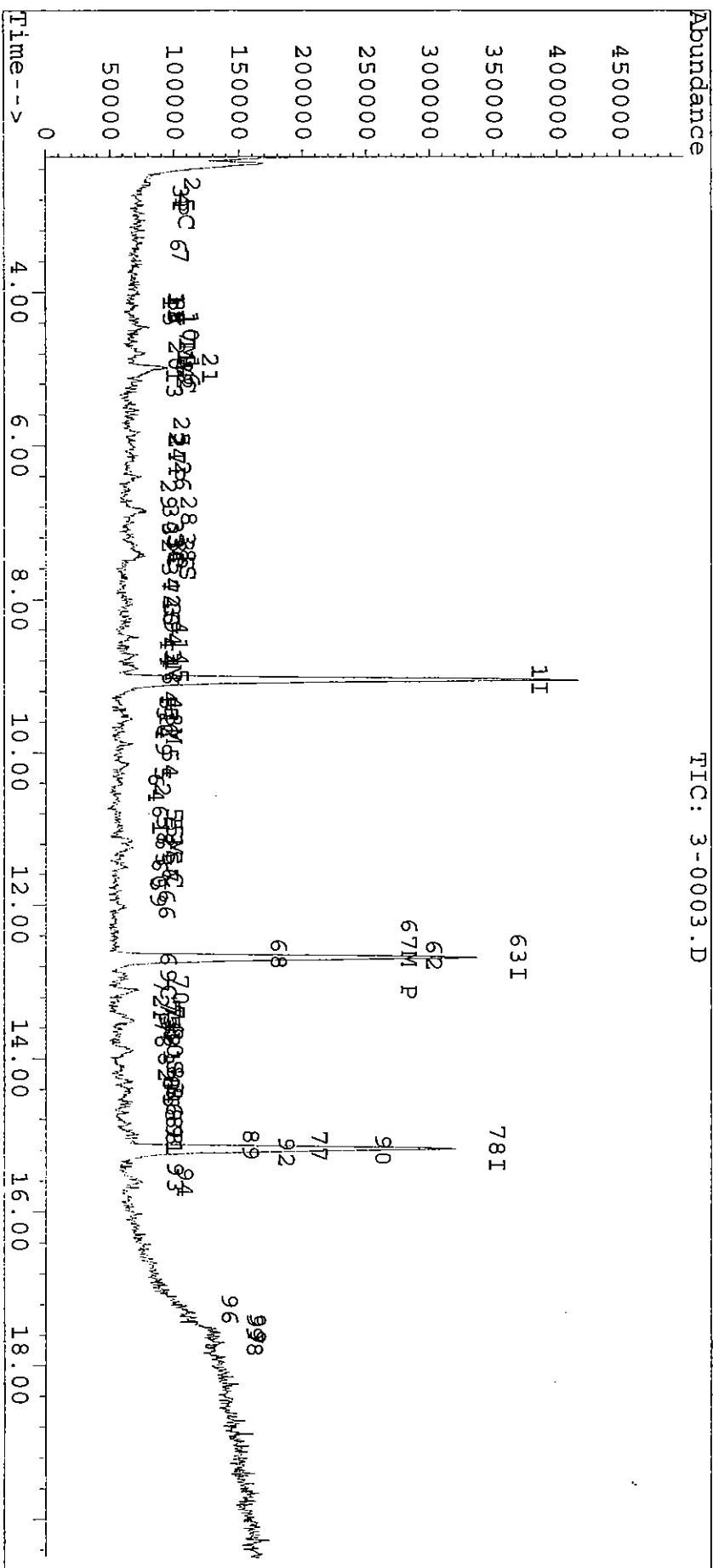
ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
82	14-di-Cl-Bz	146	15.20	15.18	0.001	146	148	9.029	0.29	0.3	1	m?
81	sec-butylbenzene		15.06	15.05	0.000	105	134	19.695	0.28	0.3	70	#
77	4-Iso-Pr-toluene		15.23	15.22	0.000	119	134	14.306	0.25	0.3	91	?
84	12-di-Cl-benzene		15.54	15.53	0.000	146	148	5.444	0.25	0.2	91	
85	n-butylbenzene		15.61	15.61	0.000	91	134	17.944	0.36	0.4	73	#
87	124-tri-Cl-Bz		17.25	17.26	0.000	180	182	3.158	0.20	0.2	85	#
88	naphthalene		17.52	17.50	0.002	128	129	4.753	0.59	0.6	81	#2
90	123-tri-Cl-Bz		17.67	17.69	-0.001	180	182	2.940	0.28	0.3	1	m
89	hx-Cl-butadiene		17.51	17.53	-0.001	225	260	2.553	0.22	0.2	44	m?/1

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

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-0003.D
 Acq On : 10 Jan 03 2:27 pm
 Sample : f=1
 Misc :
 Quant Time: Jan 13 9:28 2003
 Vial: 16
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G1044\3-002.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:26 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:30 2003
 Print Time : Mon Jan 13 10:42 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.04	9.04	0.000	96	70	768.596	10.00		0.00	
47	Cl-benzene-d5, I2	12.67	12.66	0.000	82	119	229.493	10.00		0.00	
62	1,4-DCB-d4 150 15	15.16	15.16	0.000	152	150	197.148	10.00		0.00	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.43	7.44	0.000	111	113	59.189	1.93		1.9	9.63%
29	1,2-di-Cl-ethane-	8.04	8.03	0.000	65	102	26.643	1.62		1.6	8.09%
55	toluene-d8(S2)	11.15	11.15	0.000	100	99	78.299	1.86		1.9	9.29%
70	4-Br-1-F-Bz (S3)	13.90	13.90	0.000	174	95	36.886	1.85		1.9	9.25%

Target Compounds	<<< I1 : ISTD ID = 1 >>>	Qvalue
3 di-Cl-di-F-methan	2.60 2.60 0.000 85 87 44.272 1.87 1.9 95	
4 Chloromethane	2.79 2.79 0.000 50 52 30.836 1.65 1.6 91	
9 F114 85 135	2.83 2.84 -0.002 85 135 46.094 1.87 1.9 82	
5 vinyl chloride	2.97 2.97 0.000 62 64 35.625 2.01 2.0 100	
6 bromomethane	3.39 3.38 0.000 94 96 43.488 2.52 2.5 82	
7 Chloroethane	3.53 3.54 -0.001 64 66 27.223 2.07 2.1 99	
8 tri-Cl-F-methane	4.16 4.16 0.000 101 103 56.379 1.91 1.9 87	
111 111 isopropyl alcoho	4.26 4.28 -0.002 45 43 8.181 26.30 26.3 1	
100 100 ethyl ether x5	4.45 4.45 0.000 59 74 108.089 9.87 9.9 99	
102 102 Acroolein x10	4.17 4.17 0.000 56 55 15.074 17.25 17.2 78	
119 119 methyl acetate	5.07 5.07 0.000 43 74 15.788 3.14 3.1 100	
104 104 Carbon disulfide	5.20 5.19 0.000 76 78 133.095 2.06 2.1 100	
103 103 Acrylonitrilex10	4.91 4.91 0.000 53 52 37.044 20.10 20.1 100	
95 95 Acetone x10	4.31 4.33 -0.002 43 58 41.993 28.37 28.4 90	
108 108 F-113	5.04 5.05 -0.001 151 101 41.068 2.00 2.0 97	
13 13 11-dichloroethene	4.78 4.78 0.000 61 96 61.263 2.03 2.0 95	
101 101 Acetonitrilex10	4.21 4.22 0.000 41 40 1.527 3.87 3.9 1	
109 109 Iodomethane	4.81 4.82 -0.001 142 127 55.824 2.32 2.3 97	
113 113 Tert butyl alcoh	4.86 4.87 -0.001 59 57 18.436 26.84 26.8 87	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-002.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:26 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:30 2003
 Print Time : Mon Jan 13 10:42 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	111.315	3.20	3.2	97	
112	Allyl chloride	5.08	5.08	0.000	41	76	73.643	2.71	2.7	92	?
200	200 Nitro methane x1	5.82	5.83	0.000	61	46	68.361	20.39	20.4	97	#?
10	10 t-Bu-Me-ether	6.01	6.01	0.000	73	57	73.627	1.95	1.9	94	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	45.222	1.96	2.0	94	?
98	98 Vinyl acetate x5	6.40	6.41	0.000	43	86	233.914	11.10	11.1	100	
21	21 11-dichloroethane	6.16	6.16	0.000	63	83	83.925	2.08	2.1	97	
91	91 2-butanone MEKx10	6.84	6.84	0.000	43	72	146.366	24.33	24.3	95	?
115	115 Di isoprop ether	6.85	6.85	0.000	45	87	206.054	2.36	2.4	97	?
22	22 c-12-di-Cl-ethene	6.97	6.97	0.000	96	61	46.067	2.04	2.0	95	
23	23 22-Dichloropropan	7.35	7.35	0.000	77	97	71.811	2.06	2.1	93	
24	24 Br-Cl-methane	7.19	7.19	0.000	128	130	15.248	2.10	2.1	93	
25	25 chloroform	7.27	7.27	0.000	83	85	81.604	2.50	2.5	97	
201	201 Ethyl acetate x2	7.32	7.31	0.000	43	61	44.403	4.20	4.2	88	?
116	116 ETBE	7.41	7.40	0.000	59	87	124.488	2.01	2.0	100	
117	117 Iso-butyl alcoho	7.32	7.31	0.000	43	42	44.403	22.93	22.9	1	#?
26	26 tetrahydrofuranx5	7.71	7.71	0.000	72	42	8.407	10.67	10.7	92	
34	34 111-tri-Cl-ethane	8.23	8.23	0.000	97	99	65.859	1.96	2.0	100	
30	30 12-dichloroethane	8.12	8.13	-0.002	62	64	31.555	2.35	2.3	93	
35	35 11-Di-Cl-propene	8.49	8.48	0.000	75	110	58.117	1.98	2.0	99	
36	36 benzene	8.75	8.76	-0.001	78	52	138.554	2.07	2.1	100	
37	37 CCl4	8.70	8.69	0.002	117	119	47.773	1.86	1.9	96	
97	97 thiophene	8.90	8.89	0.000	84	58	70.133	2.11	2.1	98	
118	118 TAME	9.00	9.01	0.000	73	43	97.622	2.29	2.3	92	
39	39 12-di-Cl-propane	9.50	9.50	0.000	63	76	38.931	1.99	2.0	100	
40	40 trichloroethene	9.56	9.55	0.000	130	132	41.064	1.98	2.0	94	
96	96 Me-methacrylate	9.86	9.85	0.001	69	100	19.691	2.68	2.7	78	
42	42 Br-di-Cl-methane	9.62	9.62	0.000	83	85	55.994	2.55	2.6	94	
41	41 dibromomethane	9.46	9.45	0.000	174	172	17.911	1.90	1.9	95	
45	45 c-13-di-Cl-propen	10.38	10.37	0.000	75	110	48.533	2.02	2.0	93	
92	92 2-ClEt-Vl-ether10	10.15	10.15	0.000	63	43	80.060	19.39	19.4	93	
56	56 toluene	11.22	11.23	0.000	91	92	126.101	2.01	2.0	95	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-002.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:26 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:30 2003
 Print Time : Mon Jan 13 10:42 2003
 Miscellaneous :

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

4979

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.36	11.37	0.000	69	99	35.707	3.06	3.1	93	
93	2-Hexanone x5	11.49	11.49	0.000	43	58	47.646	13.58	13.6	91	
48	112-tri-Cl-Et	11.04	11.03	0.001	97	83	21.908	2.62	2.6	95	
58	1,2-di-br-ethane	11.84	11.83	0.002	107	109	19.207	2.19	2.2	88	
51	di-Br-Cl-methane	11.58	11.57	0.001	129	127	27.173	2.20	2.2	96	
46	t-13-di-Cl-propen	10.87	10.86	0.000	75	110	30.276	1.95	2.0	99	
105	1-Chlorohexane	12.65	12.64	0.002	55	93	43.864	2.59	2.6	88	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.53	10.52	0.000	43	58	13.676	2.73	2.7	96	
49	1,3-di-Cl-propane	11.30	11.29	0.000	76	78	30.516	2.04	2.0	98	
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	40.927	2.03	2.0	96	
60	chlorobenzene	12.70	12.70	0.000	112	77	71.440	2.54	2.5	90	
61	1112-tetra-Cl-Et	12.63	12.63	0.000	131	133	28.400	2.11	2.1	100	
64	ethylbenzene	12.91	12.91	0.000	91	106	140.640	1.61	1.6	97	
65	m/p-Xylenes x2	13.11	13.11	0.000	91	106	213.036	4.35	4.4	100	
99	1-4-di-Cl-butane	13.45	13.46	0.000	55	41	32.572	2.72	2.7	94	
52	bromofom	13.20	13.23	-0.002	173	175	13.982	2.28	2.3	97	
66	styrene	13.43	13.43	0.000	104	78	75.317	2.17	2.2	97	
67	o-xylene	13.49	13.50	0.000	91	106	110.603	2.29	2.3	94	
68	1122-Tetra-Cl-Et	13.49	13.50	0.000	83	85	21.540	3.55	3.6	99	
110	t-1,4-dichloro-2	13.66	13.68	-0.001	89	53	7.032	7.71	7.7	31	
106	Cl-benzyl	15.11	15.13	-0.001	91	126	24.517	2.48	2.5	96	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.64	13.64	0.000	110	97	4.401	1.83	1.8	93	
71	isopropylbenzene	13.86	13.85	0.000	105	120	135.584	2.09	2.1	99	
72	bromobenzene	14.10	14.10	0.000	156	158	27.613	1.94	1.9	96	
73	n-propylbenzene	14.28	14.28	0.000	120	78	36.443	1.99	2.0	94	
74	2-Cl-Tl	14.38	14.38	0.000	126	128	23.738	2.05	2.0	88	
75	4-Cl-Tl	14.45	14.45	0.000	126	128	32.882	2.54	2.5	95	
76	135-tri-Me-Bz	14.56	14.56	0.000	105	120	107.323	2.05	2.1	100	
79	tert-butylbenzene	14.84	14.84	0.000	119	91	113.211	2.10	2.1	96	
78	124-tri-Me-Bz	14.95	14.94	0.000	105	120	96.787	2.18	2.2	98	

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

m
 1/13/03

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-002.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:26 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:30 2003
 Print Time : Mon Jan 13 10:42 2003
 Miscellaneous :

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

4980

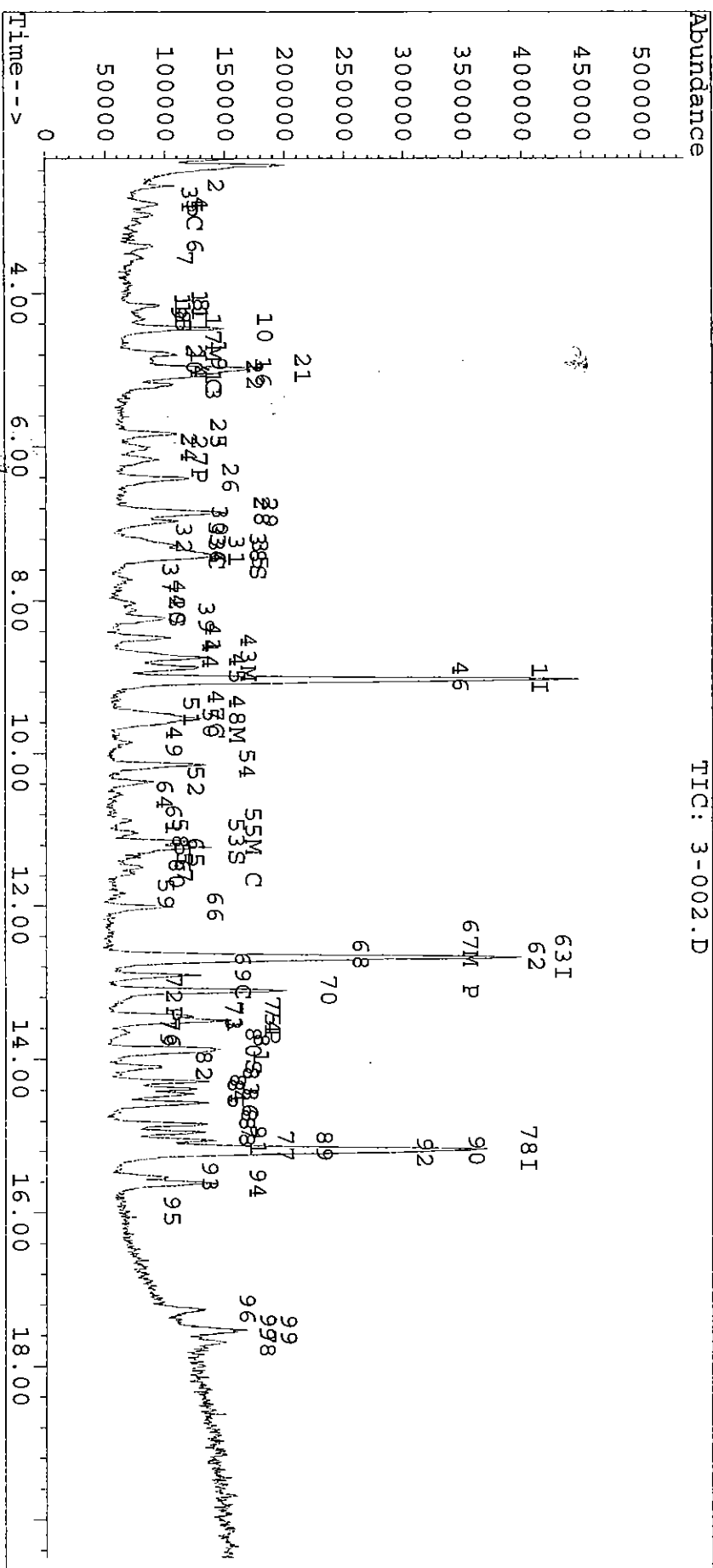
ID	Component Name	R.T.	RT0	DRRT	QIon	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
80	13-di-Cl-Bz	146	15.12	15.12	0.000	146	148	45.648	1.67	1.7	96	?
82	14-di-Cl-Bz	146	15.19	15.18	0.000	146	148	69.544	2.26	2.3	95	?
81	sec-butylbenzene		15.04	15.05	0.000	105	134	163.551	2.35	2.4	95	
77	4-iso-Pr-toluene		15.22	15.22	0.000	119	134	119.820	2.11	2.1	99	?
84	12-di-Cl-benzene		15.53	15.53	0.000	146	148	45.355	2.07	2.1	100	
85	n-butylbenzene		15.61	15.61	0.000	91	134	117.913	2.33	2.3	95	
86	12-diBr-3-Cl-Pra		15.98	16.00	-0.001	157	155	2.498	1.56	1.6	29	#
87	124-tri-Cl-Bz		17.26	17.26	0.000	180	182	29.532	1.89	1.9	93	
88	naphthalene		17.50	17.50	0.000	128	129	21.798	2.70	2.7	82	#
90	123-tri-Cl-Bz		17.70	17.69	0.000	180	182	22.060	2.11	2.1	99	
89	hx-Cl-butadiene		17.54	17.53	0.000	225	260	24.644	2.11	2.1	84	#

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

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-002.D Vial: 2
 Acq On : 10 Jan 03 3:26 pm Operator: Eddie
 Sample : f=1 Inst : GCMS-G
 Misc : Multiplr: 1.00
 Quant Time: Jan 13 9:30 2003 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G1044\3-010.D Sample : F=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 15:55 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:33 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:43 2003
 Miscellaneous :

4982

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.03	9.04	0.000	96	70	755.237	10.00		0.00	
47	47 Cl-benzene-d5, I2	12.67	12.66	0.000	82	119	225.971	10.00		0.01	
62	62 1,4-DCB-d4 150 15	15.16	15.16	0.000	152	150	188.831	10.00		0.00	
System Monitoring Compounds (Surrogate)											
27	27 Di-Br-F-Methane (	7.44	7.44	0.000	111	113	265.788	8.80	8.8	43.99%	
29	29 1,2-di-Cl-ethane-	8.02	8.03	0.000	65	102	124.715	7.71	7.7	38.53%	
55	55 toluene-d8(S2)	11.15	11.15	0.000	100	99	377.922	9.13	9.1	45.65%	
70	70 4-Br-1-F-Bz (S3)	13.90	13.90	0.000	174	95	165.570	8.67	8.7	43.37%	

