

**GC/MS Semi-Volatile Data
Standards Data**

Form 6

Initial Calibration

Instrument: GCMS_5

Level #:	Data File:	Cal Identifier:	Analysis Date/Time	Level #:	Data File:	Cal Identifier:	Analysis Date/Time
1	5M09385.	CAL BNA@50PPM	07/22/05 08:30	2	5M09386.	CAL BNA@10PPM	07/22/05 08:53
3	5M09387.	CAL BNA@25PPM	07/22/05 09:16	4	5M09388.	CAL BNA@80PPM	07/22/05 09:39
5	5M09389.	CAL BNA@120PPM	07/22/05 10:01	6	5M09390.	CAL BNA@160PPM	07/22/05 10:24
7	5M09391.	CAL BNA@200PPM	07/22/05 10:47				

Compound	Col	Mr	Fit:	RF								AvgRf	RT	Corr1	Corr2	%Rsd	Calibration Level Concentrations							
				RF1	RF2	RF3	RF4	RF5	RF6	RF7	RF8						Lvl1	Lvl2	Lvl3	Lvl4	Lvl5	Lvl6	Lvl7	Lvl8
Benzo[k]fluoranthene	1	0	Avg	1.6308	1.5111	1.6774	1.6597	1.6416	1.6263	1.4430	---	1.60	13.26	0.990	0.997	5.4	50.00	10.00	25.00	80.00	120.0	160.0	200.0	
Benzo[a]pyrene	1	0	Avg	1.4320	1.3432	1.5480	1.4982	1.5275	1.5820	1.4730	----	1.49	13.57	0.997	0.998	5.4*(30)	50.00	10.00	25.00	80.00	120.0	160.0	200.0	
Indeno[1,2,3-cd]pyrene	1	0	Avg	1.5945	1.4162	1.7632	1.6166	1.6707	1.6107	1.5992	----	1.61	14.69	0.999	0.999	6.5	50.00	10.00	25.00	80.00	120.0	160.0	200.0	
Dibenzo[a,h]anthracene	1	0	Avg	1.3175	1.1697	1.4402	1.3822	1.3923	1.2996	1.3532	----	1.34	14.72	0.998	0.998	6.5	50.00	10.00	25.00	80.00	120.0	160.0	200.0	
Benzo[g,h,i]perylene	1	0	Avg	1.2961	1.2704	1.5177	1.3809	1.3592	1.3154	1.2871	----	1.35	14.99	0.998	0.999	6.3	50.00	10.00	25.00	80.00	120.0	160.0	200.0	

Flags

a - failed the spcc criteria * - ccc compound
 b - failed the ccc criteria ** - spcc compound
 c - failed the minimum correlation coeff criteria(if applicable)

Note:

Avg Rsd: 8.15

Corr 1 = Correlation Coefficient for linear Eq.
 Corr 2 = Correlation Coefficient for quad Eq.
 Fit = Indicates whehter Avg RF, Linear, or Quadratic Curve was used for compound.

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09385.D Vial: 2
 Acq On : 22 Jul 2005 8:30 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Jul 22 12:11 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	23355	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	88320	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	52347	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	87328	40.00	ng	-0.06
77) Chrysene-d12	11.95	240	70907	40.00	ng	-0.08
88) Perylene-d12	13.54	264	56016	40.00	ng	-0.07

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.90	112	34720	44.14	ng	-0.06
Spiked Amount	200.000		Recovery	=	22.07%	
8) Phenol-d5	4.90	99	47355	41.17	ng	-0.05
Spiked Amount	200.000		Recovery	=	20.59%	
21) Nitrobenzene-d5	5.67	128	9591	24.80	ng	-0.05
Spiked Amount	100.000		Recovery	=	24.80%	
41) 2-Fluorobiphenyl	7.04	172	42278	25.84	ng	-0.05
Spiked Amount	100.000		Recovery	=	25.84%	
64) 2,4,6-Tribromophenol	8.28	330	9693	51.86	ng	-0.06
Spiked Amount	200.000		Recovery	=	25.93%	
80) Terphenyl-d14	10.75	244	42105	25.14	ng	-0.06
Spiked Amount	100.000		Recovery	=	25.14%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.10	79	50366	51.72	ng	96
3) N-Nitrosodimethylamine	2.04	74	27605	46.60	ng	95
5) Aniline	4.91	93	57814	43.79	ng	89
6) Pentachloroethane	4.94	117	14353	49.85	ng	98
7) bis(2-Chloroethyl)ether	4.98	93	38291	46.34	ng	98
9) Phenol	4.91	94	53005	43.48	ng	60
10) 2-Chlorophenol	5.01	128	41560	44.91	ng	99
11) 1,3-Dichlorobenzene	5.14	146	42174	49.26	ng	99
12) 1,4-Dichlorobenzene	5.21	146	42947	49.02	ng	100
13) 1,2-Dichlorobenzene	5.34	146	41533	49.72	ng	99
14) Benzyl alcohol	5.33	108	26925	44.06	ng	99
15) bis(2-chloroisopropyl)ethe	5.45	45	57813	46.06	ng	95
16) 2-Methylphenol	5.44	108	37366	44.23	ng	98
17) Hexachloroethane	5.62	117	17912	49.29	ng	77
18) N-Nitroso-di-n-propylamine	5.56	70	28766	43.19	ng	94
19) 3&4-Methylphenol	5.56	108	39924	44.45	ng	99
22) Nitrobenzene	5.68	77	41731	48.11	ng	99
23) Isophorone	5.88	82	76782	47.56	ng	98
24) 2-Nitrophenol	5.94	139	22756	51.12	ng	96

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09385.D Vial: 2
 Acq On : 22 Jul 2005 8:30 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 12:11 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.99	107	41629	49.17	ng	99
26) Benzoic Acid	6.07	105	19966	41.08	ng	98
27) bis(2-Chloroethoxy)methane	6.06	93	45377	49.12	ng	97
28) 2,4-Dichlorophenol	6.13	162	34895	49.42	ng	98
29) 1,2,4-Trichlorobenzene	6.19	180	36438	50.34	ng	100
30) Naphthalene	6.25	128	113670	49.11	ng	99
31) 4-Chloroaniline	6.30	127	46470	51.42	ng	98
32) Hexachlorobutadiene	6.34	225	20863	52.27	ng	98
33) 4-Chloro-3-methylphenol	6.65	107	35428	45.44	ng	98
34) 2-Methylnaphthalene	6.77	142	76447	47.82	ng	99
35) Methylnaphthalenes (Total)	6.77	142	76447	47.82	ng	99
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	37078	52.45	ng	98
38) Hexachlorocyclopentadiene	6.88	237	23632	54.28	ng	99
39) 2,4,6-Trichlorophenol	6.97	196	25325	49.71	ng	98
40) 2,4,5-Trichlorophenol	7.00	196	27732	50.04	ng	98
42) 2-Chloronaphthalene	7.14	162	73257	49.51	ng	97
43) 1,4-Dimethylnaphthalene	7.40	156	56858	50.49	ng	97
44) Dimethylnaphthalenes (Total)	7.40	156	56858	50.49	ng	97
45) Diphenyl Ether	7.20	170	47916	49.01	ng	90
46) 2-Nitroaniline	7.22	65	26263	46.60	ng	87
47) Acenaphthylene	7.46	152	116332	49.77	ng	99
48) Dimethylphthalate	7.36	163	82030	48.03	ng	100
49) 2,6-Dinitrotoluene	7.41	165	19292	49.04	ng	99
50) Acenaphthene	7.61	153	69508	48.08	ng	100
51) 3-Nitroaniline	7.55	138	21501	50.07	ng	100
52) 2,4-Dinitrophenol	7.63	184	10602	43.80	ng	90
53) Dibenzofuran	7.75	168	101145	48.06	ng	99
54) 2,4-Dinitrotoluene	7.74	165	26272	48.34	ng	92
55) 4-Nitrophenol	7.69	65	16212	47.42	ng	96
56) 2,3,4,6-Tetrachlorophenol	7.86	232	20985	48.27	ng	98
57) Fluorene	8.05	166	79949	47.15	ng	99
58) 4-Chlorophenyl-phenylether	8.06	204	39100	47.44	ng	97
59) Diethylphthalate	7.96	149	80625	46.06	ng	96
60) 4-Nitroaniline	8.08	138	22270	44.70	ng	98
62) 4,6-Dinitro-2-methylphenol	8.10	198	16287	48.51	ng	100
63) n-Nitrosodiphenylamine	8.17	169	58847	48.88	ng	98
65) 1,2-Diphenylhydrazine	8.20	77	88326	51.41	ng	99
66) 4-Bromophenyl-phenylether	8.52	248	22803	50.93	ng	94
67) Hexachlorobenzene	8.57	284	20882	49.40	ng	86
68) gamma-BHC	8.82	181	3072	9.93	ng	94
69) Pentachlorophenol	8.77	266	14920	53.84	ng	91

(#) = qualifier out of range (m) = manual integration

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 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 12:11 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.98	178	127474	50.61	ng	97
71) Anthracene	9.04	178	126461	49.40	ng	98
72) Carbazole	9.21	167	113607	48.61	ng	100
73) Heptachlor	9.49	100	3429	10.49	ng	91
74) Di-n-butylphthalate	9.62	149	138178	48.83	ng	99
75) Heptachlor epoxide	10.18	81	2196	9.67	ng	79
76) Fluoranthene	10.28	202	135723	49.43	ng	99
78) Pyrene	10.53	202	146312	51.52	ng	96
79) Benzidine	10.45	184	59401	56.65	ng	95
81) Endrin	10.98	81	1378	9.86	ng	82
82) Butylbenzylphthalate	11.35	149	60879	48.71	ng	95
83) Methoxychlor	11.99	227	13294	10.20	ng	96
84) 3,3'-Dichlorobenzidine	11.94	252	51173	62.69	ng	97
85) Benzo[a]anthracene	11.94	228	130462	50.04	ng	99
86) Chrysene	11.99	228	121552	50.83	ng	98
87) bis(2-Ethylhexyl)phthalate	12.07	149	86188	49.91	ng	98
89) Di-n-octylphthalate	12.81	149	148858	48.53	ng	97
90) Benzo[b]fluoranthene	13.15	252	107385	48.55	ng	97
91) Benzo[k]fluoranthene	13.18	252	114194	51.01	ng	97
92) Benzo[a]pyrene	13.49	252	100270	48.17	ng	96
93) Indeno[1,2,3-cd]pyrene	14.59	276	111652	49.51	ng	86
94) Dibenzo[a,h]anthracene	14.62	278	92257	49.30	ng	96
95) Benzo[g,h,i]perylene	14.88	276	90753	48.12	ng	94

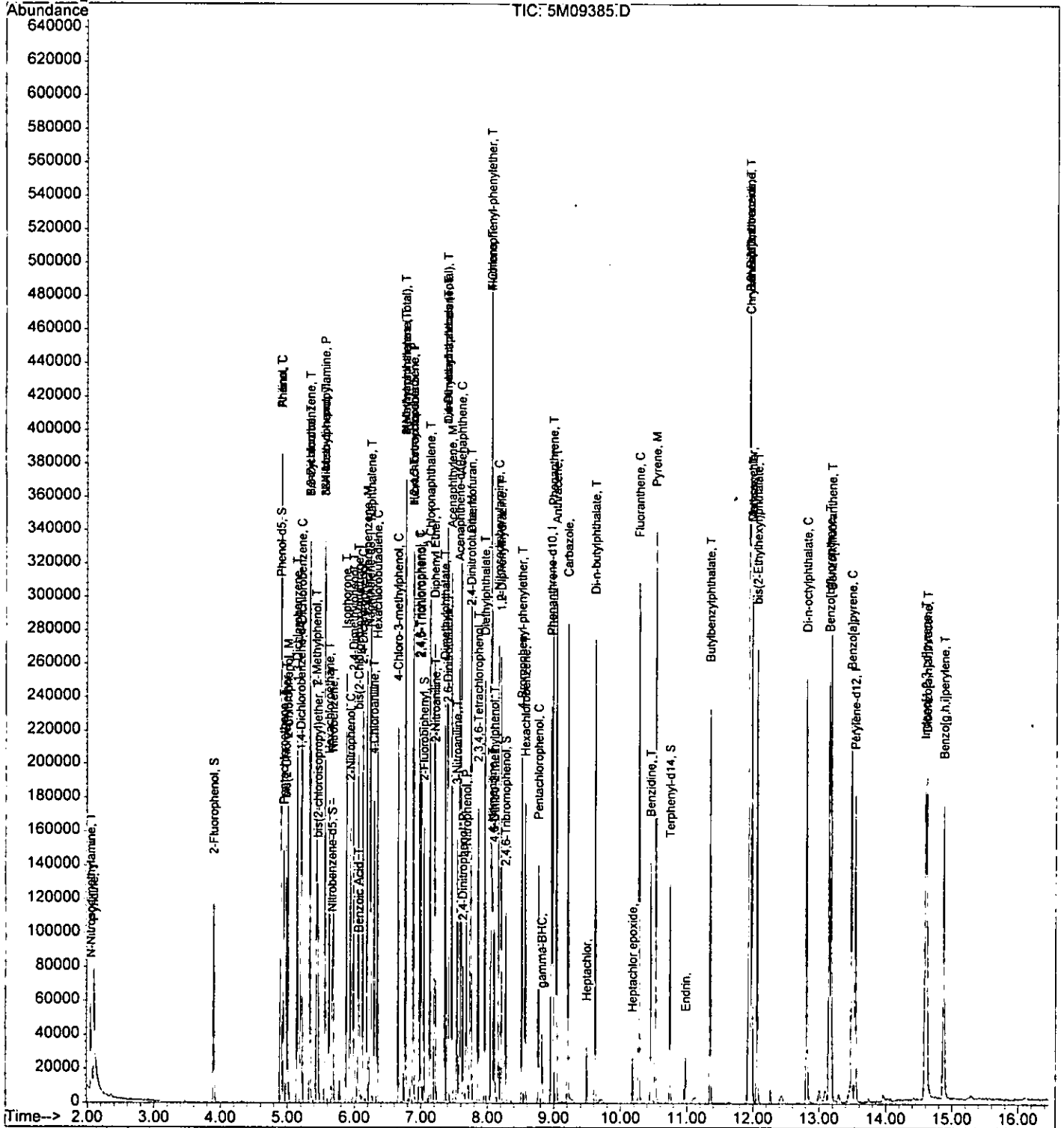
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09385.D Vial: 2
 Acq On : 22 Jul 2005 8:30 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 12:11 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09386.D Vial: 3
 Acq On : 22 Jul 2005 8:53 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:17 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	24298	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	90455	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	54766	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	92737	40.00	ng	-0.07
77) Chrysene-d12	11.95	240	79360	40.00	ng	-0.09
88) Perylene-d12	13.54	264	61273	40.00	ng	-0.08

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	6365	7.28	ng	-0.06
Spiked Amount	200.000		Recovery	=	3.64%	
8) Phenol-d5	4.89	99	9082	7.61	ng	-0.06
Spiked Amount	200.000		Recovery	=	3.81%	
21) Nitrobenzene-d5	5.67	128	1731	4.25	ng	-0.05
Spiked Amount	100.000		Recovery	=	4.25%	
41) 2-Fluorobiphenyl	7.04	172	8329	4.73	ng	-0.05
Spiked Amount	100.000		Recovery	=	4.73%	
64) 2,4,6-Tribromophenol	8.27	330	1818	9.52	ng	-0.07
Spiked Amount	200.000		Recovery	=	4.76%	
80) Terphenyl-d14	10.74	244	8221	4.33	ng	-0.07
Spiked Amount	100.000		Recovery	=	4.33%	

Target Compounds

						Qvalue
2) Pyridine	2.15	79	7384m	7.48	ng	
3) N-Nitrosodimethylamine	2.08	74	4401	7.50	ng	81
5) Aniline	4.91	93	11654	8.08	ng	85
6) Pentachloroethane	4.94	117	2849	9.20	ng	96
7) bis(2-Chloroethyl) ether	4.98	93	7502	8.67	ng	99
9) Phenol	4.90	94	10348	7.91	ng	64
10) 2-Chlorophenol	5.00	128	8079	8.45	ng	95
11) 1,3-Dichlorobenzene	5.14	146	8742	9.55	ng	97
12) 1,4-Dichlorobenzene	5.21	146	8588	9.23	ng	98
13) 1,2-Dichlorobenzene	5.33	146	8269	9.42	ng	93
14) Benzyl alcohol	5.33	108	4928	7.62	ng	96
15) bis(2-chloroisopropyl) ethe	5.45	45	11475	8.92	ng	95
16) 2-Methylphenol	5.43	108	7009	8.04	ng	99
17) Hexachloroethane	5.62	117	3663	9.51	ng	92
18) N-Nitroso-di-n-propylamine	5.55	70	5754	8.39	ng	96
19) 3&4-Methylphenol	5.56	108	7594	8.22	ng	96
22) Nitrobenzene	5.68	77	8244	9.24	ng	99
23) Isophorone	5.87	82	14886	9.14	ng	98
24) 2-Nitrophenol	5.94	139	4082	8.84	ng	95

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09386.D Vial: 3
 Acq On : 22 Jul 2005 8:53 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 11:17 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.98	107	7895	9.28	ng	98
26) Benzoic Acid	6.04	105	691	1.16	ng	86
27) bis(2-Chloroethoxy)methane	6.06	93	8993	9.45	ng	99
28) 2,4-Dichlorophenol	6.13	162	6012	8.31	ng	97
29) 1,2,4-Trichlorobenzene	6.18	180	7659	9.95	ng	96
30) Naphthalene	6.24	128	22830	9.56	ng	99
31) 4-Chloroaniline	6.29	127	8470	8.94	ng	99
32) Hexachlorobutadiene	6.34	225	4205	10.21	ng	97
33) 4-Chloro-3-methylphenol	6.65	107	6834	8.80	ng	93
34) 2-Methylnaphthalene	6.76	142	15124	9.54	ng	99
35) Methylnaphthalenes (Total)	6.76	142	15124	9.54	ng	99
37) 1,2,4,5-Tetrachlorobenzene	6.88	216	7329	9.92	ng	98
38) Hexachlorocyclopentadiene	6.88	237	3760	8.05	ng	99
39) 2,4,6-Trichlorophenol	6.97	196	4586	8.56	ng	97
40) 2,4,5-Trichlorophenol	7.00	196	5022	8.68	ng	96
42) 2-Chloronaphthalene	7.13	162	14966	9.53	ng	98
43) 1,4-Dimethylnaphthalene	7.39	156	11175	9.55	ng	96
44) Dimethylnaphthalenes (Tota	7.39	156	11175	9.55	ng	96
45) Diphenyl Ether	7.20	170	10182	9.72	ng	92
46) 2-Nitroaniline	7.21	65	5170	8.62	ng	91
47) Acenaphthylene	7.46	152	23125	9.32	ng	99
48) Dimethylphthalate	7.36	163	16138	9.12	ng	98
49) 2,6-Dinitrotoluene	7.41	165	3613	8.82	ng	82
50) Acenaphthene	7.61	153	14681	9.55	ng	97
51) 3-Nitroaniline	7.54	138	3756	7.94	ng	94
52) 2,4-Dinitrophenol	7.64	184	309	1.24	ng	20
53) Dibenzofuran	7.75	168	22268	9.94	ng	100
54) 2,4-Dinitrotoluene	7.74	165	4985	8.71	ng	93
55) 4-Nitrophenol	7.68	65	2502	6.94	ng	98
56) 2,3,4,6-Tetrachlorophenol	7.86	232	3803	8.40	ng	94
57) Fluorene	8.05	166	16907	9.77	ng	99
58) 4-Chlorophenyl-phenylether	8.05	204	8499	10.10	ng	97
59) Diethylphthalate	7.95	149	17789	9.87	ng	99
60) 4-Nitroaniline	8.07	138	4273	8.03	ng	92
62) 4,6-Dinitro-2-methylphenol	8.10	198	1965	5.57	ng	100
63) n-Nitrosodiphenylamine	8.16	169	12568	9.81	ng	98
65) 1,2-Diphenylhydrazine	8.20	77	17601	9.68	ng	97
66) 4-Bromophenyl-phenylether	8.52	248	4686	9.90	ng	97
67) Hexachlorobenzene	8.57	284	4579	10.38	ng	86
68) gamma-BHC	8.82	181	607	1.86	ng	88
69) Pentachlorophenol	8.76	266	1561	5.21	ng	97

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09386.D Vial: 3
 Acq On : 22 Jul 2005 8:53 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 11:17 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.98	178	25917	9.67	ng	99
71) Anthracene	9.03	178	26263	9.78	ng	99
72) Carbazole	9.21	167	24415	9.66	ng	99
73) Heptachlor	9.49	100	521	1.44	ng	88
74) Di-n-butylphthalate	9.61	149	28391	9.31	ng	100
75) Heptachlor epoxide	10.17	81	400	1.63	ng	96
76) Fluoranthene	10.27	202	26959	9.17	ng	100
78) Pyrene	10.53	202	29126	9.13	ng	95
79) Benzidine	10.45	184	13470	11.88	ng	95
81) Endrin	10.98	81	224	1.36	ng	# 77
82) Butylbenzylphthalate	11.35	149	11356	8.18	ng	96
83) Methoxychlor	11.98	227	2599	1.78	ng	# 87
84) 3,3'-Dichlorobenzidine	11.94	252	9203	10.69	ng	97
85) Benzo[a]anthracene	11.94	228	26478	8.86	ng	98
86) Chrysene	11.98	228	24720	9.33	ng	97
87) bis(2-Ethylhexyl)phthalate	12.06	149	16601	8.83	ng	94
89) Di-n-octylphthalate	12.81	149	28178	8.66	ng	99
90) Benzo[b]fluoranthene	13.14	252	21414	8.77	ng	97
91) Benzo[k]fluoranthene	13.17	252	23148	9.49	ng	96
92) Benzo[a]pyrene	13.48	252	20576	8.89	ng	97
93) Indeno[1,2,3-cd]pyrene	14.58	276	21695	8.74	ng	91
94) Dibenzo[a,h]anthracene	14.61	278	17918	8.74	ng	96
95) Benzo[g,h,i]perylene	14.87	276	19461	9.48	ng	91

(#) = qualifier out of range (m) = manual integration

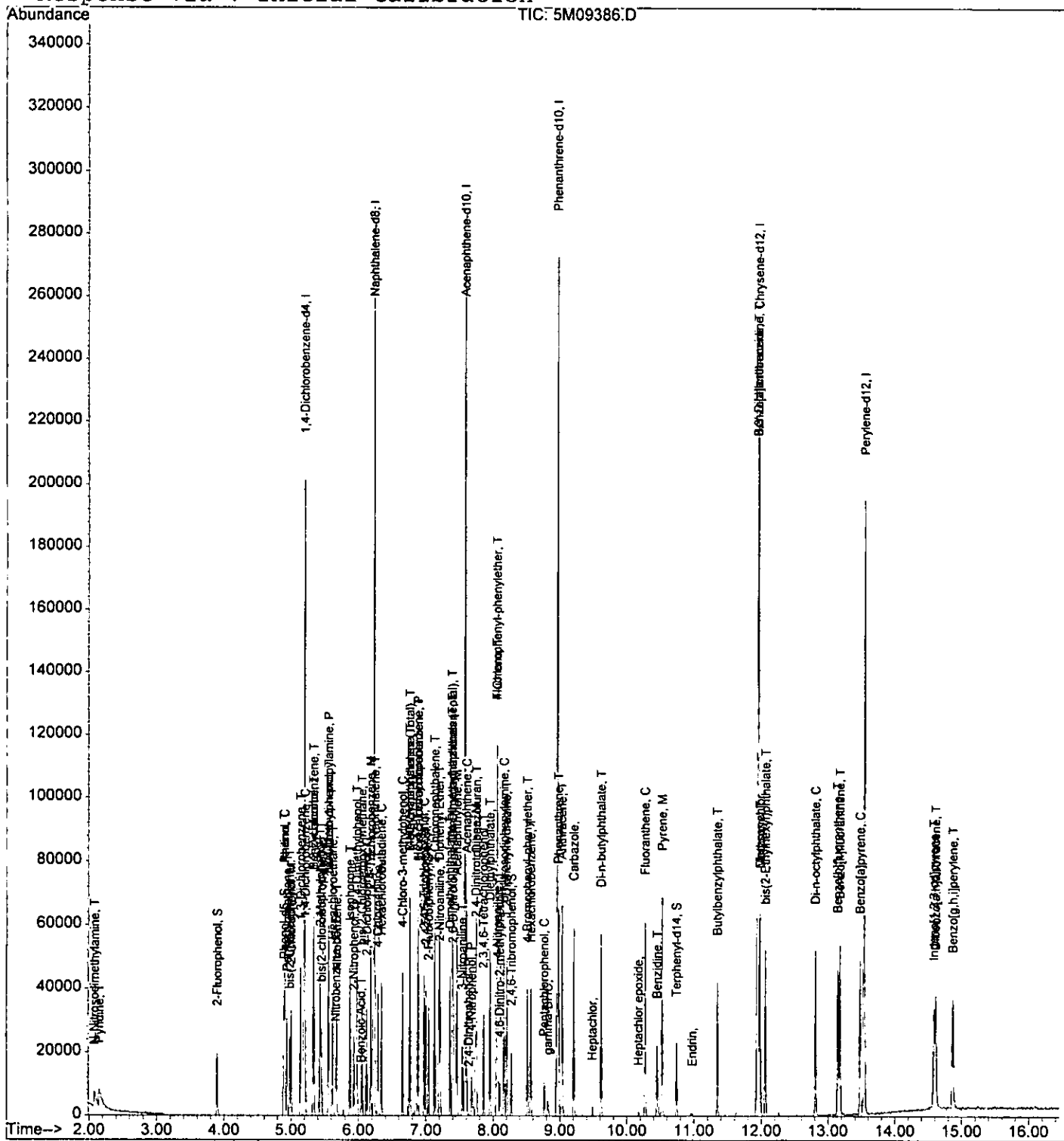
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09386.D Vial: 3
 Acq On : 22 Jul 2005 8:53 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:17 2005

9853

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09387.D Vial: 4
 Acq On : 22 Jul 2005 9:16 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 11:32 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	23586	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	91103	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	52605	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	90904	40.00	ng	-0.07
77) Chrysene-d12	11.95	240	70685	40.00	ng	-0.08
88) Perylene-d12	13.54	264	52162	40.00	ng	-0.07

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	18769	22.12	ng	-0.06
Spiked Amount	200.000		Recovery	=	11.06%	
8) Phenol-d5	4.89	99	24612	21.26	ng	-0.06
Spiked Amount	200.000		Recovery	=	10.63%	
21) Nitrobenzene-d5	5.67	128	5094	12.41	ng	-0.05
Spiked Amount	100.000		Recovery	=	12.41%	
41) 2-Fluorobiphenyl	7.04	172	22043	13.02	ng	-0.05
Spiked Amount	100.000		Recovery	=	13.02%	
64) 2,4,6-Tribromophenol	8.27	330	5138	27.45	ng	-0.07
Spiked Amount	200.000		Recovery	=	13.73%	
80) Terphenyl-d14	10.74	244	22244	13.15	ng	-0.07
Spiked Amount	100.000		Recovery	=	13.15%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.11	79	24806m	25.88	ng	
3) N-Nitrosodimethylamine	2.05	74	14519	25.48	ng	97
5) Aniline	4.91	93	30758	21.97	ng	88
6) Pentachloroethane	4.94	117	7352	24.45	ng	98
7) bis(2-Chloroethyl)ether	4.98	93	20890	24.87	ng	97
9) Phenol	4.90	94	27521	21.67	ng	63
10) 2-Chlorophenol	5.01	128	22435	24.16	ng	99
11) 1,3-Dichlorobenzene	5.14	146	23269	26.19	ng	97
12) 1,4-Dichlorobenzene	5.21	146	22855	25.30	ng	100
13) 1,2-Dichlorobenzene	5.34	146	22215	26.08	ng	99
14) Benzyl alcohol	5.33	108	14292	22.76	ng	92
15) bis(2-chloroisopropyl)ethe	5.45	45	32278	25.85	ng	93
16) 2-Methylphenol	5.43	108	20071	23.72	ng	97
17) Hexachloroethane	5.62	117	9882	26.43	ng	94
18) N-Nitroso-di-n-propylamine	5.55	70	16223	24.38	ng	98
19) 3&4-Methylphenol	5.56	108	21219	23.68	ng	99
22) Nitrobenzene	5.68	77	22854	25.44	ng	97
23) Isophorone	5.87	82	41311	25.18	ng	94
24) 2-Nitrophenol	5.94	139	11259	24.22	ng	95

(#) = qualifier out of range (m) = manual integration

hgr

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09387.D Vial: 4
 Acq On : 22 Jul 2005 9:16 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 11:32 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.98	107	21356	24.93	ng	98
26) Benzoic Acid	6.06	105	6131m	10.18	ng	
27) bis(2-Chloroethoxy)methane	6.06	93	24100	25.15	ng	100
28) 2,4-Dichlorophenol	6.13	162	18093	24.83	ng	97
29) 1,2,4-Trichlorobenzene	6.18	180	19587	25.27	ng	98
30) Naphthalene	6.24	128	61814	25.70	ng	99
31) 4-Chloroaniline	6.29	127	24620	25.79	ng	100
32) Hexachlorobutadiene	6.34	225	11336	27.33	ng	96
33) 4-Chloro-3-methylphenol	6.65	107	18911	24.17	ng	98
34) 2-Methylnaphthalene	6.76	142	42150	26.40	ng	99
35) Methylnaphthalenes (Total)	6.76	142	42150	26.40	ng	99
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	19995	28.18	ng	97
38) Hexachlorocyclopentadiene	6.88	237	11680	26.04	ng	99
39) 2,4,6-Trichlorophenol	6.97	196	13612	26.44	ng	96
40) 2,4,5-Trichlorophenol	7.00	196	13983	25.15	ng	97
42) 2-Chloronaphthalene	7.13	162	40352	26.75	ng	98
43) 1,4-Dimethylnaphthalene	7.39	156	30737	27.34	ng	99
44) Dimethylnaphthalenes (Tota	7.39	156	30737	27.34	ng	99
45) Diphenyl Ether	7.20	170	26762	26.61	ng	95
46) 2-Nitroaniline	7.21	65	14242	24.73	ng	94
47) Acenaphthylene	7.46	152	63677	26.71	ng	99
48) Dimethylphthalate	7.36	163	44085	25.93	ng	99
49) 2,6-Dinitrotoluene	7.41	165	10202	25.92	ng	93
50) Acenaphthene	7.61	153	37826	25.61	ng	99
51) 3-Nitroaniline	7.54	138	11428	25.15	ng	98
52) 2,4-Dinitrophenol	7.63	184	3431	14.29	ng	87
53) Dibenzofuran	7.75	168	55594	25.83	ng	95
54) 2,4-Dinitrotoluene	7.74	165	13404	24.37	ng	95
55) 4-Nitrophenol	7.68	65	7437	21.49	ng	97
56) 2,3,4,6-Tetrachlorophenol	7.86	232	10484	24.10	ng	100
57) Fluorene	8.05	166	44090	26.52	ng	100
58) 4-Chlorophenyl-phenylether	8.06	204	21506	26.60	ng	99
59) Diethylphthalate	7.96	149	45551	26.31	ng	96
60) 4-Nitroaniline	8.07	138	12201	23.86	ng	99
62) 4,6-Dinitro-2-methylphenol	8.10	198	6955	20.12	ng	100
63) n-Nitrosodiphenylamine	8.17	169	35141	27.97	ng	97
65) 1,2-Diphenylhydrazine	8.20	77	48294	27.10	ng	99
66) 4-Bromophenyl-phenylether	8.52	248	12025	25.91	ng	99
67) Hexachlorobenzene	8.57	284	11811	27.30	ng	89
68) gamma-BHC	8.82	181	1620	5.06	ng	92
69) Pentachlorophenol	8.76	266	5934	20.20	ng	93

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09387.D Vial: 4
 Acq On : 22 Jul 2005 9:16 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:32 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.98	178	68461	26.05	ng	98
71) Anthracene	9.04	178	70848	26.91	ng	99
72) Carbazole	9.21	167	61734	24.93	ng	99
73) Heptachlor	9.49	100	1653	4.68	ng	93
74) Di-n-butylphthalate	9.61	149	75717	25.33	ng	99
75) Heptachlor epoxide	10.17	81	1273	5.28	ng	97
76) Fluoranthene	10.28	202	72021	25.00	ng	96
78) Pyrene	10.53	202	73780	25.97	ng	97
79) Benzidine	10.45	184	28789	28.51	ng	98
81) Endrin	10.98	81	750	5.13	ng	91
82) Butylbenzylphthalate	11.35	149	32229	26.06	ng	99
83) Methoxychlor	11.98	227	6644	5.10	ng	100
84) 3,3'-Dichlorobenzidine	11.94	252	20559	26.81	ng	98
85) Benzo[a]anthracene	11.94	228	65311	24.55	ng	98
86) Chrysene	11.98	228	61876	26.22	ng	99
87) bis(2-Ethylhexyl)phthalate	12.06	149	43747	26.12	ng	97
89) Di-n-octylphthalate	12.81	149	73317	26.47	ng	98
90) Benzo[b]fluoranthene	13.14	252	51352	24.72	ng	98
91) Benzo[k]fluoranthene	13.17	252	54688	26.32	ng	96
92) Benzo[a]pyrene	13.48	252	50469	25.61	ng	98
93) Indeno[1,2,3-cd]pyrene	14.59	276	57483	27.21	ng	88
94) Dibenzo[a,h]anthracene	14.61	278	46954	26.90	ng	96
95) Benzo[g,h,i]perylene	14.87	276	49482	28.31	ng	91

(#) = qualifier out of range (m) = manual integration

Quantitation Report

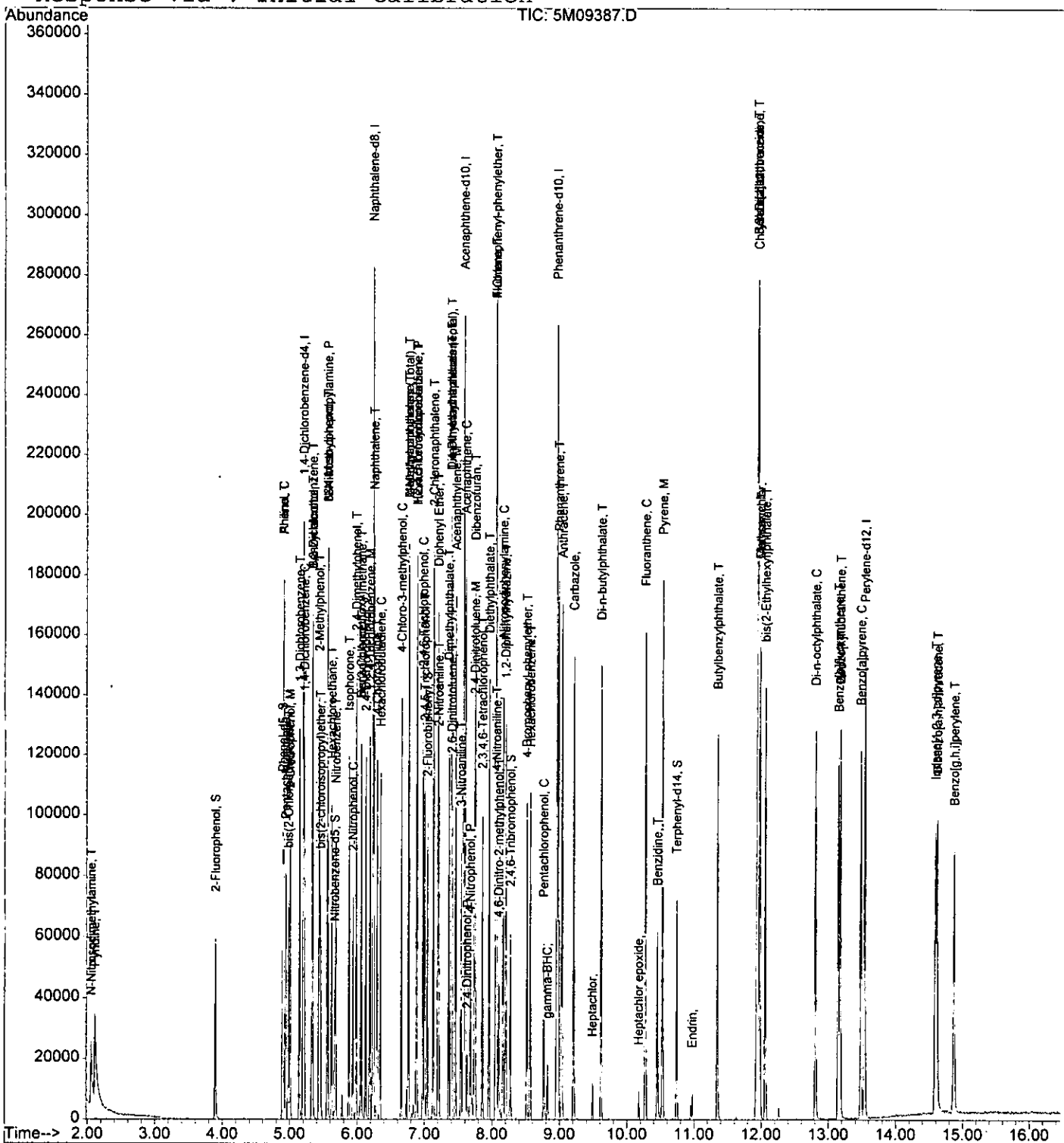
Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09387.D Vial: 4
Acq On : 22 Jul 2005 9:16
Sample : CAL BNA@25PPM
Misc : A,BNA
MS Integration Params: RTEINT.P
Quant Time: Jul 22 11:32 2005

Operator: AHD
Inst : GCMS_5
Multiplr: 1.00

3159

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09388.D Vial: 5
 Acq On : 22 Jul 2005 9:39 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

011

MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:37 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	20683	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	84396	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	49933	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	89583	40.00	ng	-0.06
77) Chrysene-d12	11.95	240	74392	40.00	ng	-0.08
88) Perylene-d12	13.54	264	54467	40.00	ng	-0.07

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	54913	73.80	ng	-0.06
Spiked Amount	200.000		Recovery	=	36.90%	
8) Phenol-d5	4.90	99	76343	75.19	ng	-0.05
Spiked Amount	200.000		Recovery	=	37.60%	
21) Nitrobenzene-d5	5.67	128	14623	38.46	ng	-0.05
Spiked Amount	100.000		Recovery	=	38.46%	
41) 2-Fluorobiphenyl	7.04	172	62049	38.61	ng	-0.05
Spiked Amount	100.000		Recovery	=	38.61%	
64) 2,4,6-Tribromophenol	8.28	330	15204	82.44	ng	-0.06
Spiked Amount	200.000		Recovery	=	41.22%	
80) Terphenyl-d14	10.74	244	68757	38.62	ng	-0.07
Spiked Amount	100.000		Recovery	=	38.62%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.09	79	72584	86.35	ng	94
3) N-Nitrosodimethylamine	2.04	74	45277	90.61	ng	93
5) Aniline	4.91	93	89277	72.72	ng	89
6) Pentachloroethane	4.94	117	20376	77.28	ng	99
7) bis(2-Chloroethyl)ether	4.98	93	56158	76.25	ng	96
9) Phenol	4.91	94	82266	73.86	ng	61
10) 2-Chlorophenol	5.01	128	61795	75.89	ng	99
11) 1,3-Dichlorobenzene	5.14	146	58858	75.53	ng	99
12) 1,4-Dichlorobenzene	5.21	146	62508	78.92	ng	99
13) 1,2-Dichlorobenzene	5.34	146	59163	79.20	ng	99
14) Benzyl alcohol	5.33	108	43261	78.57	ng	99
15) bis(2-chloroisopropyl)ethe	5.45	45	87970	80.33	ng	99
16) 2-Methylphenol	5.43	108	57711	77.77	ng	99
17) Hexachloroethane	5.62	117	25268	77.06	ng	98
18) N-Nitroso-di-n-propylamine	5.56	70	45857	78.58	ng	96
19) 3&4-Methylphenol	5.56	108	61226	77.90	ng	99
22) Nitrobenzene	5.68	77	67786	81.44	ng	99
23) Isophorone	5.88	82	119898	78.90	ng	97
24) 2-Nitrophenol	5.94	139	33397	77.55	ng	98

(#) = qualifier out of range (m) = manual integration

1-8/11

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09388.D Vial: 5
 Acq On : 22 Jul 2005 9:39 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

8912

MS Integration Params: RTEINT.P

Quant Time: Jul 22 10:37 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.99	107	64432	81.19	ng	98
26) Benzoic Acid	6.09	105	34922	62.59	ng	98
27) bis(2-Chloroethoxy)methane	6.06	93	70445	79.35	ng	99
28) 2,4-Dichlorophenol	6.13	162	54528	80.78	ng	97
29) 1,2,4-Trichlorobenzene	6.19	180	52732	73.44	ng	99
30) Naphthalene	6.24	128	173789	78.01	ng	99
31) 4-Chloroaniline	6.29	127	69715	78.82	ng	99
32) Hexachlorobutadiene	6.34	225	30412	79.15	ng	97
33) 4-Chloro-3-methylphenol	6.65	107	57270	79.02	ng	98
34) 2-Methylnaphthalene	6.77	142	119806	81.01	ng	100
35) Methylnaphthalenes (Total)	6.77	142	119806	81.01	ng	100
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	55232	82.02	ng	97
38) Hexachlorocyclopentadiene	6.88	237	34973	82.14	ng	99
39) 2,4,6-Trichlorophenol	6.97	196	39490	80.81	ng	98
40) 2,4,5-Trichlorophenol	7.00	196	42036	79.65	ng	96
42) 2-Chloronaphthalene	7.13	162	112840	78.80	ng	99
43) 1,4-Dimethylnaphthalene	7.40	156	83298	78.05	ng	96
44) Dimethylnaphthalenes (Tota	7.40	156	83298	78.05	ng	96
45) Diphenyl Ether	7.20	170	72809	76.27	ng	97
46) 2-Nitroaniline	7.22	65	42303	77.38	ng	92
47) Acenaphthylene	7.46	152	177834	78.58	ng	100
48) Dimethylphthalate	7.36	163	131779	81.65	ng	100
49) 2,6-Dinitrotoluene	7.41	165	29857	79.93	ng	99
50) Acenaphthene	7.61	153	109538	78.14	ng	98
51) 3-Nitroaniline	7.55	138	33964	78.75	ng	98
52) 2,4-Dinitrophenol	7.63	184	16704	73.32	ng	85
53) Dibenzofuran	7.75	168	160496	78.55	ng	99
54) 2,4-Dinitrotoluene	7.74	165	42299	81.03	ng	94
55) 4-Nitrophenol	7.69	65	26342	80.17	ng	99
56) 2,3,4,6-Tetrachlorophenol	7.86	232	33039	80.00	ng	99
57) Fluorene	8.05	166	130101	82.44	ng	98
58) 4-Chlorophenyl-phenylether	8.06	204	62673	81.66	ng	100
59) Diethylphthalate	7.96	149	130895	79.65	ng	97
60) 4-Nitroaniline	8.08	138	39264	80.88	ng	100
62) 4,6-Dinitro-2-methylphenol	8.10	198	26230	77.01	ng	100
63) n-Nitrosodiphenylamine	8.17	169	93386	75.43	ng	99
65) 1,2-Diphenylhydrazine	8.20	77	135131	76.95	ng	98
66) 4-Bromophenyl-phenylether	8.52	248	35519	77.67	ng	96
67) Hexachlorobenzene	8.57	284	33252	78.00	ng	83
68) gamma-BHC	8.82	181	5073	16.09	ng	91
69) Pentachlorophenol	8.76	266	24029	83.00	ng	94

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09388.D Vial: 5
 Acq On : 22 Jul 2005 9:39 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 10:37 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.98	178	198861	76.79	ng	98
71) Anthracene	9.04	178	205366	79.14	ng	98
72) Carbazole	9.21	167	192480	78.87	ng	99
73) Heptachlor	9.49	100	5254	15.08	ng	92
74) Di-n-butylphthalate	9.62	149	229034	77.76	ng	99
75) Heptachlor epoxide	10.17	81	3663	15.43	ng	87
76) Fluoranthene	10.28	202	227869	80.26	ng	100
78) Pyrene	10.53	202	228348	76.37	ng	96
79) Benzidine	10.45	184	79895	75.18	ng	95
81) Endrin	10.98	81	2423	15.74	ng	90
82) Butylbenzylphthalate	11.35	149	106257	81.65	ng	97
83) Methoxychlor	11.99	227	22243	16.21	ng	98
84) 3,3'-Dichlorobenzidine	11.94	252	66274	82.12	ng	98
85) Benzo[a]anthracene	11.94	228	218249	77.95	ng	99
86) Chrysene	11.99	228	202363	81.49	ng	100
87) bis(2-Ethylhexyl)phthalate	12.06	149	145925	82.80	ng	98
89) Di-n-octylphthalate	12.81	149	252604	87.34	ng	99
90) Benzo[b]fluoranthene	13.15	252	174306	80.35	ng	97
91) Benzo[k]fluoranthene	13.18	252	180808	83.35	ng	94
92) Benzo[a]pyrene	13.49	252	163205	79.30	ng	97
93) Indeno[1,2,3-cd]pyrene	14.59	276	176109	79.84	ng	91
94) Dibenzo[a,h]anthracene	14.62	278	150578	82.62	ng	96
95) Benzo[g,h,i]perylene	14.88	276	150430	82.43	ng	90

 (#) = qualifier out of range (m) = manual integration

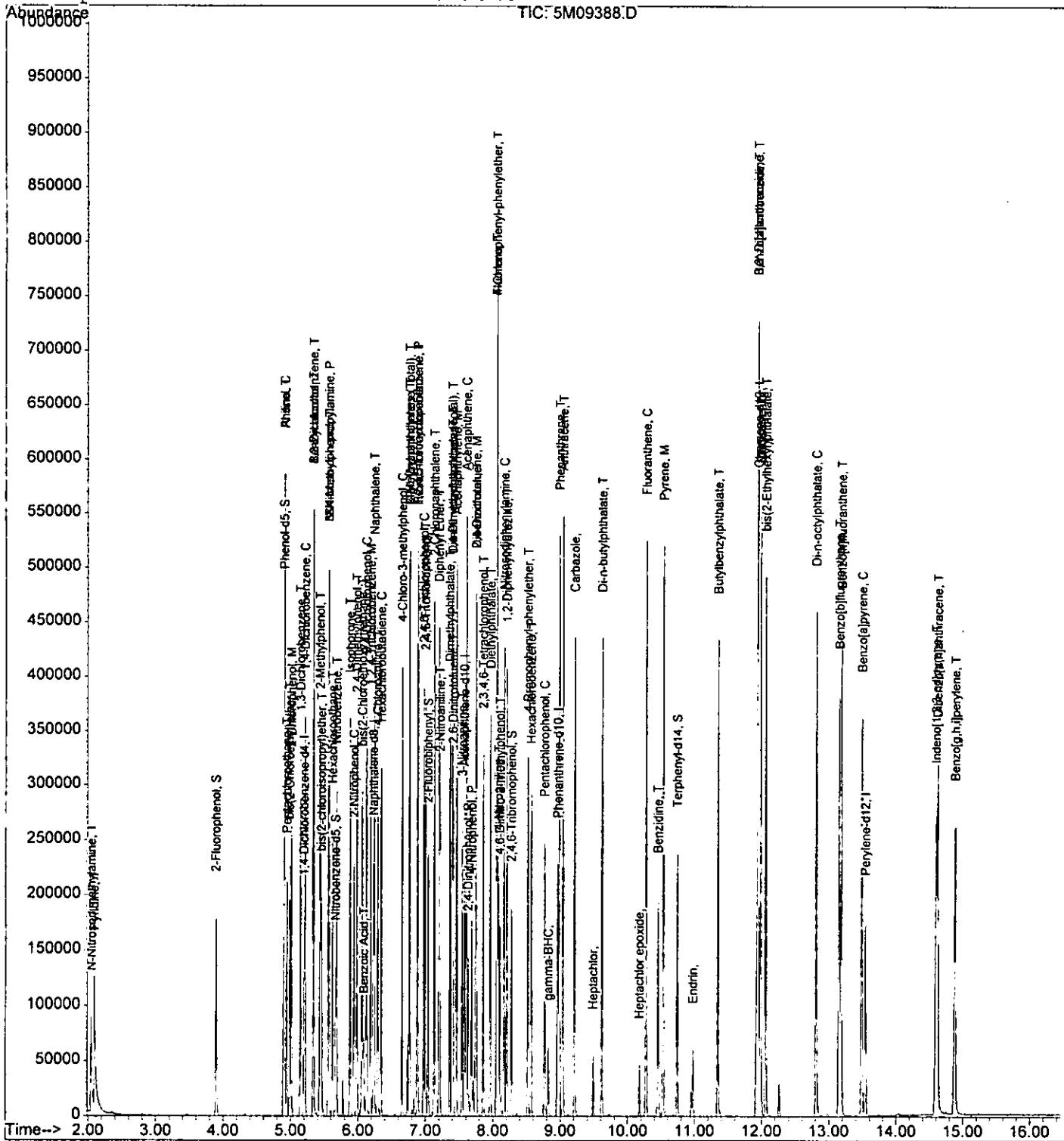
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09388.D Vial: 5
 Acq On : 22 Jul 2005 9:39 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:37 2005

7158

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09389.D Vial: 6
 Acq On : 22 Jul 2005 10:01 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:38 2005

595

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	16575	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	72739	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	48151	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	85449	40.00	ng	-0.06
77) Chrysene-d12	11.96	240	70631	40.00	ng	-0.07
88) Perylene-d12	13.54	264	54990	40.00	ng	-0.07

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	74618	125.13	ng	-0.06
Spiked Amount	200.000		Recovery	=	62.56%	
8) Phenol-d5	4.90	99	107659	132.32	ng	-0.05
Spiked Amount	200.000		Recovery	=	66.16%	
21) Nitrobenzene-d5	5.67	128	20199	61.63	ng	-0.05
Spiked Amount	100.000		Recovery	=	61.63%	
41) 2-Fluorobiphenyl	7.04	172	88572	57.15	ng	-0.05
Spiked Amount	100.000		Recovery	=	57.15%	
64) 2,4,6-Tribromophenol	8.28	330	22612	128.54	ng	-0.06
Spiked Amount	200.000		Recovery	=	64.27%	
80) Terphenyl-d14	10.74	244	108802	64.37	ng	-0.07
Spiked Amount	100.000		Recovery	=	64.37%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.08	79	94837	140.78	ng	93
3) N-Nitrosodimethylamine	2.04	74	55701	139.10	ng	91
5) Aniline	4.91	93	123731	125.76	ng	91
6) Pentachloroethane	4.94	117	25749	121.86	ng	99
7) bis(2-Chloroethyl)ether	4.98	93	77073	130.58	ng	94
9) Phenol	4.91	94	115870	129.82	ng	78
10) 2-Chlorophenol	5.01	128	85313	130.73	ng	96
11) 1,3-Dichlorobenzene	5.14	146	74216	118.85	ng	100
12) 1,4-Dichlorobenzene	5.21	146	77375	121.90	ng	99
13) 1,2-Dichlorobenzene	5.34	146	72899	121.77	ng	99
14) Benzyl alcohol	5.33	108	56832	128.80	ng	96
15) bis(2-chloroisopropyl)ethe	5.45	45	115956	132.13	ng	94
16) 2-Methylphenol	5.44	108	80165	134.80	ng	100
17) Hexachloroethane	5.62	117	32289	122.87	ng	99
18) N-Nitroso-di-n-propylamine	5.56	70	62124	132.85	ng	98
19) 3&4-Methylphenol	5.56	108	83375	132.37	ng	100
22) Nitrobenzene	5.69	77	88442	123.29	ng	96
23) Isophorone	5.88	82	173298	132.32	ng	99
24) 2-Nitrophenol	5.94	139	47616	128.28	ng	98

(#) = qualifier out of range (m) = manual integration

hgr

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09389.D Vial: 6
 Acq On : 22 Jul 2005 10:01 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 10:38 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.99	107	89278	130.52	ng	96
26) Benzoic Acid	6.10	105	58454	121.56	ng	96
27) bis(2-Chloroethoxy)methane	6.06	93	97066	126.86	ng	99
28) 2,4-Dichlorophenol	6.13	162	74883	128.71	ng	97
29) 1,2,4-Trichlorobenzene	6.19	180	72441	117.05	ng	99
30) Naphthalene	6.25	128	244630	127.40	ng	99
31) 4-Chloroaniline	6.30	127	94871	124.46	ng	100
32) Hexachlorobutadiene	6.34	225	39130	118.15	ng	99
33) 4-Chloro-3-methylphenol	6.65	107	86310	138.18	ng	99
34) 2-Methylnaphthalene	6.77	142	168257	132.00	ng	100
35) Methylnaphthalenes (Total)	6.77	142	168257	132.00	ng	100
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	74314	114.44	ng	98
38) Hexachlorocyclopentadiene	6.88	237	48239	117.49	ng	98
39) 2,4,6-Trichlorophenol	6.97	196	57261	121.51	ng	99
40) 2,4,5-Trichlorophenol	7.00	196	63072	123.94	ng	96
42) 2-Chloronaphthalene	7.13	162	168561	122.06	ng	97
43) 1,4-Dimethylnaphthalene	7.40	156	124661	121.13	ng	98
44) Dimethylnaphthalenes (Total)	7.40	156	124661	121.13	ng	98
45) Diphenyl Ether	7.20	170	106493	115.68	ng	90
46) 2-Nitroaniline	7.22	65	67024	127.13	ng	99
47) Acenaphthylene	7.47	152	258852	118.61	ng	99
48) Dimethylphthalate	7.36	163	198590	127.61	ng	99
49) 2,6-Dinitrotoluene	7.41	165	45745	126.99	ng	99
50) Acenaphthene	7.61	153	164425	121.64	ng	99
51) 3-Nitroaniline	7.55	138	50421	121.24	ng	98
52) 2,4-Dinitrophenol	7.64	184	31054	141.35	ng	76
53) Dibenzofuran	7.75	168	234562	119.05	ng	97
54) 2,4-Dinitrotoluene	7.75	165	62635	124.42	ng	96
55) 4-Nitrophenol	7.69	65	42380	133.76	ng	96
56) 2,3,4,6-Tetrachlorophenol	7.86	232	52042	130.68	ng	99
57) Fluorene	8.06	166	189503	124.52	ng	98
58) 4-Chlorophenyl-phenylether	8.06	204	90025	121.64	ng	97
59) Diethylphthalate	7.96	149	200828	126.73	ng	97
60) 4-Nitroaniline	8.09	138	59018	126.08	ng	99
62) 4,6-Dinitro-2-methylphenol	8.11	198	43850	134.96	ng	100
63) n-Nitrosodiphenylamine	8.17	169	143874	121.83	ng	97
65) 1,2-Diphenylhydrazine	8.21	77	209653	125.16	ng	94
66) 4-Bromophenyl-phenylether	8.52	248	51167	117.30	ng	91
67) Hexachlorobenzene	8.57	284	48654	119.66	ng	91
68) gamma-BHC	8.82	181	7543	25.09	ng	92
69) Pentachlorophenol	8.77	266	37918	137.31	ng	94

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09389.D Vial: 6
 Acq On : 22 Jul 2005 10:01 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:38 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Mon Jul 11 11:25:34 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.99	178	313082	126.75	ng	99
71) Anthracene	9.04	178	303812	122.74	ng	99
72) Carbazole	9.21	167	280497	120.50	ng	100
73) Heptachlor	9.49	100	8577	25.81	ng	97
74) Di-n-butylphthalate	9.62	149	344791	122.72	ng	100
75) Heptachlor epoxide	10.17	81	5760	25.43	ng	86
76) Fluoranthene	10.28	202	341094	125.96	ng	100
78) Pyrene	10.54	202	350575	123.49	ng	95
79) Benzidine	10.45	184	105478	104.54	ng	97
81) Endrin	10.98	81	3593	24.58	ng	88
82) Butylbenzylphthalate	11.36	149	162203	131.28	ng	99
83) Methoxychlor	11.99	227	32498	24.95	ng	98
84) 3,3'-Dichlorobenzidine	11.94	252	94182	122.92	ng	97
85) Benzo[a]anthracene	11.95	228	326401	122.78	ng	98
86) Chrysene	11.99	228	295658	125.40	ng	99
87) bis(2-Ethylhexyl)phthalate	12.07	149	219995	131.47	ng	97
89) Di-n-octylphthalate	12.81	149	383159	131.22	ng	99
90) Benzo[b]fluoranthene	13.16	252	265788	121.35	ng	98
91) Benzo[k]fluoranthene	13.19	252	270819	123.66	ng	95
92) Benzo[a]pyrene	13.49	252	252005	121.28	ng	98
93) Indeno[1,2,3-cd]pyrene	14.60	276	275620	123.77	ng	88
94) Dibenzo[a,h]anthracene	14.63	278	229689	124.82	ng	93
95) Benzo[g,h,i]perylene	14.89	276	224233	121.71	ng	91

(#) = qualifier out of range (m) = manual integration

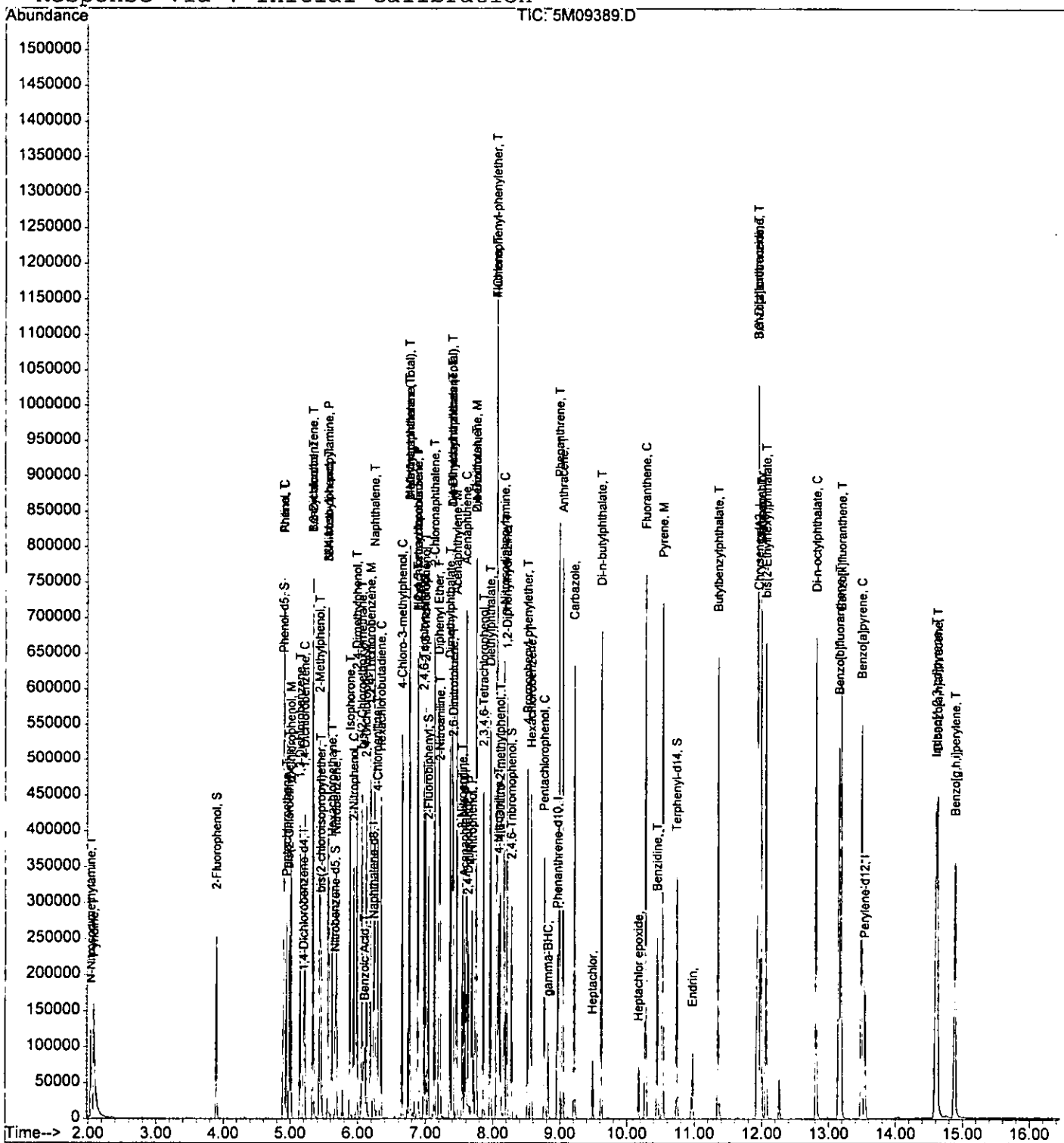
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09389.D Vial: 6
 Acq On : 22 Jul 2005 10:01 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:38 2005

8515

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09390.D Vial: 7
 Acq On : 22 Jul 2005 10:24 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:52 2005

6768

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	16184	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	72103	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	46483	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	89704	40.00	ng	-0.06
77) Chrysene-d12	11.96	240	72030	40.00	ng	-0.07
88) Perylene-d12	13.55	264	56669	40.00	ng	-0.07