Target Compounds

Qvalue

<<< I1 : ISTD ID = 1 >>>											
3	3 di-Cl-di-F-methan	2.61	2.60	0.000	85	87	235.359	10.11	10.1	99	
4	4 Chloromethane	2.79	2.79	0.000	50	52	163.038	8.87	8.9	98	
9	9 F114 85 135	2.83	2.84	-0.001	85	135	252.154	10.44	10.4	98	
5	5 vinyl chloride	2.98	2.97	0.000	62	64	183.259	10.52	10.5	99	
6	6 bromomethane	3.38	3.38	0.000	94	96	181.174	10.70	10.7	98	
7	7 Chloroethane	3.53	3.54	-0.001	64	66	133.510	10.35	10.3	99	
8	8 tri-Cl-F-methane	4.17	4.16	0.000	101	103	326.202	11.26	11.3	99	
111	111 isopropyl alcoh	4.28	4.28	0.000	45	43	34.708	113.56	113.6	1	
100	100 ethyl ether x5	4.45	4.45	0.000	59	74	509.174	47.30	47.3	99	
102	102 Acrolein x10	4.17	4.17	0.000	56	55	75.489	87.90	87.9	95	
119	119 methyl acetate	5.06	5.07	-0.001	43	74	82.815	16.77	16.8	90	
104	104 Carbon disulfide	5.19	5.19	0.000	76	78	626.926	9.87	9.9	100	
103	103 Acrylonitrilex10	4.90	4.91	0.000	53	52	178.667	98.65	98.7	100	
95	95 Acetone x10	4.33	4.33	0.000	43	58	128.289	88.19	88.2	92	
108	108 F-113	5.04	5.05	-0.001	151	101	259.145	12.84	12.8	99	
13	13 11-dichloroethene	4.78	4.78	0.000	61	96	309.126	10.41	10.4	99	
101	101 Acetonitrilex10	4.23	4.22	0.002	41	40	53.569	138.16	138.2	59	
109	109 Iodomethane	4.82	4.82	0.000	142	127	290.992	12.33	12.3	99	
113	113 Tert butyl alcoh	4.87	4.87	0.000	59	57	60.720	89.96	90.0	91	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-010.D Sample : F=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 15:55 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:33 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:43 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Q Ion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	375.753	10.99	11.0	100	
112	Allyl chloride	5.09	5.08	0.000	41	76	318.868	11.96	12.0	95	?
200	200 Nitro methane x1	5.82	5.83	-0.001	61	46	334.791	101.61	101.6	95	#?
10	10 t-Bu-Me-ether	6.01	6.01	0.000	73	57	300.489	8.08	8.1	98	
19	19 t-12-di-Cl-ethene	5.83	5.82	0.000	96	61	225.516	9.93	9.9	97	?
98	98 Vinyl acetate x5	6.41	6.41	0.000	43	86	1080.092	52.17	52.2	99	
21	21 11-dichloroethane	6.17	6.16	0.000	63	83	403.704	10.17	10.2	99	
91	91 2-butanone MEKx10	6.83	6.84	-0.001	43	72	680.153	115.05	115.1	99	?
115	115 Di isoprop ether	6.85	6.85	0.000	45	87	993.426	11.60	11.6	99	?
22	22 c-12-di-Cl-ethene	6.98	6.97	0.000	96	61	219.592	9.88	9.9	99	
23	23 22-Dichloropropan	7.36	7.35	0.000	77	97	330.092	9.65	9.6	99	
24	24 Br-Cl-methane	7.20	7.19	0.000	128	130	78.733	11.05	11.0	100	
25	25 chloroform	7.27	7.27	0.000	83	85	372.529	11.61	11.6	100	
201	201 Ethyl acetate x2	7.32	7.31	0.000	43	61	206.670	19.88	19.9	91	?
116	116 ETBE	7.40	7.40	0.000	59	87	594.002	9.75	9.7	98	
117	117 Iso-butyl alcoho	7.32	7.31	0.000	43	42	221.032	116.16	116.2	62	#?
26	26 tetrahydrofuranx5	7.72	7.71	0.000	72	42	36.840	47.60	47.6	80	
34	34 111-tri-Cl-ethane	8.23	8.23	0.000	97	99	315.664	9.56	9.6	98	
30	30 12-dichloroethane	8.14	8.13	0.000	62	64	154.021	11.67	11.7	98	
35	35 11-Di-Cl-propene	8.49	8.48	0.000	75	110	286.388	9.92	9.9	100	
36	36 benzene	8.75	8.76	-0.001	78	52	661.631	10.08	10.1	98	
37	37 CCl4	8.69	8.69	0.000	117	119	267.339	10.57	10.6	100	
97	97 thiophene	8.90	8.89	0.000	84	58	323.713	9.89	9.9	98	
118	118 TAME	9.00	9.01	0.000	73	43	406.861	9.72	9.7	98	
39	39 12-di-Cl-propane	9.50	9.50	0.000	63	76	190.695	9.93	9.9	99	
40	40 trichloroethene	9.55	9.55	0.000	130	132	207.771	10.22	10.2	98	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	65.318	9.06	9.1	99	
42	42 Br-di-Cl-methane	9.61	9.62	0.000	83	85	256.691	11.91	11.9	100	
41	41 dibromomethane	9.45	9.45	0.000	174	172	96.884	10.46	10.5	100	
45	45 c-13-di-Cl-propen	10.38	10.37	0.000	75	110	229.072	9.69	9.7	99	
92	92 2-ClEt-VI-ether10	10.15	10.15	0.000	63	43	410.701	101.25	101.3	99	
56	56 toluene	11.23	11.23	0.000	91	92	601.438	9.74	9.7	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-010.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:55 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:33 2003
 Print Time : Mon Jan 13 10:43 2003
 Miscellaneous :

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

4984

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.36	11.37	0.000	69	99	132.945	11.60	11.6	99	
93	2-Hexanone x5	11.49	11.49	0.000	43	58	218.029	63.26	63.3	93	
48	1,2-tri-Cl-Et	11.04	11.03	0.001	97	83	90.597	11.03	11.0	98	
58	1,2-di-br-ethane	11.84	11.83	0.001	107	109	94.012	10.89	10.9	98	
51	di-Br-Cl-methane	11.57	11.57	0.000	129	127	125.300	10.31	10.3	99	
46	t-13-di-Cl-propen	10.88	10.86	0.001	75	110	155.530	10.21	10.2	96	
105	1-Chlorohexane	12.64	12.64	0.000	55	93	205.359	12.34	12.3	99	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.53	10.52	0.000	43	58	74.160	15.03	15.0	95	
49	1,3-di-Cl-propane	11.30	11.29	0.001	76	78	157.929	10.71	10.7	96	
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	216.892	10.91	10.9	98	
60	chlorobenzene	12.70	12.70	0.000	112	77	352.480	12.72	12.7	100	
61	1,1,2-tetra-Cl-Et	12.63	12.63	0.000	131	133	138.607	10.48	10.5	98	
64	ethylbenzene	12.90	12.91	0.000	91	106	637.777	7.42	7.4	99	
65	m/p-Xylenes x2	13.10	13.11	0.000	91	106	1000.145	20.75	20.8	98	
99	1-4-di-Cl-butane	13.46	13.46	0.000	55	41	146.606	12.42	12.4	99	
52	bromoforn	13.22	13.23	0.000	173	175	71.698	11.86	11.9	98	
66	styrene	13.43	13.43	0.000	104	78	352.645	10.32	10.3	99	
67	o-xylene	13.50	13.50	0.000	91	106	485.194	10.18	10.2	99	
68	1,1,2-Tetra-Cl-Et	13.51	13.50	0.000	83	85	91.096	15.26	15.3	95	
110	t-1,4-dichloro-2	13.68	13.68	0.000	89	53	17.679	19.68	19.7	81	
106	Cl-benzyl	15.13	15.13	0.000	91	126	121.704	12.48	12.5	99	
<<< I3 : ISTD ID = 62 >>>											
69	1,2,3-tri-Cl-Pr	13.65	13.64	0.000	110	97	21.971	9.56	9.6	95	
71	isopropylbenzene	13.85	13.85	0.000	105	120	641.793	10.34	10.3	100	
72	bromobenzene	14.10	14.10	0.000	156	158	143.515	10.54	10.5	95	
73	n-propylbenzene	14.28	14.28	0.000	120	78	173.970	9.90	9.9	99	
74	2-Cl-Tl	14.38	14.38	0.000	126	128	106.793	9.61	9.6	93	
75	4-Cl-Tl	14.44	14.45	0.000	126	128	163.598	13.21	13.2	100	
76	1,3,5-tri-Me-Bz	14.56	14.56	0.000	105	120	509.842	10.18	10.2	98	
79	tert-butylbenzene	14.83	14.84	0.000	119	91	533.176	10.35	10.3	99	
78	1,2,4-tri-Me-Bz	14.94	14.94	0.000	105	120	424.953	9.98	10.0	96	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-010.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 15:55 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:33 2003
 Print Time : Mon Jan 13 10:43 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
80	13-di-Cl-Bz	146	15.12	15.12	0.000	146	148	238.597	9.11	9.1	94	?
82	14-di-Cl-Bz	146	15.18	15.18	0.000	146	148	305.231	10.34	10.3	92	
81	sec-butylbenzene		15.05	15.05	0.000	105	134	797.663	11.98	12.0	97	
77	4-iso-Pr-toluene		15.22	15.22	0.000	119	134	572.630	10.55	10.6	100	
84	12-di-Cl-benzene		15.53	15.53	0.000	146	148	219.614	10.49	10.5	99	
85	n-butylbenzene		15.61	15.61	0.000	91	134	579.573	11.97	12.0	100	
86	12-diBr-3-Cl-Pra		15.99	16.00	0.000	157	155	14.611	9.52	9.5	93	
87	124-tri-Cl-Bz		17.26	17.26	0.000	180	182	158.333	10.59	10.6	97	
88	naphthalene		17.50	17.50	0.000	128	129	116.373	15.07	15.1	99	
90	123-tri-Cl-Bz		17.70	17.69	0.000	180	182	121.207	12.12	12.1	97	
89	hx-Cl-butadiene		17.54	17.53	0.000	225	260	130.257	11.63	11.6	98	

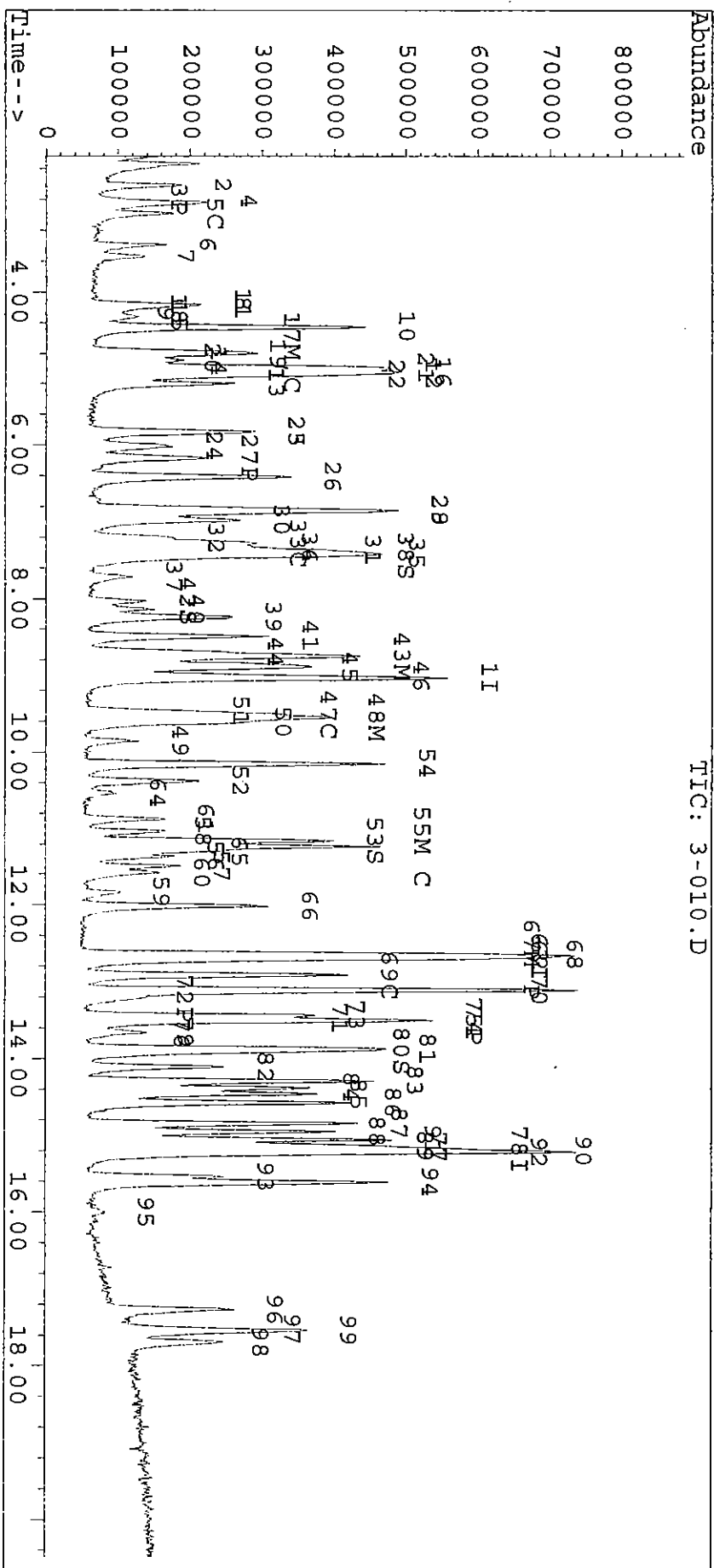
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-010.D
 Acq On : 10 Jan 03 3:55 pm
 Sample : f=1
 Misc :
 Quant Time: Jan 13 9:33 2003

Vial: 3
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G1044\3-020.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 16:24 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:35 2003
 Print Time : Mon Jan 13 10:43 2003
 Miscellaneous :

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

4987

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.04	9.04	0.000	96	70	739.905	10.00		0.00	
47	Cl-benzene-d5, I2	12.66	12.66	0.000	82	119	224.981	10.00		0.00	
62	1,4-DCB-d4 150 15	15.16	15.16	0.000	152	150	184.633	10.00		0.00	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.44	7.44	0.000	111	113	517.816	17.49		17.5	87.47%
29	1,2-di-Cl-ethane-	8.03	8.03	0.000	65	102	242.470	15.29		15.3	76.47%
55	toluene-d8(S2)	11.15	11.15	0.000	100	99	725.871	17.90		17.9	89.50%
70	4-Br-1-F-Bz (S3)	13.90	13.90	0.000	174	95	327.081	17.52		17.5	87.62%

Target Compounds

<<< I1 : ISTD ID = 1 >>>

Qvalue

3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	427.426	18.74		18.7	100	
4	Chloromethane	2.79	2.79	0.000	50	52	374.044	20.77		20.8	100	
9	F114 85 135	2.84	2.84	0.000	85	135	462.947	19.56		19.6	100	
5	vinyl chloride	2.97	2.97	0.000	62	64	344.227	20.16		20.2	100	
6	bromomethane	3.38	3.38	0.000	94	96	281.305	16.95		17.0	100	
7	Chloroethane	3.54	3.54	0.000	64	66	250.119	19.79		19.8	100	
8	tri-Cl-F-methane	4.16	4.16	0.000	101	103	612.174	21.57		21.6	100	
111	isopropyl alcoho	4.28	4.28	0.000	45	43	55.395	185.00		185.0	98	
100	ethyl ether x5	4.45	4.45	0.000	59	74	980.952	93.02		93.0	100	
102	Acrolein x10	4.17	4.17	0.000	56	55	215.710	256.38		256.4	100	
119	methyl acetate	5.07	5.07	0.000	43	74	187.169	38.69		38.7	100	
104	Carbon disulfide	5.19	5.19	0.000	76	78	1173.322	18.86		18.9	99	
103	Acrylonitrilex10	4.91	4.91	0.000	53	52	348.877	196.63		196.6	100	
95	Acetone x10	4.33	4.33	0.000	43	58	268.909	188.68		188.7	100	
108	F-113	5.05	5.05	0.000	151	101	480.691	24.32		24.3	100	
13	11-dichloroethene	4.78	4.78	0.000	61	96	575.822	19.79		19.8	99	
101	Acetonitrilex10	4.22	4.22	0.000	41	40	97.490	256.65		256.7	100	
109	Iodomethane	4.82	4.82	0.000	142	127	473.953	20.50		20.5	100	
113	Tert butyl alcoh	4.87	4.87	0.000	59	57	133.851	202.41		202.4	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-020.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 16:24 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:35 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Q1on	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	602.942	18.01	18.0	100	
112	Allyl chloride	5.08	5.08	0.000	41	76	616.004	23.57	23.6	100	?
200	200 Nitro methane x1	5.83	5.83	0.000	61	46	633.827	196.35	196.3	100	?
10	10 t-Bu-Me-ether	6.01	6.01	0.000	73	57	618.307	16.97	17.0	99	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	438.020	19.68	19.7	100	?
98	98 Vinyl acetate x5	6.41	6.41	0.000	43	86	1161.950	57.29	57.3	100	
21	21 11-dichloroethane	6.16	6.16	0.000	63	83	780.313	20.06	20.1	100	
91	91 2-butanone MEKx10	6.84	6.84	0.000	43	72	1296.553	223.87	223.9	100	?
115	115 Di Isoprop ether	6.85	6.85	0.000	45	87	1953.682	23.28	23.3	100	?
22	22 c-12-di-Cl-ethene	6.97	6.97	0.000	96	61	430.734	19.78	19.8	100	
23	23 22-Dichloropropan	7.35	7.35	0.000	77	97	619.878	18.49	18.5	100	
24	24 Br-Cl-methane	7.19	7.19	0.000	128	130	153.832	22.03	22.0	100	
25	25 chloroform	7.27	7.27	0.000	83	85	722.194	22.98	23.0	100	
201	201 Ethyl acetate x2	7.31	7.31	0.000	43	61	296.783	29.13	29.1	99	?
116	116 ETBE	7.40	7.40	0.000	59	87	1154.600	19.34	19.3	100	
117	117 Iso-butyl alcoho	7.31	7.31	0.000	43	42	297.076	159.36	159.4	100	?
26	26 tetrahydrofuranx5	7.71	7.71	0.000	72	42	75.144	99.10	99.1	100	
34	34 111-tri-Cl-ethane	8.23	8.23	0.000	97	99	604.949	18.69	18.7	100	
30	30 12-dichloroethane	8.13	8.13	0.000	62	64	298.086	23.05	23.1	100	
35	35 11-Di-Cl-propene	8.48	8.48	0.000	75	110	549.558	19.43	19.4	100	
36	36 benzene	8.76	8.76	0.000	78	52	1277.049	19.85	19.9	100	
37	37 CCl4	8.69	8.69	0.000	117	119	493.900	19.94	19.9	100	
97	97 thiophene	8.89	8.89	0.000	84	58	623.624	19.45	19.5	98	
118	118 TAME	9.01	9.01	0.000	73	43	792.558	19.33	19.3	99	
39	39 12-di-Cl-propane	9.50	9.50	0.000	63	76	376.101	19.98	20.0	100	
40	40 trichloroethene	9.55	9.55	0.000	130	132	403.682	20.26	20.3	100	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	123.371	17.47	17.5	100	
42	42 Br-di-Cl-methane	9.62	9.62	0.000	83	85	484.084	22.93	22.9	99	
41	41 dibromomethane	9.45	9.45	0.000	174	172	194.764	21.46	21.5	100	
45	45 c-13-di-Cl-propen	10.37	10.37	0.000	75	110	445.153	19.23	19.2	100	
92	92 2-ClEt-Vi-ether10	10.15	10.15	0.000	63	43	820.365	206.44	206.4	100	
56	56 toluene	11.23	11.23	0.000	91	92	1149.709	19.01	19.0	100	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-020.D
Method : C:\HPCHEM\1\METHODS\E524G003.M
Acq. Time : Jan 10 16:24 2003
Method Update: Mon Jan 13 10:38 2003
Quant. Time : Jan 13 09:35 2003
Print Time : Mon Jan 13 10:44 2003
Miscellaneous :