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	100066	195.10	ng	-0.06
Spiked Amount	200.000		Recovery	=	97.55%	
8) Phenol-d5	4.90	99	147180	206.75	ng	-0.05
Spiked Amount	200.000		Recovery	=	103.38%	
21) Nitrobenzene-d5	5.67	128	27448	88.10	ng	-0.05
Spiked Amount	100.000		Recovery	=	88.10%	
41) 2-Fluorobiphenyl	7.04	172	120250	81.88	ng	-0.05
Spiked Amount	100.000		Recovery	=	81.88%	
64) 2,4,6-Tribromophenol	8.28	330	31168	161.34	ng	-0.06
Spiked Amount	200.000		Recovery	=	80.67%	
80) Terphenyl-d14	10.75	244	134935	79.09	ng	-0.06
Spiked Amount	100.000		Recovery	=	79.09%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.08	79	118747	178.92	ng	95
3) N-Nitrosodimethylamine	2.04	74	75020	190.38	ng	92
5) Aniline	4.91	93	174254	202.54	ng	90
6) Pentachloroethane	4.94	117	33515	167.70	ng	99
7) bis(2-Chloroethyl)ether	4.99	93	102080	183.59	ng	98
9) Phenol	4.91	94	161480	205.62	ng	67
10) 2-Chlorophenol	5.01	128	117648	194.24	ng	99
11) 1,3-Dichlorobenzene	5.14	146	100071	167.62	ng	99
12) 1,4-Dichlorobenzene	5.21	146	101923	167.89	ng	100
13) 1,2-Dichlorobenzene	5.34	146	94857	163.10	ng	98
14) Benzyl alcohol	5.33	108	78566	198.48	ng	100
15) bis(2-chloroisopropyl)ethe	5.45	45	152877	179.63	ng	94
16) 2-Methylphenol	5.44	108	108981	197.98	ng	99
17) Hexachloroethane	5.62	117	40854	160.42	ng	99
18) N-Nitroso-di-n-propylamine	5.56	70	84904	194.63	ng	96
19) 3&4-Methylphenol	5.57	108	115044	197.08	ng	99
22) Nitrobenzene	5.69	77	122920	174.78	ng	98
23) Isophorone	5.88	82	230394	178.32	ng	96
24) 2-Nitrophenol	5.94	139	62278	172.67	ng	93

(#) = qualifier out of range (m) = manual integration

12.811

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09390.D Vial: 7
 Acq On : 22 Jul 2005 10:24 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 10:52 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.99	107	120175	176.17	ng	96
26) Benzoic Acid	6.12	105	80341	281.51	ng	97
27) bis(2-Chloroethoxy)methane	6.06	93	124082	164.36	ng	99
28) 2,4-Dichlorophenol	6.13	162	98356	174.23	ng	96
29) 1,2,4-Trichlorobenzene	6.19	180	95516	159.88	ng	98
30) Naphthalene	6.25	128	308044	161.97	ng	100
31) 4-Chloroaniline	6.30	127	124308	166.12	ng	98
32) Hexachlorobutadiene	6.34	225	50574	150.28	ng	99
33) 4-Chloro-3-methylphenol	6.66	107	114066	187.21	ng	94
34) 2-Methylnaphthalene	6.77	142	212720	164.71	ng	98
35) Methylnaphthalenes (Total)	6.77	142	212720	164.71	ng	98
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	99911	154.77	ng	97
38) Hexachlorocyclopentadiene	6.88	237	65910	169.29	ng	99
39) 2,4,6-Trichlorophenol	6.98	196	76436	170.59	ng	99
40) 2,4,5-Trichlorophenol	7.00	196	84222	174.77	ng	95
42) 2-Chloronaphthalene	7.13	162	210650	157.99	ng	99
43) 1,4-Dimethylnaphthalene	7.40	156	164033	163.48	ng	100
44) Dimethylnaphthalenes (Total)	7.40	156	164033	163.48	ng	100
45) Diphenyl Ether	7.20	170	145491	166.66	ng	92
46) 2-Nitroaniline	7.22	65	90374	185.19	ng	91
47) Acenaphthylene	7.47	152	339010	162.50	ng	99
48) Dimethylphthalate	7.37	163	258345	171.87	ng	99
49) 2,6-Dinitrotoluene	7.42	165	59667	172.92	ng	90
50) Acenaphthene	7.61	153	217217	169.31	ng	99
51) 3-Nitroaniline	7.55	138	64265	168.64	ng	94
52) 2,4-Dinitrophenol	7.64	184	44224	283.50	ng	93
53) Dibenzofuran	7.76	168	306754	163.07	ng	100
54) 2,4-Dinitrotoluene	7.75	165	84974	180.07	ng	93
55) 4-Nitrophenol	7.69	65	59160	209.70	ng	96
56) 2,3,4,6-Tetrachlorophenol	7.86	232	68202	182.44	ng	99
57) Fluorene	8.06	166	256658	172.22	ng	99
58) 4-Chlorophenyl-phenylether	8.06	204	123155	169.66	ng	97
59) Diethylphthalate	7.97	149	267442	173.88	ng	97
60) 4-Nitroaniline	8.10	138	81036	191.02	ng	98
62) 4,6-Dinitro-2-methylphenol	8.12	198	60326	199.61	ng	100
63) n-Nitrosodiphenylamine	8.18	169	198877	159.37	ng	98
65) 1,2-Diphenylhydrazine	8.21	77	285747	159.65	ng	97
66) 4-Bromophenyl-phenylether	8.52	248	75241	164.38	ng	97
67) Hexachlorobenzene	8.57	284	70880	162.57	ng	94
68) gamma-BHC	8.82	181	10795	34.24	ng	94
69) Pentachlorophenol	8.77	266	54471	205.67	ng	94

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09390.D Vial: 7
 Acq On : 22 Jul 2005 10:24 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 10:52 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.99	178	424063	162.40	ng	98
71) Anthracene	9.04	178	429889	163.29	ng	98
72) Carbazole	9.22	167	398300	166.02	ng	99
73) Heptachlor	9.49	100	11683	35.74	ng	94
74) Di-n-butylphthalate	9.62	149	479294	165.77	ng	100
75) Heptachlor epoxide	10.17	81	7659	33.26	ng	88
76) Fluoranthene	10.28	202	465432	165.34	ng	99
78) Pyrene	10.54	202	484679	168.68	ng	99
79) Benzidine	10.46	184	148018	135.39	ng	96
81) Endrin	10.98	81	4977	35.87	ng	86
82) Butylbenzylphthalate	11.36	149	221183	177.07	ng	96
83) Methoxychlor	12.00	227	44017	33.29	ng	97
84) 3,3'-Dichlorobenzidine	11.95	252	127321	147.52	ng	99
85) Benzo[a]anthracene	11.95	228	445548	169.65	ng	98
86) Chrysene	12.00	228	399130	163.62	ng	99
87) bis(2-Ethylhexyl)phthalate	12.07	149	300750	173.31	ng	97
89) Di-n-octylphthalate	12.81	149	525668	170.90	ng	98
90) Benzo[b]fluoranthene	13.16	252	376816	172.31	ng	98
91) Benzo[k]fluoranthene	13.19	252	368643	160.21	ng	96
92) Benzo[a]pyrene	13.50	252	358612	172.22	ng	97
93) Indeno[1,2,3-cd]pyrene	14.61	276	365114	159.85	ng	91
94) Dibenzo[a,h]anthracene	14.64	278	294594	155.13	ng	97
95) Benzo[g,h,i]perylene	14.90	276	298173	154.20	ng	92

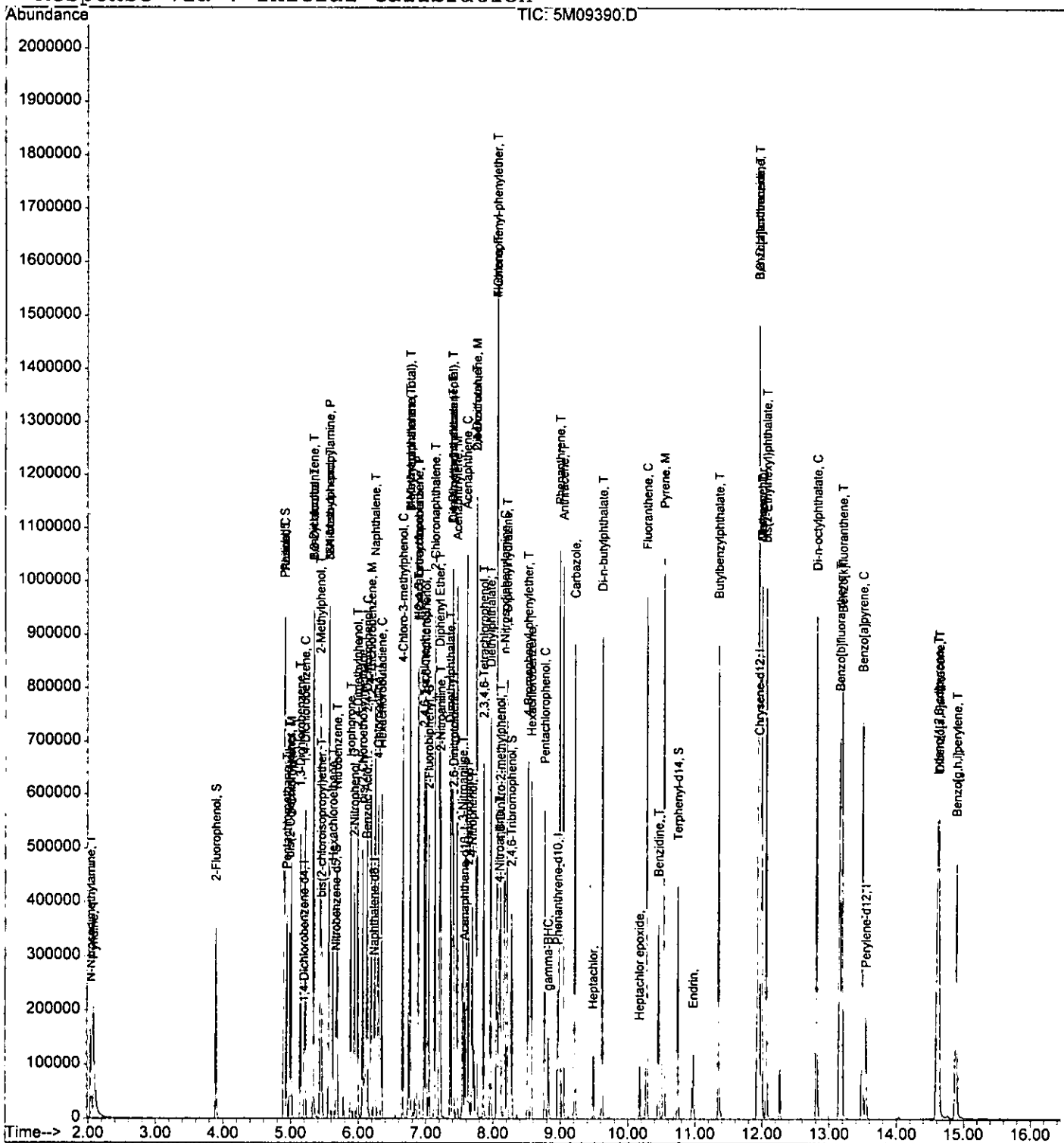
(#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09390.D Vial: 7
Acq On : 22 Jul 2005 10:24 Operator: AHD
Sample : CAL BNA@160PPM Inst : GCMS_5
Misc : A,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Jul 22 10:52 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09391.D Vial: 8
 Acq On : 22 Jul 2005 10:47 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Jul 22 11:16 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.20	152	13701	40.00	ng	-0.05
20) Naphthalene-d8	6.23	136	67172	40.00	ng	-0.05
36) Acenaphthene-d10	7.58	164	47139	40.00	ng	-0.06
61) Phenanthrene-d10	8.96	188	87189	40.00	ng	-0.06
77) Chrysene-d12	11.96	240	73027	40.00	ng	-0.07
88) Perylene-d12	13.55	264	57322	40.00	ng	-0.07

System Monitoring Compounds

4) 2-Fluorophenol	3.90	112	105940	235.38	ng	-0.06
Spiked Amount	200.000		Recovery	=	117.69%	
8) Phenol-d5	4.90	99	153727	243.23	ng	-0.05
Spiked Amount	200.000		Recovery	=	121.62%	
21) Nitrobenzene-d5	5.67	128	28790	97.55	ng	-0.05
Spiked Amount	100.000		Recovery	=	97.55%	
41) 2-Fluorobiphenyl	7.04	172	134361	89.87	ng	-0.05
Spiked Amount	100.000		Recovery	=	89.87%	
64) 2,4,6-Tribromophenol	8.28	330	35604	189.35	ng	-0.06
Spiked Amount	200.000		Recovery	=	94.68%	
80) Terphenyl-d14	10.75	244	171831	99.53	ng	-0.06
Spiked Amount	100.000		Recovery	=	99.53%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.08	79	108556m	189.48	ng	
3) N-Nitrosodimethylamine	2.04	74	73484	213.52	ng	91
5) Aniline	4.91	93	171587	225.59	ng	91
6) Pentachloroethane	4.94	117	31818	186.56	ng	99
7) bis(2-Chloroethyl)ether	4.99	93	99962	207.27	ng	98
9) Phenol	4.91	94	165555	237.72	ng	68
10) 2-Chlorophenol	5.01	128	122714	231.08	ng	98
11) 1,3-Dichlorobenzene	5.14	146	91874	180.35	ng	99
12) 1,4-Dichlorobenzene	5.21	146	97781	188.70	ng	99
13) 1,2-Dichlorobenzene	5.34	146	93299	188.88	ng	98
14) Benzyl alcohol	5.33	108	83627	239.93	ng	98
15) bis(2-chloroisopropyl)ethe	5.45	45	148683	202.22	ng	93
16) 2-Methylphenol	5.44	108	112563	232.35	ng	99
17) Hexachloroethane	5.62	117	39649	183.83	ng	99
18) N-Nitroso-di-n-propylamine	5.56	70	87873	229.65	ng	97
19) 3&4-Methylphenol	5.57	108	121798	237.29	ng	99
22) Nitrobenzene	5.69	77	125232	188.24	ng	97
23) Isophorone	5.89	82	247118	201.46	ng	97
24) 2-Nitrophenol	5.94	139	65439	192.22	ng	93

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09391.D Vial: 8
 Acq On : 22 Jul 2005 10:47 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:16 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.99	107	126108	195.16	ng	96
26) Benzoic Acid	6.13	105	94455	315.35	ng	100
27) bis(2-Chloroethoxy)methane	6.06	93	135780	192.18	ng	99
28) 2,4-Dichlorophenol	6.13	162	111292	208.52	ng	97
29) 1,2,4-Trichlorobenzene	6.19	180	102956	185.01	ng	97
30) Naphthalene	6.25	128	333769	187.99	ng	99
31) 4-Chloroaniline	6.30	127	120325	171.51	ng	99
32) Hexachlorobutadiene	6.34	225	52577	169.41	ng	99
33) 4-Chloro-3-methylphenol	6.66	107	129710	222.22	ng	95
34) 2-Methylnaphthalene	6.77	142	251185	207.75	ng	100
35) Methylnaphthalenes (Total)	6.77	142	251185	207.75	ng	100
37) 1,2,4,5-Tetrachlorobenzene	6.89	216	109890	168.78	ng	96
38) Hexachlorocyclopentadiene	6.88	237	70515	176.88	ng	99
39) 2,4,6-Trichlorophenol	6.98	196	91064	198.23	ng	99
40) 2,4,5-Trichlorophenol	7.01	196	103150	207.87	ng	98
42) 2-Chloronaphthalene	7.14	162	246360	182.58	ng	99
43) 1,4-Dimethylnaphthalene	7.40	156	194152	190.12	ng	96
44) Dimethylnaphthalenes (Total)	7.40	156	194152	190.12	ng	96
45) Diphenyl Ether	7.20	170	162742	182.56	ng	93
46) 2-Nitroaniline	7.22	65	101016	198.90	ng	92
47) Acenaphthylene	7.47	152	401564	189.31	ng	99
48) Dimethylphthalate	7.37	163	301341	195.27	ng	99
49) 2,6-Dinitrotoluene	7.42	165	70393	198.49	ng	95
50) Acenaphthene	7.61	153	246320	187.50	ng	97
51) 3-Nitroaniline	7.55	138	73497	188.48	ng	99
52) 2,4-Dinitrophenol	7.64	184	52621	294.72	ng	70
53) Dibenzofuran	7.76	168	356672	186.37	ng	100
54) 2,4-Dinitrotoluene	7.75	165	98835	202.29	ng	95
55) 4-Nitrophenol	7.70	65	69930	232.39	ng	99
56) 2,3,4,6-Tetrachlorophenol	7.86	232	82490	212.62	ng	99
57) Fluorene	8.06	166	300996	196.65	ng	98
58) 4-Chlorophenyl-phenylether	8.06	204	146858	197.51	ng	94
59) Diethylphthalate	7.97	149	307913	194.59	ng	98
60) 4-Nitroaniline	8.10	138	95140	214.23	ng	99
62) 4,6-Dinitro-2-methylphenol	8.12	198	72172	235.95	ng	100
63) n-Nitrosodiphenylamine	8.18	169	228224	188.29	ng	97
65) 1,2-Diphenylhydrazine	8.21	77	314544	180.88	ng	99
66) 4-Bromophenyl-phenylether	8.52	248	89581	200.44	ng	96
67) Hexachlorobenzene	8.58	284	81001	190.63	ng	86
68) gamma-BHC	8.82	181	12082	38.97	ng	95
69) Pentachlorophenol	8.77	266	63757	236.43	ng	93

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09391.D Vial: 8
 Acq On : 22 Jul 2005 10:47 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:16 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 10:30:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.99	178	467136	183.60	ng	98
71) Anthracene	9.05	178	496761	193.47	ng	98
72) Carbazole	9.22	167	450762	192.10	ng	100
73) Heptachlor	9.49	100	13654	42.15	ng	94
74) Di-n-butylphthalate	9.62	149	562771	199.06	ng	100
75) Heptachlor epoxide	10.17	81	9443	41.91	ng	86
76) Fluoranthene	10.28	202	536180	194.88	ng	98
78) Pyrene	10.54	202	567341	193.01	ng	95
79) Benzidine	10.46	184	157839	146.15	ng	95
81) Endrin	10.98	81	5878	40.96	ng	85
82) Butylbenzylphthalate	11.36	149	255354	198.11	ng	96
83) Methoxychlor	12.00	227	52084	38.59	ng	98
84) 3,3'-Dichlorobenzidine	11.95	252	140568	162.76	ng	97
85) Benzo[a]anthracene	11.95	228	531868	197.77	ng	97
86) Chrysene	12.00	228	468906	188.88	ng	99
87) bis(2-Ethylhexyl)phthalate	12.07	149	349595	195.99	ng	97
89) Di-n-octylphthalate	12.82	149	618336	196.50	ng	99
90) Benzo[b]fluoranthene	13.16	252	480118	214.30	ng	98
91) Benzo[k]fluoranthene	13.20	252	413578	177.65	ng	94
92) Benzo[a]pyrene	13.50	252	422176	197.91	ng	99
93) Indeno[1,2,3-cd]pyrene	14.61	276	458372	198.42	ng	91
94) Dibenzo[a,h]anthracene	14.64	278	387841	202.94	ng	94
95) Benzo[g,h,i]perylene	14.90	276	368899	189.75	ng	94

(#) = qualifier out of range (m) = manual integration

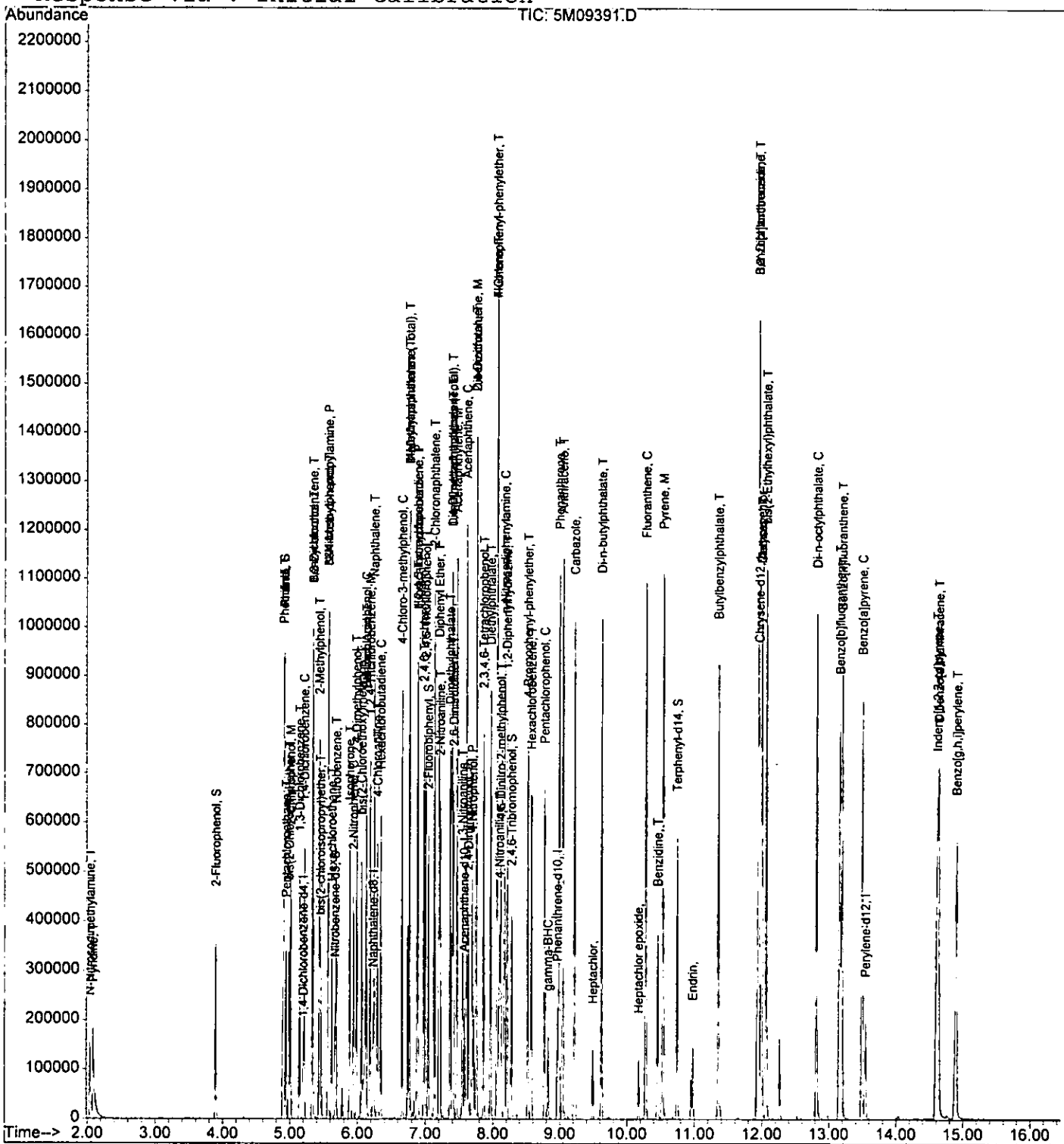
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09391.D Vial: 8
 Acq On : 22 Jul 2005 10:47 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Jul 22 11:16 2005

9255

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05299.D Vial: 2
 Acq On : 3 Aug 2005 8:52 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	39673	40.00	ng	0.00
19) Naphthalene-d8	5.94	136	128050	40.00	ng	0.00
35) Acenaphthene-d10	7.53	164	71936	40.00	ng	0.00
59) Phenanthrene-d10	9.13	188	122199	40.00	ng	0.00
72) Chrysene-d12	12.33	240	86303	40.00	ng	0.00
81) Perylene-d12	14.18	264	51188	40.00	ng	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.80	112	57467	50.98	ng	0.00
Spiked Amount				200.000		
			Recovery	=	25.49%	
7) Phenol-d5	4.65	99	77347	51.08	ng	0.00
Spiked Amount				200.000		
			Recovery	=	25.54%	
20) Nitrobenzene-d5	5.38	128	16723	25.97	ng	0.00
Spiked Amount				100.000		
			Recovery	=	25.97%	
40) 2-Fluorobiphenyl	6.87	172	63978	26.92	ng	0.00
Spiked Amount				100.000		
			Recovery	=	26.92%	
62) 2,4,6-Tribromophenol	8.36	332	29747	53.59	ng	0.00
Spiked Amount				200.000		
			Recovery	=	26.80%	
75) Terphenyl-d14	11.02	244	61668	25.52	ng	0.00
Spiked Amount				100.000		
			Recovery	=	25.52%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.32	79	85909	56.33	ng	98
3) N-Nitrosodimethylamine	2.27	74	45215	53.36	ng	99
5) Aniline	4.67	93	88026	53.54	ng	39
6) bis(2-Chloroethyl)ether	4.74	93	68671	54.94	ng	94
8) Phenol	4.66	94	87209	53.20	ng	67
9) 2-Chlorophenol	4.77	128	63997	50.95	ng	81
10) 1,3-Dichlorobenzene	4.89	146	66100	49.05	ng	99
11) 1,4-Dichlorobenzene	4.95	146	64865	49.07	ng	98
12) 1,2-Dichlorobenzene	5.07	146	67933	53.46	ng	98
13) Benzyl alcohol	5.06	108	40415	51.35	ng	96
14) bis(2-chloroisopropyl)ethe	5.17	45	149247	49.20	ng	97
15) 2-Methylphenol	5.14	108	51097	47.99	ng	100
16) Hexachloroethane	5.34	117	32835	52.87	ng	74
17) N-Nitroso-di-n-propylamine	5.27	70	55752	51.13	ng	89
18) 3&4-Methylphenol	5.27	108	53725	49.52	ng	100
21) Nitrobenzene	5.39	77	68884	49.98	ng	85
22) Isophorone	5.58	82	129469	50.63	ng	97
23) 2-Nitrophenol	5.64	139	34081	48.75	ng	82
24) 2,4-Dimethylphenol	5.68	107	67135	51.63	ng	95

(#) = qualifier out of range (m) = manual integration

11.811

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05299.D Vial: 2
 Acq On : 3 Aug 2005 8:52 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.76	105	10522	51.87	ng	94
26) bis(2-Chloroethoxy)methane	5.76	93	77750	50.26	ng	98
27) 2,4-Dichlorophenol	5.83	162	56526	52.67	ng	94
28) 1,2,4-Trichlorobenzene	5.90	180	59328	51.69	ng	98
29) Naphthalene	5.96	128	155594	53.46	ng	99
30) 4-Chloroaniline	6.00	127	59255	54.18	ng	98
31) Hexachlorobutadiene	6.05	225	36507	49.63	ng	93
32) 4-Chloro-3-methylphenol	6.40	107	59253	50.26	ng	83
33) 2-Methylnaphthalene	6.55	142	103012	50.51	ng	97
34) Methylnaphthalene (Total)	6.55	142	103012	50.51	ng	97
36) 1,2,4,5-Tetrachlorobenzene	6.69	216	64403	53.10	ng	97
37) Hexachlorocyclopentadiene	6.68	237	45464	53.89	ng	98
38) 2,4,6-Trichlorophenol	6.79	196	45343	52.16	ng	99
39) 2,4,5-Trichlorophenol	6.82	196	46633	52.50	ng	98
41) 2-Chloronaphthalene	6.99	162	109150	52.36	ng	97
42) 2-Nitroaniline	7.08	65	61585	50.86	ng	99
43) 1,4-Dimethylnaphthalene	7.30	156	74815	52.31	ng	93
44) Dimethylnaphthalene (Total)	7.30	156	74815	52.31	ng	93
45) Diphenyl Ether	7.07	170	93430	53.16	ng	85
46) Acenaphthylene	7.38	152	175347	54.93	ng	99
47) Dimethylphthalate	7.26	163	133883	51.70	ng	100
48) 2,6-Dinitrotoluene	7.32	165	33039	53.50	ng	93
49) Acenaphthene	7.56	153	108815	52.25	ng	98
50) 3-Nitroaniline	7.49	138	29542	55.96	ng	84
51) 2,4-Dinitrophenol	7.59	184	12946	44.55	ng	80
52) Dibenzofuran	7.73	168	149063	53.38	ng	89
53) 2,4-Dinitrotoluene	7.72	165	43321	53.55	ng	93
54) 4-Nitrophenol	7.65	65	33300	50.25	ng	100
55) Fluorene	8.09	166	109614	53.69	ng	98
56) 4-Chlorophenyl-phenylether	8.10	204	61868	54.75	ng	93
57) Diethylphthalate	7.99	149	142915	52.65	ng	99
58) 4-Nitroaniline	8.12	138	34476	50.16	ng	84
60) 4,6-Dinitro-2-methylphenol	8.15	198	20728	54.40	ng	100
61) n-Nitrosodiphenylamine	8.22	169	86656	54.29	ng	99
63) 1,2-Diphenylhydrazine	8.27	77	152252	53.55	ng	85
64) 4-Bromophenyl-phenylether	8.63	248	43454	53.73	ng	96
65) Hexachlorobenzene	8.69	284	59455	54.70	ng	98
66) Pentachlorophenol	8.92	266	29346	56.42	ng	92
67) Phenanthrene	9.15	178	166352	53.07	ng	99
68) Anthracene	9.21	178	170892	53.44	ng	99
69) Carbazole	9.41	167	160300	54.76	ng	99

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05299.D Vial: 2
 Acq On : 3 Aug 2005 8:52 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	236389	51.66	ng	98
71) Fluoranthene	10.54	202	174359	53.10	ng	90
73) Pyrene	10.81	202	176886	52.73	ng	85
74) Benzidine	10.73	184	50425	57.56	ng	99
76) Butylbenzylphthalate	11.67	149	82010	50.68	ng	91
77) 3,3'-Dichlorobenzidine	12.31	252	34075	47.38	ng	99
78) Benzo[a]anthracene	12.32	228	139158	50.57	ng	100
79) Chrysene	12.36	228	123049	50.29	ng	98
80) bis(2-Ethylhexyl)phthalate	12.46	149	109441	52.05	ng	94
82) Di-n-octylphthalate	13.31	149	145189	52.88	ng	99
83) Benzo[b]fluoranthene	13.71	252	119383	55.94	ng	96
84) Benzo[k]fluoranthene	13.74	252	93154	49.57	ng	95
85) Benzo[a]pyrene	14.11	252	90928	51.41	ng	98
86) Indeno[1,2,3-cd]pyrene	15.42	276	80045	49.64	ng	85
87) Dibenzo[a,h]anthracene	15.45	278	66699	50.55	ng	95
88) Benzo[g,h,i]perylene	15.71	276	63755	49.40	ng	92