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

4989

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	CO,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.37	11.37	0.000	69	99	165.590	14.75	14.8	100	
93	2-Hexanone x5	11.49	11.49	0.000	43	58	368.204	109.05	109.1	99	
48	112-tri-Cl-Et	11.03	11.03	0.000	97	83	172.036	21.38	21.4	100	
58	1,2-di-br-ethane	11.83	11.83	0.000	107	109	191.133	22.61	22.6	100	
51	di-Br-Cl-methane	11.57	11.57	0.000	129	127	243.477	20.44	20.4	100	
46	t-13-di-Cl-propen	10.86	10.86	0.000	75	110	301.394	20.19	20.2	100	
105	1-Chlorohexane	12.64	12.64	0.000	55	93	374.248	22.95	22.9	100	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.52	10.52	0.000	43	58	143.757	29.26	29.3	99	
49	1,3-di-Cl-propane	11.29	11.29	0.000	76	78	310.529	21.15	21.1	100	
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	417.298	21.08	21.1	100	
60	chlorobenzene	12.70	12.70	0.000	112	77	668.143	24.21	24.2	100	
61	1112-tetra-Cl-Et	12.63	12.63	0.000	131	133	283.855	21.56	21.6	100	
64	ethylbenzene	12.91	12.91	0.000	91	106	1239.073	14.48	14.5	98	
65	m/p-Xylenes x2	13.11	13.11	0.000	91	106	1911.139	39.83	39.8	100	
99	1-4-di-Cl-butane	13.46	13.46	0.000	55	41	276.558	23.53	23.5	99	
52	bromoforn	13.23	13.23	0.000	173	175	143.599	23.85	23.8	100	
66	styrene	13.43	13.43	0.000	104	78	677.932	19.93	19.9	100	
67	o-xylene	13.50	13.50	0.000	91	106	915.641	19.30	19.3	100	
68	1122-Tetra-Cl-Et	13.50	13.50	0.000	83	85	173.152	29.14	29.1	100	
110	t-1,4-dichloro-2	13.68	13.68	0.000	89	53	27.009	30.20	30.2	94	
106	Cl-benzyl	15.13	15.13	0.000	91	126	238.713	24.59	24.6	99	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.64	13.64	0.000	110	97	45.443	20.21	20.2	100	
71	isopropylbenzene	13.85	13.85	0.000	105	120	1217.921	20.07	20.1	100	
72	bromobenzene	14.10	14.10	0.000	156	158	285.095	21.41	21.4	100	
73	n-propylbenzene	14.28	14.28	0.000	120	78	335.858	19.55	19.5	100	
74	2-Cl-Tl	14.38	14.38	0.000	126	128	201.349	18.53	18.5	100	
75	4-Cl-Tl	14.45	14.45	0.000	126	128	289.530	23.91	23.9	93	
76	135-tri-Me-Bz	14.56	14.56	0.000	105	120	969.387	19.79	19.8	100	
79	tert-butylbenzene	14.84	14.84	0.000	119	91	1020.985	20.26	20.3	99	
78	124-tri-Me-Bz	14.94	14.94	0.000	105	120	847.796	20.36	20.4	100	

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

m
1/13/03

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-020.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 16:24 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:35 2003
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.12	15.12	0.000	146	148	400.250	15.63	15.6	84
82	14-di-Cl-Bz	146	15.18	15.18	0.000	146	148	644.124	22.31	22.3	100
81	sec-butylbenzene		15.05	15.05	0.000	105	134	1465.228	22.51	22.5	97
77	4-iso-Pr-toluene		15.22	15.22	0.000	119	134	1085.210	20.45	20.5	100
84	12-di-Cl-benzene		15.53	15.53	0.000	146	148	410.082	20.03	20.0	100
85	n-butylbenzene		15.61	15.61	0.000	91	134	1096.665	23.17	23.2	100
86	12-diBr-3-Cl-Pra		16.00	16.00	0.000	157	155	31.395	20.93	20.9	100
87	124-tri-Cl-Bz		17.26	17.26	0.000	180	182	322.231	22.03	22.0	100
88	naphthalene		17.50	17.50	0.000	128	129	245.906	32.56	32.6	100
90	123-tri-Cl-Bz		17.69	17.69	0.000	180	182	239.739	24.52	24.5	100
89	hx-Cl-butadiene		17.53	17.53	0.000	225	260	244.690	22.34	22.3	100

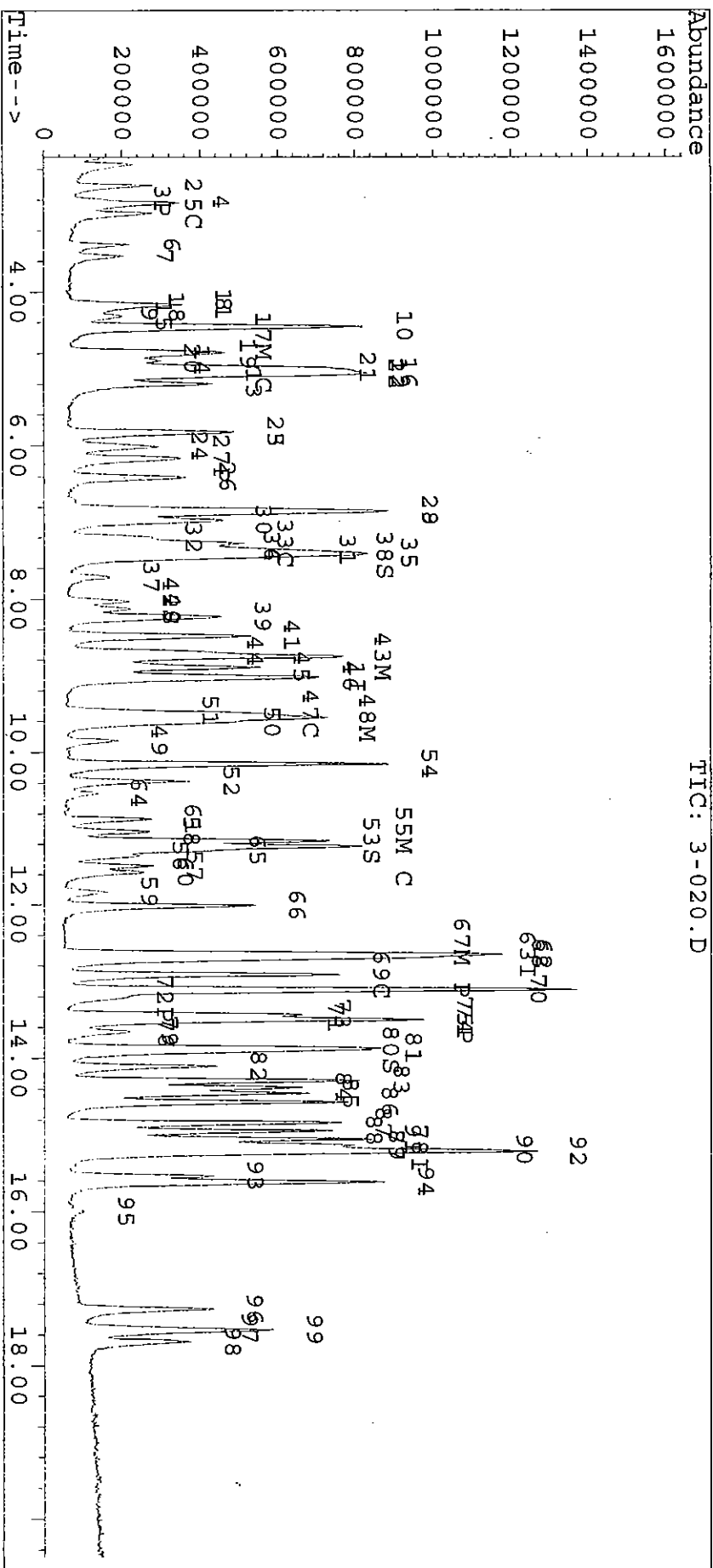
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-020.D
 Acq On : 10 Jan 03 4:24 pm
 Sample : f=1
 Misc :
 Quant Time: Jan 13 9:35 2003

Vial: 4
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G1044\3-040.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 16:54 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:38 2003
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.04	9.04	0.000	96	70	721.219	10.00		0.00	
47	Cl-benzene-d5, I2	12.67	12.66	0.001	82	119	233.528	10.00		0.01	
62	1,4-DCB-d4 150 15	15.18	15.16	0.001	152	150	176.207	10.00		0.02	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.44	7.44	0.000	111	113	1038.870	36.01	36.0	180.04%	
29	1,2-di-Cl-ethane-	8.02	8.03	0.000	65	102	475.685	30.78	30.8	153.91%	
55	toluene-d8(S2)	11.15	11.15	0.000	100	99	1455.135	36.81	36.8	184.07%	
70	4-Br-1-F-Bz (S3)	13.90	13.90	0.000	174	95	627.145	35.21	35.2	176.03%	

Target Compounds

<<< I1 : ISTD ID = 1 >>>

Qvalue

3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	820.884	36.93	36.9	99	
4	Chloromethane	2.79	2.79	0.000	50	52	803.175	45.76	45.8	100	
9	F114 85 135	2.83	2.84	-0.002	85	135	909.235	39.41	39.4	94	
5	vinyl chloride	2.97	2.97	0.000	62	64	682.967	41.04	41.0	99	
6	bromomethane	3.37	3.38	0.000	94	96	570.773	35.29	35.3	96	
7	Chloroethane	3.52	3.54	-0.002	64	66	505.034	40.99	41.0	99	
8	tri-Cl-F-methane	4.16	4.16	0.000	101	103	1192.828	43.11	43.1	100	
111	isopropyl alcoho	4.26	4.28	-0.002	45	43	131.600	450.89	450.9	1	#
100	ethyl ether x5	4.45	4.45	0.000	59	74	1975.186	192.15	192.1	99	
102	Acrolein x10	4.17	4.17	0.000	56	55	340.415	415.08	415.1	92	?
119	methyl acetate	5.07	5.07	0.000	43	74	271.933	57.66	57.7	96	?
104	Carbon disulfide	5.19	5.19	0.000	76	78	2357.195	38.87	38.9	100	
103	Acrylonitrilex10	4.90	4.91	0.000	53	52	701.313	405.50	405.5	98	
95	Acetone x10	4.32	4.33	0.000	43	58	465.240	334.90	334.9	98	
108	F-113	5.03	5.05	-0.002	151	101	925.581	48.03	48.0	100	
13	11-dichloroethene	4.77	4.78	0.000	61	96	1147.783	40.46	40.5	99	
101	Acetonitrilex10	4.22	4.22	0.000	41	40	189.257	511.14	511.1	80	#
109	Iodomethane	4.81	4.82	0.000	142	127	954.242	42.35	42.3	99	
113	Tert butyl alcoh	4.87	4.87	0.000	59	57	240.444	373.03	373.0	96	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-040.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 16:54 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:38 2003
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	1052.979	32.26	32.3	100	
112	Allyl chloride	5.08	5.08	0.000	41	76	1168.854	45.89	45.9	99	?
200	200 Nitro methane x1	5.82	5.83	0.000	61	46	1264.975	402.01	402.0	100	?
10	10 t-Bu-Me-ether	6.01	6.01	0.000	73	57	1211.982	34.13	34.1	98	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	867.973	40.01	40.0	99	?
98	98 Vinyl acetate x5	6.41	6.41	0.000	43	86	4543.111	229.80	229.8	99	
21	21 11-dichloroethane	6.15	6.16	0.000	63	83	1539.657	40.61	40.6	98	
91	91 2-butanone MEKx10	6.84	6.84	0.000	43	72	2708.302	479.74	479.7	99	?
115	115 Di Isoprop ether	6.85	6.85	0.000	45	87	3898.884	47.66	47.7	100	
22	22 c-12-di-Cl-ethene	6.97	6.97	0.000	96	61	857.123	40.38	40.4	99	
23	23 22-Dichloropropan	7.36	7.35	0.001	77	97	1207.116	36.94	36.9	99	
24	24 Br-Cl-methane	7.19	7.19	0.000	128	130	312.235	45.88	45.9	99	
25	25 chloroform	7.27	7.27	0.000	83	85	1453.363	47.45	47.4	99	
201	201 Ethyl acetate x2	7.31	7.31	0.000	43	61	873.148	87.93	87.9	83	?
116	116 ETBE	7.40	7.40	0.000	59	87	2264.664	38.92	38.9	99	
117	117 Iso-butyl alcoho	7.31	7.31	0.000	43	42	880.890	484.77	484.8	63	#?
26	26 tetrahydrofuranx5	7.71	7.71	0.000	72	42	146.600	198.35	198.3	94	
34	34 111-tri-Cl-ethane	8.24	8.23	0.000	97	99	1193.974	37.85	37.8	97	
30	30 12-dichloroethane	8.13	8.13	0.000	62	64	600.265	47.63	47.6	99	
35	35 11-Di-Cl-propene	8.48	8.48	0.000	75	110	1091.787	39.61	39.6	99	
36	36 benzene	8.75	8.76	0.000	78	52	2547.937	40.64	40.6	100	
37	37 CCl4	8.69	8.69	0.000	117	119	981.263	40.64	40.6	100	
97	97 thiophene	8.90	8.89	0.000	84	58	1239.916	39.68	39.7	99	
118	118 TAME	9.01	9.01	0.000	73	43	1521.011	38.05	38.1	100	
39	39 12-di-Cl-propane	9.50	9.50	0.000	63	76	741.594	40.42	40.4	100	
40	40 trichloroethene	9.55	9.55	0.000	130	132	804.090	41.41	41.4	100	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	250.724	36.43	36.4	97	
42	42 Br-di-Cl-methane	9.61	9.62	0.000	83	85	955.698	46.44	46.4	98	
41	41 dibromomethane	9.45	9.45	0.000	174	172	383.350	43.33	43.3	99	
45	45 c-13-di-Cl-propen	10.37	10.37	0.000	75	110	890.451	39.45	39.5	99	
92	92 2-ClEt-VI-ether10	10.15	10.15	0.000	63	43	1655.105	427.28	427.3	98	
56	56 toluene	11.23	11.23	0.000	91	92	2291.459	38.88	38.9	99	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-040.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 16:54 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:38 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	Co,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.37	11.37	0.000	69	99	512.644	46.85	46.8	99	
93	2-Hexanone x5	11.48	11.49	-0.001	43	58	831.419	252.62	252.6	99	
48	112-tri-Cl-Et	11.04	11.03	0.000	97	83	347.876	44.36	44.4	94	
58	1,2-di-br-ethane	11.84	11.83	0.001	107	109	371.448	45.07	45.1	100	
51	di-Br-Cl-methane	11.58	11.57	0.000	129	127	489.254	42.14	42.1	100	
46	t-13-di-Cl-propen	10.87	10.86	0.000	75	110	594.704	40.88	40.9	96	
105	1-Chlorohexane	12.64	12.64	0.000	55	93	719.249	45.24	45.2	99	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.53	10.52	0.000	43	58	303.184	59.45	59.4	96	
49	1,3-di-Cl-propane	11.30	11.29	0.000	76	78	604.220	39.64	39.6	100	
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	825.820	40.18	40.2	99	
60	chlorobenzene	12.71	12.70	0.000	112	77	1330.525	46.45	46.4	99	
61	1112-tetra-Cl-Et	12.63	12.63	0.000	131	133	562.377	41.15	41.1	100	
64	ethylbenzene	12.90	12.91	0.000	91	106	2446.417	27.53	27.5	98	
65	m/p-Xylenes x2	13.10	13.11	0.000	91	106	3754.507	75.38	75.4	99	
99	1-4-di-Cl-butane	13.46	13.46	0.000	55	41	532.573	43.66	43.7	96	
52	bromoforn	13.22	13.23	0.000	173	175	281.887	45.10	45.1	96	
66	styrene	13.43	13.43	0.000	104	78	1346.269	38.14	38.1	98	
67	o-xylene	13.50	13.50	0.000	91	106	1764.503	35.83	35.8	100	
68	1122-Tetra-Cl-Et	13.51	13.50	0.000	83	85	326.262	52.90	52.9	100	
110	t-1,4-dichloro-2	13.67	13.68	0.000	89	53	54.200	58.39	58.4	95	
106	Cl-benzyl	15.13	15.13	0.000	91	126	457.227	45.37	45.4	99	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.65	13.64	0.000	110	97	89.331	41.64	41.6	94	
71	isopropylbenzene	13.86	13.85	0.000	105	120	2372.165	40.96	41.0	99	
72	bromobenzene	14.11	14.10	0.000	156	158	554.412	43.62	43.6	98	
73	n-propylbenzene	14.29	14.28	0.000	120	78	643.879	39.27	39.3	98	
74	2-Cl-Tl	14.38	14.38	0.000	126	128	427.470	41.22	41.2	95	
75	4-Cl-Tl	14.45	14.45	0.000	126	128	594.245	51.42	51.4	98	
76	135-tri-Me-Bz	14.57	14.56	0.000	105	120	1885.346	40.33	40.3	99	
79	tert-butylbenzene	14.84	14.84	0.000	119	91	1994.231	41.47	41.5	88	
78	124-tri-Me-Bz	14.94	14.94	0.000	105	120	1639.260	41.24	41.2	99	

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

m
1/13/02

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-040.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 16:54 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:38 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:44 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
80	13-di-Cl-Bz	146	15.13	15.12	0.000	146	148	844.281	34.55	34.6	94	?
82	14-di-Cl-Bz	146	15.19	15.18	0.000	146	148	1193.857	43.33	43.3	92	
81	sec-butylbenzene		15.05	15.05	0.000	105	134	2859.247	46.03	46.0	96	
77	4-iso-Pr-toluene		15.22	15.22	0.000	119	134	2064.029	40.76	40.8	99	
84	12-di-Cl-benzene		15.54	15.53	0.000	146	148	812.246	41.57	41.6	97	
85	n-butylbenzene		15.61	15.61	0.000	91	134	2099.124	46.47	46.5	99	
86	12-diBr-3-Cl-Pra		15.99	16.00	0.000	157	155	62.914	43.94	43.9	93	
87	124-tri-Cl-Bz		17.27	17.26	0.000	180	182	604.462	43.31	43.3	100	
88	naphthalene		17.50	17.50	0.000	128	129	459.400	63.75	63.7	100	
90	123-tri-Cl-Bz		17.69	17.69	0.000	180	182	448.256	48.03	48.0	98	
89	hx-Cl-butadiene		17.55	17.53	0.001	225	260	466.707	44.65	44.6	99	

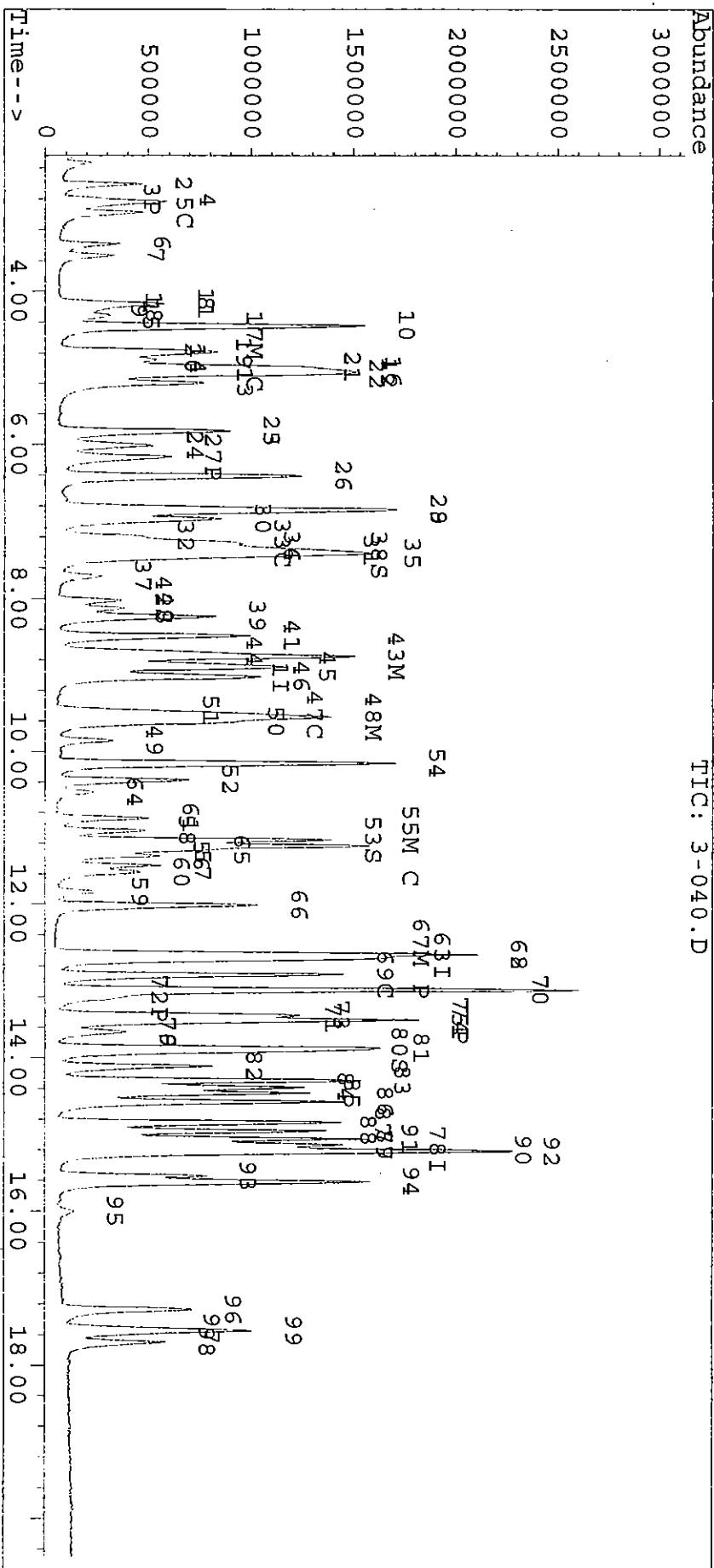
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-040.D
 Acq On : 10 Jan 03 4:54 pm
 Sample : f=1
 Misc :
 Quant Time: Jan 13 9:38 2003