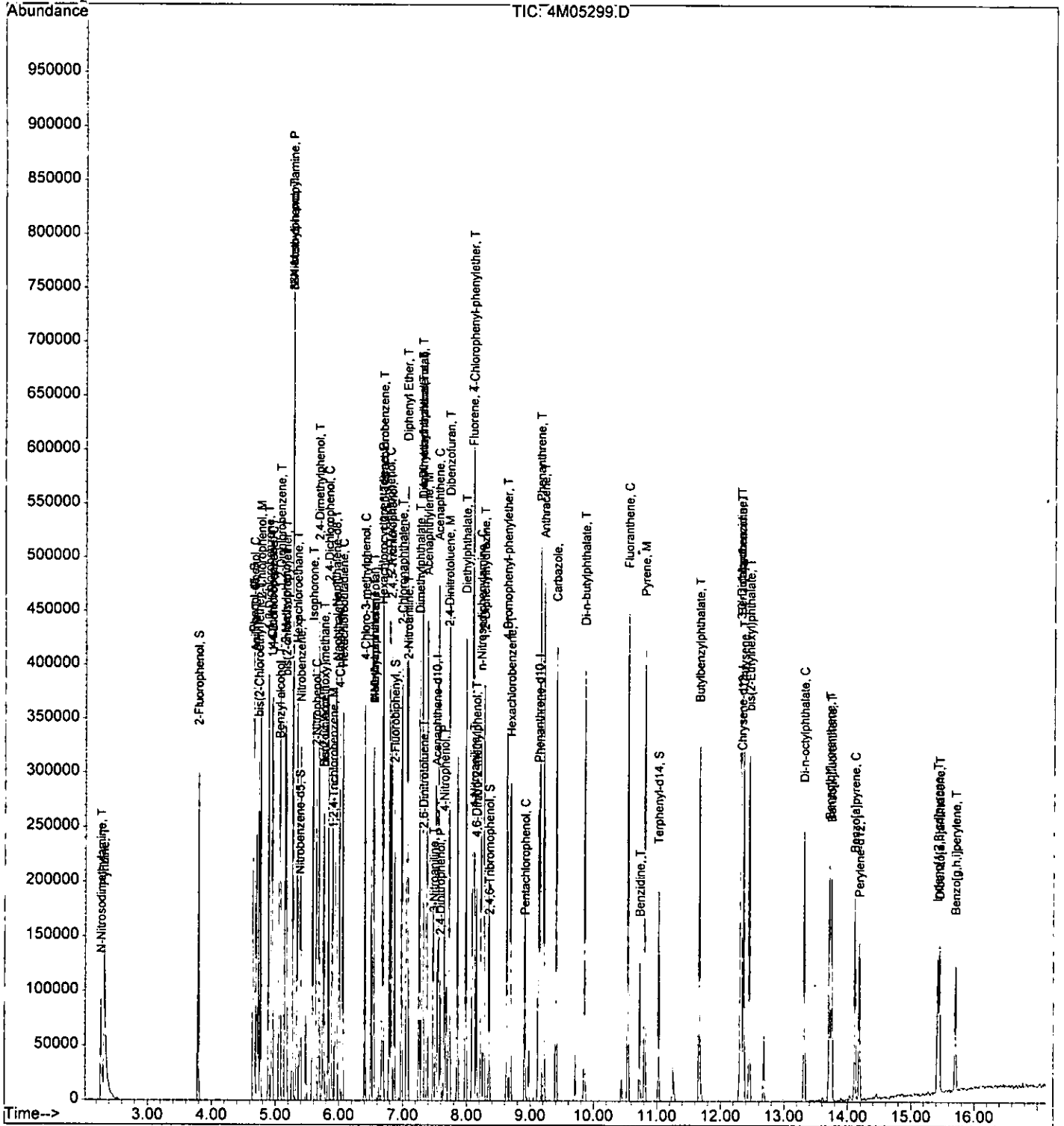
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05299.D Vial: 2
 Acq On : 3 Aug 2005 8:52 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05300.D Vial: 3
 Acq On : 3 Aug 2005 9:19 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 9:36 2005 Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	30220	40.00	ng	-0.09
19) Naphthalene-d8	5.94	136	95624	40.00	ng	-0.09
35) Acenaphthene-d10	7.53	164	54722	40.00	ng	-0.10
59) Phenanthrene-d10	9.13	188	96248	40.00	ng	-0.11
72) Chrysene-d12	12.33	240	78377	40.00	ng	-0.13
81) Perylene-d12	14.18	264	54571	40.00	ng	-0.14

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
4) 2-Fluorophenol	3.79	112	8338	9.53	ng	-0.10
Spiked Amount 200.000			Recovery =	4.77%		
7) Phenol-d5	4.64	99	10969	9.48	ng	-0.09
Spiked Amount 200.000			Recovery =	4.74%		
20) Nitrobenzene-d5	5.38	128	2557	5.63	ng	-0.08
Spiked Amount 100.000			Recovery =	5.63%		
40) 2-Fluorobiphenyl	6.87	172	10608	5.66	ng	-0.10
Spiked Amount 100.000			Recovery =	5.66%		
62) 2,4,6-Tribromophenol	8.35	332	4477	9.99	ng	-0.10
Spiked Amount 200.000			Recovery =	5.00%		
75) Terphenyl-d14	11.02	244	10503	5.04	ng	-0.12
Spiked Amount 100.000			Recovery =	5.04%		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.40	79	8457	6.91	ng	96
3) N-Nitrosodimethylamine	2.32	74	3898	5.41	ng	96
5) Aniline	4.67	93	9168	7.14	ng	21
6) bis(2-Chloroethyl)ether	4.73	93	9317	10.20	ng	92
8) Phenol	4.65	94	12224	9.24	ng	58
9) 2-Chlorophenol	4.77	128	9655	10.22	ng	85
10) 1,3-Dichlorobenzene	4.89	146	10533	10.34	ng	98
11) 1,4-Dichlorobenzene	4.95	146	10279	9.92	ng	95
12) 1,2-Dichlorobenzene	5.07	146	10736	10.59	ng	98
13) Benzyl alcohol	5.05	108	5367	8.97	ng	57
14) bis(2-chloroisopropyl)ethe	5.16	45	22155	9.44	ng	97
15) 2-Methylphenol	5.14	108	8915	11.79	ng	95
16) Hexachloroethane	5.34	117	4685	10.30	ng	75
17) N-Nitroso-di-n-propylamine	5.27	70	8441	11.04	ng	84
18) 3&4-Methylphenol	5.27	108	9426	11.94	ng	100
21) Nitrobenzene	5.39	77	10526	10.29	ng	88
22) Isophorone	5.58	82	18903	10.28	ng	94
23) 2-Nitrophenol	5.63	139	4617	8.87	ng	68
24) 2,4-Dimethylphenol	5.68	107	10772	11.71	ng	98

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Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05300.D Vial: 3
 Acq On : 3 Aug 2005 9:19 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 9:36 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.68	105	667	3.45	ng	# 1
26) bis(2-Chloroethoxy)methane	5.75	93	11781	10.03	ng	96
27) 2,4-Dichlorophenol	5.83	162	8385	10.45	ng	90
28) 1,2,4-Trichlorobenzene	5.89	180	8537	9.72	ng	94
29) Naphthalene	5.96	128	23646	9.92	ng	100
30) 4-Chloroaniline	6.00	127	7807	7.77	ng	98
31) Hexachlorobutadiene	6.06	225	5680	9.96	ng	92
32) 4-Chloro-3-methylphenol	6.40	107	8708	10.30	ng	90
33) 2-Methylnaphthalene	6.54	142	17387	11.59	ng	93
34) Methylnaphthalene(Total)	6.54	142	17387	11.59	ng	93
36) 1,2,4,5-Tetrachlorobenzene	6.69	216	10414	10.53	ng	99
37) Hexachlorocyclopentadiene	6.68	237	5209	7.81	ng	99
38) 2,4,6-Trichlorophenol	6.79	196	6851	10.29	ng	97
39) 2,4,5-Trichlorophenol	6.82	196	6856	10.12	ng	95
41) 2-Chloronaphthalene	6.98	162	18177	11.06	ng	94
42) 2-Nitroaniline	7.08	65	9539	10.79	ng	90
43) 1,4-Dimethylnaphthalene	7.30	156	12250	11.06	ng	92
44) Dimethylnaphthalene(Total)	7.30	156	12250	11.06	ng	92
45) Diphenyl Ether	7.07	170	15349	14.25	ng	87
46) Acenaphthylene	7.38	152	27020	10.65	ng	97
47) Dimethylphthalate	7.26	163	21942	11.63	ng	98
48) 2,6-Dinitrotoluene	7.31	165	4630	10.81	ng	80
49) Acenaphthene	7.56	153	18118	11.02	ng	98
50) 3-Nitroaniline	7.47	138	4410	9.74	ng	78
51) 2,4-Dinitrophenol	0.00	184	0	N.D.		
52) Dibenzofuran	7.73	168	24244	11.10	ng	93
53) 2,4-Dinitrotoluene	7.72	165	6255	10.57	ng	96
54) 4-Nitrophenol	7.65	65	3874	8.14	ng	77
55) Fluorene	8.09	166	18461	11.50	ng	95
56) 4-Chlorophenyl-phenylether	8.10	204	10165	11.84	ng	89
57) Diethylphthalate	7.99	149	22817	11.85	ng	97
58) 4-Nitroaniline	8.11	138	5773	12.47	ng	85
60) 4,6-Dinitro-2-methylphenol	8.15	198	791	2.14	ng	100
61) n-Nitrosodiphenylamine	8.22	169	13807	10.48	ng	99
63) 1,2-Diphenylhydrazine	8.26	77	21999	9.55	ng	96
64) 4-Bromophenyl-phenylether	8.63	248	6454	9.96	ng	89
65) Hexachlorobenzene	8.68	284	8652	9.85	ng	70
66) Pentachlorophenol	8.92	266	1923	4.10	ng	96
67) Phenanthrene	9.15	178	28026	10.79	ng	98
68) Anthracene	9.21	178	28354	10.72	ng	99
69) Carbazole	9.41	167	24952	10.68	ng	97

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05300.D Vial: 3
 Acq On : 3 Aug 2005 9:19 Operator: AHD
 Sample : CAL BNA@10PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 9:36 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	47520	14.87	ng	99
71) Fluoranthene	10.54	202	29760	11.65	ng	87
73) Pyrene	10.81	202	31795	10.49	ng	86
74) Benzidine	10.73	184	5048	8.06	ng	93
76) Butylbenzylphthalate	11.67	149	14139	9.93	ng	95
77) 3,3'-Dichlorobenzidine	12.31	252	8466	9.36	ng	90
78) Benzo[a]anthracene	12.32	228	26060	10.35	ng	99
79) Chrysene	12.36	228	24048	10.30	ng	98
80) bis(2-Ethylhexyl)phthalate	12.45	149	17483	8.99	ng	95
82) Di-n-octylphthalate	13.32	149	23454	8.49	ng	96
83) Benzo[b]fluoranthene	13.71	252	21994	10.52	ng	94
84) Benzo[k]fluoranthene	13.74	252	22340	11.16	ng	97
85) Benzo[a]pyrene	14.11	252	19538	10.46	ng	97
86) Indeno[1,2,3-cd]pyrene	15.42	276	17539	8.04	ng	91
87) Dibenzo[a,h]anthracene	15.45	278	14381	8.20	ng	92
88) Benzo[g,h,i]perylene	15.71	276	14447	8.01	ng	94

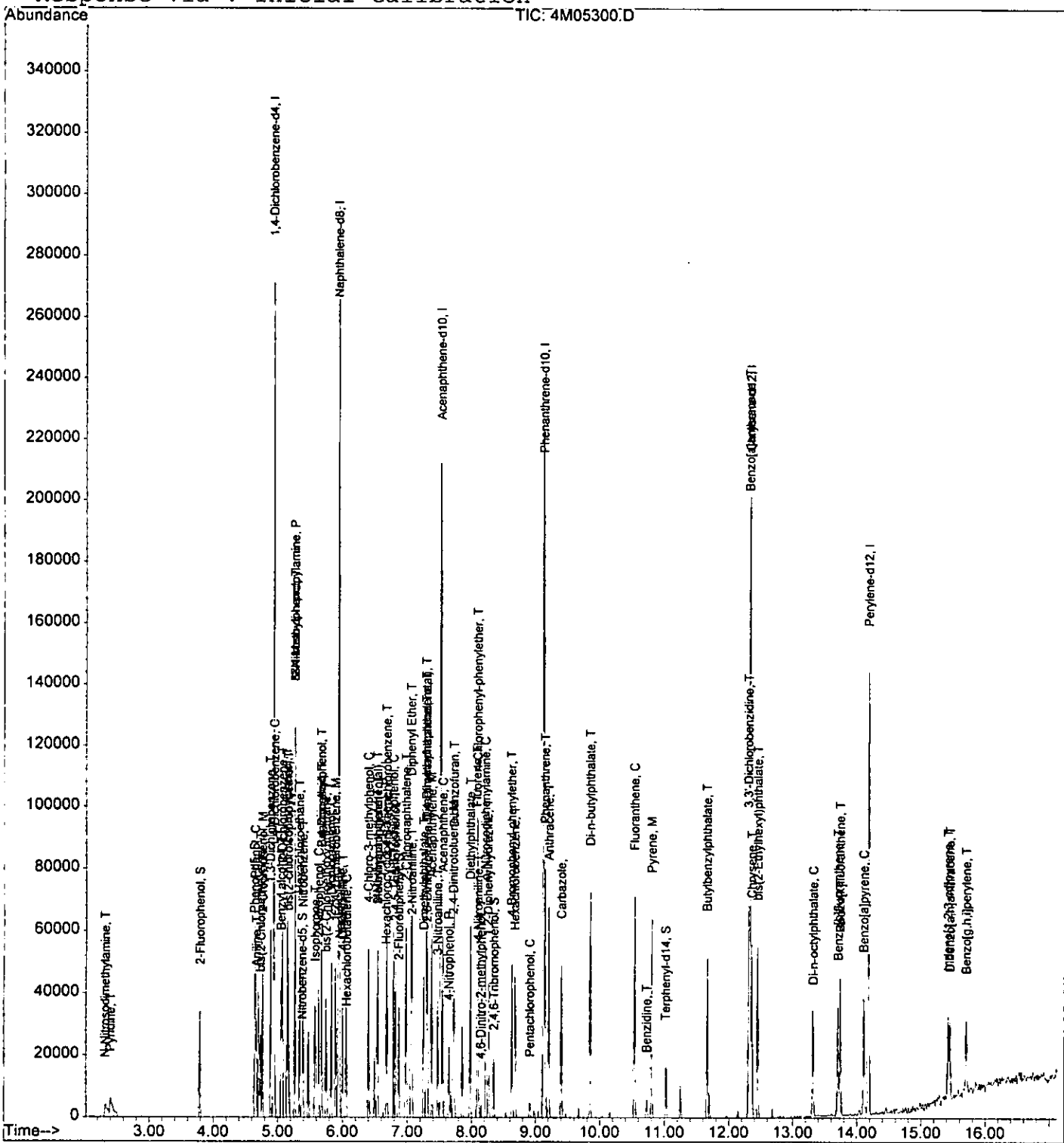
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05300.D Vial: 3
Acq On : 3 Aug 2005 9:19 Operator: AHD
Sample : CAL BNA@10PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 9:36 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05301.D Vial: 4
 Acq On : 3 Aug 2005 9:43 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 10:00 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	44723	40.00	ng	-0.09
19) Naphthalene-d8	5.94	136	140434	40.00	ng	-0.09
35) Acenaphthene-d10	7.52	164	75558	40.00	ng	-0.11
59) Phenanthrene-d10	9.12	188	127553	40.00	ng	-0.12
72) Chrysene-d12	12.33	240	90498	40.00	ng	-0.13
81) Perylene-d12	14.18	264	57551	40.00	ng	-0.14
System Monitoring Compounds						
4) 2-Fluorophenol	3.80	112	30986	23.94	ng	-0.10
Spiked Amount	200.000		Recovery	=	11.97%	
7) Phenol-d5	4.64	99	42217	24.66	ng	-0.09
Spiked Amount	200.000		Recovery	=	12.33%	
20) Nitrobenzene-d5	5.38	128	8126	12.18	ng	-0.08
Spiked Amount	100.000		Recovery	=	12.18%	
40) 2-Fluorobiphenyl	6.87	172	32098	12.39	ng	-0.10
Spiked Amount	100.000		Recovery	=	12.39%	
62) 2,4,6-Tribromophenol	8.35	332	14338	24.14	ng	-0.11
Spiked Amount	200.000		Recovery	=	12.07%	
75) Terphenyl-d14	11.02	244	31540	13.11	ng	-0.12
Spiked Amount	100.000		Recovery	=	13.11%	
Target Compounds						
2) Pyridine	2.33	79	44735	24.70	ng	97
3) N-Nitrosodimethylamine	2.27	74	25639	24.03	ng	93
5) Aniline	4.68	93	42578	22.42	ng	31
6) bis(2-Chloroethyl)ether	4.74	93	35042	25.92	ng	83
8) Phenol	4.65	94	45814	23.41	ng	58
9) 2-Chlorophenol	4.77	128	36133	25.84	ng	90
10) 1,3-Dichlorobenzene	4.89	146	41882	27.78	ng	99
11) 1,4-Dichlorobenzene	4.95	146	40304	26.27	ng	96
12) 1,2-Dichlorobenzene	5.07	146	39655	26.42	ng	99
13) Benzyl alcohol	5.05	108	23115	26.11	ng	69
14) bis(2-chloroisopropyl)ethe	5.17	45	85170	24.52	ng	96
15) 2-Methylphenol	5.15	108	32360	28.92	ng	99
16) Hexachloroethane	5.34	117	18337	27.24	ng	55
17) N-Nitroso-di-n-propylamine	5.27	70	31218	27.59	ng	85
18) 3&4-Methylphenol	5.27	108	35346	30.26	ng	95
21) Nitrobenzene	5.39	77	42030	27.97	ng	89
22) Isophorone	5.58	82	70827	26.22	ng	89
23) 2-Nitrophenol	5.64	139	19430	25.41	ng	92
24) 2,4-Dimethylphenol	5.68	107	38860	28.77	ng	98

(#) = qualifier out of range (m) = manual integration

4M05301.D 4M_0803.M

Thu Aug 11 17:09:17 2005

RPT1

Page 1

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05301.D Vial: 4
 Acq On : 3 Aug 2005 9:43 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 10:00 2005 Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.74	105	5249	18.49	ng	95
26) bis(2-Chloroethoxy)methane	5.75	93	43983	25.51	ng	98
27) 2,4-Dichlorophenol	5.83	162	31352	26.61	ng	89
28) 1,2,4-Trichlorobenzene	5.89	180	33616	26.06	ng	96
29) Naphthalene	5.95	128	81143	23.18	ng	99
30) 4-Chloroaniline	6.00	127	31873	21.60	ng	95
31) Hexachlorobutadiene	6.06	225	21704	25.93	ng	98
32) 4-Chloro-3-methylphenol	6.40	107	33315	26.83	ng	89
33) 2-Methylnaphthalene	6.54	142	56869	25.81	ng	96
34) Methylnaphthalene(Total)	6.54	142	56869	25.81	ng	96
36) 1,2,4,5-Tetrachlorobenzene	6.69	216	36052	26.40	ng	97
37) Hexachlorocyclopentadiene	6.68	237	24503	26.62	ng	94
38) 2,4,6-Trichlorophenol	6.79	196	27067	29.44	ng	97
39) 2,4,5-Trichlorophenol	6.82	196	24429	26.12	ng	97
41) 2-Chloronaphthalene	6.99	162	58848	25.92	ng	95
42) 2-Nitroaniline	7.08	65	34503	28.28	ng	88
43) 1,4-Dimethylnaphthalene	7.30	156	42475	27.78	ng	96
44) Dimethylnaphthalene(Total)	7.30	156	42475	27.78	ng	96
45) Diphenyl Ether	7.07	170	52976	35.62	ng	80
46) Acenaphthylene	7.38	152	91287	26.07	ng	98
47) Dimethylphthalate	7.26	163	72655	27.89	ng	99
48) 2,6-Dinitrotoluene	7.31	165	16532	27.95	ng	77
49) Acenaphthene	7.56	153	59672	26.29	ng	99
50) 3-Nitroaniline	7.48	138	15681	25.08	ng	86
51) 2,4-Dinitrophenol	7.59	184	3598	9.87	ng	76
52) Dibenzofuran	7.73	168	81724	27.11	ng	95
53) 2,4-Dinitrotoluene	7.72	165	23099	28.27	ng	99
54) 4-Nitrophenol	7.65	65	15714	23.92	ng	83
55) Fluorene	8.09	166	59425	26.81	ng	97
56) 4-Chlorophenyl-phenylether	8.10	204	33506	28.26	ng	95
57) Diethylphthalate	7.99	149	75778	28.51	ng	98
58) 4-Nitroaniline	8.11	138	17644	27.60	ng	88
60) 4,6-Dinitro-2-methylphenol	8.15	198	8275	16.89	ng	100
61) n-Nitrosodiphenylamine	8.22	169	45156	25.86	ng	97
63) 1,2-Diphenylhydrazine	8.26	77	78290	25.63	ng	99
64) 4-Bromophenyl-phenylether	8.63	248	23004	26.79	ng	98
65) Hexachlorobenzene	8.68	284	30228	25.96	ng	79
66) Pentachlorophenol	8.91	266	11493	18.49	ng	96
67) Phenanthrene	9.15	178	87333	25.38	ng	99
68) Anthracene	9.21	178	90771	25.91	ng	98
69) Carbazole	9.40	167	78660	25.41	ng	95

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05301.D Vial: 4
 Acq On : 3 Aug 2005 9:43 Operator: AHD
 Sample : CAL BNA@25PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 10:00 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.85	149	124432	29.37	ng	99
71) Fluoranthene	10.54	202	87980	25.99	ng	86
73) Pyrene	10.80	202	91576	26.17	ng	97
74) Benzidine	10.73	184	22721	31.41	ng	91
76) Butylbenzylphthalate	11.67	149	42230	25.68	ng	90
77) 3,3'-Dichlorobenzidine	12.31	252	19350	18.52	ng	95
78) Benzo[a]anthracene	12.31	228	75324	25.92	ng	98
79) Chrysene	12.36	228	66545	24.68	ng	98
80) bis(2-Ethylhexyl)phthalate	12.45	149	56542	25.19	ng	99
82) Di-n-octylphthalate	13.31	149	74305	25.52	ng	98
83) Benzo[b]fluoranthene	13.71	252	58175	26.38	ng	95
84) Benzo[k]fluoranthene	13.74	252	57814	27.38	ng	98
85) Benzo[a]pyrene	14.11	252	50576	25.67	ng	97
86) Indeno[1,2,3-cd]pyrene	15.42	276	44676	19.43	ng	83
87) Dibenzo[a,h]anthracene	15.45	278	36253	19.59	ng	92
88) Benzo[g,h,i]perylene	15.71	276	34490	18.14	ng	88

 (#) = qualifier out of range (m) = manual integration

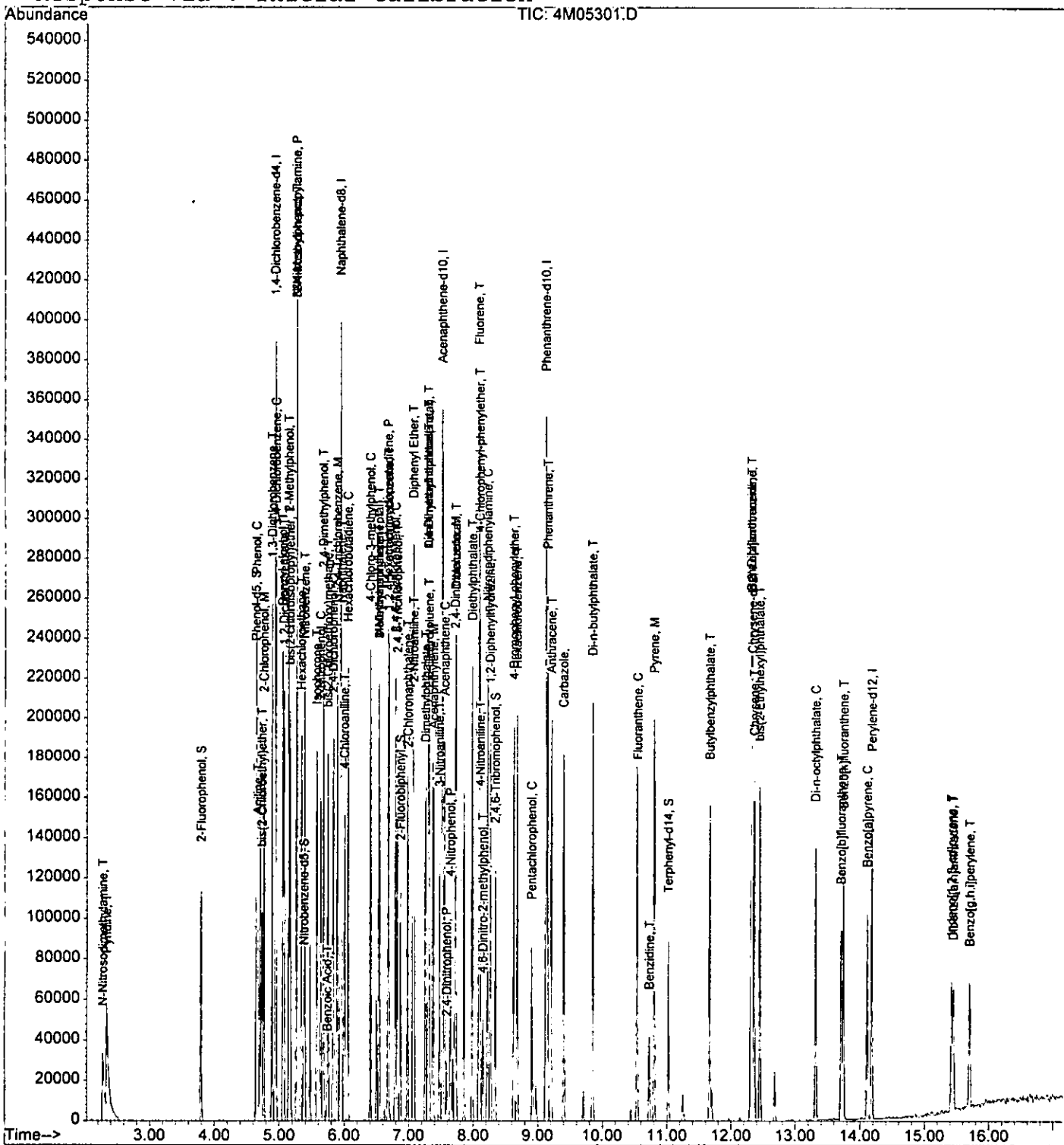
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05301.D Vial: 4
Acq On : 3 Aug 2005 9:43 Operator: AHD
Sample : CAL BNA@25PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 10:00 2005

6755

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05302.D Vial: 5
 Acq On : 3 Aug 2005 10:07 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 10:24 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Thu Jul 28 12:16:29 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	38055	40.00	ng	-0.09
19) Naphthalene-d8	5.94	136	125850	40.00	ng	-0.09
35) Acenaphthene-d10	7.53	164	72993	40.00	ng	-0.10
59) Phenanthrene-d10	9.13	188	125355	40.00	ng	-0.11
72) Chrysene-d12	12.33	240	82228	40.00	ng	-0.13
81) Perylene-d12	14.18	264	47754	40.00	ng	-0.14

System Monitoring Compounds

4) 2-Fluorophenol	3.79	112	90601	82.27	ng	-0.10
Spiked Amount	200.000		Recovery	=	41.14%	
7) Phenol-d5	4.65	99	123684	84.92	ng	-0.08
Spiked Amount	200.000		Recovery	=	42.46%	
20) Nitrobenzene-d5	5.38	128	25680	42.95	ng	-0.08
Spiked Amount	100.000		Recovery	=	42.95%	
40) 2-Fluorobiphenyl	6.87	172	95265	38.08	ng	-0.10
Spiked Amount	100.000		Recovery	=	38.08%	
62) 2,4,6-Tribromophenol	8.35	332	46606	79.83	ng	-0.10
Spiked Amount	200.000		Recovery	=	39.92%	
75) Terphenyl-d14	11.03	244	99121	45.33	ng	-0.11
Spiked Amount	100.000		Recovery	=	45.33%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.32	79	110193	71.49	ng	98
3) N-Nitrosodimethylamine	2.27	74	72835	80.23	ng	97
5) Aniline	4.67	93	148960	92.17	ng	45
6) bis(2-Chloroethyl)ether	4.74	93	96140	83.57	ng	97
8) Phenol	4.66	94	129307	77.65	ng	67
9) 2-Chlorophenol	4.77	128	92547	77.77	ng	73
10) 1,3-Dichlorobenzene	4.89	146	98289	76.63	ng	98
11) 1,4-Dichlorobenzene	4.96	146	97548	74.73	ng	97
12) 1,2-Dichlorobenzene	5.07	146	101025	79.10	ng	97
13) Benzyl alcohol	5.06	108	69570	92.37	ng	82
14) bis(2-chloroisopropyl)ethe	5.16	45	243948	82.54	ng	98
15) 2-Methylphenol	5.15	108	80543	84.60	ng	99
16) Hexachloroethane	5.34	117	49119	85.77	ng	85
17) N-Nitroso-di-n-propylamine	5.27	70	88409	91.83	ng	93
18) 3&4-Methylphenol	5.28	108	78992	79.48	ng	96
21) Nitrobenzene	5.40	77	105499	78.36	ng	91
22) Isophorone	5.58	82	194956	80.54	ng	96
23) 2-Nitrophenol	5.65	139	60041	87.61	ng	73
24) 2,4-Dimethylphenol	5.68	107	94399	77.99	ng	88

(#) = qualifier out of range (m) = manual integration

hmr

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05302.D Vial: 5
 Acq On : 3 Aug 2005 10:07 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 10:24 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)

Title : @GCMS_4,mg,625,8270

Last Update : Thu Jul 28 12:16:29 2005

Response via : Initial Calibration

DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.79	105	19128	75.18	ng	93
26) bis(2-Chloroethoxy)methane	5.76	93	131119	84.85	ng	99
27) 2,4-Dichlorophenol	5.83	162	82482	78.11	ng	100
28) 1,2,4-Trichlorobenzene	5.90	180	93946	81.28	ng	99
29) Naphthalene	5.96	128	249345	79.48	ng	99
30) 4-Chloroaniline	6.00	127	97373	73.65	ng	100
31) Hexachlorobutadiene	6.06	225	57036	76.03	ng	98
32) 4-Chloro-3-methylphenol	6.40	107	94746	85.15	ng	80
33) 2-Methylnaphthalene	6.55	142	168274	85.21	ng	98
34) Methylnaphthalene (Total)	6.55	142	168274	85.21	ng	98
36) 1,2,4,5-Tetrachlorobenzene	6.69	216	94256	71.44	ng	97
37) Hexachlorocyclopentadiene	6.68	237	72477	81.51	ng	99
38) 2,4,6-Trichlorophenol	6.79	196	68532	77.17	ng	97
39) 2,4,5-Trichlorophenol	6.82	196	73846	81.72	ng	98
41) 2-Chloronaphthalene	6.98	162	169634	77.35	ng	99
42) 2-Nitroaniline	7.09	65	99115	84.08	ng	66
43) 1,4-Dimethylnaphthalene	7.30	156	111509	75.48	ng	92
44) Dimethylnaphthalene (Total)	7.30	156	111509	75.48	ng	92
45) Diphenyl Ether	7.07	170	135235	94.12	ng	92
46) Acenaphthylene	7.38	152	254362	75.19	ng	97
47) Dimethylphthalate	7.26	163	199926	79.45	ng	98
48) 2,6-Dinitrotoluene	7.32	165	51641	90.39	ng	83
49) Acenaphthene	7.56	153	164163	74.87	ng	99
50) 3-Nitroaniline	7.49	138	43368	71.80	ng	95
51) 2,4-Dinitrophenol	7.59	184	24618	69.91	ng	49
52) Dibenzofuran	7.73	168	227751	78.20	ng	90
53) 2,4-Dinitrotoluene	7.73	165	69951	88.61	ng	93
54) 4-Nitrophenol	7.66	65	59695	94.05	ng	86
55) Fluorene	8.10	166	167020	78.00	ng	96
56) 4-Chlorophenyl-phenylether	8.10	204	87824	76.66	ng	85
57) Diethylphthalate	7.99	149	221442	86.24	ng	96
58) 4-Nitroaniline	8.13	138	56719	91.83	ng	92
60) 4,6-Dinitro-2-methylphenol	8.16	198	40159	83.39	ng	100
61) n-Nitrosodiphenylamine	8.23	169	133816	77.99	ng	98
63) 1,2-Diphenylhydrazine	8.27	77	251635	83.83	ng	92
64) 4-Bromophenyl-phenylether	8.64	248	71616	84.88	ng	85
65) Hexachlorobenzene	8.69	284	93205	81.44	ng	93
66) Pentachlorophenol	8.92	266	52709	86.29	ng	96
67) Phenanthrene	9.15	178	265349	78.46	ng	100
68) Anthracene	9.21	178	263765	76.60	ng	99
69) Carbazole	9.41	167	241236	79.31	ng	98