Vial: 5
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



Data Filename: C:\HPCHEM\1\DATA\03G1044\3-080.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 17:23 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:39 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:45 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.04	9.04	0.000	96	70	703.721	10.00		0.00	
47	Cl-benzene-d5, I2	12.67	12.66	0.000	82	119	241.678	10.00		0.01	
62	1,4-DCB-d4 150 15	15.18	15.16	0.002	152	150	164.785	10.00			
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.44	7.44	0.000	111	113	1963.588	69.75		348.75%	
29	1,2-di-Cl-ethane-	8.02	8.03	0.000	65	102	894.560	59.33		296.64%	
55	toluene-d8(S2)	11.16	11.15	0.000	100	99	2774.904	71.95		359.74%	
70	4-Br-1-F-Bz (S3)	13.91	13.90	0.000	174	95	1159.540	69.60		348.02%	

Target Compounds	<<< I1 : ISTD ID = 1 >>>	Qvalue
3 di-Cl-di-F-methan	2.60 2.60 0.000 85 87 1551.879 71.55	99
4 Chloromethane	2.78 2.79 -0.002 50 52 1514.667 88.45	97
9 F114 85 135	2.83 2.84 -0.002 85 135 1714.061 76.15	97
5 vinyl chloride	2.96 2.97 0.000 62 64 1258.237 77.49	99
6 bromomethane	3.38 3.38 0.000 94 96 1116.110 70.72	96
7 Chloroethane	3.52 3.54 -0.002 64 66 952.606 79.23	100
8 tri-Cl-F-methane	4.15 4.16 0.000 101 103 2238.212 82.90	100
111 111 isopropyl alcoho	4.27 4.28 0.000 45 43 244.204 857.50	59
100 100 ethyl ether x5	4.44 4.45 -0.001 59 74 3742.221 373.10	100
102 102 Acrolein x10	4.17 4.17 0.000 56 55 601.157 751.24	90
119 119 methyl acetate	5.07 5.07 0.000 43 74 696.519 151.37	95
104 104 Carbon disulfide	5.18 5.19 0.000 76 78 4372.134 73.88	100
103 103 Acrylonitrilex10	4.90 4.91 0.000 53 52 1341.637 795.03	98
95 95 Acetone x10	4.32 4.33 0.000 43 58 912.905 673.49	97
108 108 F-113	5.03 5.05 -0.002 151 101 1749.444 93.05	100
13 13 11-dichloroethene	4.78 4.78 0.000 61 96 2146.767 77.56	99
101 101 Acetonitrilex10	4.21 4.22 0.000 41 40 353.648 978.88	83
109 109 Iodomethane	4.81 4.82 -0.001 142 127 1666.625 75.80	99
113 113 Tert butyl alcoh	4.87 4.87 0.000 59 57 485.913 772.60	85

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-080.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 17:23 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:39 2003
 Print Time : Mon Jan 13 10:45 2003
 Miscellaneous :

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

4998

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.98	4.98	0.000	84	49	1493.871	46.91	46.9	100	
112	Allyl chloride	5.08	5.08	0.000	41	76	2056.270	82.74	82.7	98	?
200	200 Nitro methane x1	5.82	5.83	0.000	61	46	2385.881	777.09	777.1	99	?
10	10 t-Bu-Me-ether	6.01	6.01	0.000	73	57	2415.631	69.73	69.7	98	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	1629.626	76.99	77.0	98	?
98	98 Vinyl acetate x5	6.40	6.41	0.000	43	86	6984.572	362.07	362.1	100	
21	21 11-dichloroethane	6.16	6.16	0.000	63	83	2853.804	77.14	77.1	98	
91	91 2-butanone MEKx10	6.84	6.84	0.000	43	72	5159.574	936.67	936.7	98	?
115	115 Di isoprop ether	6.85	6.85	0.000	45	87	7414.434	92.88	92.9	100	?
22	22 c-12-di-Cl-ethene	6.98	6.97	0.000	96	61	1613.152	77.88	77.9	99	
23	23 22-Dichloropropan	7.36	7.35	0.000	77	97	2269.373	71.18	71.2	100	
24	24 Br-Cl-methane	7.19	7.19	0.000	128	130	593.248	89.34	89.3	100	
25	25 chloroform	7.27	7.27	0.000	83	85	2739.754	91.67	91.7	100	
201	201 Ethyl acetate x2	7.32	7.31	0.000	43	61	1526.995	157.60	157.6	87	?
116	116 ETBE	7.41	7.40	0.000	59	87	4329.036	76.24	76.2	99	
117	117 Iso-butyl alcoho	7.32	7.31	0.000	43	42	1546.333	872.13	872.1	70	#?
26	26 tetrahydrofuranx5	7.71	7.71	0.000	72	42	287.877	399.18	399.2	97	
34	34 111-tri-Cl-ethane	8.24	8.23	0.000	97	99	2324.189	75.51	75.5	99	
30	30 12-dichloroethane	8.13	8.13	0.000	62	64	1155.412	93.96	94.0	99	
35	35 11-Di-Cl-propene	8.48	8.48	0.000	75	110	2099.663	78.07	78.1	99	
36	36 benzene	8.75	8.76	-0.001	78	52	4710.056	76.99	77.0	100	
37	37 CCl4	8.69	8.69	0.000	117	119	1852.391	78.63	78.6	100	
97	97 thiophene	8.89	8.89	0.000	84	58	2337.761	76.68	76.7	98	
118	118 TAME	9.01	9.01	0.000	73	43	3045.504	78.09	78.1	100	
39	39 12-di-Cl-propane	9.50	9.50	0.000	63	76	1426.638	79.70	79.7	100	
40	40 trichloroethene	9.55	9.55	0.000	130	132	1515.012	79.96	80.0	99	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	519.513	77.36	77.4	98	
42	42 Br-di-Cl-methane	9.61	9.62	0.000	83	85	1823.535	90.81	90.8	99	
41	41 dibromomethane	9.46	9.45	0.000	174	172	714.384	82.76	82.8	98	
45	45 c-13-di-Cl-propen	10.38	10.37	0.000	75	110	1727.952	78.47	78.5	100	
92	92 2-ClEt-Vl-ether10	10.16	10.15	0.000	63	43	3234.891	855.88	855.9	97	
56	56 toluene	11.23	11.23	0.000	91	92	4425.814	76.96	77.0	98	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-080.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Jan 10 17:23 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : Jan 13 09:39 2003
 Print Time : Mon Jan 13 10:45 2003
 Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.37	11.37	0.000	69	99	828.103	77.56	77.6	98	
93	2-Hexanone x5	11.49	11.49	0.000	43	58	1601.350	498.66	498.7	98	
48	112-tri-Cl-Et	11.04	11.03	0.000	97	83	664.699	86.86	86.9	92	
58	1,2-di-br-ethane	11.83	11.83	0.000	107	109	723.238	89.94	89.9	100	
51	di-Br-Cl-methane	11.57	11.57	0.000	129	127	967.601	85.41	85.4	99	
46	t-13-di-Cl-propen	10.88	10.86	0.001	75	110	1109.294	78.14	78.1	96	
105	1-Chlorohexane	12.65	12.64	0.001	55	93	1336.615	86.16	86.2	99	
<<< I2 : ISTD ID = 47 >>>											
54	MIBK	10.53	10.52	0.000	43	58	587.112	111.24	111.2	97	
49	1,3-di-Cl-propane	11.30	11.29	0.001	76	78	1155.285	73.25	73.2	99	
59	tetra-Cl-ethene	12.01	12.01	0.000	166	168	1643.233	77.26	77.3	99	
60	chlorobenzene	12.70	12.70	0.000	112	77	2470.738	83.34	83.3	99	
61	1112-tetra-Cl-Et	12.64	12.63	0.000	131	133	1108.773	78.39	78.4	99	
64	ethylbenzene	12.91	12.91	0.000	91	106	4804.958	52.25	52.3	97	
65	m/p-Xylenes x2	13.11	13.11	0.000	91	106	7222.435	140.12	140.1	99	
99	1-4-di-Cl-butane	13.47	13.46	0.001	55	41	1001.628	79.35	79.3	98	
52	bromoform	13.24	13.23	0.000	173	175	544.898	84.24	84.2	95	
66	styrene	13.44	13.43	0.000	104	78	2563.056	70.16	70.2	100	
67	o-xylene	13.51	13.50	0.001	91	106	3287.299	64.50	64.5	100	
68	1122-Tetra-Cl-Et	13.51	13.50	0.000	83	85	625.224	97.95	98.0	98	
110	t-1,4-dichloro-2	13.68	13.68	0.000	89	53	103.110	107.34	107.3	94	
106	Cl-benzyl	15.14	15.13	0.001	91	126	866.531	83.08	83.1	99	
<<< I3 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.65	13.64	0.000	110	97	172.722	86.09	86.1	95	
71	isopropylbenzene	13.87	13.85	0.001	105	120	4528.346	83.61	83.6	99	
72	bromobenzene	14.11	14.10	0.000	156	158	1030.644	86.70	86.7	98	
73	n-propylbenzene	14.30	14.28	0.001	120	78	1212.584	79.08	79.1	97	
74	2-Cl-Tl	14.39	14.38	0.000	126	128	745.993	76.92	76.9	99	
75	4-Cl-Tl	14.46	14.45	0.000	126	128	1075.809	99.53	99.5	92	
76	135-tri-Me-Bz	14.58	14.56	0.001	105	120	3580.223	81.88	81.9	99	
79	tert-butylbenzene	14.85	14.84	0.000	119	91	3739.615	83.15	83.2	99	
78	124-tri-Me-Bz	14.96	14.94	0.001	105	120	3054.447	82.17	82.2	99	

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

m
1/13/03

Data Filename: C:\HPCHEM\1\DATA\03G1044\3-080.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 17:23 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : Jan 13 09:39 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:45 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	Q10n	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
80	13-di-Cl-Bz	146	15.14	15.12	0.002	146	148	1658.291	72.57	72.6	95	?
82	14-di-Cl-Bz	146	15.20	15.18	0.001	146	148	2207.723	85.68	85.7	93	
81	sec-butylbenzene		15.06	15.05	0.000	105	134	5499.576	94.67	94.7	96	
77	4-Iso-Pr-toluene		15.23	15.22	0.001	119	134	3804.782	80.34	80.3	100	
84	12-di-Cl-benzene		15.54	15.53	0.000	146	148	1524.040	83.41	83.4	99	
85	n-butylbenzene		15.61	15.61	0.000	91	134	3991.637	94.49	94.5	99	
86	12-diBr-3-Cl-Pra		15.99	16.00	0.000	157	155	120.318	89.86	89.9	91	
87	124-tri-Cl-Bz		17.26	17.26	0.000	180	182	1187.497	90.98	91.0	100	
88	naphthalene		17.50	17.50	0.000	128	129	922.945	136.94	136.9	100	
90	123-tri-Cl-Bz		17.69	17.69	0.000	180	182	845.940	96.93	96.9	100	
89	hx-Cl-butadiene		17.54	17.53	0.000	225	260	859.594	87.93	87.9	98	

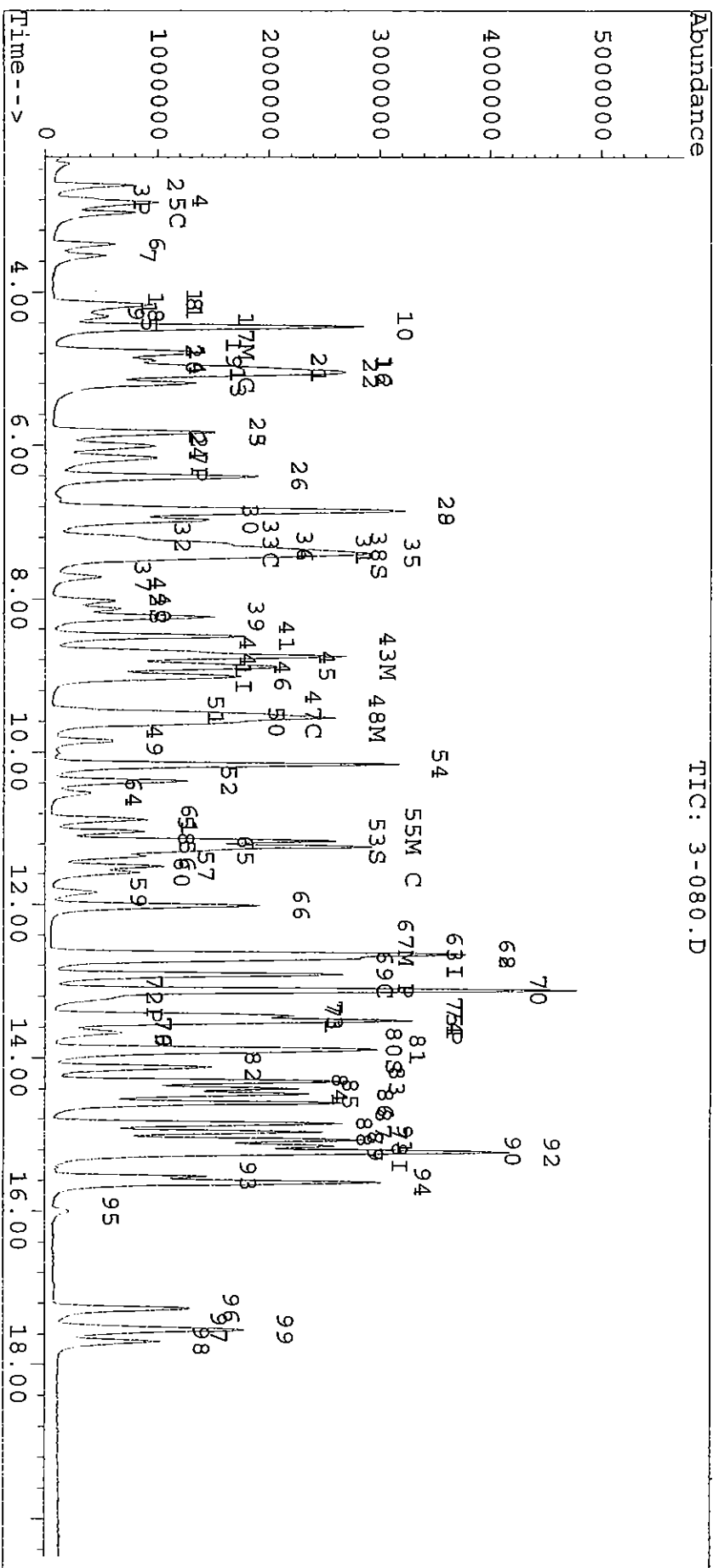
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G1044\3-080.D
 Acq On : 10 Jan 03 5:23 pm
 Sample : f=1
 Misc :
 Quant Time: Jan 13 9:39 2003

Vial: 6
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



5002

```
Dev (Min)
0.00
0.00
0.00
```

%Recovery

4	87.02%
2	85.82%
5	87.70%
5	87.89%

Qualitative	Quantitative
1. Qualitative 2. Quantitative	1. Qualitative 2. Quantitative

2	100
9	98
0	97
6	99
3	94
4	99
7	99
4	47
8	100
7	99
7	99
6	99
2	99
6	99
6	96
7	98
0	99
8	80
1	99
0	100

•

Data Filename: C:\HPCHEM\1\DATA\03G1044\G1044Q01.D Sample : f=1 ccv/icv
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 18:50 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 09:57 2003 Operator: Eddie
 Quant. Time : Jan 13 10:36 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:37 2003
 Miscellaneous :

5003

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	4.97	4.98	-0.001	84	49	397.006	11.83	11.8	100	
112	Allyl chloride	5.07	5.08	-0.001	41	76	588.305	17.03	17.0	97	?
200	200 Nitro methane x1	5.82	5.83	-0.001	61	46	610.779	175.74	175.7	95	#?
10	10 t-Bu-Me-ether	6.00	6.01	-0.001	73	57	557.717	16.56	16.6	96	
19	19 t-12-di-Cl-ethene	5.82	5.82	0.000	96	61	406.048	16.96	17.0	95	?
98	98 Vinyl acetate x5	6.41	6.41	0.000	43	86	1965.632	91.24	91.2	100	
21	21 11-dichloroethane	6.15	6.16	-0.001	63	83	750.915	17.96	18.0	99	
91	91 2-butanone MEKx10	6.83	6.84	-0.001	43	72	1231.875	164.89	164.9	97	?
115	115 Di isoprop ether	6.85	6.85	0.000	45	87	1808.509	17.14	17.1	99	?
22	22 c-12-di-Cl-ethene	6.97	6.97	0.000	96	61	395.661	16.89	16.9	97	
23	23 22-Dichloropropan	7.35	7.35	0.000	77	97	576.991	15.99	16.0	99	
24	24 Br-Cl-methane	7.18	7.19	-0.001	128	130	141.700	16.86	16.9	99	
25	25 chloroform	7.27	7.27	0.000	83	85	689.236	16.48	16.5	100	
201	201 Ethyl acetate x2	7.31	7.31	0.000	43	61	378.636	35.55	35.6	90	?
116	116 ETBE	7.40	7.40	0.000	59	87	1052.083	16.37	16.4	100	
117	117 Iso-butyl alcoho	7.31	7.31	0.000	43	42	392.518	184.17	184.2	64	#?
26	26 tetrahydrofuranx5	7.71	7.71	0.000	72	42	69.559	85.20	85.2	99	
34	34 111-tri-Cl-ethane	8.23	8.23	0.000	97	99	564.989	16.22	16.2	100	
30	30 12-dichloroethane	8.13	8.13	0.000	62	64	282.242	17.32	17.3	99	
35	35 11-Di-Cl-propene	8.48	8.48	0.000	75	110	497.493	16.34	16.3	100	
36	36 benzene	8.75	8.76	-0.001	78	52	1199.382	17.33	17.3	100	
37	37 CCl4	8.68	8.69	0.000	117	119	455.994	17.10	17.1	99	
97	97 thiophene	8.89	8.89	0.000	84	58	583.305	16.95	16.9	99	
118	118 TAME	9.00	9.01	0.000	73	43	702.458	15.87	15.9	100	
39	39 12-di-Cl-propane	9.49	9.50	0.000	63	76	351.243	17.34	17.3	99	
40	40 trichloroethene	9.54	9.55	-0.001	130	132	377.871	17.62	17.6	98	
96	96 Me-methacrylate	9.85	9.85	0.000	69	100	125.425	17.85	17.8	98	
42	42 Br-di-Cl-methane	9.61	9.62	0.000	83	85	450.244	15.94	15.9	96	
41	41 dibromomethane	9.45	9.45	0.000	174	172	180.293	18.45	18.5	98	
45	45 c-13-di-Cl-propen	10.37	10.37	0.000	75	110	412.841	16.57	16.6	99	
92	92 2-ClBt-Vi-ether10	10.15	10.15	0.000	63	43	750.880	175.75	175.8	99	
56	56 toluene	11.23	11.23	0.000	91	92	1093.591	16.81	16.8	98	

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

Data Filename: C:\HPCHEM\1\DATA\03G1044\G1044Q01.D Sample : F=1 CCV/ICV
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 18:50 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 09:57 2003 Operator: Eddie
 Quant. Time : Jan 13 10:36 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:37 2003
 Miscellaneous :

5004

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.36	11.37	0.000	69	99	232.775	17.29	17.3	100	
93	2-Hexanone x5	11.48	11.49	-0.001	43	58	392.829	86.81	86.8	97	
48	112-tri-Cl-Et	11.03	11.03	0.000	97	83	161.722	16.95	17.0	98	
58	1,2-di-br-ethane	11.82	11.83	0.000	107	109	172.988	16.98	17.0	98	
51	di-Br-Cl-methane	11.57	11.57	0.000	129	127	224.132	17.48	17.5	99	
46	t-13-di-Cl-propen	10.87	10.86	0.000	75	110	282.573	17.60	17.6	100	
105	1-Chlorohexane	12.64	12.64	0.000	55	93	359.945	18.07	18.1	96	
<<< 12 : ISTD ID = 47 >>>											
54	MIBK	10.52	10.52	0.000	43	58	140.136	18.90	18.9	97	
49	1,3-di-Cl-propane	11.29	11.29	0.000	76	78	281.641	18.31	18.3	100	
59	tetra-Cl-ethene	12.00	12.01	0.000	166	168	377.073	18.21	18.2	100	
60	chlorobenzene	12.70	12.70	0.000	112	77	623.775	18.14	18.1	99	
61	1112-tetra-Cl-Et	12.62	12.63	0.000	131	133	260.584	18.92	18.9	100	
64	ethylbenzene	12.90	12.91	0.000	91	106	1145.522	17.29	17.3	99	
65	m/p-Xylenes x2	13.10	13.11	0.000	91	106	1801.521	35.76	35.8	100	
99	1-4-di-Cl-butane	13.46	13.46	0.000	55	41	249.550	16.44	16.4	98	
52	bromoform	13.22	13.23	0.000	173	175	130.519	18.57	18.6	98	
66	styrene	13.43	13.43	0.000	104	78	627.907	17.65	17.7	98	
67	o-xylene	13.50	13.50	0.000	91	106	854.318	17.23	17.2	100	
68	1122-Tetra-Cl-Et	13.51	13.50	0.000	83	85	157.341	17.29	17.3	97	
110	t-1,4-dichloro-2	13.67	13.68	0.000	89	53	29.653	20.23	20.2	82	
106	Cl-benzyl	15.12	15.13	0.000	91	126	217.674	18.94	18.9	99	
<<< 13 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.64	13.64	0.000	110	97	40.148	16.92	16.9	97	
71	isopropylbenzene	13.86	13.85	0.000	105	120	1123.964	17.55	17.6	99	
72	bromobenzene	14.10	14.10	0.000	156	158	256.336	18.24	18.2	98	
73	n-propylbenzene	14.29	14.28	0.000	120	78	307.368	17.81	17.8	100	
74	2-Cl-Tl	14.37	14.38	0.000	126	128	196.033	17.09	17.1	94	
75	4-Cl-Tl	14.45	14.45	0.000	126	128	271.926	17.35	17.3	99	
76	135-tri-Me-Bz	14.57	14.56	0.000	105	120	876.887	16.96	17.0	99	
79	tert-butylbenzene	14.84	14.84	0.000	119	91	953.072	17.96	18.0	87	
78	124-tri-Me-Bz	14.94	14.94	0.000	105	120	768.449	17.48	17.5	99	

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

m
1/13/03

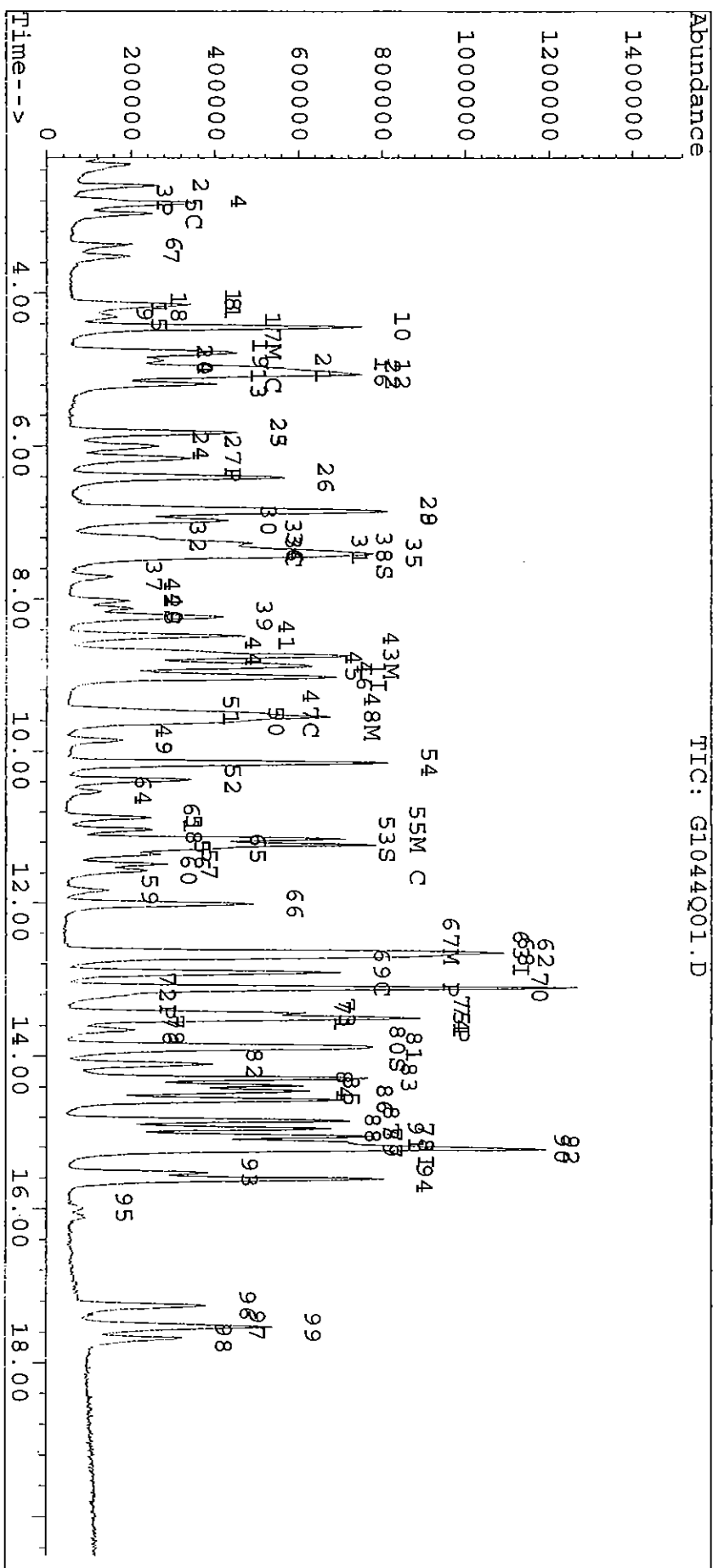
Data Filename: C:\HPCHEM\1\DATA\03G1044\G1044Q01.D Sample : F=1 CCV/LCV
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Jan 10 18:50 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 09:57 2003 Operator: Eddie
 Quant. Time : Jan 13 10:36 2003 Multiplr: 1.000000
 Print Time : Mon Jan 13 10:37 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note	
80	13-di-Cl-Bz	146	15.11	15.12	0.000	146	148	402.440	17.97	18.0	90	?
82	14-di-Cl-Bz	146	15.19	15.18	0.000	146	148	541.239	16.63	16.6	93	
81	sec-butylbenzene		15.04	15.05	0.000	105	134	1390.850	17.92	17.9	98	
77	4-iso-Pr-toluene		15.22	15.22	0.000	119	134	1009.580	18.03	18.0	98	
84	12-di-Cl-benzene		15.52	15.53	0.000	146	148	377.204	17.46	17.5	99	
85	n-butylbenzene		15.60	15.61	0.000	91	134	1017.452	17.34	17.3	100	
86	12-diBr-3-Cl-Pra		15.97	16.00	-0.001	157	155	28.653	18.10	18.1	87	
87	124-tri-Cl-Bz		17.24	17.26	0.000	180	182	286.395	16.93	16.9	99	
88	naphthalene		17.49	17.50	0.000	128	129	236.255	18.23	18.2	100	
90	123-tri-Cl-Bz		17.68	17.69	0.000	180	182	211.274	17.95	17.9	100	
89	hx-Cl-butadiene		17.52	17.53	0.000	225	260	228.481	17.77	17.8	99	

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

Data File : C:\HPCHEM\1\DATA\03G1044\G1044Q01.D Vial: 9
 Acq On : 10 Jan 03 6:50 pm Operator: Eddie
 Sample : f=1 ccv/icv Inst : GCMS-G
 Misc : Multiplr: 1.00
 Quant Time: Jan 13 10:36 2003 Quant Results File: quant.res

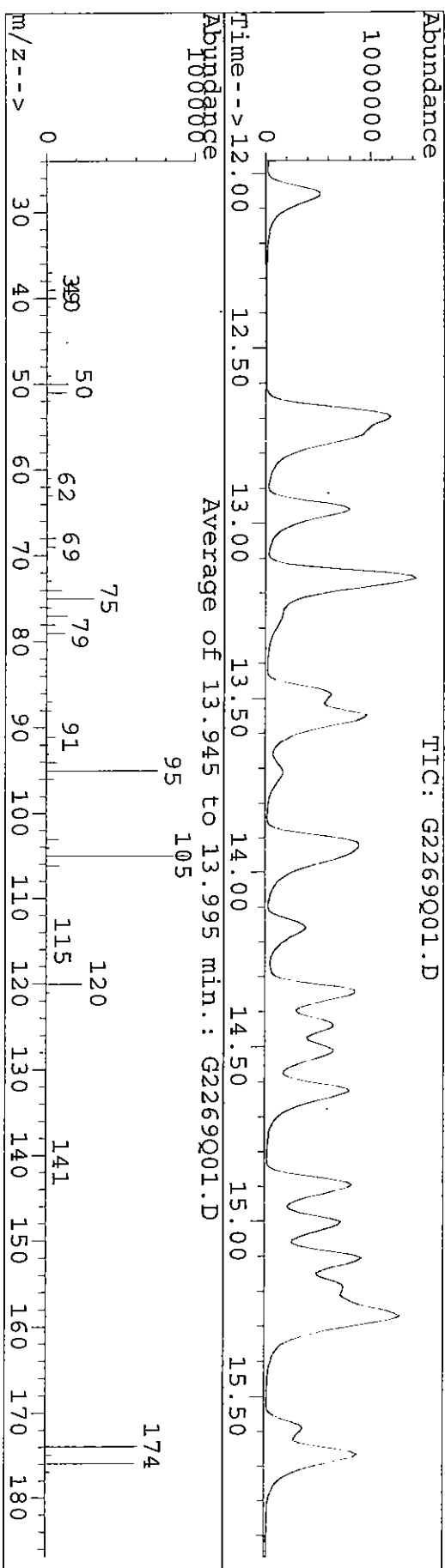
Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P & Ch Lab** EPA 524.2
 Last Update : Mon Jan 13 09:57:17 2003
 Response via : Multiple Level Calibration



Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :

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

Method : C:\HPCHEM\1\METHODS\E524G003.M
 Title : **Applied P &h Lab** EPA 524.2



Peak Apex is scan: 1443

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	19.6	14722	PASS
75	95	30	60	42.3	31794	PASS
95	95	100	100	100.0	75222	PASS
96	95	5	9	6.7	5047	PASS
173	174	0	2	0.0	0	PASS
174	95	50	100	82.8	62279	PASS
175	174	5	9	7.5	4689	PASS
176	174	95	101	96.3	60000	PASS
177	176	5	9	6.8	4089	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: 032964

Project ID: JPL

BFB Inj. Date: 04/30/03

Batch No: 03G2269

BFB Inj. Time: 17:40

Sequence No: 03G2269

Project No: 04-4428.10

Instrument ID: G

GC Column: DB-VEX

Data File Name: G2269Q01

Heated Purge: (Y/N) N

Column ID: 0.45 mm

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

#	Client Sample No	Lab Sample ID	Data File Name	Date Analyzed	Time Analyzed
1	03G2269-CCV-01	03G2269-CCV-01	G2269Q01	04/30/03	17:40
2	03G2269-LCS-01	03G2269-LCS-01	G2269L01	04/30/03	18:09
3	MW-17-2MS	03-2933-3MS	G2269M01	04/30/03	18:38
4	MW-17-2MSD	03-2933-3MSD	G2269N01	04/30/03	19:08
5	03G2269-MB-01	03G2269-MB-01	G2269K02	04/30/03	21:03
6	DUPE-4-2Q03	03-2964-1	2964-01	05/01/03	00:53
7	EB-7-4/29/03	03-2964-2	2964-02	05/01/03	01:21
8	MW-24-1	03-2964-3	2964-03	05/01/03	01:50
9	MW-24-2	03-2964-4	2964-04	05/01/03	02:19
10	MW-24-3	03-2964-5	2964-05	05/01/03	02:48
11	MW-24-4	03-2964-6	2964-06	05/01/03	03:17
12	MW-24-5	03-2964-7	2964-07	05/01/03	03:46
13	TB-7-4/29/03	03-2964-8	2964-08	05/01/03	04:15
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :

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

5009

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

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

Compound	AVGRF	CCRF	%Dev	Area%	Dev(min)
1 I	1 Fluorobenzene I1	1	1.000	1.000	0.0
2	3 di-Cl-di-F-methane	85	0.308	0.255	17.3
3 P	4 Chloromethane	50	0.243	0.207	15.1
4	9 F114 85 135		0.319	0.329	-3.0
5 C	5 vinyl chloride	62	0.231	0.210	9.2
6	6 bromomethane	94	0.227	0.195	14.2
7	7 Chloroethane	64	0.171	0.166	3.1
8	8 tri-Cl-F-methane	101	0.390	0.412	-5.7
9	111 isopropyl alcohol x10	10	0.005	0.004	12.3
10	100 ethyl ether x5		0.143	0.115	19.2
11	102 Acrolein x10		0.011	0.012	-1.7
12	119 methyl acetate		0.128	0.104	18.4
13	104 Carbon disulfide		0.834	0.651	21.9
14	103 Acrylonitrile x10		0.024	0.020	14.3
15	95 Acetone x10		0.019	0.017	8.9
16	108 F-113		0.300	0.346	-15.4
17 M,C	13 11-dichloroethene	61	0.392	0.380	3.1
18	101 Acetonitrile x10		0.006	0.000	95.5
19	109 Iodomethane		0.311	0.352	-13.3
20	113 Tert butyl alcohol x10		0.009	0.008	16.1
21	18 methylene chloride	49	0.643	0.381	40.7
22	112 Allyl chloride		0.434	0.407	6.1
23	200 Nitro methane x10		0.044	0.040	8.2
24	10 t-Bu-Me-ether	73	0.498	0.387	22.1
25	19 t-12-di-Cl-ethene	96	0.301	0.274	8.8
26	98 Vinyl acetate x5		0.270	0.239	11.5
27 P	21 11-dichloroethane	63	0.525	0.543	-3.5

(#) = Out of Range

G2269Q01.D E524G003.M Wed May 28 16:50:52 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :

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

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

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

Compound	AVGRF	CCRF	%Dev Area	%Dev	Area	Dev(min)
28	91 2-butanone MEKx10	0.094	0.080	14.3	94	0.00
29	115 Di isoprop ether	1.125	1.196	-6.3	93	0.00
30	22 c-12-di-Cl-ethene	0.294	0.277	5.8	98	0.00
31	23 22-Dichloropropane	0.453	0.454	-0.2	112	0.00
32	24 Br-Cl-methane	0.094	0.092	2.3	91	0.00
33	25 chloroform	0.525	0.481	8.3	102	0.00
34	201 Ethyl acetate x2	0.134	0.119	11.3	122	0.00
35	116 ETBE	0.807	0.690	14.4	91	0.00
36	117 Iso-butyl alcohol X10	0.027	0.024#	9.6	124	0.00
37	26 tetrahydrofuranx5	0.010	0.009#	12.9	91	0.00
38	27 Di-Br-F-Methane (S1)	0.400	0.340	14.9	100	0.00
39	34 111-tri-Cl-ethane	0.437	0.427	2.4	108	0.00
40	30 12-dichloroethane	0.175	0.180	-3.2	92	0.00
41	35 11-Di-Cl-propene	0.382	0.385	-0.9	107	0.00
42	29 1,2-di-Cl-ethane-d4 [Sur	0.165	0.141	14.7	89	0.00
43	36 benzene	0.869	0.876	-0.8	105	0.00
44	37 CCl4	0.335	0.357	-6.6	110	0.00
45	97 thiophene	0.432	0.401	7.2	98	0.00
46	118 TAME	0.556	0.471	15.3	90	0.00
47	39 12-di-Cl-propane	0.254	0.242	5.0	98	0.00
48	40 trichloroethene	0.269	0.271	-0.7	102	0.00
49	96 Me-methacrylate	0.095	0.079	17.0	98	0.00
50	42 Br-di-Cl-methane	0.355	0.307	13.4	97	0.00
51	41 dibromomethane	0.123	0.117	4.9	91	0.00
52	45 c-13-di-Cl-propene	0.313	0.290	7.3	99	0.00
53	55 toluene-d8(S2)	0.500	0.505	-1.0	106	0.00
54	92 2-ClEt-Vi-ether10	0.054	0.019#	64.3#	36	0.00

(#) = Out of Range

G2269Q01.D E524G003.M Wed May 28 16:50:57 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :

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

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

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

Compound	AVGRF	CCRF	%Dev Area	Dev(min)
55 M C 56 toluene	91	9	0.817	0.806
56 107 Et methacrylate	0.169	0.157	1.3	107
57 93 2-Hexanone x5	0.057	0.052	7.0	145
58 48 112-tri-Cl-Et	97	8	0.141	0.103
59 58 1,2-di-br-ethane	107	109	8.0	108
60 51 di-Br-Cl-methane	129	12	0.114	0.112
61 46 t-13-di-Cl-propene	75	11	0.161	0.147
62 105 1-Chlorohexane	0.202	0.187	8.7	92
	0.310	0.277	7.3	94
			10.8	113
				0.00
63 I 47 Cl-benzene-d5, I2	1.000	1.000	0.0	99
64 54 MIBK	0.297	0.301	-1.5	93
65 49 1,3-di-Cl-propane	76	78	0.654	0.637
66 59 tetra-Cl-ethene	166	16	2.6	92
67 M P 60 chlorobenzene	112	7	-12.1	106
68 61 1112-tetra-Cl-Et	131	13	-9.7	107
69 C 64 ethylbenzene	91	10	-7.3	99
70 65 m/p-Xylenes x2	2.816	3.096	-10.0	112
71 99 1-4-di-Cl-butane	2.141	2.328	-8.7	109
72 P 52 bromoform	0.645	0.575	-10.8	93
73 66 styrene	173	17	0.575	0.00
74 67 o-xylene	104	7	-6.7	91
75 P 68 1122-Tetra-Cl-Et	91	10	-6.7	106
76 110 t-1,4-dichloro-2-butene	83	8	-7.9	111
77 106 Cl-benzyl	0.387	0.364	5.9	94
	0.081	0.069	14.4	114
	0.572	0.565	1.2	106
				0.00
78 I 62 1,4-DCB-d4	150	152	0.0	102
79 69 123-tri-Cl-Pr	110	9	0.000	0.00
	0.122	0.109	10.6	90
				0.00

(#) = Out of Range

G2269Q01.D E524G003.M

Wed May 28 16:51:01 2003

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :

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

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

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

	Compound	AvGRF	CCRF	%Dev	Area% Dev(min)
80 S	70 4-Br-1-F-Bz (S3)	174 9	0.893	0.921	-3.1 106 0.00
81	71 isopropylbenzene	105 12	3.286	3.857	-17.4 119 0.00
82	72 bromobenzene	156 15	0.721	0.740	-2.5 97 0.00
83	73 n-propylbenzene	120 7	0.886	1.037	-17.1 116 0.00
84	74 2-Cl-Tl	126 128	0.589	0.606	-3.0 113 0.00
85	75 4-Cl-Tl	126 128	0.804	0.773	4.0 100 0.00
86	76 135-tri-Me-Bz	105 12	2.653	2.961	-11.6 115 0.00
87	79 tert-butylbenzene	119 9	2.724	3.089	-13.4 114 0.00
88	78 124-tri-Me-Bz	105 12	2.256	2.498	-10.7 111 0.00
89	80 13-di-Cl-Bz	146 148	1.149	1.104	4.0 104 0.00
90	82 14-di-Cl-Bz	146 148	1.670	1.730	-3.6 101 0.00
91	81 sec-butylbenzene	105 13	3.983	4.446	-11.6 114 0.00
92	77 4-iso-Pr-toluene	119 13	2.874	3.318	-15.5 115 0.00
93	84 12-di-Cl-benzene	146 14	1.109	1.043	6.0 95 0.00
94	85 n-butylbenzene	91 13	3.012	3.287	-9.1 113 0.00
95	86 12-diBr-3-Cl-Pra	157 15	0.081	0.073	10.1 87 0.00
96	87 124-tri-Cl-Bz	180 18	0.792	0.814	-2.8 95 0.00
97	88 naphthalene	128 12	0.665	0.571	14.1 87 0.00
98	90 123-tri-Cl-Bz	180 18	0.604	0.592	2.0 93 0.00
99	89 hx-Cl-butadiene	225 26	0.621	0.797	-28.4# 122 0.00

(#) = Out of Range
 G2269Q01.D E524G003.M SPCC's out = 0 CCC's out = 0
 Wed May 28 16:51:05 2003