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05302.D Vial: 5
 Acq On : 3 Aug 2005 10:07 Operator: AHD
 Sample : CAL BNA@80PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 10:24 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)

Title : @GCMS_4,mg,625,8270

Last Update : Thu Jul 28 12:16:29 2005

Response via : Initial Calibration

DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	358569	86.13	ng	99
71) Fluoranthene	10.54	202	262150	78.80	ng	99
73) Pyrene	10.81	202	262559	82.58	ng	93
74) Benzidine	10.73	184	80751	122.85	ng	99
76) Butylbenzylphthalate	11.67	149	126733	84.82	ng	99
77) 3,3'-Dichlorobenzidine	12.31	252	53730	56.60	ng	98
78) Benzo[a]anthracene	12.32	228	212410	80.43	ng	98
79) Chrysene	12.37	228	188595	76.97	ng	99
80) bis(2-Ethylhexyl)phthalate	12.45	149	163909	80.37	ng	93
82) Di-n-octylphthalate	13.32	149	228212	94.45	ng	98
83) Benzo[b]fluoranthene	13.71	252	161705	88.38	ng	97
84) Benzo[k]fluoranthene	13.75	252	147295	84.08	ng	98
85) Benzo[a]pyrene	14.12	252	134825	82.45	ng	93
86) Indeno[1,2,3-cd]pyrene	15.42	276	120320	63.06	ng	93
87) Dibenzo[a,h]anthracene	15.45	278	98006	63.83	ng	97
88) Benzo[g,h,i]perylene	15.71	276	95806	60.74	ng	96

 (#) = qualifier out of range (m) = manual integration

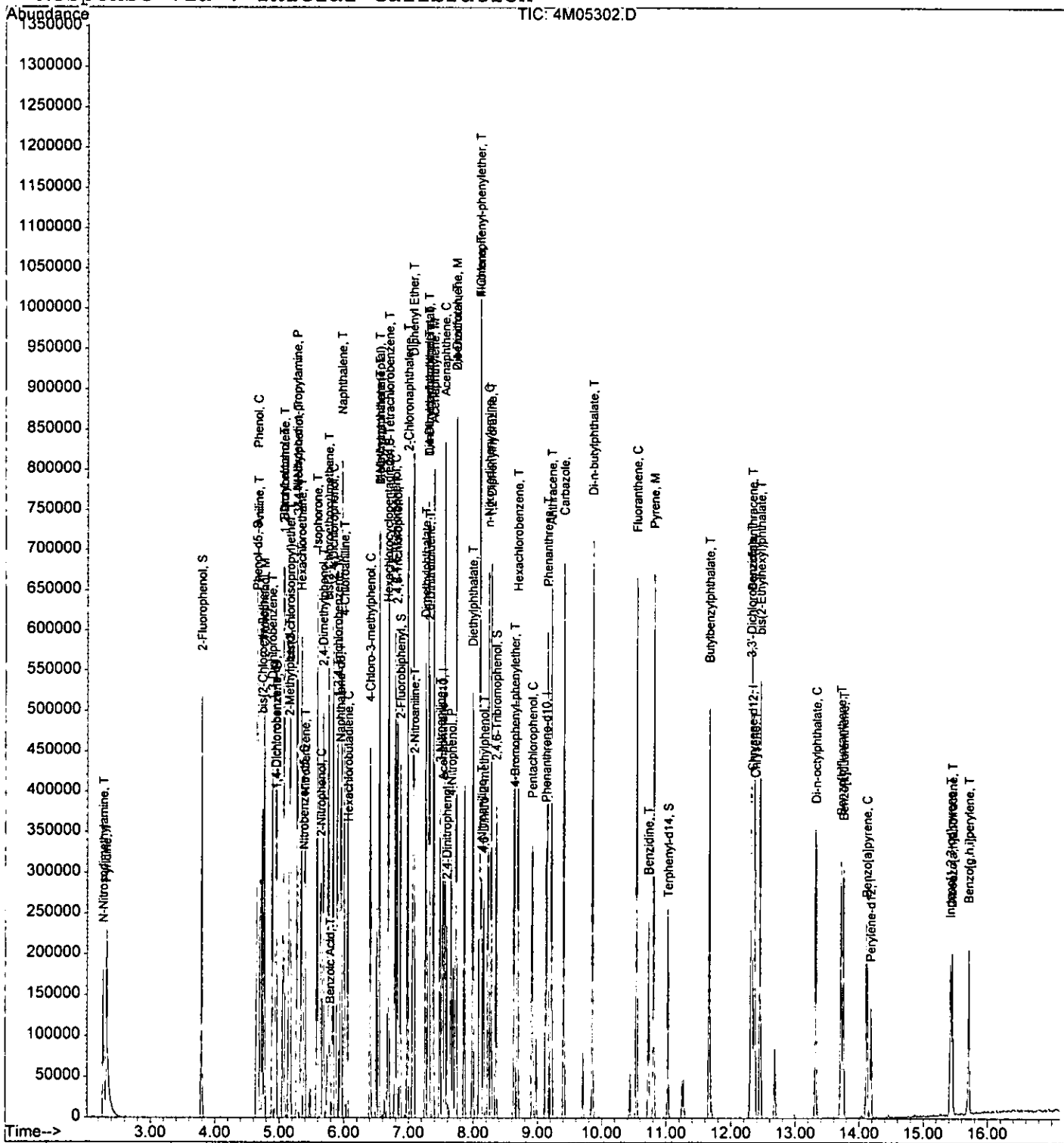
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05302.D Vial: 5
Acq On : 3 Aug 2005 10:07 Operator: AHD
Sample : CAL BNA@80PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 10:24 2005

7755

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05303.D Vial: 6
 Acq On : 3 Aug 2005 10:31 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:27 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	43595	40.00	ng	0.00
19) Naphthalene-d8	5.94	136	141452	40.00	ng	0.00
35) Acenaphthene-d10	7.53	164	87716	40.00	ng	0.00
59) Phenanthrene-d10	9.13	188	158300	40.00	ng	0.00
72) Chrysene-d12	12.34	240	111899	40.00	ng	0.00
81) Perylene-d12	14.18	264	77716	40.00	ng	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
4) 2-Fluorophenol	3.79	112	140357	113.30	ng	0.00
Spiked Amount				200.000		
			Recovery	=	56.65%	
7) Phenol-d5	4.65	99	199212	119.72	ng	0.00
Spiked Amount				200.000		
			Recovery	=	59.86%	
20) Nitrobenzene-d5	5.38	128	41633	58.52	ng	0.00
Spiked Amount				100.000		
			Recovery	=	58.52%	
40) 2-Fluorobiphenyl	6.87	172	150039	51.77	ng	0.00
Spiked Amount				100.000		
			Recovery	=	51.77%	
62) 2,4,6-Tribromophenol	8.35	332	82577	114.84	ng	0.00
Spiked Amount				200.000		
			Recovery	=	57.42%	
75) Terphenyl-d14	11.03	244	182824	58.35	ng	0.00
Spiked Amount				100.000		
			Recovery	=	58.35%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.31	79	219757	131.13	ng	96
3) N-Nitrosodimethylamine	2.27	74	119165	127.98	ng	98
5) Aniline	4.67	93	226924	125.62	ng	38
6) bis(2-Chloroethyl)ether	4.75	93	157466	114.64	ng	85
8) Phenol	4.67	94	230585	128.02	ng	92
9) 2-Chlorophenol	4.78	128	166226	120.43	ng	91
10) 1,3-Dichlorobenzene	4.90	146	172119	116.23	ng	98
11) 1,4-Dichlorobenzene	4.96	146	170951	117.69	ng	100
12) 1,2-Dichlorobenzene	5.07	146	145977	104.55	ng	96
13) Benzyl alcohol	5.06	108	92629	107.10	ng	64
14) bis(2-chloroisopropyl)ethe	5.18	45	401940	120.58	ng	98
15) 2-Methylphenol	5.15	108	142079	121.43	ng	99
16) Hexachloroethane	5.34	117	79343	116.27	ng	92
17) N-Nitroso-di-n-propylamine	5.28	70	129025	107.68	ng	93
18) 3&4-Methylphenol	5.28	108	118175	99.12	ng	99
21) Nitrobenzene	5.40	77	194253	127.59	ng	91
22) Isophorone	5.59	82	346664	122.72	ng	98
23) 2-Nitrophenol	5.65	139	101508	131.45	ng	90
24) 2,4-Dimethylphenol	5.69	107	170047	118.38	ng	95

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05303.D Vial: 6
 Acq On : 3 Aug 2005 10:31 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:27 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.81	105	31896	142.33	ng	97
26) bis(2-Chloroethoxy)methane	5.76	93	181806	106.39	ng	100
27) 2,4-Dichlorophenol	5.84	162	141402	119.27	ng	92
28) 1,2,4-Trichlorobenzene	5.90	180	149157	117.64	ng	97
29) Naphthalene	5.96	128	350413	108.99	ng	99
30) 4-Chloroaniline	6.00	127	138148	114.35	ng	99
31) Hexachlorobutadiene	6.06	225	96115	118.29	ng	97
32) 4-Chloro-3-methylphenol	6.41	107	159089	122.16	ng	95
33) 2-Methylnaphthalene	6.55	142	256347	113.78	ng	95
34) Methylnaphthalene (Total)	6.55	142	256347	113.78	ng	95
36) 1,2,4,5-Tetrachlorobenzene	6.70	216	154331	104.35	ng	99
37) Hexachlorocyclopentadiene	6.68	237	119473	116.14	ng	98
38) 2,4,6-Trichlorophenol	6.80	196	112867	106.47	ng	96
39) 2,4,5-Trichlorophenol	6.83	196	126199	116.52	ng	98
41) 2-Chloronaphthalene	6.98	162	260696	102.55	ng	97
42) 2-Nitroaniline	7.09	65	157940	106.97	ng	94
43) 1,4-Dimethylnaphthalene	7.30	156	184732	105.92	ng	89
44) Dimethylnaphthalene (Total)	7.30	156	184732	105.92	ng	89
45) Diphenyl Ether	7.07	170	218268	101.85	ng	99
46) Acenaphthylene	7.38	152	407682	104.74	ng	98
47) Dimethylphthalate	7.27	163	345526	109.43	ng	100
48) 2,6-Dinitrotoluene	7.33	165	90234	119.83	ng	99
49) Acenaphthene	7.56	153	265459	104.54	ng	96
50) 3-Nitroaniline	7.49	138	64247	99.81	ng	85
51) 2,4-Dinitrophenol	7.60	184	54038	152.52	ng	82
52) Dibenzofuran	7.74	168	359778	105.65	ng	93
53) 2,4-Dinitrotoluene	7.73	165	109896	111.40	ng	68
54) 4-Nitrophenol	7.67	65	112018	138.61	ng	81
55) Fluorene	8.10	166	249580	100.26	ng	99
56) 4-Chlorophenyl-phenylether	8.11	204	141171	102.45	ng	95
57) Diethylphthalate	8.00	149	339070	102.43	ng	98
58) 4-Nitroaniline	8.14	138	95829	114.34	ng	82
60) 4,6-Dinitro-2-methylphenol	8.17	198	76287	154.56	ng	100
61) n-Nitrosodiphenylamine	8.24	169	227302	109.92	ng	99
63) 1,2-Diphenylhydrazine	8.27	77	406099	110.25	ng	95
64) 4-Bromophenyl-phenylether	8.64	248	112110	107.01	ng	95
65) Hexachlorobenzene	8.69	284	152560	108.36	ng	81
66) Pentachlorophenol	8.92	266	95934	142.38	ng	98
67) Phenanthrene	9.16	178	423390	104.26	ng	99
68) Anthracene	9.22	178	450464	108.74	ng	99
69) Carbazole	9.42	167	439141	115.80	ng	100

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05303.D Vial: 6
 Acq On : 3 Aug 2005 10:31 Operator: AHD
 Sample : CAL BNA@120PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:27 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	588405	99.26	ng	99
71) Fluoranthene	10.55	202	481345	113.16	ng	88
73) Pyrene	10.82	202	494469	113.68	ng	83
74) Benzidine	10.73	184	143442	126.28	ng	94
76) Butylbenzylphthalate	11.68	149	254657	121.37	ng	78
77) 3,3'-Dichlorobenzidine	12.32	252	103172	110.63	ng	98
78) Benzo[a]anthracene	12.33	228	415702	116.51	ng	98
79) Chrysene	12.37	228	357197	112.59	ng	99
80) bis(2-Ethylhexyl)phthalate	12.45	149	323842	118.79	ng	94
82) Di-n-octylphthalate	13.32	149	506209	121.44	ng	99
83) Benzo[b]fluoranthene	13.72	252	383401	118.33	ng	96
84) Benzo[k]fluoranthene	13.76	252	290468	101.81	ng	96
85) Benzo[a]pyrene	14.12	252	308441	114.85	ng	99
86) Indeno[1,2,3-cd]pyrene	15.43	276	305592	124.82	ng	91
87) Dibenzo[a,h]anthracene	15.46	278	247734	123.66	ng	94
88) Benzo[g,h,i]perylene	15.72	276	249663	127.41	ng	96

(#) = qualifier out of range (m) = manual integration

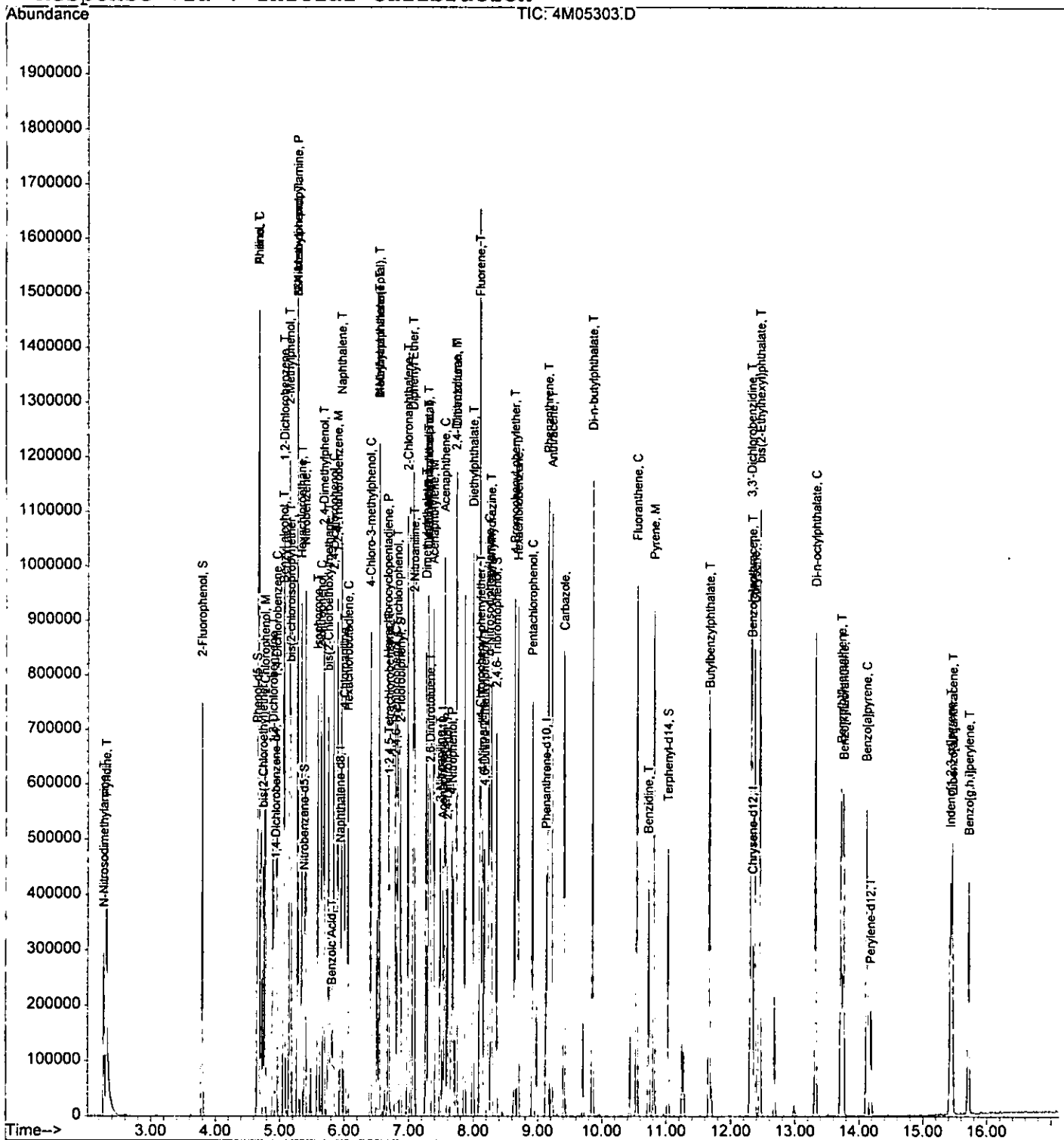
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05303.D Vial: 6
Acq On : 3 Aug 2005 10:31 Operator: AHD
Sample : CAL BNA@120PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 11:27 2005

6755

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05304.D Vial: 7
 Acq On : 3 Aug 2005 10:55 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	40792	40.00	ng	0.00
19) Naphthalene-d8	5.94	136	142226	40.00	ng	0.00
35) Acenaphthene-d10	7.53	164	86087	40.00	ng	0.00
59) Phenanthrene-d10	9.13	188	163050	40.00	ng	0.00
72) Chrysene-d12	12.34	240	110455	40.00	ng	0.00
81) Perylene-d12	14.19	264	70365	40.00	ng	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)	Recovery
4) 2-Fluorophenol	3.80	112	193404	166.85	ng	0.00	83.43%
Spiked Amount 200.000							
7) Phenol-d5	4.66	99	243184	156.19	ng	0.00	78.10%
Spiked Amount 200.000							
20) Nitrobenzene-d5	5.39	128	56513	79.01	ng	0.00	79.01%
Spiked Amount 100.000							
40) 2-Fluorobiphenyl	6.87	172	198110	69.65	ng	0.00	69.65%
Spiked Amount 100.000							
62) 2,4,6-Tribromophenol	8.37	332	110701	149.47	ng	0.00	74.74%
Spiked Amount 200.000							
75) Terphenyl-d14	11.03	244	241751	78.17	ng	0.00	78.17%
Spiked Amount 100.000							

Target Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)	Qvalue
2) Pyridine	2.32	79	268527	171.24	ng	98	98
3) N-Nitrosodimethylamine	2.28	74	148938	170.95	ng	96	96
5) Aniline	4.68	93	284099	168.07	ng	49	49
6) bis(2-Chloroethyl)ether	4.75	93	199542	155.26	ng	97	97
8) Phenol	4.67	94	234696	139.25	ng	96	96
9) 2-Chlorophenol	4.78	128	204098	158.02	ng	85	85
10) 1,3-Dichlorobenzene	4.90	146	215307	155.38	ng	98	98
11) 1,4-Dichlorobenzene	4.96	146	211667	155.74	ng	98	98
12) 1,2-Dichlorobenzene	5.07	146	168640	129.08	ng	98	98
13) Benzyl alcohol	5.07	108	128406	158.66	ng	78	78
14) bis(2-chloroisopropyl)ethe	5.18	45	503118	161.30	ng	99	99
15) 2-Methylphenol	5.15	108	151504	138.39	ng	98	98
16) Hexachloroethane	5.34	117	92445	144.78	ng	99	99
17) N-Nitroso-di-n-propylamine	5.29	70	177850	158.63	ng	85	85
18) 3&4-Methylphenol	5.29	108	167473	150.13	ng	98	98
21) Nitrobenzene	5.40	77	202908	132.55	ng	88	88
22) Isophorone	5.59	82	452112	159.18	ng	95	95
23) 2-Nitrophenol	5.65	139	116626	150.21	ng	98	98
24) 2,4-Dimethylphenol	5.69	107	198361	137.34	ng	95	95

(#) = qualifier out of range (m) = manual integration

h2811

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05304.D Vial: 7
 Acq On : 3 Aug 2005 10:55 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.83	105	43024	190.94	ng	95
26) bis(2-Chloroethoxy)methane	5.77	93	267640	155.76	ng	99
27) 2,4-Dichlorophenol	5.84	162	164714	138.18	ng	98
28) 1,2,4-Trichlorobenzene	5.90	180	179648	140.92	ng	97
29) Naphthalene	5.96	128	428360	132.51	ng	99
30) 4-Chloroaniline	6.01	127	157833	129.94	ng	100
31) Hexachlorobutadiene	6.06	225	120957	148.05	ng	98
32) 4-Chloro-3-methylphenol	6.41	107	195809	149.54	ng	90
33) 2-Methylnaphthalene	6.55	142	302128	133.38	ng	96
34) Methylnaphthalene (Total)	6.55	142	302128	133.38	ng	96
36) 1,2,4,5-Tetrachlorobenzene	6.70	216	197469	136.05	ng	98
37) Hexachlorocyclopentadiene	6.69	237	158016	156.52	ng	95
38) 2,4,6-Trichlorophenol	6.80	196	145976	140.31	ng	97
39) 2,4,5-Trichlorophenol	6.83	196	151856	142.86	ng	98
41) 2-Chloronaphthalene	6.99	162	349123	139.94	ng	94
42) 2-Nitroaniline	7.10	65	223056	153.94	ng	73
43) 1,4-Dimethylnaphthalene	7.31	156	233893	136.64	ng	93
44) Dimethylnaphthalene (Total)	7.31	156	233893	136.64	ng	93
45) Diphenyl Ether	7.08	170	284091	135.07	ng	83
46) Acenaphthylene	7.39	152	516513	135.21	ng	98
47) Dimethylphthalate	7.27	163	456312	147.26	ng	99
48) 2,6-Dinitrotoluene	7.33	165	105977	143.40	ng	77
49) Acenaphthene	7.57	153	350074	140.47	ng	98
50) 3-Nitroaniline	7.50	138	81599	129.17	ng	97
51) 2,4-Dinitrophenol	7.60	184	73583	211.62	ng	74
52) Dibenzofuran	7.74	168	423440	126.70	ng	90
53) 2,4-Dinitrotoluene	7.74	165	128888	133.12	ng	68
54) 4-Nitrophenol	7.67	65	134390	169.44	ng	96
55) Fluorene	8.10	166	307000	125.66	ng	99
56) 4-Chlorophenyl-phenylether	8.11	204	169261	125.16	ng	91
57) Diethylphthalate	8.01	149	478528	147.30	ng	98
58) 4-Nitroaniline	8.15	138	124486	151.34	ng	81
60) 4,6-Dinitro-2-methylphenol	8.18	198	101177	199.01	ng	100
61) n-Nitrosodiphenylamine	8.24	169	270712	127.10	ng	95
63) 1,2-Diphenylhydrazine	8.28	77	542952	143.11	ng	87
64) 4-Bromophenyl-phenylether	8.64	248	146951	136.19	ng	98
65) Hexachlorobenzene	8.70	284	204535	141.04	ng	96
66) Pentachlorophenol	8.92	266	125946	181.47	ng	95
67) Phenanthrene	9.16	178	559269	133.71	ng	99
68) Anthracene	9.22	178	550726	129.07	ng	99
69) Carbazole	9.42	167	514996	131.84	ng	97

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05304.D Vial: 7
 Acq On : 3 Aug 2005 10:55 Operator: AHD
 Sample : CAL BNA@160PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 10:51:44 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	805630	131.94	ng	99
71) Fluoranthene	10.55	202	592322	135.19	ng	94
73) Pyrene	10.82	202	608835	141.81	ng	90
74) Benzidine	10.74	184	172809	154.13	ng	92
76) Butylbenzylphthalate	11.68	149	328092	158.41	ng	90
77) 3,3'-Dichlorobenzidine	12.32	252	121977	132.51	ng	97
78) Benzo[a]anthracene	12.33	228	517393	146.90	ng	99
79) Chrysene	12.38	228	463554	148.02	ng	99
80) bis(2-Ethylhexyl)phthalate	12.45	149	432716	160.80	ng	95
82) Di-n-octylphthalate	13.32	149	635547	168.40	ng	99
83) Benzo[b]fluoranthene	13.72	252	442884	150.96	ng	97
84) Benzo[k]fluoranthene	13.76	252	372061	144.03	ng	99
85) Benzo[a]pyrene	14.13	252	365711	150.41	ng	95
86) Indeno[1,2,3-cd]pyrene	15.43	276	341022	153.84	ng	89
87) Dibenzo[a,h]anthracene	15.46	278	279686	154.19	ng	96
88) Benzo[g,h,i]perylene	15.72	276	271068	152.79	ng	96

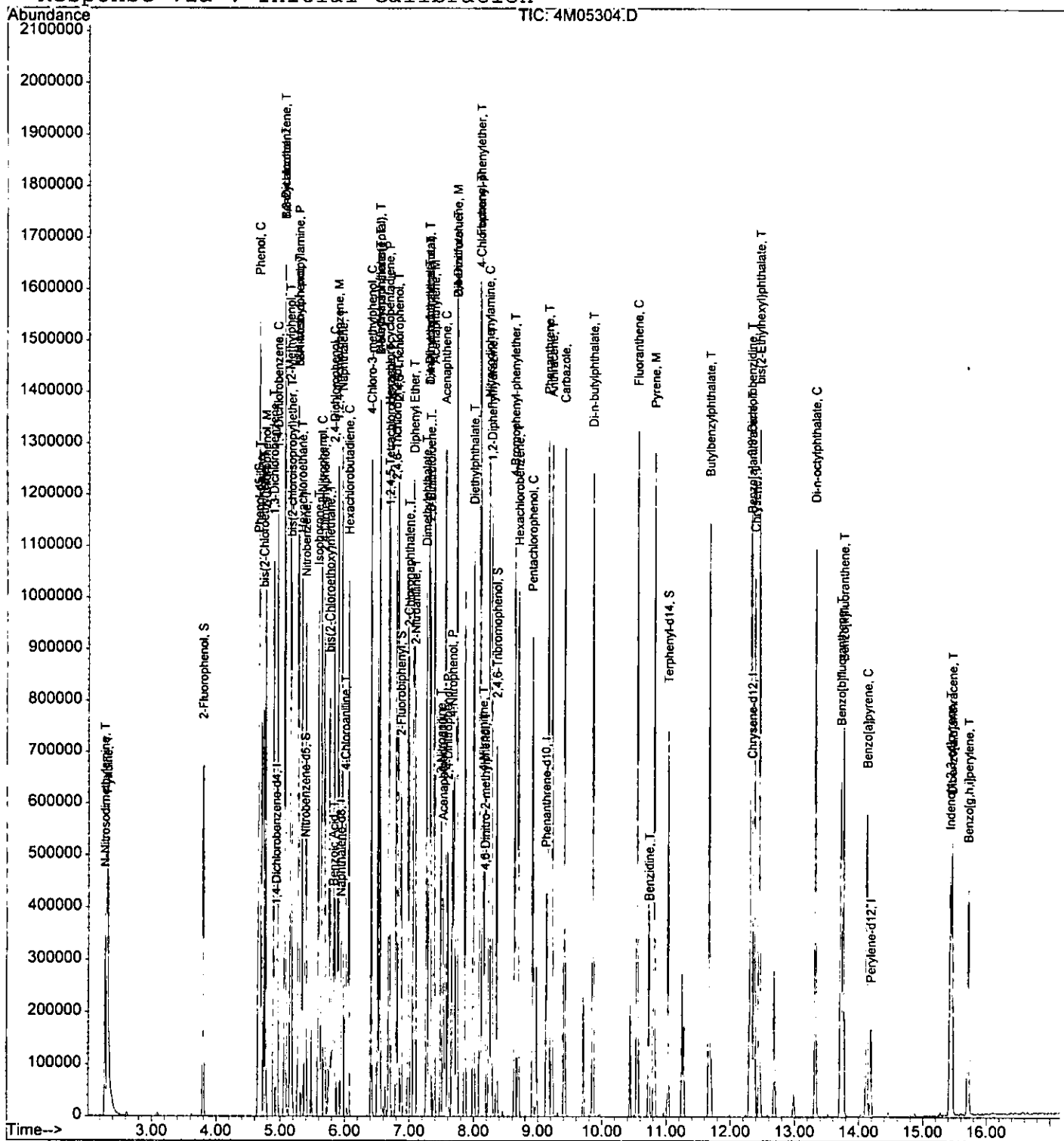
(#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05304.D Vial: 7
Acq On : 3 Aug 2005 10:55 Operator: AHD
Sample : CAL BNA@160PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 11:28 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05305.D Vial: 8
 Acq On : 3 Aug 2005 11:19 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 3 12:15 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 11:27:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	4.94	152	44370	40.00	ng	0.00
19) Naphthalene-d8	5.94	136	145276	40.00	ng	0.00
35) Acenaphthene-d10	7.53	164	91941	40.00	ng	0.00
59) Phenanthrene-d10	9.13	188	168689	40.00	ng	0.00
72) Chrysene-d12	12.34	240	98308	40.00	ng	0.01
81) Perylene-d12	14.19	264	56071	40.00	ng	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)	Recovery
4) 2-Fluorophenol	3.81	112	239180	189.70	ng	0.01	94.85%
Spiked Amount 200.000							
7) Phenol-d5	4.67	99	300243	177.28	ng	0.01	88.64%
Spiked Amount 200.000							
20) Nitrobenzene-d5	5.39	128	70739	96.82	ng	0.01	96.82%
Spiked Amount 100.000							
40) 2-Fluorobiphenyl	6.87	172	239754	78.93	ng	0.00	78.93%
Spiked Amount 100.000							
62) 2,4,6-Tribromophenol	8.37	332	137559	179.52	ng	0.01	89.76%
Spiked Amount 200.000							
75) Terphenyl-d14	11.04	244	287293	104.38	ng	0.01	104.38%
Spiked Amount 100.000							