Continuing Calibration Concentration Summary

Data File G2269Q01.D
Method File E524G003

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
1 Fluorobenzene l1 1	10	10.00	ppb	0.00	761786
3 di-Cl-di-F-methane 85 87	20	17.84	ppb	10.82	388177
4 Chloromethane 50 52	20	16.98	ppb	15.10	314773
9 F114 85 135	20	20.61	ppb	3.04	501305
5 vinyl chloride 62 64	20	18.16	ppb	9.19	319243
6 bromomethane 94 96	20	19.26	ppb	3.68	297323
7 Chloroethane 64 66	20	19.38	ppb	3.10	252239
8 tri-Cl-F-methane 101 103	20	21.14	ppb	5.68	628206
111 isopropyl alcohol x10	200	175.40	ppb	12.30	60291
100 ethyl ether x5	100	80.76	ppb	19.24	876837
102 Acrolein x10	200	203.31	ppb	1.65	176116
119 methyl acetate	20	17.07	ppb	14.66	158971
104 Carbon disulfide	20	15.62	ppb	21.90	992000
103 Acrylonitrile x10	200	171.37	ppb	14.32	311929
95 Acetone x10	200	202.99	ppb	1.50	263169
108 F-113	20	23.08	ppb	15.41	527111
13 11-dichloroethene 61 96	20	19.38	ppb	3.09	578845
101 Acetonitrile x10	200	6.47	ppb	96.76	4137
109 Iodomethane	20	22.19	ppb	10.97	536775
113 Tert butyl alcohol x10	200	178.57	ppb	10.72	117653
18 methylene chloride 49 84	20	19.35	ppb	3.27	581064
112 Allyl chloride	20	18.78	ppb	6.08	620506
200 Nitro methane x10	200	183.54	ppb	8.23	610000
10 t-Bu-Me-ether 73 57	20	18.30	ppb	8.48	590288
19 t-12-di-Cl-ethene 96 61	20	18.25	ppb	8.77	417743
98 Vinyl acetate x5	100	88.47	ppb	11.53	1822747
21 11-dichloroethane 63 83	20	20.70	ppb	3.52	827680
91 2-butanone MEK x10	200	171.45	ppb	14.27	1224926
115 Di isoprop ether	20	18.06	ppb	9.69	1822892
22 c-12-di-Cl-ethene 96 61	20	18.83	ppb	5.84	421869
23 22-Dichloropropane 77 97	20	20.04	ppb	0.18	691258
24 Br-Cl-methane 128 130	20	17.36	ppb	13.21	139523
25 chloroform 83 85	20	18.34	ppb	8.31	733492
201 Ethyl acetate x2	40	35.50	ppb	11.25	361550
116 ETBE	20	17.11	ppb	14.44	1051771
117 Iso-butyl alcohol X10	200	180.84	ppb	9.58	368565
26 tetrahydrofuran x5	100	87.15	ppb	12.85	68033
27 Di-Br-F-Methane (S1) 111 1	20	19.21	ppb	3.93	518608
34 111-tri-Cl-ethane 97 99	20	19.53	ppb	2.37	650600
30 12-dichloroethane 64 62	20	17.61	ppb	11.94	274524
35 11-Di-Cl-propene 75 110	20	20.17	ppb	0.86	587246
29 1,2-di-Cl-ethane-d4 [Surr] 10	20	17.06	ppb	14.70	214731
36 benzene 78 52	20	20.17	ppb	0.84	1335059
37 CCl4 117 119	20	21.31	ppb	6.56	543484

97 thiophene	20	18.57	ppb	7.16	611092
118 TAME	20	16.95	ppb	15.27	717138
39 12-di-Cl-propane 63 76	20	19.01	ppb	4.96	368246
40 trichloroethene 130 132	20	20.14	ppb	0.71	413159
96 Me-methacrylate	20	17.96	ppb	10.20	120737
42 Br-di-Cl-methane 83 85	20	17.33	ppb	13.37	468055
41 dibromomethane 174 172	20	19.01	ppb	4.94	177648
45 c-13-di-Cl-propene 75 110	20	18.54	ppb	7.29	441720
55 toluene-d8(S2) 100 99	20	20.20	ppb	1.02	768828
92 2-ClEt-Vi-ether10	200	71.42	ppb	64.29	291782

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
56 toluene 91 92	20	19.73	ppb	1.33	1227802
107 Et methacrylate	20	18.60	ppb	6.98	239507
93 2-Hexanone x5	100	92.03	ppb	7.97	398233
48 112-tri-Cl-Et 97 83	20	17.24	ppb	13.79	157257
58 1,2-di-br-ethane 107 109	20	17.46	ppb	12.68	170146
51 di-Br-Cl-methane 129 127	20	18.25	ppb	8.73	223865
46 t-13-di-cl-propene 75 110	20	18.55	ppb	7.26	284786
105 1-Chlorohexane	20	22.36	ppb	11.78	421400
47 Cl-benzene-d5, I2	10	10.00	ppb	0.00	223184
54 MIBK	20	19.12	ppb	4.40	134404
49 1,3-di-cl-propane 76 78	20	19.48	ppb	2.59	284260
59 tetra-Cl-ethene 166 168	20	22.42	ppb	12.08	440272
60 chlorobenzene 112 77	20	21.94	ppb	9.70	715660
61 1112-tetra-Cl-Et 131 133	20	21.46	ppb	7.31	280352
64 ethylbenzene 91 106	20	21.99	ppb	9.95	1381912

<u>Compound Name</u>	<u>Amount</u>	<u>Actual</u>	<u>Units</u>	<u>%Dev</u>	<u>Target Response</u>
65 m/p-Xylenes x2	40	43.49	ppb	8.73	2077922
99 1-4-di-Cl-butane	20	17.83	ppb	10.83	256743
52 bromoform 173 175	20	19.63	ppb	1.86	130445
66 styrene 104 78	20	21.34	ppb	6.71	720039
67 o-xylene 91 106	20	21.58	ppb	7.88	1014307
68 1122-Tetra-Cl-Et 83 85	20	18.82	ppb	5.91	162432
110 t-1,4-dichloro-2-butene	20	22.58	ppb	12.90	30780
106 Cl-benzyl	20	23.54	ppb	17.69	252253
62 1,4-DCB-d4 150 152 I3	10	10.00	ppb	0.00	187712
69 123-tri-Cl-Pr 110 97	20	17.87	ppb	10.65	40845
70 4-Br-1-F-Bz (S3) 174 95	20	20.61	ppb	3.07	345743
71 isopropylbenzene 105 120	20	23.48	ppb	17.38	1448076
72 bromobenzene 156 158	20	20.51	ppb	2.54	277685
73 n-propylbenzene 120 78	20	23.42	ppb	17.12	389498
74 2-Cl-TI 126 128	20	20.60	ppb	2.98	227537
75 4-Cl-TI 126 128	20	19.21	ppb	3.96	290072
76 135-tri-Me-Bz 105 120	20	22.32	ppb	11.60	1111705
79 tert-butylbenzene 119 91	20	22.68	ppb	13.42	1159830
78 124-tri-Me-Bz 105 120	20	22.15	ppb	10.73	937730
80 13-di-Cl-Bz 146 148	20	19.21	ppb	3.97	414282
82 14-di-Cl-Bz 146 148	20	20.72	ppb	3.61	649550
81 sec-butylbenzene 105 134	20	22.32	ppb	11.61	1669005

77 4-iso-Pr-toluene	119 134	20	23.09	ppb	15.46	1245699
84 12-di-Cl-benzene	146 148	20	18.81	ppb	5.95	391500
85 n-butylbenzene	91 134	20	21.83	ppb	9.14	1233921
86 12-diBr-3-Cl-Pra	157 155	20	17.99	ppb	10.06	27436
87 124-tri-Cl-Bz	180 182	20	18.69	ppb	6.53	305769
88 naphthalene	128 129	20	17.18	ppb	14.09	214523
90 123-tri-Cl-Bz	180 182	20	19.61	ppb	1.96	222391

Ave.% Dev	10.12
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Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269Q01.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 17:40 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:32 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:32 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
Internal Standards											
1	Fluorobenzene I1	9.07	9.03	0.004	96	70	761.786	10.00		0.04	
47	Cl-benzene-d5, I2	12.71	12.66	0.004	82	119	223.184	10.00		0.05	
62	1,4-DCB-d4 150 15	15.23	15.15	0.005	152	150	187.712	10.00		0.08	
System Monitoring Compounds (Surrogate)											
27	Di-Br-F-Methane (	7.48	7.44	0.003	111	113	518.608	19.21		19.2	%Recovery
29	1,2-di-Cl-ethane-	8.06	8.02	0.002	65	102	214.731	17.06		17.1	
55	toluene-d8(S2)	11.20	11.15	0.003	100	99	768.828	20.20		20.2	
70	4-Br-1-F-Bz (S3)	13.96	13.90	0.004	174	95	345.743	20.61		20.6	103.07%

Target Compounds											
<<< I1 : ISTD ID = 1 >>>											
3	di-Cl-di-F-methan	2.60	2.60	0.000	85	87	388.177	17.84		17.8	99
4	Chloromethane	2.78	2.78	0.000	50	52	314.773	16.98		17.0	98
9	F114 85 135	2.83	2.83	0.000	85	135	501.305	20.61		20.6	96
5	vinyl chloride	2.97	2.96	0.001	62	64	319.243	18.16		18.2	98
6	bromomethane	3.38	3.37	0.001	94	96	297.323	19.26		19.3	95
7	Chloroethane	3.53	3.52	0.002	64	66	252.239	19.38		19.4	98
8	tri-Cl-F-methane	4.17	4.14	0.003	101	103	628.206	21.14		21.1	98
111	isopropyl alcoho	4.29	4.27	0.002	45	43	60.291	175.40		175.4	1
100	ethyl ether x5	4.47	4.44	0.004	59	74	876.837	80.76		80.8	99
102	Acrolein x10	4.18	4.15	0.004	56	55	176.116	203.31		203.3	95
119	methyl acetate	5.09	5.06	0.004	43	74	158.971	17.07		17.1	95
104	Carbon disulfide	5.22	5.18	0.004	76	78	992.000	15.62		15.6	100
103	Acrylonitrilex10	4.91	4.89	0.003	53	52	311.929	171.37		171.4	99
95	Acetone x10	4.33	4.31	0.003	43	58	263.169	202.99		203.0	99
108	F-113	5.06	5.02	0.004	151	101	527.111	23.08		23.1	98
13	11-dichloroethene	4.79	4.76	0.004	61	96	578.845	19.38		19.4	99
101	Acetonitrilex10	4.08	4.20	-0.013	41	40	4.137	6.47		6.5	96
109	Iodomethane	4.83	4.80	0.003	142	127	536.775	22.19		22.2	99
113	Tert butyl alcoh	4.88	4.86	0.002	59	57	117.653	178.57		178.6	99

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269Q01.D Sample : f=1
 Method : C:\HPCHEM\1\METHODS\E524G003.M Inst. : GCMS-G
 Acq. Time : Apr 30 17:40 2003 RF via : Multiple Level Calibration
 Method Update: Mon Jan 13 10:38 2003 Operator: Eddie
 Quant. Time : May 01 11:32 2003 Multiplr: 1.000000
 Print Time : Thu May 01 11:33 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	QI	RF/1000	C0,ppb	C,ppb	Quality	Note
18	18 methylene chlorid	5.00	4.97	0.004	84	49	581.064	19.35	19.3	97	
112	Allyl chloride	5.10	5.07	0.004	41	76	620.506	18.78	18.8	94	?
200	200 Nitro methane x1	5.85	5.82	0.004	61	46	610.000	183.54	183.5	97	#?
10	10 t-Bu-Me-ether	6.03	6.00	0.004	73	57	590.288	18.30	18.3	97	
19	19 t-12-di-Cl-ethene	5.85	5.82	0.004	96	61	417.743	18.25	18.2	99	?
98	98 Vinyl acetate x5	6.44	6.41	0.004	43	86	1822.747	88.47	88.5	97	
21	21 11-dichloroethane	6.19	6.15	0.004	63	83	827.680	20.70	20.7	99	
91	91 2-butanone MEKx10	6.87	6.83	0.004	43	72	1224.926	171.45	171.5	97	?
115	115 Di isoprop ether	6.90	6.85	0.005	45	87	1822.892	18.06	18.1	98	?
22	22 c-12-di-Cl-ethene	7.02	6.97	0.005	96	61	421.869	18.83	18.8	99	
23	23 22-Dichloropropan	7.39	7.35	0.004	77	97	691.258	20.04	20.0	99	
24	24 Br-Cl-methane	7.24	7.18	0.006	128	130	139.523	17.36	17.4	98	
25	25 chloroform	7.30	7.27	0.004	83	85	733.492	18.34	18.3	99	
201	201 Ethyl acetate x2	7.35	7.31	0.004	43	61	361.550	35.50	35.5	95	?
116	116 ETBE	7.45	7.40	0.005	59	87	1051.771	17.11	17.1	100	
117	117 Iso-butyl alcoho	7.35	7.31	0.004	43	42	368.565	180.84	180.8	77	#?
26	26 tetrahydrofuranx5	7.74	7.71	0.003	72	42	68.033	87.15	87.1	97	
34	34 111-tri-Cl-ethane	8.27	8.23	0.004	97	99	650.600	19.53	19.5	99	
30	30 12-dichloroethane	8.17	8.13	0.004	62	64	274.524	17.61	17.6	99	
35	35 11-Di-Cl-propene	8.52	8.48	0.004	75	110	587.246	20.17	20.2	100	
36	36 benzene	8.79	8.75	0.004	78	52	1335.059	20.17	20.2	98	
37	37 CCl4	8.73	8.68	0.005	117	119	543.484	21.31	21.3	99	
97	97 thiophene	8.93	8.89	0.004	84	58	611.092	18.57	18.6	97	
118	118 TAME	9.04	9.00	0.004	73	43	717.138	16.95	16.9	97	
39	39 12-di-Cl-propane	9.54	9.49	0.005	63	76	368.246	19.01	19.0	98	
40	40 trichloroethene	9.60	9.54	0.006	130	132	413.159	20.14	20.1	100	
96	96 Me-methacrylate	9.89	9.85	0.004	69	100	120.737	17.96	18.0	100	
42	42 Br-di-Cl-methane	9.65	9.61	0.004	83	85	468.055	17.33	17.3	97	
41	41 dibromomethane	9.49	9.45	0.004	174	172	177.648	19.01	19.0	99	
45	45 c-13-di-Cl-propen	10.42	10.37	0.006	75	110	441.720	18.54	18.5	98	
92	92 2-ClEt-Vi-ether10	10.19	10.15	0.005	63	43	291.782	71.42	71.4	99	
56	56 toluene	11.28	11.23	0.006	91	92	1227.802	19.73	19.7	100	

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

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Method : C:\HPCHEM\1\METHODS\E524G003.M
 Acq. Time : Apr 30 17:40 2003
 Method Update: Mon Jan 13 10:38 2003
 Quant. Time : May 01 11:32 2003
 Print Time : Thu May 01 11:33 2003
 Miscellaneous :

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

5018

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
107	Et methacrylate	11.42	11.36	0.007	69	99	239.507	18.60	18.6	97	
93	2-Hexanone x5	11.54	11.48	0.007	43	58	398.233	92.03	92.0	96	
48	112-tri-Cl-Et	11.08	11.03	0.006	97	83	157.257	17.24	17.2	98	
58	1,2-di-br-ethane	11.88	11.82	0.007	107	109	170.146	17.46	17.5	100	
51	di-Br-Cl-methane	11.62	11.57	0.006	129	127	223.865	18.25	18.3	96	
46	t-13-di-Cl-propen	10.92	10.87	0.005	75	110	284.786	18.55	18.5	99	
105	1-Chlorohexane	12.69	12.64	0.006	55	93	421.400	22.36	22.4	99	
<<< 12 : ISTD ID = 47 >>>											
54	MIBK	10.59	10.52	0.005	43	58	134.404	19.12	19.1	98	
49	1,3-di-Cl-propane	11.34	11.29	0.004	76	78	284.260	19.48	19.5	99	
59	tetra-Cl-ethene	12.05	12.00	0.004	166	168	440.272	22.42	22.4	99	
60	chlorobenzene	12.76	12.70	0.004	112	77	715.660	21.94	21.9	97	
61	1112-tetra-Cl-Et	12.68	12.62	0.005	131	133	280.352	21.46	21.5	100	
64	ethylbenzene	12.96	12.90	0.005	91	106	1381.912	21.99	22.0	99	
65	m/p-Xylenes x2	13.16	13.10	0.005	91	106	2077.922	43.49	43.5	98	
99	1-4-di-Cl-butane	13.51	13.46	0.004	55	41	256.743	17.83	17.8	100	
52	bromoforn	13.27	13.22	0.004	173	175	130.445	19.63	19.6	92	
66	styrene	13.49	13.43	0.005	104	78	720.039	21.34	21.3	98	
67	o-xylene	13.55	13.50	0.004	91	106	1014.307	21.58	21.6	99	
68	1122-Tetra-Cl-Et	13.57	13.51	0.005	83	85	162.432	18.82	18.8	98	
110	t-1,4-dichloro-2	13.73	13.67	0.005	89	53	30.780	22.58	22.6	75	
106	Cl-benzyl	15.19	15.12	0.005	91	126	252.253	23.54	23.5	100	
<<< 13 : ISTD ID = 62 >>>											
69	123-tri-Cl-Pr	13.70	13.64	0.004	110	97	40.845	17.87	17.9	90	
71	isopropylbenzene	13.91	13.86	0.004	105	120	1448.076	23.48	23.5	99	
72	bromobenzene	14.16	14.10	0.004	156	158	277.685	20.51	20.5	96	
73	n-propylbenzene	14.35	14.29	0.004	120	78	389.498	23.42	23.4	99	
74	2-Cl-Tl	14.44	14.37	0.004	126	128	227.537	20.60	20.6	98	
75	4-Cl-Tl	14.51	14.45	0.004	126	128	290.072	19.21	19.2	95	
76	135-tri-Me-Bz	14.63	14.57	0.004	105	120	1111.705	22.32	22.3	99	
79	tert-butylbenzene	14.90	14.84	0.004	119	91	1159.830	22.68	22.7	98	
78	124-tri-Me-Bz	15.00	14.94	0.004	105	120	937.730	22.15	22.1	98	

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

75
 5/1/03

Data Filename: C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
Method : C:\HPCHEM\1\METHODS\E524G003.M
Acq. Time : Apr 30 17:40 2003
Method Update: Mon Jan 13 10:38 2003
Quant. Time : May 01 11:32 2003
Print Time : Thu May 01 11:33 2003
Miscellaneous :

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

ID	Component Name	R.T.	RT0	DRRT	Qion	Q1	RF/1000	C0,ppb	C,ppb	Quality	Note
80	13-di-Cl-Bz	146	15.18	15.11	0.004	146	148	414.282	19.21	19.2	90
82	14-di-Cl-Bz	146	15.25	15.19	0.004	146	148	649.550	20.72	20.7	94
81	sec-butylbenzene		15.11	15.04	0.004	105	134	1669.005	22.32	22.3	99
77	4-iso-Pr-toluene		15.28	15.22	0.004	119	134	1245.699	23.09	23.1	99
84	12-di-Cl-benzene		15.59	15.52	0.004	146	148	391.500	18.81	18.8	99
85	n-butylbenzene		15.67	15.60	0.004	91	134	1233.921	21.83	21.8	100
86	12-diBr-3-Cl-Pra		16.04	15.97	0.004	157	155	27.436	17.99	18.0	94
87	124-tri-Cl-Bz		17.32	17.24	0.005	180	182	305.769	18.69	18.7	98
88	naphthalene		17.55	17.49	0.004	128	129	214.523	17.18	17.2	99
90	123-tri-Cl-Bz		17.74	17.68	0.004	180	182	222.391	19.61	19.6	98
89	hx-Cl-butadiene		17.58	17.52	0.004	225	260	299.222	24.22	24.2	96

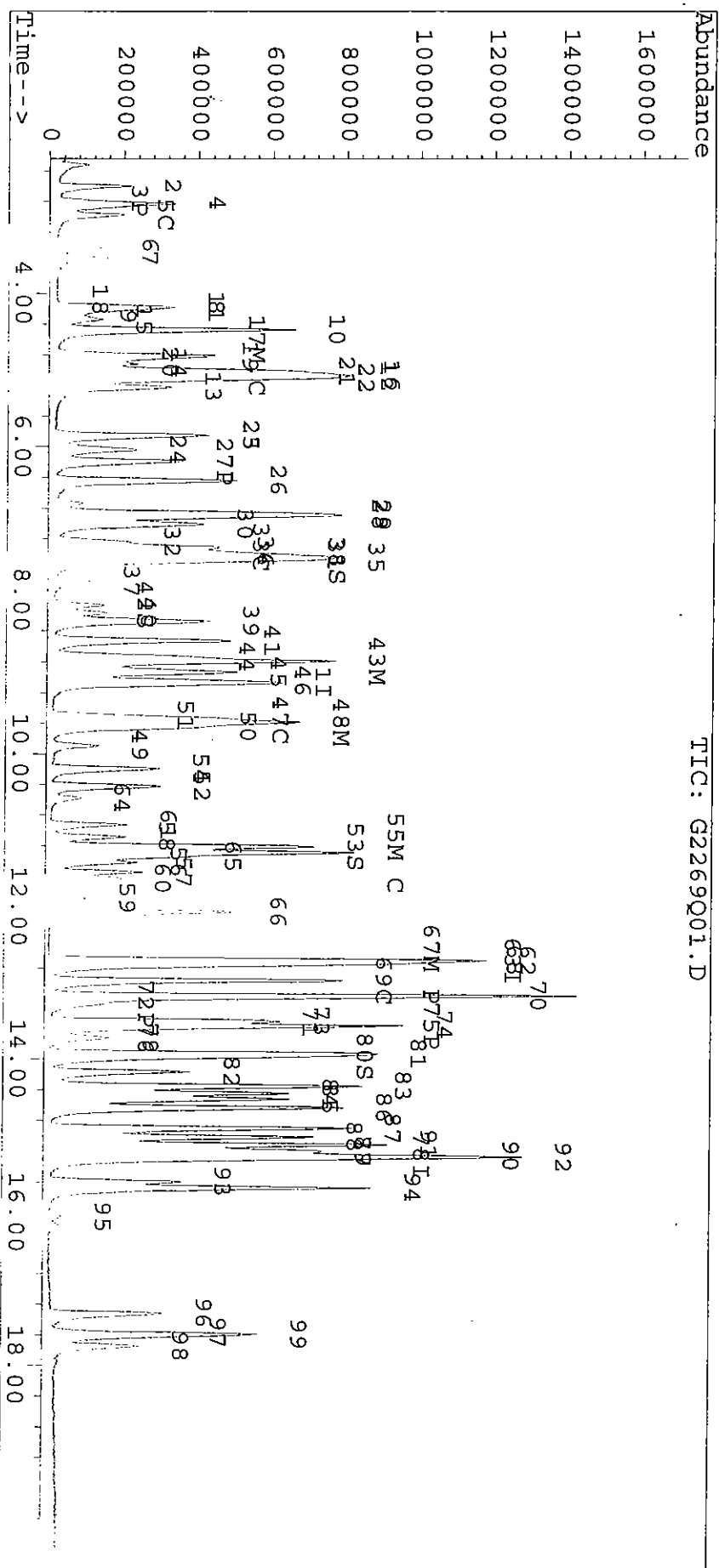
= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2269\G2269Q01.D
 Acq On : 30 Apr 03 5:40 pm
 Sample : f=1
 Misc :
 Quant Time: May 1 11:32 2003