Target Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)	Qvalue
2) Pyridine	2.32	79	306077	179.45	ng	98	98
3) N-Nitrosodimethylamine	2.29	74	197613	193.22	ng	100	100
5) Aniline	4.69	93	327233	177.98	ng	38	38
6) bis(2-Chloroethyl)ether	4.75	93	227081	162.44	ng	98	98
8) Phenol	4.69	94	324044	176.76	ng	95	95
9) 2-Chlorophenol	4.78	128	246298	175.32	ng	83	83
10) 1,3-Dichlorobenzene	4.90	146	264954	175.79	ng	98	98
11) 1,4-Dichlorobenzene	4.96	146	254561	172.19	ng	99	99
12) 1,2-Dichlorobenzene	5.08	146	189354	133.25	ng	95	95
13) Benzyl alcohol	5.08	108	147574	167.64	ng	74	74
14) bis(2-chloroisopropyl)ethe	5.18	45	604247	178.11	ng	98	98
15) 2-Methylphenol	5.16	108	182284	153.07	ng	99	99
16) Hexachloroethane	5.34	117	113972	164.10	ng	89	89
17) N-Nitroso-di-n-propylamine	5.29	70	197477	161.93	ng	94	94
18) 3&4-Methylphenol	5.29	108	206177	169.92	ng	99	99
21) Nitrobenzene	5.40	77	243861	155.96	ng	89	89
22) Isophorone	5.60	82	569789	196.40	ng	96	96
23) 2-Nitrophenol	5.65	139	145240	183.14	ng	97	97
24) 2,4-Dimethylphenol	5.69	107	255860	173.43	ng	92	92

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Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05305.D Vial: 8
 Acq On : 3 Aug 2005 11:19 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 12:15 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 11:27:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.83	105	45688	178.52	ng	96
26) bis(2-Chloroethoxy)methane	5.77	93	314216	179.03	ng	98
27) 2,4-Dichlorophenol	5.84	162	202995	166.71	ng	96
28) 1,2,4-Trichlorobenzene	5.90	180	222136	170.58	ng	96
29) Naphthalene	5.96	128	505454	153.07	ng	99
30) 4-Chloroaniline	6.01	127	174670	140.78	ng	99
31) Hexachlorobutadiene	6.07	225	143761	172.27	ng	98
32) 4-Chloro-3-methylphenol	6.41	107	250514	187.31	ng	92
33) 2-Methylnaphthalene	6.55	142	349764	151.16	ng	96
34) Methylnaphthalene(Total)	6.55	142	349764	151.16	ng	96
36) 1,2,4,5-Tetrachlorobenzene	6.70	216	241157	155.57	ng	97
37) Hexachlorocyclopentadiene	6.68	237	184176	170.81	ng	97
38) 2,4,6-Trichlorophenol	6.80	196	180363	162.32	ng	95
39) 2,4,5-Trichlorophenol	6.83	196	188314	165.88	ng	99
41) 2-Chloronaphthalene	6.99	162	407102	152.79	ng	95
42) 2-Nitroaniline	7.10	65	276838	178.89	ng	80
43) 1,4-Dimethylnaphthalene	7.31	156	285049	155.93	ng	95
44) Dimethylnaphthalene(Total)	7.31	156	285049	155.93	ng	95
45) Diphenyl Ether	7.07	170	336052	149.60	ng	98
46) Acenaphthylene	7.40	152	636399	155.99	ng	99
47) Dimethylphthalate	7.27	163	577125	174.38	ng	99
48) 2,6-Dinitrotoluene	7.33	165	135733	171.97	ng	78
49) Acenaphthene	7.57	153	431925	162.27	ng	99
50) 3-Nitroaniline	7.50	138	90808	134.60	ng	92
51) 2,4-Dinitrophenol	7.61	184	102289	275.44	ng	87
52) Dibenzofuran	7.74	168	496356	139.06	ng	88
53) 2,4-Dinitrotoluene	7.75	165	175249	169.48	ng	61
54) 4-Nitrophenol	7.68	65	197080	232.66	ng	92
55) Fluorene	8.10	166	372934	142.93	ng	100
56) 4-Chlorophenyl-phenylether	8.11	204	195906	135.64	ng	91
57) Diethylphthalate	8.01	149	579412	167.00	ng	99
58) 4-Nitroaniline	8.16	138	161988	184.39	ng	79
60) 4,6-Dinitro-2-methylphenol	8.18	198	122882	203.63	ng	100
61) n-Nitrosodiphenylamine	8.24	169	351406	159.48	ng	97
63) 1,2-Diphenylhydrazine	8.29	77	660144	168.19	ng	87
64) 4-Bromophenyl-phenylether	8.64	248	187050	167.55	ng	98
65) Hexachlorobenzene	8.70	284	250158	166.73	ng	94
66) Pentachlorophenol	8.92	266	163353	205.68	ng	96
67) Phenanthrene	9.16	178	706348	163.23	ng	100
68) Anthracene	9.23	178	692541	156.88	ng	100
69) Carbazole	9.42	167	648285	160.42	ng	97

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05305.D Vial: 8
 Acq On : 3 Aug 2005 11:19 Operator: AHD
 Sample : CAL BNA@200PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 3 12:15 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 11:27:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.86	149	976894	154.64	ng	99
71) Fluoranthene	10.56	202	696818	153.72	ng	95
73) Pyrene	10.82	202	708018	185.28	ng	92
74) Benzidine	10.73	184	208054	194.29	ng	93
76) Butylbenzylphthalate	11.68	149	368550	199.93	ng	86
77) 3,3'-Dichlorobenzidine	12.32	252	121437	148.22	ng	98
78) Benzo[a]anthracene	12.33	228	558194	178.07	ng	99
79) Chrysene	12.39	228	514143	184.46	ng	99
80) bis(2-Ethylhexyl)phthalate	12.46	149	482403	201.41	ng	92
82) Di-n-octylphthalate	13.33	149	691110	229.80	ng	99
83) Benzo[b]fluoranthene	13.72	252	460181	196.85	ng	96
84) Benzo[k]fluoranthene	13.77	252	357910m	173.88	ng	
85) Benzo[a]pyrene	14.12	252	359176	185.38	ng	99
86) Indeno[1,2,3-cd]pyrene	15.43	276	315081	178.37	ng	87
87) Dibenzo[a,h]anthracene	15.46	278	259370	179.44	ng	94
88) Benzo[g,h,i]perylene	15.72	276	246975	174.69	ng	95

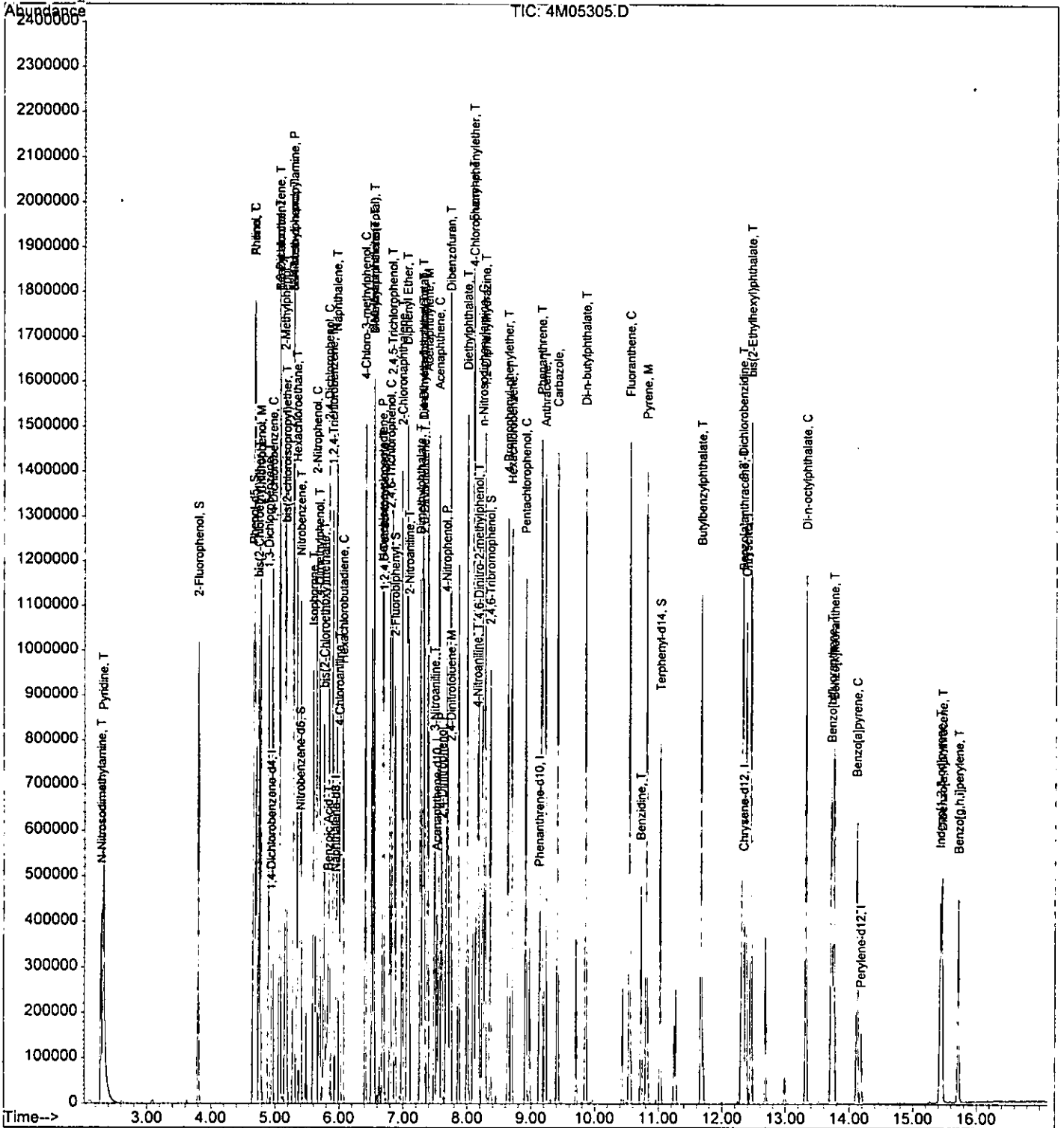
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05305.D Vial: 8
Acq On : 3 Aug 2005 11:19 Operator: AHD
Sample : CAL BNA@200PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 3 12:15 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/4/05 6:44:00 AM

Data File: 5M09736.D
Method: 8270

Instrument: GCMS_5

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
1,4-Dichlorobenzene-d4	1	0	I	5.11	40.00	40			0.000	0.00		
Pyridine	1	0		1.98	50.55	50			1.668	1.686	1.10	
N-Nitrosodimethylamine	1	0		2.02	1.63	50			1.014	0.033	96.74	
2-Fluorophenol	1	0	S	3.79	49.44	50			1.347	1.332	1.12	
Aniline	1	0		4.82	44.87	50			2.261	2.029	10.26	
Pentachloroethane	1	0		4.85	52.46	50			0.493	0.517	4.92	
bis(2-Chloroethyl)ether	1	0		4.89	49.30	50			1.415	1.395	1.40	
Phenol-d5	1	0	S	4.81	43.95	50			1.970	1.732	12.10	
Phenol	1	0	CC	4.82	47.73	50	20		2.088	1.993	4.54	
2-Chlorophenol	1	0		4.92	45.31	50			1.585	1.436	9.38	
1,3-Dichlorobenzene	1	0		5.05	51.23	50			1.466	1.502	2.46	
1,4-Dichlorobenzene	1	0	CC	5.12	50.79	50	20		1.501	1.524	1.58	
1,2-Dichlorobenzene	1	0		5.25	50.30	50			1.431	1.439	0.60	
Benzyl alcohol	1	0		5.24	45.05	50			1.047	0.943	9.90	
bis(2-chloroisopropyl)ether	1	0		5.37	49.88	50			2.150	2.145	0.24	
2-Methylphenol	1	0		5.35	45.61	50			1.447	1.320	8.78	
Hexachloroethane	1	0		5.53	51.57	50			0.622	0.642	3.14	
N-Nitroso-di-n-propylamine	1	0	CP	5.47	45.25	50	0.05		1.141	1.032	9.50	
3&4-Methylphenol	1	0		5.48	45.39	50			1.538	1.396	9.22	
Naphthalene-d8	1	0	I	6.14	40.00	40				0.000	0.00	
Nitrobenzene-d5	1	0	S	5.58	25.19	25			0.175	0.176	0.76	
Nitrobenzene	1	0		5.60	52.44	50			0.393	0.412	4.88	
Isophorone	1	0		5.80	48.26	50			0.731	0.706	3.48	
2-Nitrophenol	1	0	CC	5.85	51.26	50	20		0.202	0.207	2.52	
2,4-Dimethylphenol	1	0		5.90	48.30	50			0.383	0.370	3.40	
Benzoic Acid	1	0		5.99	45.30	50			0.220	0.199	9.40	
bis(2-Chloroethoxy)methane	1	0		5.98	51.04	50			0.418	0.427	2.08	
2,4-Dichlorophenol	1	0	CC	6.04	48.76	50	20		0.320	0.312	2.48	
1,2,4-Trichlorobenzene	1	0		6.11	52.07	50			0.328	0.341	4.14	
Naphthalene	1	0		6.16	50.75	50			1.048	1.064	1.50	
4-Chloroaniline	1	0		6.21	46.26	50			0.409	0.379	7.48	
Hexachlorobutadiene	1	0	CC	6.26	51.21	50	20		0.181	0.185	2.42	
4-Chloro-3-methylphenol	1	0	CC	6.58	47.39	50	20		0.353	0.335	5.22	
2-Methylnaphthalene	1	0		6.68	48.80	50			0.724	0.707	2.40	
Methylnaphthalenes	1	0		6.68	48.80							
Acenaphthene-d10	1	0	I	7.48	40.00	40				0.000	0.00	
1,2,4,5-Tetrachlorobenzene	1	0		6.80	50.13	50			0.540	0.542	0.26	
Hexachlorocyclopentadiene	1	0	CP	6.79	52.75	50	0.05		0.333	0.351	5.50	
2,4,6-Trichlorophenol	1	0	CC	6.89	49.28	50	20		0.389	0.384	1.44	
2,4,5-Trichlorophenol	1	0		6.92	50.18	50			0.423	0.425	0.36	
2-Fluorobiphenyl	1	0	S	6.95	26.44	25			1.250	1.323	5.76	
2-Chloronaphthalene	1	0		7.04	49.69	50			1.131	1.124	0.62	
1,4-Dimethylnaphthalene	1	0		7.30	49.64	50			0.860	0.854	0.72	
Dimethylnaphthalenes	1	0		7.30	49.64							
Diphenyl Ether	1	0		7.11	67.95	50			0.747	1.015	35.90	
2-Nitroaniline	1	0		7.13	55.58	50			0.431	0.479	11.16	
Acenaphthylene	1	0		7.37	51.52	50			1.786	1.841	3.04	
Dimethylphthalate	1	0		7.27	48.57	50			1.305	1.268	2.86	
2,6-Dinitrotoluene	1	0		7.32	47.54	50			0.301	0.286	4.92	
Acenaphthene	1	0	CC	7.51	49.04	50	20		1.105	1.084	1.92	
3-Nitroaniline	1	0		7.45	50.37	50			0.328	0.331	0.74	
2,4-Dinitrophenol	1	0	CP	7.54	48.96	50	0.05		0.185	0.181	2.08	
Dibenzofuran	1	0		7.65	49.97	50			1.608	1.607	0.06	
2,4-Dinitrotoluene	1	0		7.64	47.65	50			0.415	0.396	4.70	
4-Nitrophenol	1	0	CP	7.59	52.50	50	0.05		0.261	0.274	5.00	
2,3,4,6-Tetrachlorophenol	1	0		7.76	46.20	50			0.332	0.307	7.60	
Fluorene	1	0		7.95	47.27	50			1.296	1.225	5.46	
4-Chlorophenyl-phenylether	1	0		7.96	46.93	50			0.630	0.591	6.14	
Diethylphthalate	1	0		7.86	47.09	50			1.338	1.260	5.82	
4-Nitroaniline	1	0		7.97	46.29	50			0.381	0.352	7.42	
Phenanthrene-d10	1	0	I	8.85	40.00	40				0.000	0.00	
4,6-Dinitro-2-methylphenol	1	0		8.00	47.57	50			0.154	0.146	4.86	
n-Nitrosodiphenylamine	1	0	CC	8.07	47.89	50	20		0.551	0.528	4.22	
2,4,6-Tribromophenol	1	0	S	8.17	46.73	50			0.086	0.080	6.54	
1,2-Diphenylhydrazine	1	0		8.10	52.67	50			0.787	0.829	5.34	
4-Bromophenyl-phenylether	1	0		8.41	48.99	50			0.205	0.201	2.02	
Hexachlorobenzene	1	0		8.46	48.66	50			0.194	0.188	2.68	
gamma-BHC	1	0		8.71	10.08	10			0.142	0.143	0.80	
Pentachlorophenol	1	0	CC	8.66	50.15	50	20		0.127	0.127	0.30	
Phenanthrene	1	0		8.87	49.09	50			1.154	1.133	1.82	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 1 of 2

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Form7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/4/05 6:44:00 AM

Data File: 5M09736.D
Method: 8270

Instrument: GCMS_5

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
Anthracene	1	0		8.93	49.96	50			1.172	1.172	0.08	
Carbazole	1	0		9.10	50.16	50			1.070	1.074	0.32	
Heptachlor	1	0		9.37	10.98	10			0.150	0.164	9.80	
Di-n-butylphthalate	1	0		9.51	49.65	50			1.296	1.287	0.70	
Heptachlor_epoxide	1	0		10.05	10.42	10			0.104	0.108	4.20	
Fluoranthene	1	0	CC	10.15	46.08	50	20		1.258	1.159	7.84	
Chrysene-d12	1	0	I	11.83	40.00	40				0.000	0.00	
Pyrene	1	0		10.41	48.21	50			1.602	1.545	3.58	
Benzidine	1	0		10.33	38.32	50			0.592	0.453	23.36	
Terphenyl-d14	1	0	S	10.62	24.46	25			0.945	0.925	2.16	
Endrin	1	0		10.85	10.89	10			0.079	0.086	8.90	
Butylbenzylphthalate	1	0		11.23	50.07	50			0.705	0.706	0.14	
Methoxychlor	1	0		11.85	9.30	10			0.735	0.684	7.00	
3,3'-Dichlorobenzidine	1	0		11.82	40.50	50			0.460	0.373	19.00	
Benzo[a]anthracene	1	0		11.82	48.05	50			1.471	1.413	3.90	
Chrysene	1	0		11.85	46.09	50			1.349	1.243	7.82	
bis(2-Ethylhexyl)phthalate	1	0		11.94	50.83	50			0.974	0.990	1.66	
Perylene-d12	1	0	I	13.41	40.00	40				0.000	0.00	
Di-n-octylphthalate	1	0	CC	12.69	50.50	50	20		2.190	2.212	1.00	
Benzo[b]fluoranthene	1	0		13.02	51.01	50			1.579	1.611	2.02	
Benzo[k]fluoranthene	1	0		13.05	47.12	50			1.599	1.506	5.76	
Benzo[a]pyrene	1	0	CC	13.35	50.05	50	20		1.486	1.488	0.10	
Indeno[1,2,3-cd]pyrene	1	0		14.44	51.44	50			1.610	1.657	2.88	
Dibenzo[a,h]anthracene	1	0		14.46	51.16	50			1.336	1.367	2.32	
Benzo[g,h,i]perylene	1	0		14.71	51.67	50			1.347	1.392	3.34	
Diaminotoluene Dihydrochloride	1	1E		0.00	0.00	50				0.000	100.00	
Toluene Diisocyanate	1	1E		0.00	0.00	50				0.000	100.00	
2,4 Diaminotoluene	1	1E		0.00	0.00	50				0.000	100.00	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 2 of 2

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF.

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09736.D Vial: 2
 Acq On : 4 Aug 2005 6:44 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 4 7:10 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.11	152	20348	40.00	ng	-0.14
20) Naphthalene-d8	6.14	136	77780	40.00	ng	-0.14
36) Acenaphthene-d10	7.48	164	45021	40.00	ng	-0.15
61) Phenanthrene-d10	8.85	188	77436	40.00	ng	-0.18
77) Chrysene-d12	11.83	240	61871	40.00	ng	-0.21
88) Perylene-d12	13.41	264	46212	40.00	ng	-0.21
System Monitoring Compounds						
4) 2-Fluorophenol	3.79	112	33880	49.44	ng	-0.18
Spiked Amount	200.000		Recovery	=	24.72%	
8) Phenol-d5	4.81	99	44045	43.95	ng	-0.14
Spiked Amount	200.000		Recovery	=	21.98%	
21) Nitrobenzene-d5	5.58	128	8580	25.19	ng	-0.14
Spiked Amount	100.000		Recovery	=	25.19%	
41) 2-Fluorobiphenyl	6.95	172	37213	26.44	ng	-0.14
Spiked Amount	100.000		Recovery	=	26.44%	
64) 2,4,6-Tribromophenol	8.17	330	7744	46.73	ng	-0.17
Spiked Amount	200.000		Recovery	=	23.37%	
80) Terphenyl-d14	10.62	244	35755	24.46	ng	-0.19
Spiked Amount	100.000		Recovery	=	24.46%	
Target Compounds						
2) Pyridine	1.98	79	42892	50.55	ng	90
3) N-Nitrosodimethylamine	2.02	74	840	1.63	ng	62
5) Aniline	4.82	93	51611	44.87	ng	94
6) Pentachloroethane	4.85	117	13160	52.46	ng	100
7) bis(2-Chloroethyl)ether	4.89	93	35492	49.30	ng	95
9) Phenol	4.82	94	50694	47.73	ng	87
10) 2-Chlorophenol	4.92	128	36528	45.31	ng	97
11) 1,3-Dichlorobenzene	5.05	146	38213	51.23	ng	99
12) 1,4-Dichlorobenzene	5.12	146	38773	50.79	ng	99
13) 1,2-Dichlorobenzene	5.25	146	36610	50.30	ng	96
14) Benzyl alcohol	5.24	108	23985	45.05	ng	99
15) bis(2-chloroisopropyl)ethe	5.37	45	54557	49.88	ng	96
16) 2-Methylphenol	5.35	108	33574	45.61	ng	99
17) Hexachloroethane	5.53	117	16329	51.57	ng	89
18) N-Nitroso-di-n-propylamine	5.47	70	26261	45.25	ng	99
19) 3&4-Methylphenol	5.48	108	35519	45.39	ng	97
22) Nitrobenzene	5.60	77	40054	52.44	ng	99
23) Isophorone	5.80	82	68622	48.26	ng	97
24) 2-Nitrophenol	5.85	139	20095	51.26	ng	99

(#) = qualifier out of range (m) = manual integration

h811

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09736.D Vial: 2
 Acq On : 4 Aug 2005 6:44 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 4 7:10 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.90	107	36013	48.30	ng	98
26) Benzoic Acid	5.99	105	19393	45.30	ng	96
27) bis(2-Chloroethoxy)methane	5.98	93	41524	51.04	ng	99
28) 2,4-Dichlorophenol	6.04	162	30317	48.76	ng	96
29) 1,2,4-Trichlorobenzene	6.11	180	33194	52.07	ng	99
30) Naphthalene	6.16	128	103429	50.75	ng	99
31) 4-Chloroaniline	6.21	127	36815	46.26	ng	100
32) Hexachlorobutadiene	6.26	225	18002	51.21	ng	99
33) 4-Chloro-3-methylphenol	6.58	107	32542	47.39	ng	99
34) 2-Methylnaphthalene	6.68	142	68701	48.80	ng	99
35) Methylnaphthalenes (Total)	6.68	142	68701	48.80	ng	99
37) 1,2,4,5-Tetrachlorobenzene	6.80	216	30477	50.13	ng	98
38) Hexachlorocyclopentadiene	6.79	237	19752	52.75	ng	99
39) 2,4,6-Trichlorophenol	6.89	196	21596	49.28	ng	98
40) 2,4,5-Trichlorophenol	6.92	196	23913	50.18	ng	99
42) 2-Chloronaphthalene	7.04	162	63234	49.69	ng	99
43) 1,4-Dimethylnaphthalene	7.30	156	48071	49.64	ng	99
44) Dimethylnaphthalenes (Total)	7.30	156	48071	49.64	ng	99
45) Diphenyl Ether	7.11	170	57133	67.95	ng	90
46) 2-Nitroaniline	7.13	65	26938	55.58	ng	94
47) Acenaphthylene	7.37	152	103584	51.52	ng	100
48) Dimethylphthalate	7.27	163	71348	48.57	ng	100
49) 2,6-Dinitrotoluene	7.32	165	16083	47.54	ng	98
50) Acenaphthene	7.51	153	60981	49.04	ng	99
51) 3-Nitroaniline	7.45	138	18605	50.37	ng	100
52) 2,4-Dinitrophenol	7.54	184	10192	48.96	ng	93
53) Dibenzofuran	7.65	168	90442	49.97	ng	100
54) 2,4-Dinitrotoluene	7.64	165	22273	47.65	ng	92
55) 4-Nitrophenol	7.59	65	15436	52.50	ng	96
56) 2,3,4,6-Tetrachlorophenol	7.76	232	17274	46.20	ng	99
57) Fluorene	7.95	166	68930	47.27	ng	100
58) 4-Chlorophenyl-phenylether	7.96	204	33268	46.93	ng	99
59) Diethylphthalate	7.86	149	70897	47.09	ng	97
60) 4-Nitroaniline	7.97	138	19834	46.29	ng	96
62) 4,6-Dinitro-2-methylphenol	8.00	198	14162	47.57	ng	100
63) n-Nitrosodiphenylamine	8.07	169	51126	47.89	ng	97
65) 1,2-Diphenylhydrazine	8.10	77	80229	52.67	ng	96
66) 4-Bromophenyl-phenylether	8.41	248	19452	48.99	ng	97
67) Hexachlorobenzene	8.46	284	18240	48.66	ng	89
68) gamma-BHC	8.71	181	2766	10.08	ng	96
69) Pentachlorophenol	8.66	266	12323	50.15	ng	94

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09736.D Vial: 2
 Acq On : 4 Aug 2005 6:44 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 4 7:10 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.87	178	109626	49.09	ng	98
71) Anthracene	8.93	178	113403	49.96	ng	99
72) Carbazole	9.10	167	103947	50.16	ng	99
73) Heptachlor	9.37	100	3184	10.98	ng	97
74) Di-n-butylphthalate	9.51	149	124580	49.65	ng	100
75) Heptachlor epoxide	10.05	81	2099	10.42	ng	94
76) Fluoranthene	10.15	202	112187	46.08	ng	98
78) Pyrene	10.41	202	119473	48.21	ng	98
79) Benzidine	10.33	184	35064	38.32	ng	99
81) Endrin	10.85	81	1329	10.89	ng	93
82) Butylbenzylphthalate	11.23	149	54600	50.07	ng	97
83) Methoxychlor	11.85	227	10579	9.30	ng	100
84) 3,3'-Dichlorobenzidine	11.82	252	28844	40.50	ng	97
85) Benzo[a]anthracene	11.82	228	109308	48.05	ng	99
86) Chrysene	11.85	228	96161	46.09	ng	99
87) bis(2-Ethylhexyl)phthalate	11.94	149	76599	50.83	ng	95
89) Di-n-octylphthalate	12.69	149	127777	50.50	ng	98
90) Benzo[b]fluoranthene	13.02	252	93071	51.01	ng	98
91) Benzo[k]fluoranthene	13.05	252	87017	47.12	ng	95
92) Benzo[a]pyrene	13.35	252	85938	50.05	ng	99
93) Indeno[1,2,3-cd]pyrene	14.44	276	95694	51.44	ng	85
94) Dibenzo[a,h]anthracene	14.46	278	78992	51.16	ng	98
95) Benzo[g,h,i]perylene	14.71	276	80392	51.67	ng	92

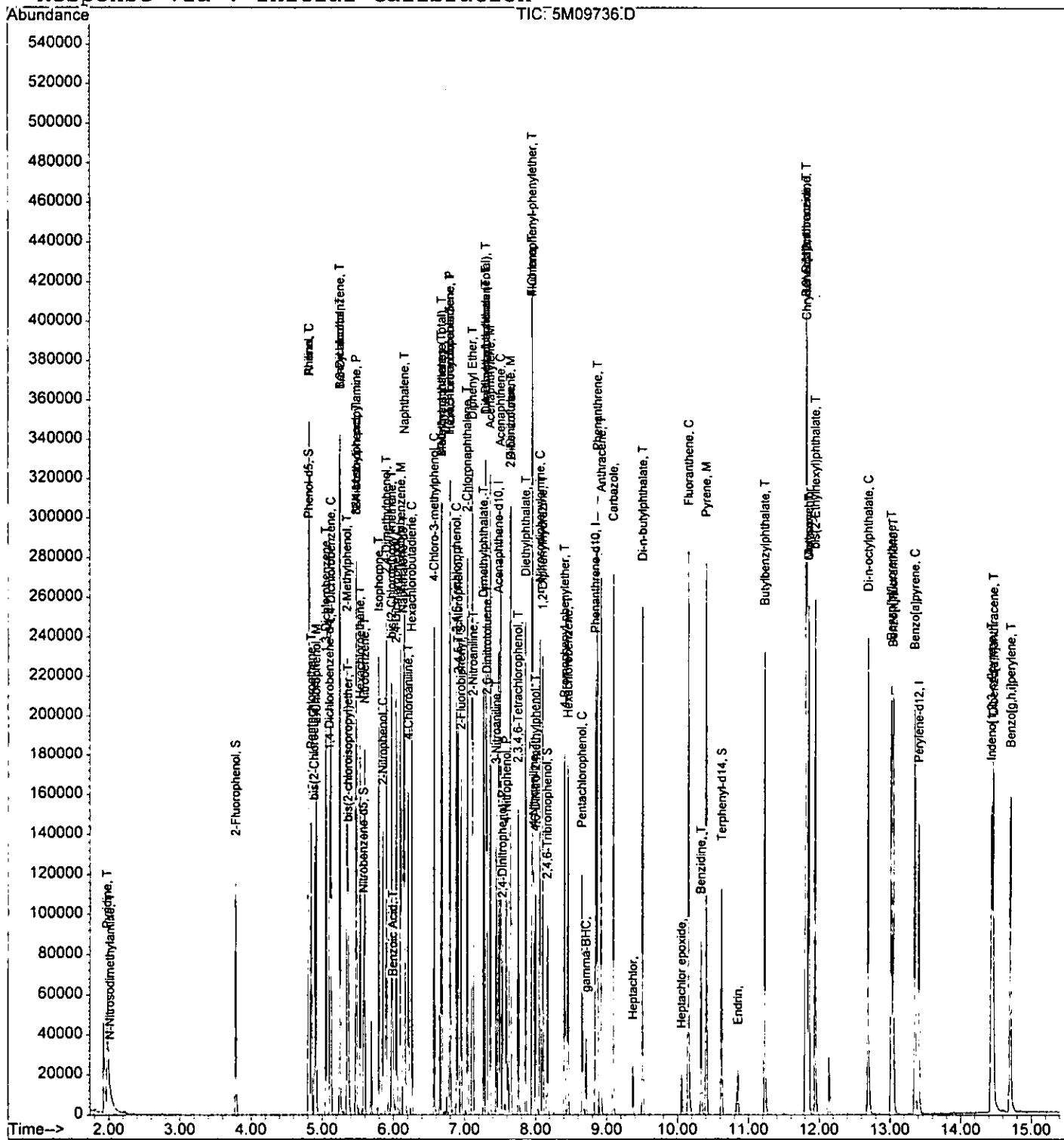
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09736.D Vial: 2
Acq On : 4 Aug 2005 6:44 Operator: AHD
Sample : CAL BNA@50PPM Inst : GCMS_5
Misc : A,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 4 7:10 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/5/05 6:42:00 AM