Vial: 17
 Operator: Eddie
 Inst : GCMS-G
 Multiplr: 1.00
 Quant Results File: quant.res

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



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: 032964

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

CCV Data File: G2269Q01

Instrument ID: G

Batch No: 03G2269

#	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		04/30/03 17:40	761786 9.07	223184 12.71	187712 15.23
	CCV Upper Limit			1523572 9.57	446368 13.21	375424 15.73
	CCV Lower Limit			380893 8.57	111592 12.21	93856 14.73
1	03G2269-LCS-01	03G2269-LCS-01	04/30/03 18:09	760366 9.08	211140 12.72	183003 15.23
2	MW-17-2MS	03-2933-3MS	04/30/03 18:38	774675 9.07	217575 12.72	187626 15.24
3	MW-17-2MSD	03-2933-3MSD	04/30/03 19:08	799033 9.08	232426 12.73	196889 15.24
4	03G2269-MB-01	03G2269-MB-01	04/30/03 21:03	730711 9.12	192175 12.76	163983 15.28
5	DUPE-4-2Q03	03-2964-1	05/01/03 00:53	785372 9.11	227250 12.74	180327 15.25
6	EB-7-4/29/03	03-2964-2	05/01/03 01:21	748622 9.11	208208 12.74	177365 15.24
7	MW-24-1	03-2964-3	05/01/03 01:50	764286 9.11	212917 12.74	169783 15.25
8	MW-24-2	03-2964-4	05/01/03 02:19	732476 9.11	207779 12.75	169276 15.26
9	MW-24-3	03-2964-5	05/01/03 02:48	799254 9.10	217705 12.74	179025 15.25
10	MW-24-4	03-2964-6	05/01/03 03:17	817500 9.11	230007 12.75	176463 15.25
11	MW-24-5	03-2964-7	05/01/03 03:46	777713 9.12	215822 12.76	170404 15.25
12	TB-7-4/29/03	03-2964-8	05/01/03 04:15	780494 9.12	224518 12.76	175459 15.25
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						

IS-1 = FLUOROBENZENE

IS-2 = CHLOROBENZENE-D5

IS-3 = 1,4-DICHLOROBENZENE-D4

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

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

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

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

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

* Values outside of QC limits

Applied P & Ch Laboratory

100 Magnolia Ave. Chino CA 91710

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

VOC Analysis General Logbook

 Sample # 0361044 Batch # 0361044 Matrix: W Date: 1/10/03 Analyst: E. Li

 IS/Surrogate: GC14507/14508 Methanol(mark-M): _____ PEG (mark-PEG): _____ Defoaming(mark-DF): _____

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

	Type	Sample ID	Method	$V/X=f_1$	$V_1/V_i=f_2$	$V_{123}/V_{inj}=f_3$	F	A-#	Datafile	Note	pH
2378	SP	G1044P01	E524G-	21725=	/ =	/ =	1		G1044P01	1/10/03 1.51pH	
2379	Calib	3-0003	603	/ =	/ =	/ =			3-0003	GC14720	
2380		3-002		/ =	/ =	/ =			3-002		
2381		3-010		/ =	/ =	/ =			3-010		
2382		3-020		/ =	/ =	/ =			3-020		
2383		3-040		/ =	/ =	/ =			3-040		
2384		3-080		/ =	/ =	/ =			3-080		
2384	CCV	G1044Q01		/ =	/ =	/ =			G1044Q01	GC14721 CCV/ICV/IFD	
2385	MS	M01		/ =	/ =	/ =			M01	1011-03	C2
2386	VCS	L01		/ =	/ =	/ =			L01		
2387	MSD	N01		/ =	/ =	/ =			N01	1011-03	C2
2388	MSB	K01		/ =	/ =	/ =			K01		
2389	Sample	1084-03		/ =	/ =	/ =			1084-03		C2
2390		6844-02A		/ =	/ =	/ =			6844-02A		
2391		6844-01A		/ =	/ =	/ =			↓ -01A		
2392		1017-1		/ =	/ =	/ =			1017-1		
2393		1011-02		/ =	/ =	/ =			1011-02		
2394		3		/ =	/ =	/ =			3		
2395		4		/ =	/ =	/ =			4		
2396		5		/ =	/ =	/ =			↓ 5		
2397		1047-01		/ =	/ =	/ =			1047-01		
2398		↓ -02		/ =	/ =	/ =			↓ -02		
2399		1051-2		/ =	/ =	/ =			1051-02		
2400		↓ 3		/ =	/ =	/ =			↓ 3		
2401		1084-1		/ =	/ =	/ =			1084-01		
2402		↓ 2		/ =	/ =	/ =			↓ 2		
2403		↓ 4		/ =	/ =	/ =			↓ 4		
2404				/ =	/ =	/ =					
2405				/ =	/ =	/ =					
2406				/ =	/ =	/ =					
2407				/ =	/ =	/ =					
2408				/ =	/ =	/ =					

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	2386	GC-14721	$200 \times 2.5 / X = 20$ ppb		GC-	$x / X =$ ppb
MS/MSD	2385/2387	GC-	$x / X =$ ppb		GC-	$x / X =$ ppb

Note/Anomaly:

5022

Sample # 0221 2269 Batch # 07672269 Matrix: W Date: 4-30-07 Analyst: Eddie
 IS/Surrogate: GC-15114/15115 Methanol (mark-M): _____ PEG (mark-PEG): _____ Defoaming (mark-DF): _____
☒ Init. Batch ☒ Initial Batch; ☐ Middle Batch; ☐ Final Batch. ☐ Internal Study Datafile Path: _____

otnote/Anomaly:

Level C Data Package Deliverables

8270-SIM



Applied P & Ch Laboratory

13760 Magnolia Ave. Chino, CA 91710

Telephone (909)590-1828

Fax (909)590-1498

Applied P & Ch Laboratory
Organic Analysis Results for Method 8270-SIM

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 04/29/2003
Project ID: JPL	Service ID: 32964	Collected by:
Sample ID: MW-24-1	Lab Sample ID: 03-2964-3	Received Date: 04/29/2003
Sample Type: Field Sample	Sample Matrix: Water	Moisture %: -
Anal. Method: 8270-SIM	Prep. Method: 3520	Instrument ID: GC/MS: M
Batch No: 03G2304	Prep. Date: 05/01/03	Anal. Date: 05/02/03
Data File Name: 2964-03	Prep. No: 1 of 1	Anal. Time: 13:12
Extract Vol. 1.0 mL	Sample Amount: 1000 mL	Dilution Factor: 1

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	1,4-DIOXANE	123-91-1	µg/L	1	3.6	
Internal Standard				Control Limit, %	IS Rec.%	
1	1,4-DIOXANE-D8	17647-74-4		50-200	78	
# of out-of-control					0	

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 8270-SIM

Client Name: GEOFON, Inc.	Project No: 04-4428.10	Collection Date: 05/01/2003
Project ID: JPL	Service ID: 32964	Collected by:
Sample ID: 03G2304-MB-01	Lab Sample ID: 03G2304-MB-01	Received Date: 05/01/2003
Sample Type: Method Blank	Sample Matrix: Water	Moisture %: -
Anal. Method: 8270-SIM	Prep. Method: 3520	Instrument ID: GC/MS: M
Batch No: 03G2304	Prep. Date: 05/01/03	Anal. Date: 05/02/03
Data File Name: G2304K01	Prep. No: 1 of 1	Anal. Time: 11:34
Extract Vol. 1.0 mL	Sample Amount: 1000 mL	Dilution Factor: 1

#	Component Name	CAS No	Unit	RL	Result	Qualifier
1	1,4-DIOXANE	123-91-1	µg/L	1	<1	U
Internal Standard				Control Limit, %	IS Rec.%	
1	1,4-DIOXANE-D8	17647-74-4		50-200	75	
# 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-3C

Applied P & Ch Laboratory

Lab Control Spike/Lab Control Spike Duplicate Recovery for Method 8270-SIM

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32964

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

Batch No: 03G2304

LCS Filename: G2304L01

Date Analyzed: 050203

Time Analyzed: 11:53

LCSD Filename: G2304J01

Date Analyzed: 050203

Time Analyzed: 12:12

Spiked Components	Unit	Spike Added	Concentration		LCS Rec% #	QC Limit, % REC
			Unspiked	LCS		
1,4-DIOXANE	µg/L	20	0	19.5	98	40-140
# of Out-of-control					0	

Spiked Components	Unit	Spike Added	LCSD Concentration	LCSD Rec% #	RPD% #	QC Limit, %	
						RPD	REC
1,4-DIOXANE	µg/L	20	19.7	99	1	30	40-140
# 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-4B

Applied P & Ch Laboratory

Method Blank Summary for Method 8270-SIM

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 32964

Project ID: JPL

Project No: 04-4428.10

Analysis Date: 05/02/03

Sample Matrix: Water

Analysis Time: 11:34

Sample ID: 03G2304-MB-01

Batch No: 03G2304

Instrument ID: GC/MS: M

Lab Sample ID: 03G2304-MB-01

Data File Name: G2304K01

GC Column: DB-5.625

Column ID: 0.25 mm

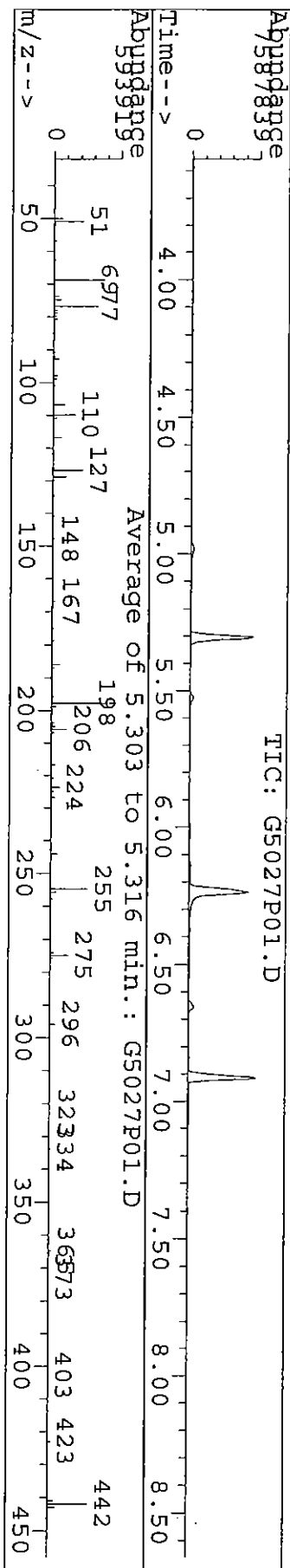
This Method Blank applies to the following samples and QC samples:

#	Client Sample No	Lab Sample ID	Sample Type	Data Filename	Analysis Date	Analysis Time
1	03G2304-LCS-01	03G2304-LCS-01	Lab Control Spike	G2304L01	05/02/03	11:53
2	03G2304-LSD-01	03G2304-LSD-01	Lab Control Spike Duplicate	G2304J01	05/02/03	12:12
3	MW-24-1	03-2964-3	Field Sample	2964-03	05/02/03	13:12
4						
5						
6						
7						
8						
9						
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						
21						
22						
23						
24						
25						

Data File : C:\HPCHEM\1\DATA\02G5027\G5027P01.D
 Acq On : 17 Dec 02 2:19 pm
 Sample : ##02g5027,w dftpp gc14349
 Misc :

Vial: 99
 Operator: Andy Huang
 Inst : GC/MS - M
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\DIOSIM06.M
 Title : * Applied P & Ch Lab * GC/MS 8270



Peak Apex is scan: AVERAGE

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
51	198	30	80	44.9	254307	PASS
68	69	0	2	0.0	0	PASS
69	198	1	100	78.3	443243	PASS
70	69	0	2	0.3	1277	PASS
127	198	25	75	46.2	261139	PASS
197	198	0	1	0.0	0	PASS
198	198	100	100	100.0	565803	PASS
199	198	5	9	6.7	37981	PASS
275	198	10	30	27.4	155040	PASS
365	198	1	100	3.7	20870	PASS
441	443	1	99	78.6	52501	PASS
442	198	40	110	61.8	349717	PASS
443	442	15	24	19.1	66784	PASS

5B
SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name : APPLIED P & CH LAB Contract: _____
 Lab Code: _____ Case No.: _____ SAS No.: _____ SDG No.: 03-2964
 Lab File ID: G5027P01 DFTPP Injection Date: 12/17/02
 Instrument ID: GCMS-M DFTPP Injection Time: 1419

m/e	ION ABUNDANCE CRITERIA	%RELATIVE ABUNDANCE
51	30.0 - 60.0% of mass 198	44.9
68	Less than 2.0% of mass 69	0.0 (0.0)1
69	Mass 69 relative abundance	78.3
70	Less than 2.0% of mass 69	0.2 (0.3)1
127	40.0 - 60.0% of mass 198	46.2
197	Less than 1.0% of mass 198	0.0
198	Base Peak, 100 % relative abundance	100.0
199	5.0 - 9.0% of mass 198	6.7
275	10.0 - 30.0% of mass 198	27.4
365	Greater than 0.75% of mass 198	3.7
441	Present, but less than mass 443	9.3 (78.6)3
442	40.0 - 100.0% of mass 198	61.8
443	17.0 - 23.0% of mass 442	11.8 (19.1)2

1-Value is % mass 69

2-Value is % mass 442

3-Value is % mass 443

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	SSTD040	DS006-40	DS006-040.D	12/17/2002	1438
02	SSTD030	DS006-30	DS006-030.D	12/17/2002	1457
03	SSTD020	DS006-20	DS006-020.D	12/17/2002	1516
04	SSTD010	DS006-10	DS006-10.D	12/17/2002	1536
05	SSTD001	DS006-01	DS006-01.D	12/17/2002	1555
06					
07					
08					
09					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					

Method : C:\HPCHEM\1\METHODS\DIOSIM06.M
 Title : * Applied P & Ch Lab * GC/MS 8270
 Last Update : Tue Dec 17 16:08:23 2002
 Response via : Initial Calibration

Calibration Files
 30 =M06-030.D 20 =M06-020.D 10 =M06-010.D
 1 =M06-001.D 40 =M06-040.D

Compound		30	20	10	1	40	Avg	%RSD

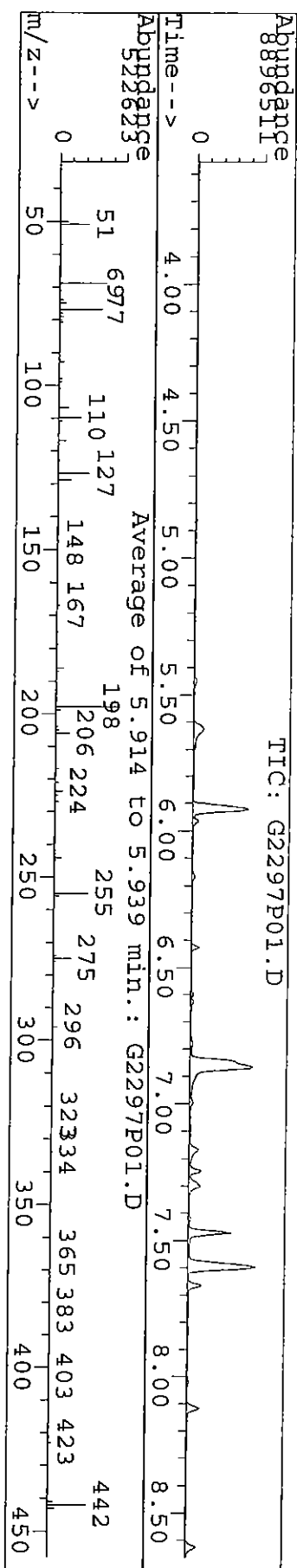
1) I	1 1,4-Dioxane-d8	-----ISTD-----						
2)	2 1,4-Dioxane	1.151	1.165	1.216	1.379	1.165	1.215	7.81

(#) = Out of Range
 DIOSIM06.M Tue Dec 17 16:08:40 2002 5972

Data File : C:\HPCHEM\1\DATA\03G2297\G2297P01.D
 Acq On : 2 May 03 9:57 am
 Sample : ##03g2297,s dftpp gcl4349
 Misc :

Vial: 99
 Operator: Andy Huang
 Inst : GC/MS - M
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\DIOSIM06.M
 Title : * Applied P & Ch Lab * GC/MS 8270



Peak Apex is scan: AVERAGE

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result
51	198	30	80	45.2	224750	PASS
68	69	0	2	0.0	0	PASS
69	198	1	100	72.5	360752	PASS
70	69	0	2	0.9	3341	PASS
127	198	25	75	49.4	245959	PASS
197	198	0	1	0.0	0	PASS
198	198	100	100	100.0	497757	PASS
199	198	5	9	7.0	34603	PASS
275	198	10	30	27.4	136388	PASS
365	198	1	100	3.4	16958	PASS
441	443	1	99	79.6	45081	PASS
442	198	40	110	60.1	298989	PASS
443	442	15	24	18.9	56603	PASS

FORM-5B

Applied P & Ch Laboratory

Semivolatile Organic Instrument Performance Check for Method 8270-SIM

Decafluorotriphenylphosphine (DFTPP), Part II

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 032964

Project ID: JPL

DFTPP Inj. Date: 05/02/03

Batch No: 03G2304

DFTPP Inj. Time: 09:57

Sequence No: 03G2297

Project No: 04-4428.10

Instrument ID: M

GC Column: DB-5.625

Data File Name: G2297P01

Column ID: 0.25 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	03G2304-CCV-01	03G2304-CCV-01	G2297Q01	05/02/03	10:16
2	03G2304-MB-01	03G2304-MB-01	G2304K01	05/02/03	11:34
3	03G2304-LCS-01	03G2304-LCS-01	G2304L01	05/02/03	11:53
4	03G2304-LSD-01	03G2304-LSD-01	G2304J01	05/02/03	12:12
5	MW-24-1	03-2964-3	2964-03	05/02/03	13:12
6					
7					
8					
9					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					

Evaluate Continuing Calibration Report

Data File : C:\HPCHEM\1\DATA\03G2297\G2297Q01.D
 Acq On : 2 May 03 10:16 am
 Sample : gcl4554
 Misc :
 Vial: 100
 Operator: Andy Huang
 Inst : GC/MS - M
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\DIOSIM06.M
 Title : * Applied P & Ch Lab * GC/MS 8270
 Last Update : Fri May 02 10:23:53 2003
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 200%

Compound		AvgRRF	CCRF	%Dev Area% Dev(min)	
1	1 1,4-Dioxane-d8	1.000	1.000	0.0	101 0.00
2	2 1,4-Dioxane	1.215	1.171	3.6	102 0.00

(#) = Out of Range
 G2297Q01.D DIOSIM06.M
 SPC's out = 0 CCC's out = 0
 Fri May 02 11:32:39 2003 5972

Data Filename: C:\HPCHEM\1\DATA\03G2297\G2297Q01.D Sample : gcl4554
 Method : C:\HPCHEM\1\METHODS\DIOSIM06.M Inst. : GC/MS - M
 Acq. Time : May 2 10:16 2003 RF via : Multiple Level Calibration
 Method Update: Fri May 02 10:23 2003 Operator: Andy Huang
 Quant. Time : May 02 11:32 2003 Multiplr: 1.000000
 Print Time : Fri May 02 11:32 2003
 Miscellaneous :

ID	Component Name	R.T.	RT0	DRRT	QIon	Q1	RF/1000	C0,ppm	C,ppb	Quality	Note
Internal Standards											
1	1,4-Dioxane-d8	5.34	5.34	0.000	96	66	471.664	20.00		Dev(Min)	0.00
System Monitoring Compounds (Surrogate)											
Target Compounds											
<<< I1 : ISTD ID = 1 >>>											
2	1,4-Dioxane	5.40	5.40	0.000	88	58	552.289	19.27	19272.4	85	Qvalue