Data File: 5M09779.D
Method: 8270

Instrument: GCMS_5

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
1,4-Dichlorobenzene-d4	1	0	I	5.10	40.00	40				0.000	0.00	
Pyridine	1	0		1.97	48.38	50			1.668	1.614	3.24	
N-Nitrosodimethylamine	1	0		1.92	46.19	50			1.014	0.937	7.62	
2-Fluorophenol	1	0	S	3.78	48.00	50			1.347	1.293	4.00	
Aniline	1	0		4.81	45.79	50			2.261	2.071	8.42	
Pentachloroethane	1	0		4.84	50.80	50			0.493	0.501	1.60	
bis(2-Chloroethyl)ether	1	0		4.89	50.33	50			1.415	1.425	0.66	
Phenol-d5	1	0	S	4.80	43.95	50			1.970	1.732	12.10	
Phenol	1	0	CC	4.81	45.96	50	20		2.088	1.919	8.08	
2-Chlorophenol	1	0		4.91	45.40	50			1.585	1.439	9.20	
1,3-Dichlorobenzene	1	0		5.05	52.12	50			1.466	1.529	4.24	
1,4-Dichlorobenzene	1	0	CC	5.12	51.33	50	20		1.501	1.541	2.66	
1,2-Dichlorobenzene	1	0		5.25	50.89	50			1.431	1.456	1.78	
Benzyl alcohol	1	0		5.24	46.53	50			1.047	0.974	6.94	
bis(2-chloroisopropyl)ether	1	0		5.36	52.05	50			2.150	2.238	4.10	
2-Methylphenol	1	0		5.34	49.97	50			1.447	1.446	0.06	
Hexachloroethane	1	0		5.52	54.55	50			0.622	0.679	9.10	
N-Nitroso-di-n-propylamine	1	0	CP	5.46	49.33	50	0.05		1.141	1.125	1.34	
3&4-Methylphenol	1	0		5.47	48.00	50			1.538	1.477	4.00	
Naphthalene-d8	1	0	I	6.14	40.00	40				0.000	0.00	
Nitrobenzene-d5	1	0	S	5.58	23.99	25			0.175	0.168	4.04	
Nitrobenzene	1	0		5.59	50.50	50			0.393	0.397	1.00	
Isophorone	1	0		5.78	49.15	50			0.731	0.719	1.70	
2-Nitrophenol	1	0	CC	5.84	51.04	50	20		0.202	0.206	2.08	
2,4-Dimethylphenol	1	0		5.90	48.67	50			0.383	0.373	2.66	
Benzoic Acid	1	0		5.98	35.96	50			0.220	0.158	28.08	
bis(2-Chloroethoxy)methane	1	0		5.97	48.33	50			0.418	0.404	3.34	
2,4-Dichlorophenol	1	0	CC	6.04	46.88	50	20		0.320	0.300	6.24	
1,2,4-Trichlorobenzene	1	0		6.09	51.24	50			0.328	0.336	2.48	
Naphthalene	1	0		6.15	50.95	50			1.048	1.068	1.90	
4-Chloroaniline	1	0		6.20	50.29	50			0.409	0.412	0.58	
Hexachlorobutadiene	1	0	CC	6.25	49.75	50	20		0.181	0.180	0.50	
4-Chloro-3-methylphenol	1	0	CC	6.57	47.30	50	20		0.353	0.334	5.40	
2-Methylnaphthalene	1	0		6.67	47.66	50			0.724	0.690	4.68	
Methylnaphthalenes	1	0		6.67	47.66							
Acenaphthene-d10	1	0	I	7.47	40.00	40				0.000	0.00	
1,2,4,5-Tetrachlorobenzene	1	0		6.79	51.15	50			0.540	0.553	2.30	
Hexachlorocyclopentadiene	1	0	CP	6.79	55.61	50	0.05		0.333	0.370	11.22	
2,4,6-Trichlorophenol	1	0	CC	6.89	49.79	50	20		0.389	0.388	0.42	
2,4,5-Trichlorophenol	1	0		6.91	52.09	50			0.423	0.441	4.18	
2-Fluorobiphenyl	1	0	S	6.95	26.26	25			1.250	1.313	5.04	
2-Chloronaphthalene	1	0		7.03	53.47	50			1.131	1.209	6.94	
1,4-Dimethylnaphthalene	1	0		7.30	52.10	50			0.860	0.897	4.20	
Dimethylnaphthalenes	1	0		7.30	52.10							
Diphenyl Ether	1	0		7.10	71.06	50			0.747	1.062	42.12	
2-Nitroaniline	1	0		7.11	57.16	50			0.431	0.492	14.32	
Acenaphthylene	1	0		7.36	51.96	50			1.786	1.856	3.92	
Dimethylphthalate	1	0		7.26	51.05	50			1.305	1.333	2.10	
2,6-Dinitrotoluene	1	0		7.31	50.91	50			0.301	0.306	1.82	
Acenaphthene	1	0	CC	7.50	53.72	50	20		1.105	1.187	7.44	
3-Nitroaniline	1	0		7.44	54.69	50			0.328	0.359	9.38	
2,4-Dinitrophenol	1	0	CP	7.53	46.65	50	0.05		0.185	0.173	6.70	
Dibenzofuran	1	0		7.64	53.76	50			1.608	1.729	7.52	
2,4-Dinitrotoluene	1	0		7.64	51.00	50			0.415	0.424	2.00	
4-Nitrophenol	1	0	CP	7.58	53.76	50	0.05		0.261	0.281	7.52	
2,3,4,6-Tetrachlorophenol	1	0		7.75	50.18	50			0.332	0.333	0.36	
Fluorene	1	0		7.94	49.48	50			1.296	1.282	1.04	
4-Chlorophenyl-phenylether	1	0		7.95	50.24	50			0.630	0.633	0.48	
Diethylphthalate	1	0		7.85	50.17	50			1.338	1.342	0.34	
4-Nitroaniline	1	0		7.96	46.01	50			0.381	0.350	7.98	
Phenanthrene-d10	1	0	I	8.84	40.00	40				0.000	0.00	
4,6-Dinitro-2-methylphenol	1	0		7.99	46.35	50			0.154	0.143	7.30	
n-Nitrosodiphenylamine	1	0	CC	8.06	47.71	50	20		0.551	0.526	4.58	
2,4,6-Tribromophenol	1	0	S	8.16	47.62	50			0.086	0.082	4.76	
1,2-Diphenylhydrazine	1	0		8.09	50.91	50			0.787	0.801	1.82	
4-Bromophenyl-phenylether	1	0		8.40	47.96	50			0.205	0.197	4.08	
Hexachlorobenzene	1	0		8.45	48.28	50			0.194	0.187	3.44	
gamma-BHC	1	0		8.71	9.49	10			0.142	0.134	5.10	
Pentachlorophenol	1	0	CC	8.65	45.22	50	20		0.127	0.115	9.56	
Phenanthrene	1	0		8.86	48.55	50			1.154	1.120	2.90	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 1 of 2

** - No limit specified in method

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/5/05 6:42:00 AM

Data File: 5M09779.D
Method: 8270

Instrument: GCMS_5

79554

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
Anthracene	1	0		8.92	47.92	50			1.172	1.124	4.16	
Carbazole	1	0		9.09	48.09	50			1.070	1.029	3.82	
Heptachlor	1	0		9.36	9.52	10			0.150	0.143	4.80	
Di-n-butylphthalate	1	0		9.50	47.88	50			1.296	1.241	4.24	
Heptachlor epoxide	1	0		10.05	9.91	10			0.104	0.103	0.90	
Fluoranthene	1	0	CC	10.14	46.62	50	20		1.258	1.173	6.76	
Chrysene-d12	1	0	I	11.82	40.00	40				0.000	0.00	
Pyrene	1	0		10.40	50.34	50			1.602	1.613	0.68	
Benzidine	1	0		10.33	44.85	50			0.592	0.531	10.30	
Terphenyl-d14	1	0	S	10.61	24.43	25			0.945	0.923	2.28	
Endrin	1	0		10.84	9.50	10			0.079	0.075	5.00	
Butylbenzylphthalate	1	0		11.22	49.86	50			0.705	0.703	0.28	
Methoxychlor	1	0		11.85	9.47	10			0.735	0.697	5.30	
3,3'-Dichlorobenzidine	1	0		11.81	46.04	50			0.460	0.424	7.92	
Benzo[a]anthracene	1	0		11.81	48.82	50			1.471	1.436	2.36	
Chrysene	1	0		11.85	47.60	50			1.349	1.284	4.80	
bis(2-Ethylhexyl)phthalate	1	0		11.94	48.33	50			0.974	0.942	3.34	
Perylene-d12	1	0	I	13.40	40.00	40				0.000	0.00	
Di-n-octylphthalate	1	0	CC	12.68	50.89	50	20		2.190	2.229	1.78	
Benzo[b]fluoranthene	1	0		13.01	52.94	50			1.579	1.672	5.88	
Benzo[k]fluoranthene	1	0		13.04	50.40	50			1.599	1.612	0.80	
Benzo[a]pyrene	1	0	CC	13.34	49.83	50	20		1.486	1.481	0.34	
Indeno[1,2,3-cd]pyrene	1	0		14.42	53.98	50			1.610	1.738	7.96	
Dibenzo[a,h]anthracene	1	0		14.45	53.82	50			1.336	1.439	7.64	
Benzo[g,h,i]perylene	1	0		14.69	54.65	50			1.347	1.472	9.30	
Diaminotoluene Dihydrochloride	1	1E		0.00	0.00	50				0.000	100.00	
Toluene Diisocyanate	1	1E		0.00	0.00	50				0.000	100.00	
2,4 Diaminotoluene	1	1E		0.00	0.00	50				0.000	100.00	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound 1 - Internal Standard
* - Failed the C or P Criteria

Page 2 of 2

** - No limit specified in method

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09779.D Vial: 2
 Acq On : 5 Aug 2005 6:42 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 7:13 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	17569	40.00	ng	-0.15
20) Naphthalene-d8	6.14	136	70683	40.00	ng	-0.14
36) Acenaphthene-d10	7.47	164	38532	40.00	ng	-0.16
61) Phenanthrene-d10	8.84	188	69550	40.00	ng	-0.19
77) Chrysene-d12	11.82	240	55217	40.00	ng	-0.22
88) Perylene-d12	13.40	264	40060	40.00	ng	-0.22

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.78	112	28403	48.00	ng	-0.19
Spiked Amount 200.000			Recovery =	24.00%		
8) Phenol-d5	4.80	99	38027	43.95	ng	-0.15
Spiked Amount 200.000			Recovery =	21.98%		
21) Nitrobenzene-d5	5.58	128	7425	23.99	ng	-0.14
Spiked Amount 100.000			Recovery =	23.99%		
41) 2-Fluorobiphenyl	6.95	172	31623	26.26	ng	-0.14
Spiked Amount 100.000			Recovery =	26.26%		
64) 2,4,6-Tribromophenol	8.16	330	7088	47.62	ng	-0.18
Spiked Amount 200.000			Recovery =	23.81%		
80) Terphenyl-d14	10.61	244	31867	24.43	ng	-0.20
Spiked Amount 100.000			Recovery =	24.43%		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	1.97	79	35445	48.38	ng	97
3) N-Nitrosodimethylamine	1.92	74	20580	46.19	ng	89
5) Aniline	4.81	93	45478	45.79	ng	91
6) Pentachloroethane	4.84	117	11004	50.80	ng	99
7) bis(2-Chloroethyl)ether	4.89	93	31290	50.33	ng	97
9) Phenol	4.81	94	42149	45.96	ng	75
10) 2-Chlorophenol	4.91	128	31603	45.40	ng	94
11) 1,3-Dichlorobenzene	5.05	146	33571	52.12	ng	99
12) 1,4-Dichlorobenzene	5.12	146	33833	51.33	ng	99
13) 1,2-Dichlorobenzene	5.25	146	31978	50.89	ng	98
14) Benzyl alcohol	5.24	108	21389	46.53	ng	99
15) bis(2-chloroisopropyl)ethe	5.36	45	49153	52.05	ng	96
16) 2-Methylphenol	5.34	108	31760	49.97	ng	98
17) Hexachloroethane	5.52	117	14913	54.55	ng	97
18) N-Nitroso-di-n-propylamine	5.46	70	24716	49.33	ng	98
19) 3&4-Methylphenol	5.47	108	32434	48.00	ng	97
22) Nitrobenzene	5.59	77	35057	50.50	ng	98
23) Isophorone	5.78	82	63510	49.15	ng	95
24) 2-Nitrophenol	5.84	139	18182	51.04	ng	92

(#) = qualifier out of range (m) = manual integration

18/11

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09779.D Vial: 2
 Acq On : 5 Aug 2005 6:42 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 7:13 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.90	107	32982	48.67	ng	99
26) Benzoic Acid	5.98	105	13987	35.96	ng	97
27) bis(2-Chloroethoxy)methane	5.97	93	35732	48.33	ng	100
28) 2,4-Dichlorophenol	6.04	162	26489	46.88	ng	98
29) 1,2,4-Trichlorobenzene	6.09	180	29685	51.24	ng	99
30) Naphthalene	6.15	128	94365	50.95	ng	100
31) 4-Chloroaniline	6.20	127	36372	50.29	ng	99
32) Hexachlorobutadiene	6.25	225	15891	49.75	ng	97
33) 4-Chloro-3-methylphenol	6.57	107	29512	47.30	ng	94
34) 2-Methylnaphthalene	6.67	142	60973	47.66	ng	97
35) Methylnaphthalenes (Total)	6.67	142	60973	47.66	ng	97
37) 1,2,4,5-Tetrachlorobenzene	6.79	216	26618	51.15	ng	97
38) Hexachlorocyclopentadiene	6.79	237	17821	55.61	ng	98
39) 2,4,6-Trichlorophenol	6.89	196	18674	49.79	ng	99
40) 2,4,5-Trichlorophenol	6.91	196	21249	52.09	ng	96
42) 2-Chloronaphthalene	7.03	162	58240	53.47	ng	98
43) 1,4-Dimethylnaphthalene	7.30	156	43182	52.10	ng	95
44) Dimethylnaphthalenes (Tota	7.30	156	43182	52.10	ng	95
45) Diphenyl Ether	7.10	170	51138	71.06	ng	94
46) 2-Nitroaniline	7.11	65	23712	57.16	ng	99
47) Acenaphthylene	7.36	152	89397	51.96	ng	99
48) Dimethylphthalate	7.26	163	64183	51.05	ng	99
49) 2,6-Dinitrotoluene	7.31	165	14741	50.91	ng	90
50) Acenaphthene	7.50	153	57174	53.72	ng	98
51) 3-Nitroaniline	7.44	138	17290	54.69	ng	95
52) 2,4-Dinitrophenol	7.53	184	8312	46.65	ng	83
53) Dibenzofuran	7.64	168	83284	53.76	ng	97
54) 2,4-Dinitrotoluene	7.64	165	20401	51.00	ng	98
55) 4-Nitrophenol	7.58	65	13530	53.76	ng	88
56) 2,3,4,6-Tetrachlorophenol	7.75	232	16056	50.18	ng	99
57) Fluorene	7.94	166	61758	49.48	ng	98
58) 4-Chlorophenyl-phenylether	7.95	204	30480	50.24	ng	96
59) Diethylphthalate	7.85	149	64641	50.17	ng	99
60) 4-Nitroaniline	7.96	138	16871	46.01	ng	98
62) 4,6-Dinitro-2-methylphenol	7.99	198	12394	46.35	ng	100
63) n-Nitrosodiphenylamine	8.06	169	45739	47.71	ng	99
65) 1,2-Diphenylhydrazine	8.09	77	69658	50.91	ng	99
66) 4-Bromophenyl-phenylether	8.40	248	17105	47.96	ng	94
67) Hexachlorobenzene	8.45	284	16256	48.28	ng	90
68) gamma-BHC	8.71	181	2337	9.49	ng	96
69) Pentachlorophenol	8.65	266	9981	45.22	ng	94

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09779.D Vial: 2
 Acq On : 5 Aug 2005 6:42 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 5 7:13 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.86	178	97375	48.55	ng	99
71) Anthracene	8.92	178	97685	47.92	ng	98
72) Carbazole	9.09	167	89501	48.09	ng	100
73) Heptachlor	9.36	100	2479	9.52	ng	94
74) Di-n-butylphthalate	9.50	149	107903	47.88	ng	99
75) Heptachlor epoxide	10.05	81	1793	9.91	ng	94
76) Fluoranthene	10.14	202	101945	46.62	ng	97
78) Pyrene	10.40	202	111326	50.34	ng	94
79) Benzidine	10.33	184	36624	44.85	ng	98
81) Endrin	10.84	81	1034	9.50	ng	85
82) Butylbenzylphthalate	11.22	149	48527	49.86	ng	95
83) Methoxychlor	11.85	227	9616	9.47	ng	100
84) 3,3'-Dichlorobenzidine	11.81	252	29266	46.04	ng	98
85) Benzo[a]anthracene	11.81	228	99123	48.82	ng	99
86) Chrysene	11.85	228	88636	47.60	ng	99
87) bis(2-Ethylhexyl)phthalate	11.94	149	64994	48.33	ng	98
89) Di-n-octylphthalate	12.68	149	111635	50.89	ng	99
90) Benzo[b]fluoranthene	13.01	252	83733	52.94	ng	98
91) Benzo[k]fluoranthene	13.04	252	80697	50.40	ng	94
92) Benzo[a]pyrene	13.34	252	74171	49.83	ng	99
93) Indeno[1,2,3-cd]pyrene	14.42	276	87042	53.98	ng	93
94) Dibenzo[a,h]anthracene	14.45	278	72039	53.82	ng	97
95) Benzo[g,h,i]perylene	14.69	276	73715	54.65	ng	94

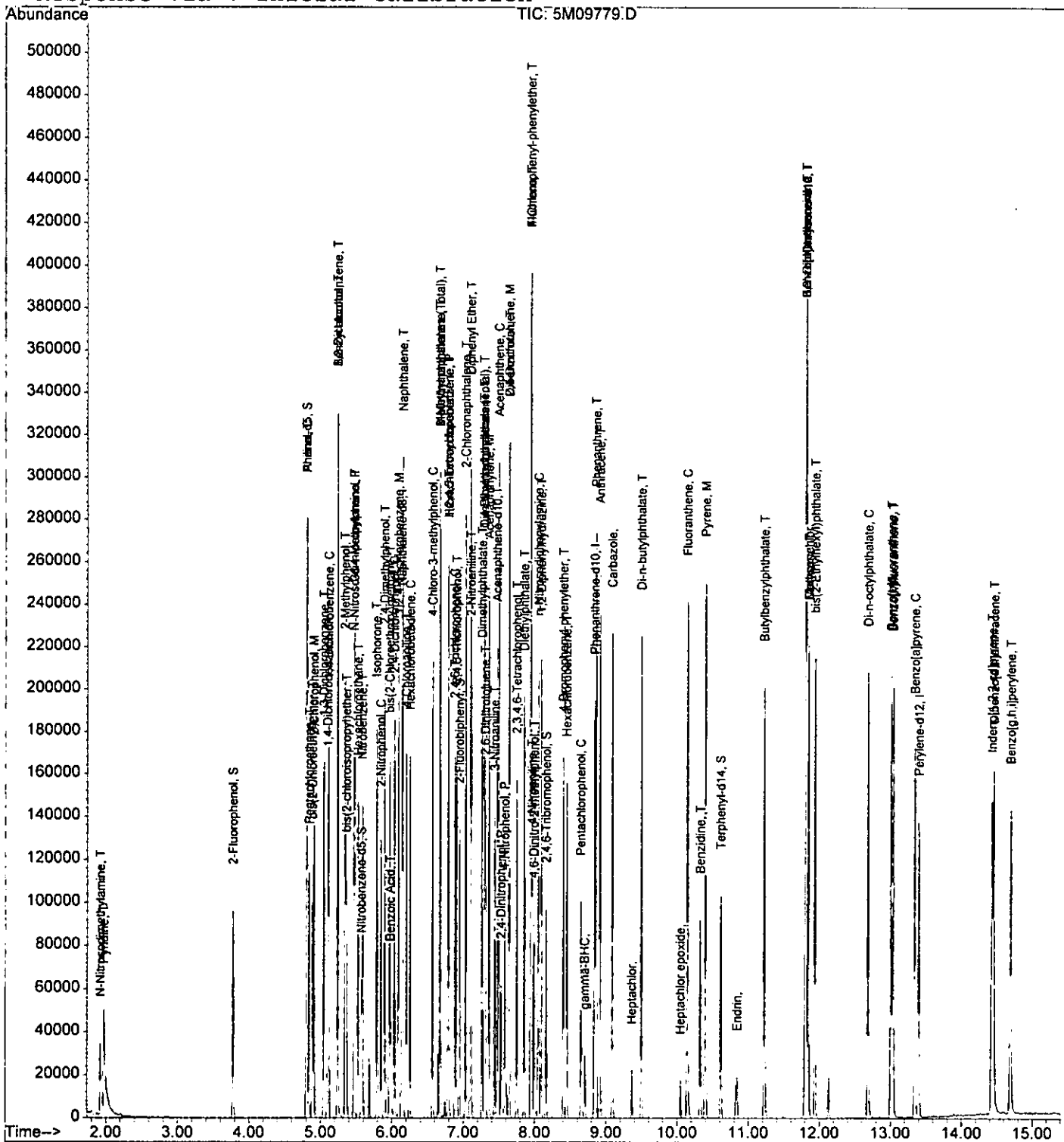
(#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09779.D Vial: 2
Acq On : 5 Aug 2005 6:42 Operator: AHD
Sample : CAL BNA@50PPM Inst : GCMS_5
Misc : A,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 5 7:13 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/5/05 8:18:00 AM

Data File: 4M05385.D
Method: 8270

Instrument: GCMS_4

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
1,4-Dichlorobenzene-d4	1	0	I	4.91	40.00	40				0.000	0.00	
Pyridine	1	0		2.29	55.41	50			1.515	1.679	10.82	
N-Nitrosodimethylamine	1	0		2.24	42.75	50			0.917	0.784	14.50	
2-Fluorophenol	1	0	S	3.76	52.42	50			1.128	1.183	4.84	
Aniline	1	0		4.64	57.54	50			1.631	1.878	15.08	
bis(2-Chloroethyl)ether	1	0		4.70	55.92	50			1.226	1.372	11.84	
Phenol-d5	1	0	S	4.62	52.81	50			1.502	1.586	5.62	
Phenol	1	0	CC	4.63	54.87	50	20		1.625	1.783	9.74	
2-Chlorophenol	1	0		4.73	49.00	50			1.244	1.219	2.00	
1,3-Dichlorobenzene	1	0		4.85	50.78	50			1.335	1.356	1.56	
1,4-Dichlorobenzene	1	0	CC	4.93	51.63	50	20		1.306	1.349	3.26	
1,2-Dichlorobenzene	1	0		5.04	53.05	50			1.281	1.359	6.10	
Benzyl alcohol	1	0		5.03	50.02	50			0.775	0.775	0.04	
bis(2-chloroisopropyl)ether	1	0		5.13	49.71	50			3.011	2.993	0.58	
2-Methylphenol	1	0		5.12	49.59	50			1.038	1.029	0.82	
Hexachloroethane	1	0		5.30	55.71	50			0.610	0.680	11.42	
N-Nitroso-di-n-propylamine	1	0	CP	5.23	47.76	50	0.05		1.070	1.021	4.48	
3&4-Methylphenol	1	0		5.24	47.11	50			1.070	1.008	5.78	
Naphthalene-d8	1	0	I	5.91	40.00	40				0.000	0.00	
Nitrobenzene-d5	1	0	S	5.34	24.93	25			0.200	0.200	0.28	
Nitrobenzene	1	0		5.36	47.46	50			0.417	0.396	5.08	
Isophorone	1	0		5.55	51.57	50			0.797	0.822	3.14	
2-Nitrophenol	1	0	CC	5.61	50.51	50	20		0.216	0.218	1.02	
2,4-Dimethylphenol	1	0		5.65	46.13	50			0.398	0.368	7.74	
Benzoic Acid	1	0		5.72	59.37	50			0.069	0.082	18.74	
bis(2-Chloroethoxy)methane	1	0		5.72	52.22	50			0.476	0.497	4.44	
2,4-Dichlorophenol	1	0	CC	5.79	47.83	50	20		0.327	0.313	4.34	
1,2,4-Trichlorobenzene	1	0		5.87	49.64	50			0.351	0.348	0.72	
Naphthalene	1	0		5.93	55.23	50			0.879	0.971	10.46	
4-Chloroaniline	1	0		5.97	58.79	50			0.342	0.402	17.58	
Hexachlorobutadiene	1	0	CC	6.03	51.81	50	20		0.225	0.233	3.62	
4-Chloro-3-methylphenol	1	0	CC	6.37	43.83	50	20		0.365	0.320	12.34	
2-Methylnaphthalene	1	0		6.50	51.18	50			0.615	0.629	2.36	
Methylnaphthalene	1	0		6.50	51.18	50						
Acenaphthene-d10	1	0	I	7.48	40.00	40				0.000	0.00	
1,2,4,5-Tetrachlorobenzene	1	0		6.65	56.47	50			0.653	0.738	12.94	
Hexachlorocyclopentadiene	1	0	CP	6.64	53.10	50	0.05		0.459	0.488	6.20	
2,4,6-Trichlorophenol	1	0	CC	6.76	55.86	50	20		0.470	0.526	11.72	
2,4,5-Trichlorophenol	1	0		6.79	52.94	50			0.482	0.510	5.88	
2-Fluorobiphenyl	1	0	S	6.84	25.60	25			1.282	1.312	2.40	
2-Chloronaphthalene	1	0		6.94	52.54	50			1.120	1.177	5.08	
2-Nitroaniline	1	0		7.04	51.67	50			0.663	0.685	3.34	
1,4-Dimethylnaphthalene	1	0		7.27	53.91	50			0.770	0.831	7.82	
Dimethylnaphthalene	1	0		7.27	53.91	50						
Diphenyl Ether	1	0		7.03	52.83	50			0.977	1.033	5.66	
Acenaphthylene	1	0		7.34	55.59	50			1.719	1.911	11.18	
Dimethylphthalate	1	0		7.23	48.93	50			1.413	1.383	2.14	
2,6-Dinitrotoluene	1	0		7.28	51.01	50			0.337	0.343	2.02	
Acenaphthene	1	0	CC	7.51	53.60	50	20		1.127	1.208	7.20	
3-Nitroaniline	1	0		7.44	60.90	50			0.294	0.358	21.80	
2,4-Dinitrophenol	1	0	CP	7.55	33.44	50	0.05		0.191	0.128	33.12	
Dibenzofuran	1	0		7.69	54.59	50			1.485	1.622	9.18	
2,4-Dinitrotoluene	1	0		7.69	51.76	50			0.440	0.456	3.52	
4-Nitrophenol	1	0	CP	7.61	47.30	50	0.05		0.377	0.357	5.40	
Fluorene	1	0		8.05	55.37	50			1.092	1.210	10.74	
4-Chlorophenyl-phenylether	1	0		8.05	49.79	50			0.656	0.653	0.42	
Diethylphthalate	1	0		7.94	49.05	50			1.474	1.446	1.90	
4-Nitroaniline	1	0		8.07	49.93	50			0.378	0.377	0.14	
Phenanthrene-d10	1	0	I	9.08	40.00	40				0.000	0.00	
4,6-Dinitro-2-methylphenol	1	0		8.12	46.38	50			0.144	0.133	7.24	
n-Nitrosodiphenylamine	1	0	CC	8.19	55.45	50	20		0.507	0.563	10.90	
2,4,6-Tribromophenol	1	0	S	8.31	55.61	50			0.179	0.199	11.22	
1,2-Diphenylhydrazine	1	0		8.23	54.01	50			0.910	0.983	8.02	
4-Bromophenyl-phenylether	1	0		8.60	53.99	50			0.259	0.279	7.98	
Hexachlorobenzene	1	0		8.65	56.28	50			0.347	0.391	12.56	
Pentachlorophenol	1	0	CC	8.87	44.31	50	20		0.189	0.168	11.38	
Phenanthrene	1	0		9.11	54.18	50			0.999	1.083	8.36	
Anthracene	1	0		9.17	54.83	50			1.015	1.113	9.66	
Carbazole	1	0		9.36	54.75	50			0.931	1.020	9.50	
Di-n-butylphthalate	1	0		9.81	53.98	50			1.362	1.470	7.96	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 1 of 2

** - No limit specified in method

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Form 7

Continuing Calibration

Calibration Name: CAL_BNA@50PPM
 Cont Calibration Date/Time 8/5/05 8:18:00 AM

Data File: 4M05385.D
 Method: 8270

Instrument: GCMS_4

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
Fluoranthene	1	0	CC	10.50	52.26	50	20		1.039	1.086	4.52	
Chrysene-d12	1	0	I	12.28	40.00	40				0.000	0.00	
Pyrene	1	0		10.76	48.62	50			1.538	1.496	2.76	
Benzidine	1	0		10.68	55.26	50			0.434	0.479	10.52	
Terphenyl-d14	1	0	S	10.98	22.65	25			1.127	1.021	9.40	
Butylbenzylphthalate	1	0		11.62	49.36	50			0.750	0.740	1.28	
3,3'-Dichlorobenzidine	1	0		12.27	65.41	50			0.303	0.396	30.82	
Benzo[a]anthracene	1	0		12.27	50.94	50			1.255	1.279	1.88	
Chrysene	1	0		12.32	51.03	50			1.121	1.145	2.06	
bis(2-Ethylhexyl)phthalate	1	0		12.41	52.13	50			0.976	1.017	4.26	
Perylene-d12	1	0	I	14.13	40.00	40				0.000	0.00	
Di-n-octylphthalate	1	0	CC	13.27	44.72	50	20		2.191	1.960	10.56	
Benzo[b]fluoranthene	1	0		13.66	46.51	50			1.664	1.548	6.98	
Benzo[k]fluoranthene	1	0		13.70	49.27	50			1.441	1.420	1.46	
Benzo[a]pyrene	1	0	CC	14.07	48.72	50	20		1.368	1.333	2.56	
Indeno[1,2,3-cd]pyrene	1	0		15.36	58.85	50			1.241	1.460	17.70	
Dibenzo[a,h]anthracene	1	0		15.40	57.76	50			1.016	1.174	15.52	
Benzo[g,h,i]perylene	1	0		15.65	62.63	50			0.990	1.240	25.26	
2,4 Diaminotoluene	1	1E		0.00	0.00	50				0.000	100.00	
Heptachlor_epoxide	1	1E		0.00	0.00	10				0.000	100.00	
Methoxychlor	1	1E		0.00	0.00	10				0.000	100.00	
Diaminotoluene Dihydrochloride	1	1E		0.00	0.00	50				0.000	100.00	
Toluene Diisocyanate	1	1E		0.00	0.00	50				0.000	100.00	
Endrin	1	1E		0.00	0.00	10				0.000	100.00	
2,3,4,6-Tetrachlorophenol	1	1E		0.00	0.00	50				0.000	100.00	
gamma-BHC	1	1E		0.00	0.00	10				0.000	100.00	
Pentachloroethane	1	1E		0.00	0.00	50				0.000	100.00	
Heptachlor	1	1E		0.00	0.00	10				0.000	100.00	