%Recovery

= qualifier out of range, m = manual integration, ? = RT coelution, * = DRRT > 0.06
 5972

Quantitation Report

Data File : C:\HPCHEM\1\DATA\03G2297\G2297Q01.D

Vial: 100
Operator: Andy Huang

Acq On : 2 May 03 10:16 am

Inst : GC/MS - M

Sample : gcl4554

Multiplr: 1.00

Misc :
Quant Time: May 2 11:32 2003

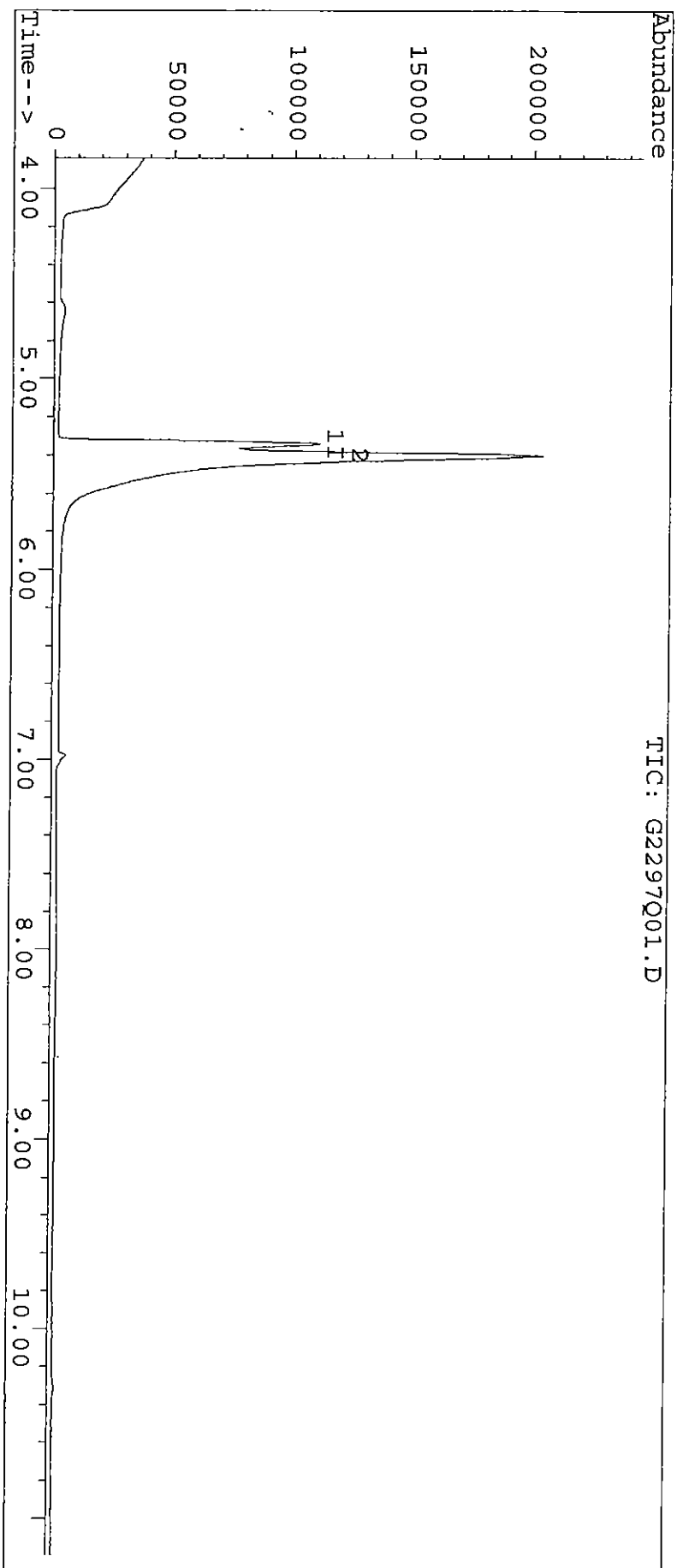
Quant Results File: quant.res

Method : C:\HPCHEM\1\METHODS\DIOSIM06.M

Title : * Applied P & Ch Lab * GC/MS 8270

Last Update : Fri May 02 10:23:53 2003

Response via : Multiple Level Calibration



FORM-8B

Applied P & Ch Laboratory

Internal Standard Area and RT Summary for Method 8270-SIM

Client Name: GEOFON, Inc.

Contract No:

Lab Code: APCL

Case No:

SAS No:

Service ID: 032964

Project ID: JPL

Project No: 04-4428.10

Sample Matrix: Water

CCV Data File: G2297Q01

Instrument ID: M

Batch No: 03G2304

#	Client	Lab	Analysis	IS-1	
	Sample No	Sample ID	Date & Time	Area #	RT #
12 Hour CCV STD			05/02/03 10:16	471664	5.34
CCV Upper Limit				943328	5.84
CCV Lower Limit				235832	4.84
1	03G2304-MB-01	03G2304-MB-01	05/02/03 11:34	354962	5.35
2	03G2304-LCS-01	03G2304-LCS-01	05/02/03 11:53	320271	5.40
3	03G2304-LSD-01	03G2304-LSD-01	05/02/03 12:12	283812	5.39
4	MW-24-1	03-2964-3	05/02/03 13:12	365843	5.46
5					
6					
7					
8					
9					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					

IS-1 = 1,4-DIOXANE-D8

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

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

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

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

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

* Values outside of QC limits

Applied P & Ch Laboratory

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Organic Sample Preparation Logbook

Batch # 0392304 Matrix: W Target method: 1,4-Dioxane EXT Method: 2570C

Date: 5-1-2

Solvent: Chel Lot # 007471 Exchange Solvent: NA Lot # NA

Analyst: ML

Lot #: Surr.: GC NA Conc.: NA ppm Na₂SO₄: Recov 78

Op. #	Sample Type	Sample ID #	Smpl Amnt (mL or g)	Surr. Vol mL	Extract sub-ID	Final Vol. mL	Dilution F=V/X	Note & Anomaly
3131	MB	03-2304-4	1000.0	(V _A)	1,4-Dioxane	1.00	0.001	
3132	LCS	201	1000.0					
3133	LCSD	-201	1000.0					
3134	Sample-1	03-2933-5	1000.0					
3135	MS							
3136	MSD							
3137	Sample-2	03-2940-2	1000.0	NA	1,4-Dioxane	1.00	0.001	
3138	Sample-3	03-2964-3	1000.0					
3139	Sample-4							
3140	Sample-5							
3141	Sample-6							
3142	Sample-7							
3143	Sample-8							
3144	Sample-9							
3145	Sample-10							
3146	Sample-11							
3147	Sample-12							
3148	Sample-13							
3149	Sample-14							
3150	Sample-15							
3151	Sample-16							
3152	Sample-17							
3153	Sample-18							
3154	Sample-19							
3155	Sample-20							
3156	MTX Dup.							
3157	LCS'							
3158	LCSD'							
3159	MS'							
3160	MSD'							

Type	Op #	STD Lot #	$C_{std}(ppm) \times V_{std}(mL) / X(g \text{ or } mL) = T$				Op #	STD Lot #	$C_{std}(ppm) \times V_{std}(mL) / X(g \text{ or } mL) = T$							
LCS/LCSD	3132	GC-14768	1000	x	20.0	x	1.00	3133	GC-14768	1000	x	20.0	x	1.00	/X=	ppb
MS/MSD		GC-	1000	x		x	/X=		GC-	1000	x		x	/X=	ppb	
LCS'/LCSD'		GC-	1000	x		x	/X=		GC-	1000	x		x	/X=	ppb	
MS'/MSD'		GC-	1000	x		x	/X=		GC-	1000	x		x	/X=	ppb	

Cur

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Organic Sample Preparation

Extraction method: 1 3510-separate funnel; 2 3520-L-L continue extraction; 3 3550-Ultrasonic; 4 LUFF-Shaker; 5 3540-Soxhlet;
6 Solid phase Extraction; 7 Microextraction; 8 SFE; 9 Dilution; 10 Concentration; 11 Other, specify.

Service ID: 2933 Batch # 0362304 Matrix W GC, GC/MS Method 14-Dioxane Ext Method 3520C

Sample Extraction

OP. by HL Date 5/1/03 Surro Lot GC-NA Conc. NA PP
LCS Lot GC-14768 Conc. 20.0 PP MS Lot GC-14768 Conc. 20.0 PP
Ext. solv. Which Lot # 027471 Exchange solv. NA Lot # NA

Sample ID.

Sample Matrix

Sample Amount, X (g/mL)

Any TCLP, EP, WET, SPLP ?

Surr. STD used, mL

Spike STD used, mL

[Extracted at pH]

Solvent amount, mL

Final volume, V, mL

Extraction DF $f_c = V/X$

Sub-ID of extracts

Anomaly Footnote¹:

Sample Cleanup

Cleanup method:

Cleanup DF, f_c

Preparation DF, $f_1 = f_c f_c$

Op. by _____ Date 1/1 (For details, see Cleanup worksheet)

Pre-injection*:

2nd dilution, f_2 (1) Op. by _____ Date 1/1 IS Lot: GC Conc. _____ PP

V_i , μ L (40) _____

Single Extract GC, GC/MS Analysis:

$V_2 = V_i/f_2$, μ L (40/ f_2) _____

$V_{\text{solvent}} = V_i - V_2$, μ L _____

Double Extract GC, GC/MS Analysis:

$V_{>} = \frac{1}{2} V_i/f_2$, μ L (20/ f_2) _____

$V_{<} = \frac{1}{2} \frac{V_i}{f_2} \frac{f_{<}}{f_{>}}$, μ L ($\frac{20}{f_2} \frac{f_{<}}{f_{>}}$) _____

$V_{\text{solvent}} = V_i - V_{>} - V_{<}$ _____

IS volume, μ L (10) _____

Final Conc. of IS (40 ppm) _____

Total F = 2 $f_1 f_2$ or $f_1 f_2$ _____

Extraction Anomaly Footnote:

- Difficult to concentrate to 1.0 mL.
- Extract is dark color
- Extract is sticky
- Heterogenous matrix.
- Too much precipitate during extraction.
- There is an oil layer above the sample.
- Mixture of soil & water.
- BN-extract is interfered, A-extract only.
- Sample has strong stink
- Some solvent suspected
- special color, specify

For double extract analysis (ABN), $f_{>} = \max[f_A, f_B]$ and $f_{<} = \min[f_A, f_B]$, where f_A, f_B are f_1 for A and BN extracts, respectively.
The values in () is the typical case for 625/8270 ABN analysis.

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

Tel: (909) 590-1626 Fax: (909) 590-1498

Organic Sample Preparation

Extraction method: 1 3510-separate funnel; 2 3520-L-L continue extraction; 3 3550-Ultrasonic; 4 LUFT-Shaker; 5 3540-Soxhlet;
6 Solid phase Extraction; 7 Microextraction; 8 SFE; 9 Dilution; 10 Concentration; 11 Other, specify.

Service ID: 2964 Batch # 352304 Matrix W GC, GC/MS Method 14-Dioxane Ext Method 3520C

Sample Extraction

Op. by HL Date 5/1/03 Surro Lot GC-NA Conc. NA pp
LCS Lot GC-14768 Conc. 20.0 pp MS Lot GC-14768 Conc. 20.0 pp
Ext. solv. Which Lot # 027471 Exchange solv. NA Lot # NA

Sample ID. -3

Sample Matrix W

Sample Amount, X (g/mL) 1000

Any TCLP, EP, WET, SPLP ?

Surr. STD used, mL NA

Spike STD used, mL

[Extracted at pH] 1

Solvent amount, mL 200.0 X

Final volume, V, mL 1.00

Extraction DF $f_e = V/X$ 200/

Sub-ID of extracts 14-Dioxane

Anomaly Footnote† :

Sample Cleanup

Cleanup method:

up DF, f_c

Preparation DF, $f_1 = f_e f_c$

Op. by _____ Date 1/1 (For details, see Cleanup worksheet)

Pre-injection*:

2nd dilution, f_2 (1) _____ Op. by _____ Date 1/1 IS Lot: GC- Conc. _____ pp

V_i , μL (40) _____

Single Extract GC, GC/MS Analysis:

$V_2 = V_i/f_2$, μL (40/ f_2) _____

$V_{\text{solvent}} = V_i - V_2$, μL _____

Double Extract GC, GC/MS Analysis:

$V_> = \frac{1}{2} V_i/f_2$, μL (20/ f_2) _____

$V_< = \frac{1}{2} V_i/f_2$, μL (20/ f_2) _____

$V_{\text{solvent}} = V_i - V_> - V_<$ _____

IS volume, μL (10) _____

Final Conc. of IS (40 ppm) _____

Total F = 2 $f_1 f_2$ or $f_1 f_2$ _____

† Extraction Anomaly Footnote:

1. Difficult to concentrate to 1.0 mL.
2. Extract is dark color
3. Extract is sticky
4. Heterogenous matrix.
5. Too much precipitate during extraction.
6. There is an oil layer above the sample.
7. Mixture of soil & water.
8. BN-extract is interfered, A-extract only.
9. Sample has strong stink
10. Some solvent suspected
11. special color, specify

* For double extract analysis (ABN), $f_2 = \max[f_A, f_B]$ and $f_1 = \min[f_A, f_B]$, where f_A, f_B are f_1 for A and BN extracts, respectively. The values in () is the typical case for 625/8270 ABN analysis.

Analysis Anomaly Record

Batch #: 0392304 Method: 1.4 Dioxane

Matrix: w Date: 5-1-03

Samples Involved: 2933, 2940, 2964

• Anomaly for MS/MSD/MD

- ☐ 1. MS/MSD out of control limit, sample matrix interference suspected
- ☐ 2. MS or MSD out of control limit, but $\overline{MS}$ is O.K.
- ☐ 3. One of MS/MSD, and $\overline{MS}$ out of control limit, but not enough sample for re-analysis
- ☒ 4. Not enough sample for MS/MSD, LCS/LCSD may be used
- ☐ 5. Spiked samples contain high concentration of analytes ($>4 \times$ spiked concentration)
- ☐ 6. MS/MSD out of control limit, but post spike is O.K.
- ☐ 7. MS/MSD out of control limit, due to the heterogenous matrix
- ☐ 8. MS/MSD out of control limit, due to the very special matrix
- ☐ 9. Not enough sample for matrix duplicate (MD)

• Anomaly for Holding time (HT)

- ☐ 10. HT has been exceeded when received, authorized by the Client to carry on with the analyses
- ☐ 11. Extraction HT was passed, authorized by the Client to carry on with the analyses
- ☐ 12. Analysis HT was passed, authorized by the Client to carry on with the analyses

• Surrogates Anomaly

- ☐ 13. Surrogate recovery out of control limit (see Level-1 Part III), sample matrix interference suspected
- ☐ 14. Surrogate recovery out of control limit (see Level-1 Part III), but not enough sample for re-analysis
- ☐ 15. Surrogate recovery out of control limit, due to very special matrix
- ☐ 16. Surrogate diluted out (for highly contaminated samples)
- ☐ 17. Lower surrogate recovery due to cleanup process

• Chromatogram Pattern Anomaly

- ☐ 18. Not a typical gasoline chromatogram pattern
- ☐ 19. Not a typical diesel chromatogram pattern
- ☐ 20. Not a typical diesel chromatogram pattern, but similar to heavy-oil or motor oil
- ☐ 21. Not a typical kerosene chromatogram pattern
- ☐ 22. Not a typical jet fuel chromatogram pattern
- ☐ 23. Not a complete PCB chromatogram pattern, interference/degradation suspected

• Other anomaly

Problem Description: _____

Reason: _____

Analyst: HL Date: 5-1-03

Note: _____

Applied P & Ch Laboratory

13760 Magnolia Ave. Chino CA 91710

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

Semi-VOC Analysis Logbook

M LOG-136

Sequence # 0362297

Starting Date: 5-2-03 Time 9:57am Analyst: Jmr

Seq. type: ☐ Cal. ☒ Ini. Batch ☐ Middle ☐ Final ☐ Continue ☐ Study

Datafile Path: C:\HPCHEM\1\DATA\0362297

Routine Maintenance: ☐ Replace Septum ☒ Replace Liner ☐ Replace Seal ☐ Cut Guid Column ☒ Cut Column ☐ Others

Op. #	Batch-No	MTX	S Type*	Sample ID	Method**	f ₂	F	A-#	Datafile	OK ?†	Note††
5321			SP	62297P01	PFTPP625			99	62297P01	-	5-2-03 9:57am
5322			CCV	Q01	D10S1M06			100	Q01	-	GC14554
5323	0362297	S	MB	P01			0.0333	1	P01	-	
5324			LCS	L01				2	L01	-	
5325			LCSD	J01	Jmr			3	J01	-	
5326	0362304	W	MB	2933G230401	5/1/03		0.001	4	G2304P01	-	
5327			LCS	G2304L01				5	L01	-	
5328			LCSD	J01				6	J01	-	
5329				2933-05				7	2933-05	-	
5330				2940-02				8	2940-02	-	
5331				2964-03				9	2964-03	-	
5332	0362297	S		2997-02			0.0333	10	2997-02	-	
5333				-040				11	-040	-	
5334				2929-11				12	2929-11	-	
5335				2920-03				13	2920-03	-	
5336				-10				14	-10	-	
5337				-12				15	-12	-	
5338				-23				16	-23	-	
5339			MS	G2297M01				17	G2297M01	-	\$2920-03
5340			MSM	N01				18	N01	-	1
5341											
5342											
5343											
5344											
5345											
5346											
5347											
5348											
5349											
5350											

Footnote/Anomaly:

* Sample type such as, SP(BFB, or DFTPP), CCV, CVA, CVB, LCS, LCSD, MS, MSD, MD, CLS etc. For field samples leave as blank.

** Method name: [MMMM][S][nnn]. [MMMM] - method code, per SOP C-88. Special codes: DSL2: Diesel #2; FP30: Fuel Finger Print (C8-C30); FP40: Fuel Finger Print (C8-C40); EGLY: Ethylene Glycol; FORM: Formaldehyde; MEON: Methyl Alcohol. [S] - APCL system code; [nnn] - initial calibration number.

† Specify the result of the injection is accepted (✓), or denied (X), or need Re-run (R) or use as a reference data (ref).

†† Lot # for CCV/Closing, Time for SP (e.g. DFTPP) must be recorded.

Semi-VOC Analysis Logbook

M LOG-063

Sample # 026502

Starting Date: 12-17-02 Time 2:19 pm Analyst: J.Y.

Sample type: ☐ Cal. ☒ Ini. Batch ☐ Middle ☐ Final ☐ Continue ☐ Study

Datafile Path: _____

Line Maintenance: ☐ Replace Septum ☒ Replace Liner ☐ Replace Seal ☐ Cut Guid Column ☐ Cut Column ☐ Others _____

	Batch-No	MTX	S Type*	Sample ID	Method**	f ₂	F	A-#	Datafile	OK ?†	Note††
			SP	G5027P01	DFTPP625			99	G5027P01	✓	12-17-02 2:19 pm
			Cali-	M06-040	D10S1M06			1	M06-040	✓	60ppm G414554
				-030				2	-030	✓	30
				-020				3	-020	✓	20
				-010				4	-010	✓	10
				-001				5	-001	✓	1
			CCV	G5027Q01				100	G5027Q01	✓	G414554
	026502	W	MB	K01			0.001	6	K01	✓	
			LCS	L01				7	L01	✓	
			LCSD	J01				8	J01	✓	
			MS	M01				9	M01	✓	\$6556-03
			MSD	N01				10	N01	✓	1
			MS'	M02			0.000962	11	M02	✓	\$6587-01
			MSD'	N02				12	N02	✓	1
				6587-01				13	6587-01	✓	
				-02				14	-02	✓	
				-03				15	-03	✓	
				6556-01			0.002	16	6556-01	✓	
				-02			0.001	17	-02	✓	
				-03				18	-03	✓	
				6588-01			0.000962	19	6588-01	✓	
				6603-01				20	6603-01	✓	
				-02				21	-02	✓	
				6558-01				22	6558-01	✓	
				-03			0.001	23	-03	✓	
				6586-02				24	6586-02	✓	
				-04				25	-04	✓	
				-05				26	-05	✓	
				-06				27	-06	✓	Reverse scale
				-08				28	-08	✓	↓

Footnote/Anomaly:

* Sample type such as, SP(BFB, or DFTPP), CCV, CVA, CVB, LCS, LCSD, MS, MSD, MD, CLS etc. For field samples leave as blank.

** Method name: [MMMM][S][nnn]. [MMMM] - method code, per SOP C-88. Special codes: DSL2: Diesel #2; FP30: Fuel Finger Print (C8-C30); FP40: Fuel Finger Print (C8-C40); EGLY: Ethylene Glycol; FORM: Formaldehyde; MEOR: Methyl Alcohol. [S] - APCL system code; [nnn] - initial calibration number.

† Specify the result of the injection is accepted (✓), or denied (X), or need Re-run (R) or use as a reference data (ref).

†† Lot # for CCV/Closing, Time for SP (e.g. DFTPP) must be recorded.

5045