CC - Continuing Calibration Check Compound
 N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
 * - Failed the C or P Criteria

Page 2 of 2

** - No limit specified in method

Note:

8260/8270 limits are compared against the %DIFF/R.F.
 624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
 524.2 limits are compared against the %DIFF

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05385.D Vial: 2
 Acq On : 5 Aug 2005 8:18 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 8:36 2005 Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.91	152	33586	40.00	ng	-0.04
19) Naphthalene-d8	5.91	136	104620	40.00	ng	-0.04
35) Acenaphthene-d10	7.48	164	54282	40.00	ng	-0.05
59) Phenanthrene-d10	9.08	188	87118	40.00	ng	-0.06
72) Chrysene-d12	12.28	240	65220	40.00	ng	-0.06
81) Perylene-d12	14.13	264	48868	40.00	ng	-0.06
System Monitoring Compounds						
4) 2-Fluorophenol	3.76	112	49658	52.42	ng	-0.04
Spiked Amount	200.000		Recovery	=	26.21%	
7) Phenol-d5	4.62	99	66603	52.81	ng	-0.04
Spiked Amount	200.000		Recovery	=	26.41%	
20) Nitrobenzene-d5	5.34	128	13057	24.93	ng	-0.04
Spiked Amount	100.000		Recovery	=	24.93%	
40) 2-Fluorobiphenyl	6.84	172	44528	25.60	ng	-0.04
Spiked Amount	100.000		Recovery	=	25.60%	
62) 2,4,6-Tribromophenol	8.31	332	21684	55.61	ng	-0.05
Spiked Amount	200.000		Recovery	=	27.81%	
75) Terphenyl-d14	10.98	244	41622	22.65	ng	-0.05
Spiked Amount	100.000		Recovery	=	22.65%	
Target Compounds						
2) Pyridine	2.29	79	70490	55.41	ng	Qvalue 93
3) N-Nitrosodimethylamine	2.24	74	32905	42.75	ng	88
5) Aniline	4.64	93	78823	57.54	ng	46
6) bis(2-Chloroethyl)ether	4.70	93	57587	55.92	ng	96
8) Phenol	4.63	94	74872	54.87	ng	61
9) 2-Chlorophenol	4.73	128	51192	49.00	ng	77
10) 1,3-Dichlorobenzene	4.85	146	56933	50.78	ng	98
11) 1,4-Dichlorobenzene	4.93	146	56633	51.63	ng	96
12) 1,2-Dichlorobenzene	5.04	146	57063	53.05	ng	98
13) Benzyl alcohol	5.03	108	32557	50.02	ng	87
14) bis(2-chloroisopropyl)ethe	5.13	45	125670	49.71	ng	98
15) 2-Methylphenol	5.12	108	43203	49.59	ng	99
16) Hexachloroethane	5.30	117	28539	55.71	ng	90
17) N-Nitroso-di-n-propylamine	5.23	70	42885	47.76	ng	92
18) 3&4-Methylphenol	5.24	108	42336	47.11	ng	96
21) Nitrobenzene	5.36	77	51761	47.46	ng	81
22) Isophorone	5.55	82	107455	51.57	ng	96
23) 2-Nitrophenol	5.61	139	28501	50.51	ng	75
24) 2,4-Dimethylphenol	5.65	107	48079	46.13	ng	97

(#) = qualifier out of range (m) = manual integration

h811

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05385.D Vial: 2
 Acq On : 5 Aug 2005 8:18 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 5 8:36 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.72	105	10747	59.37	ng	99
26) bis(2-Chloroethoxy)methane	5.72	93	65009	52.22	ng	98
27) 2,4-Dichlorophenol	5.79	162	40946	47.83	ng	97
28) 1,2,4-Trichlorobenzene	5.87	180	45574	49.64	ng	90
29) Naphthalene	5.93	128	126927	55.23	ng	99
30) 4-Chloroaniline	5.97	127	52532	58.79	ng	97
31) Hexachlorobutadiene	6.03	225	30523	51.81	ng	98
32) 4-Chloro-3-methylphenol	6.37	107	41834	43.83	ng	82
33) 2-Methylnaphthalene	6.50	142	82301	51.18	ng	97
34) Methylnaphthalene (Total)	6.50	142	82301	51.18	ng	97
36) 1,2,4,5-Tetrachlorobenzene	6.65	216	50044	56.47	ng	98
37) Hexachlorocyclopentadiene	6.64	237	33099	53.10	ng	96
38) 2,4,6-Trichlorophenol	6.76	196	35657	55.86	ng	99
39) 2,4,5-Trichlorophenol	6.79	196	34619	52.94	ng	98
41) 2-Chloronaphthalene	6.94	162	79861	52.54	ng	92
42) 2-Nitroaniline	7.04	65	46497	51.67	ng	85
43) 1,4-Dimethylnaphthalene	7.27	156	56356	53.91	ng	89
44) Dimethylnaphthalene (Total)	7.27	156	56356	53.91	ng	89
45) Diphenyl Ether	7.03	170	70070	52.83	ng	77
46) Acenaphthylene	7.34	152	129700	55.59	ng	99
47) Dimethylphthalate	7.23	163	93854	48.93	ng	98
48) 2,6-Dinitrotoluene	7.28	165	23293	51.01	ng	87
49) Acenaphthene	7.51	153	81962	53.60	ng	99
50) 3-Nitroaniline	7.44	138	24258	60.90	ng	91
51) 2,4-Dinitrophenol	7.55	184	8659	33.44	ng	90
52) Dibenzofuran	7.69	168	110042	54.59	ng	90
53) 2,4-Dinitrotoluene	7.69	165	30908	51.76	ng	88
54) 4-Nitrophenol	7.61	65	24207	47.30	ng	100
55) Fluorene	8.05	166	82072	55.37	ng	94
56) 4-Chlorophenyl-phenylether	8.05	204	44302	49.79	ng	84
57) Diethylphthalate	7.94	149	98105	49.05	ng	99
58) 4-Nitroaniline	8.07	138	25606	49.93	ng	92
60) 4,6-Dinitro-2-methylphenol	8.12	198	14497	46.38	ng	100
61) n-Nitrosodiphenylamine	8.19	169	61276	55.45	ng	99
63) 1,2-Diphenylhydrazine	8.23	77	106996	54.01	ng	87
64) 4-Bromophenyl-phenylether	8.60	248	30404	53.99	ng	81
65) Hexachlorobenzene	8.65	284	42569	56.28	ng	98
66) Pentachlorophenol	8.87	266	18261	44.31	ng	98
67) Phenanthrene	9.11	178	117901	54.18	ng	98
68) Anthracene	9.17	178	121161	54.83	ng	99
69) Carbazole	9.36	167	111031	54.75	ng	99

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05385.D Vial: 2
 Acq On : 5 Aug 2005 8:18 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 8:36 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.81	149	160120	53.98	ng	99
71) Fluoranthene	10.50	202	118297	52.26	ng	89
73) Pyrene	10.76	202	121958	48.62	ng	87
74) Benzidine	10.68	184	39073	55.26	ng	96
76) Butylbenzylphthalate	11.62	149	60361	49.36	ng	87
77) 3,3'-Dichlorobenzidine	12.27	252	32265	65.41	ng	97
78) Benzo[a]anthracene	12.27	228	104278	50.94	ng	99
79) Chrysene	12.32	228	93313	51.03	ng	99
80) bis(2-Ethylhexyl)phthalate	12.41	149	82924	52.13	ng	94
82) Di-n-octylphthalate	13.27	149	119715	44.72	ng	97
83) Benzo[b]fluoranthene	13.66	252	94543	46.51	ng	97
84) Benzo[k]fluoranthene	13.70	252	86743	49.27	ng	99
85) Benzo[a]pyrene	14.07	252	81404	48.72	ng	94
86) Indeno[1,2,3-cd]pyrene	15.36	276	89201	58.85	ng	92
87) Dibenzo[a,h]anthracene	15.40	278	71694	57.76	ng	98
88) Benzo[g,h,i]perylene	15.65	276	75771	62.63	ng	94

 (#) = qualifier out of range (m) = manual integration

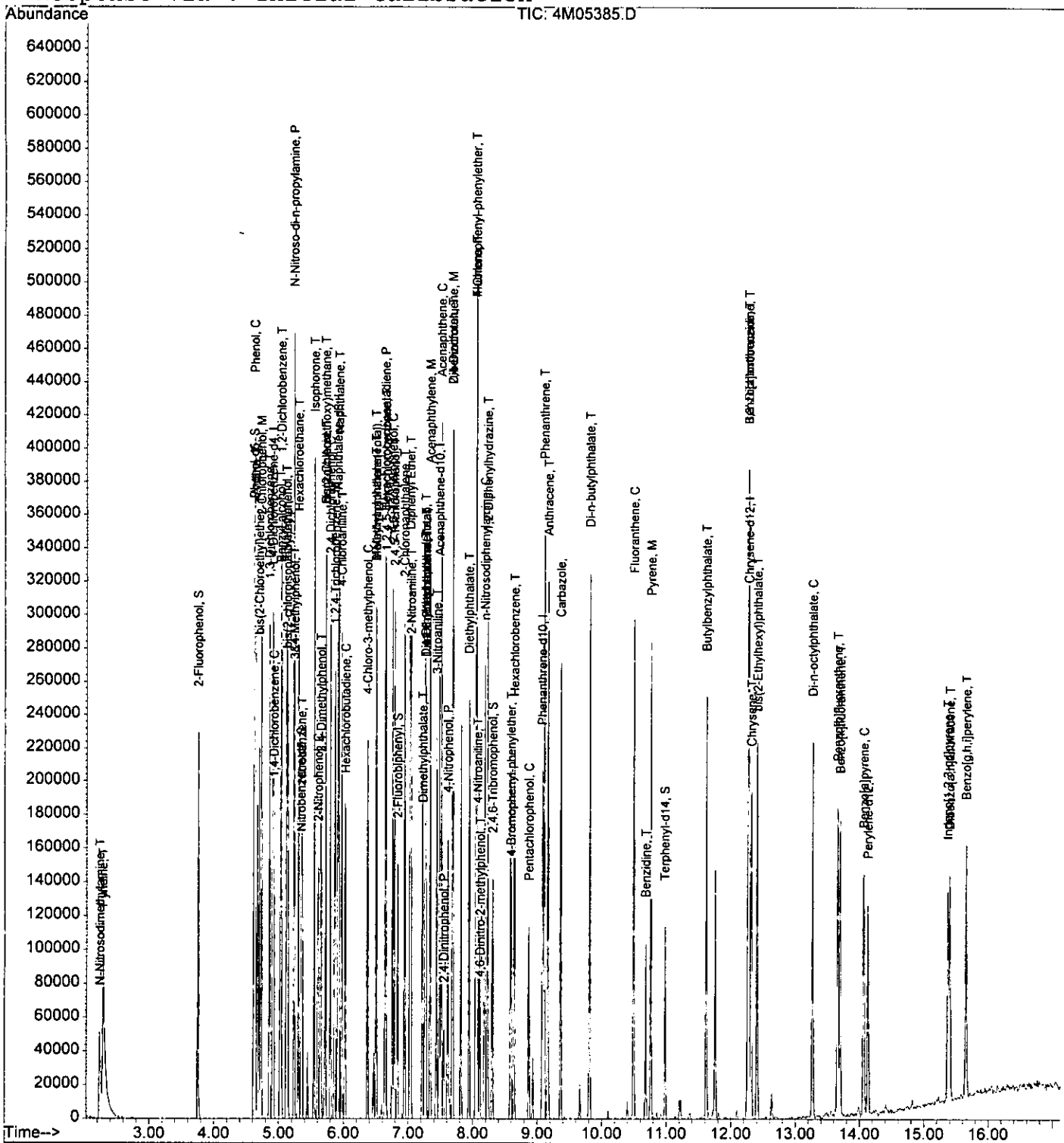
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05385.D Vial: 2
 Acq On : 5 Aug 2005 8:18 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 8:36 2005

9.065

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration



Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/8/05 6:40:00 AM

Data File: 5M09827.D
Method: 8270

Instrument: GCMS_5

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
1,4-Dichlorobenzene-d4	1	0	I	5.10	40.00	40			0.000	0.00		
Pyridine	1	0		1.95	48.27	50			1.668	1.610	3.46	
N-Nitrosodimethylamine	1	0		1.91	45.61	50			1.014	0.925	8.78	
2-Fluorophenol	1	0	S	3.77	49.47	50			1.347	1.333	1.06	
Aniline	1	0		4.80	47.59	50			2.261	2.152	4.82	
Pentachloroethane	1	0		4.83	51.74	50			0.493	0.510	3.48	
bis(2-Chloroethyl)ether	1	0		4.88	49.18	50			1.415	1.392	1.64	
Phenol-d5	1	0	S	4.80	42.72	50			1.970	1.683	14.56	
Phenol	1	0	CC	4.81	47.44	50	20		2.088	1.981	5.12	
2-Chlorophenol	1	0		4.90	46.70	50			1.585	1.480	6.60	
1,3-Dichlorobenzene	1	0		5.04	51.34	50			1.466	1.506	2.68	
1,4-Dichlorobenzene	1	0	CC	5.11	50.07	50	20		1.501	1.503	0.14	
1,2-Dichlorobenzene	1	0		5.23	49.50	50			1.431	1.416	1.00	
Benzyl alcohol	1	0		5.23	48.01	50			1.047	1.005	3.98	
bis(2-chloroisopropyl)ether	1	0		5.35	51.80	50			2.150	2.227	3.60	
2-Methylphenol	1	0		5.34	47.18	50			1.447	1.366	5.64	
Hexachloroethane	1	0		5.52	50.23	50			0.622	0.625	0.46	
N-Nitroso-di-n-propylamine	1	0	CP	5.46	48.28	50	0.05		1.141	1.102	3.44	
3&4-Methylphenol	1	0		5.47	44.85	50			1.538	1.380	10.30	
Naphthalene-d8	1	0	I	6.13	40.00	40			0.000	0.00		
Nitrobenzene-d5	1	0	S	5.57	24.98	25			0.175	0.175	0.08	
Nitrobenzene	1	0		5.58	50.87	50			0.393	0.400	1.74	
Isophorone	1	0		5.78	48.68	50			0.731	0.712	2.64	
2-Nitrophenol	1	0	CC	5.84	50.71	50	20		0.202	0.204	1.42	
2,4-Dimethylphenol	1	0		5.89	46.70	50			0.383	0.358	6.60	
Benzoic Acid	1	0		5.99	36.52	50			0.220	0.161	26.96	
bis(2-Chloroethoxy)methane	1	0		5.96	53.36	50			0.418	0.447	6.72	
2,4-Dichlorophenol	1	0	CC	6.03	46.99	50	20		0.320	0.300	6.02	
1,2,4-Trichlorobenzene	1	0		6.09	49.66	50			0.328	0.326	0.68	
Naphthalene	1	0		6.15	48.69	50			1.048	1.021	2.62	
4-Chloroaniline	1	0		6.20	51.06	50			0.409	0.418	2.12	
Hexachlorobutadiene	1	0	CC	6.24	51.47	50	20		0.181	0.186	2.94	
4-Chloro-3-methylphenol	1	0	CC	6.56	43.88	50	20		0.353	0.310	12.24	
2-Methylnaphthalene	1	0		6.67	47.15	50			0.724	0.683	5.70	
Methylnaphthalenes	1	0		6.67	47.15							
Acenaphthene-d10	1	0	I	7.47	40.00	40			0.000	0.00		
1,2,4,5-Tetrachlorobenzene	1	0		6.79	49.75	50			0.540	0.537	0.50	
Hexachlorocyclopentadiene	1	0	CP	6.78	55.88	50	0.05		0.333	0.372	11.76	
2,4,6-Trichlorophenol	1	0	CC	6.87	48.23	50	20		0.389	0.376	3.54	
2,4,5-Trichlorophenol	1	0		6.90	46.34	50			0.423	0.392	7.32	
2-Fluorobiphenyl	1	0	S	6.94	25.37	25			1.250	1.269	1.48	
2-Chloronaphthalene	1	0		7.03	50.71	50			1.131	1.147	1.42	
1,4-Dimethylnaphthalene	1	0		7.29	48.87	50			0.860	0.841	2.26	
Dimethylnaphthalenes	1	0		7.29	48.87							
Diphenyl Ether	1	0		7.10	65.92	50			0.747	0.985	31.84	
2-Nitroaniline	1	0		7.11	55.08	50			0.431	0.474	10.16	
Acenaphthylene	1	0		7.35	47.10	50			1.786	1.682	5.80	
Dimethylphthalate	1	0		7.26	48.28	50			1.305	1.260	3.44	
2,6-Dinitrotoluene	1	0		7.30	48.57	50			0.301	0.292	2.86	
Acenaphthene	1	0	CC	7.49	50.84	50	20		1.105	1.123	1.68	
3-Nitroaniline	1	0		7.44	49.97	50			0.328	0.328	0.06	
2,4-Dinitrophenol	1	0	CP	7.52	47.10	50	0.05		0.185	0.174	5.80	
Dibenzofuran	1	0		7.64	48.65	50			1.608	1.565	2.70	
2,4-Dinitrotoluene	1	0		7.63	46.10	50			0.415	0.383	7.80	
4-Nitrophenol	1	0	CP	7.57	49.64	50	0.05		0.261	0.259	0.72	
2,3,4,6-Tetrachlorophenol	1	0		7.75	46.00	50			0.332	0.306	8.00	
Fluorene	1	0		7.94	47.54	50			1.296	1.232	4.92	
4-Chlorophenyl-phenylether	1	0		7.94	47.73	50			0.630	0.601	4.54	
Diethylphthalate	1	0		7.85	47.87	50			1.338	1.281	4.26	
4-Nitroaniline	1	0		7.96	44.90	50			0.381	0.342	10.20	
Phenanthrene-d10	1	0	I	8.83	40.00	40			0.000	0.00		
4,6-Dinitro-2-methylphenol	1	0		7.99	48.78	50			0.154	0.150	2.44	
n-Nitrosodiphenylamine	1	0	CC	8.05	48.93	50	20		0.551	0.540	2.14	
2,4,6-Tribromophenol	1	0	S	8.16	48.36	50			0.086	0.083	3.28	
1,2-Diphenylhydrazine	1	0		8.08	54.78	50			0.787	0.862	9.56	
4-Bromophenyl-phenylether	1	0		8.40	47.23	50			0.205	0.194	5.54	
Hexachlorobenzene	1	0		8.45	47.42	50			0.194	0.184	5.16	
gamma-BHC	1	0		8.70	10.38	10			0.142	0.147	3.80	
Pentachlorophenol	1	0	CC	8.64	48.61	50	20		0.127	0.123	2.78	
Phenanthrene	1	0		8.85	48.13	50			1.154	1.111	3.74	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 1 of 2

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM Data File: 5M09827.D
 Cont Calibration Date/Time 8/8/05 6:40:00 AM Method: 8270

Instrument: GCMS_5

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
Anthracene	1	0		8.90	47.90	50			1.172	1.123	4.20	
Carbazole	1	0		9.08	48.82	50			1.070	1.045	2.36	
Heptachlor	1	0		9.35	10.30	10			0.150	0.154	3.00	
Di-n-butylphthalate	1	0		9.49	50.12	50			1.296	1.299	0.24	
Heptachlor_epoxide	1	0		10.04	9.82	10			0.104	0.102	1.80	
Fluoranthene	1	0	CC	10.13	48.03	50	20		1.258	1.208	3.94	
Chrysene-d12	1	0	I	11.81	40.00	40				0.000	0.00	
Pyrene	1	0		10.39	51.04	50			1.602	1.635	2.08	
Benzidine	1	0		10.31	50.70	50			0.592	0.600	1.40	
Terphenyl-d14	1	0	S	10.60	25.02	25			0.945	0.946	0.08	
Endrin	1	0		10.83	10.65	10			0.079	0.084	6.50	
Butylbenzylphthalate	1	0		11.21	51.36	50			0.705	0.724	2.72	
Methoxychlor	1	0		11.84	10.05	10			0.735	0.739	0.50	
3,3'-Dichlorobenzidine	1	0		11.79	50.67	50			0.460	0.467	1.34	
Benzo[a]anthracene	1	0		11.79	49.33	50			1.471	1.451	1.34	
Chrysene	1	0		11.84	49.45	50			1.349	1.334	1.10	
bis(2-Ethylhexyl)phthalate	1	0		11.93	51.74	50			0.974	1.008	3.48	
Perylene-d12	1	0	I	13.39	40.00	40				0.000	0.00	
Di-n-octylphthalate	1	0	CC	12.67	50.45	50	20		2.190	2.210	0.90	
Benzo[b]fluoranthene	1	0		13.00	50.52	50			1.579	1.596	1.04	
Benzo[k]fluoranthene	1	0		13.03	46.72	50			1.599	1.494	6.56	
Benzo[a]pyrene	1	0	CC	13.33	49.53	50	20		1.486	1.472	0.94	
Indeno[1,2,3-cd]pyrene	1	0		14.41	51.89	50			1.610	1.671	3.78	
Dibenzo[a,h]anthracene	1	0		14.44	51.83	50			1.336	1.385	3.66	
Benzo[g,h,i]perylene	1	0		14.68	51.27	50			1.347	1.381	2.54	
Diaminotoluene Dihydrochloride	1	1E		0.00	0.00	50				0.000	100.00	
Toluene Diisocyanate	1	1E		0.00	0.00	50				0.000	100.00	
2,4 Diaminotoluene	1	1E		0.00	0.00	50				0.000	100.00	

CC - Continuing Calibration Check Compound
 N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
 * - Failed the C or P Criteria

Page 2 of 2

Note:

8260/8270 limits are compared against the %DIFF/R.F.
 624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
 524.2 limits are compared against the %DIFF

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09827.D Vial: 2
 Acq On : 8 Aug 2005 6:40 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 8 7:16 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	33792	40.00	ng	-0.15
20) Naphthalene-d8	6.13	136	125969	40.00	ng	-0.15
36) Acenaphthene-d10	7.47	164	69689	40.00	ng	-0.17
61) Phenanthrene-d10	8.83	188	114519	40.00	ng	-0.20
77) Chrysene-d12	11.81	240	88593	40.00	ng	-0.23
88) Perylene-d12	13.39	264	67613	40.00	ng	-0.23

System Monitoring Compounds

4) 2-Fluorophenol	3.77	112	56303	49.47	ng	-0.20
Spiked Amount	200.000		Recovery	=	24.74%	
8) Phenol-d5	4.80	99	71095	42.72	ng	-0.15
Spiked Amount	200.000		Recovery	=	21.36%	
21) Nitrobenzene-d5	5.57	128	13780	24.98	ng	-0.15
Spiked Amount	100.000		Recovery	=	24.98%	
41) 2-Fluorobiphenyl	6.94	172	55262	25.37	ng	-0.15
Spiked Amount	100.000		Recovery	=	25.37%	
64) 2,4,6-Tribromophenol	8.16	330	11852	48.36	ng	-0.19
Spiked Amount	200.000		Recovery	=	24.18%	
80) Terphenyl-d14	10.60	244	52371	25.02	ng	-0.21
Spiked Amount	100.000		Recovery	=	25.02%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	1.95	79	68024	48.27	ng	91
3) N-Nitrosodimethylamine	1.91	74	39089	45.61	ng	92
5) Aniline	4.80	93	90913	47.59	ng	89
6) Pentachloroethane	4.83	117	21557	51.74	ng	99
7) bis(2-Chloroethyl)ether	4.88	93	58805	49.18	ng	96
9) Phenol	4.81	94	83680	47.44	ng	83
10) 2-Chlorophenol	4.90	128	62529	46.70	ng	97
11) 1,3-Dichlorobenzene	5.04	146	63602	51.34	ng	99
12) 1,4-Dichlorobenzene	5.11	146	63472	50.07	ng	100
13) 1,2-Dichlorobenzene	5.23	146	59821	49.50	ng	99
14) Benzyl alcohol	5.23	108	42447	48.01	ng	92
15) bis(2-chloroisopropyl)ethe	5.35	45	94077	51.80	ng	96
16) 2-Methylphenol	5.34	108	57680	47.18	ng	97
17) Hexachloroethane	5.52	117	26411	50.23	ng	88
18) N-Nitroso-di-n-propylamine	5.46	70	46530	48.28	ng	97
19) 3&4-Methylphenol	5.47	108	58288	44.85	ng	99
22) Nitrobenzene	5.58	77	62933	50.87	ng	98
23) Isophorone	5.78	82	112097	48.68	ng	92
24) 2-Nitrophenol	5.84	139	32192	50.71	ng	97

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09827.D Vial: 2
 Acq On : 8 Aug 2005 6:40 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 7:16 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) 2,4-Dimethylphenol	5.89	107	56402	46.70	ng	98
26) Benzoic Acid	5.99	105	25319	36.52	ng	94
27) bis(2-Chloroethoxy)methane	5.96	93	70311	53.36	ng	99
28) 2,4-Dichlorophenol	6.03	162	47316	46.99	ng	97
29) 1,2,4-Trichlorobenzene	6.09	180	51270	49.66	ng	99
30) Naphthalene	6.15	128	160709	48.69	ng	99
31) 4-Chloroaniline	6.20	127	65814	51.06	ng	98
32) Hexachlorobutadiene	6.24	225	29301	51.47	ng	98
33) 4-Chloro-3-methylphenol	6.56	107	48794	43.88	ng	90
34) 2-Methylnaphthalene	6.67	142	107491	47.15	ng	99
35) Methylnaphthalenes (Total)	6.67	142	107491	47.15	ng	99
37) 1,2,4,5-Tetrachlorobenzene	6.79	216	46818	49.75	ng	97
38) Hexachlorocyclopentadiene	6.78	237	32392	55.88	ng	99
39) 2,4,6-Trichlorophenol	6.87	196	32717	48.23	ng	95
40) 2,4,5-Trichlorophenol	6.90	196	34188	46.34	ng	98
42) 2-Chloronaphthalene	7.03	162	99890	50.71	ng	97
43) 1,4-Dimethylnaphthalene	7.29	156	73259	48.87	ng	100
44) Dimethylnaphthalenes (Total)	7.29	156	73259	48.87	ng	100
45) Diphenyl Ether	7.10	170	85800	65.92	ng	94
46) 2-Nitroaniline	7.11	65	41323	55.08	ng	97
47) Acenaphthylene	7.35	152	146562	47.10	ng	99
48) Dimethylphthalate	7.26	163	109773	48.28	ng	98
49) 2,6-Dinitrotoluene	7.30	165	25439	48.57	ng	99
50) Acenaphthene	7.49	153	97850	50.84	ng	98
51) 3-Nitroaniline	7.44	138	28568	49.97	ng	99
52) 2,4-Dinitrophenol	7.52	184	15178	47.10	ng	91
53) Dibenzofuran	7.64	168	136307	48.65	ng	97
54) 2,4-Dinitrotoluene	7.63	165	33353	46.10	ng	92
55) 4-Nitrophenol	7.57	65	22595	49.64	ng	94
56) 2,3,4,6-Tetrachlorophenol	7.75	232	26621	46.00	ng	99
57) Fluorene	7.94	166	107321	47.54	ng	99
58) 4-Chlorophenyl-phenylether	7.94	204	52377	47.73	ng	94
59) Diethylphthalate	7.85	149	111551	47.87	ng	98
60) 4-Nitroaniline	7.96	138	29779	44.90	ng	94
62) 4,6-Dinitro-2-methylphenol	7.99	198	21479	48.78	ng	100
63) n-Nitrosodiphenylamine	8.05	169	77241	48.93	ng	99
65) 1,2-Diphenylhydrazine	8.08	77	123404	54.78	ng	97
66) 4-Bromophenyl-phenylether	8.40	248	27731	47.23	ng	99
67) Hexachlorobenzene	8.45	284	26287	47.42	ng	86
68) gamma-BHC	8.70	181	4212	10.38	ng	95
69) Pentachlorophenol	8.64	266	17664	48.61	ng	92

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09827.D Vial: 2
 Acq On : 8 Aug 2005 6:40 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 7:16 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Phenanthrene	8.85	178	158972	48.13	ng	99
71) Anthracene	8.90	178	160781	47.90	ng	98
72) Carbazole	9.08	167	149619	48.82	ng	99
73) Heptachlor	9.35	100	4417	10.30	ng	95
74) Di-n-butylphthalate	9.49	149	185971	50.12	ng	99
75) Heptachlor epoxide	10.04	81	2925	9.82	ng	91
76) Fluoranthene	10.13	202	172928	48.03	ng	95
78) Pyrene	10.39	202	181091	51.04	ng	97
79) Benzidine	10.31	184	66423	50.70	ng	98
81) Endrin	10.83	81	1860	10.65	ng	90
82) Butylbenzylphthalate	11.21	149	80205	51.36	ng	94
83) Methoxychlor	11.84	227	16367	10.05	ng	99
84) 3,3'-Dichlorobenzidine	11.79	252	51680	50.67	ng	99
85) Benzo[a]anthracene	11.79	228	160700	49.33	ng	99
86) Chrysene	11.84	228	147755	49.45	ng	99
87) bis(2-Ethylhexyl)phthalate	11.93	149	111650	51.74	ng	97
89) Di-n-octylphthalate	12.67	149	186771	50.45	ng	99
90) Benzo[b]fluoranthene	13.00	252	134882	50.52	ng	99
91) Benzo[k]fluoranthene	13.03	252	126248	46.72	ng	96
92) Benzo[a]pyrene	13.33	252	124426	49.53	ng	97
93) Indeno[1,2,3-cd]pyrene	14.41	276	141245	51.89	ng	92
94) Dibenzo[a,h]anthracene	14.44	278	117094	51.83	ng	96
95) Benzo[g,h,i]perylene	14.68	276	116715	51.27	ng	94

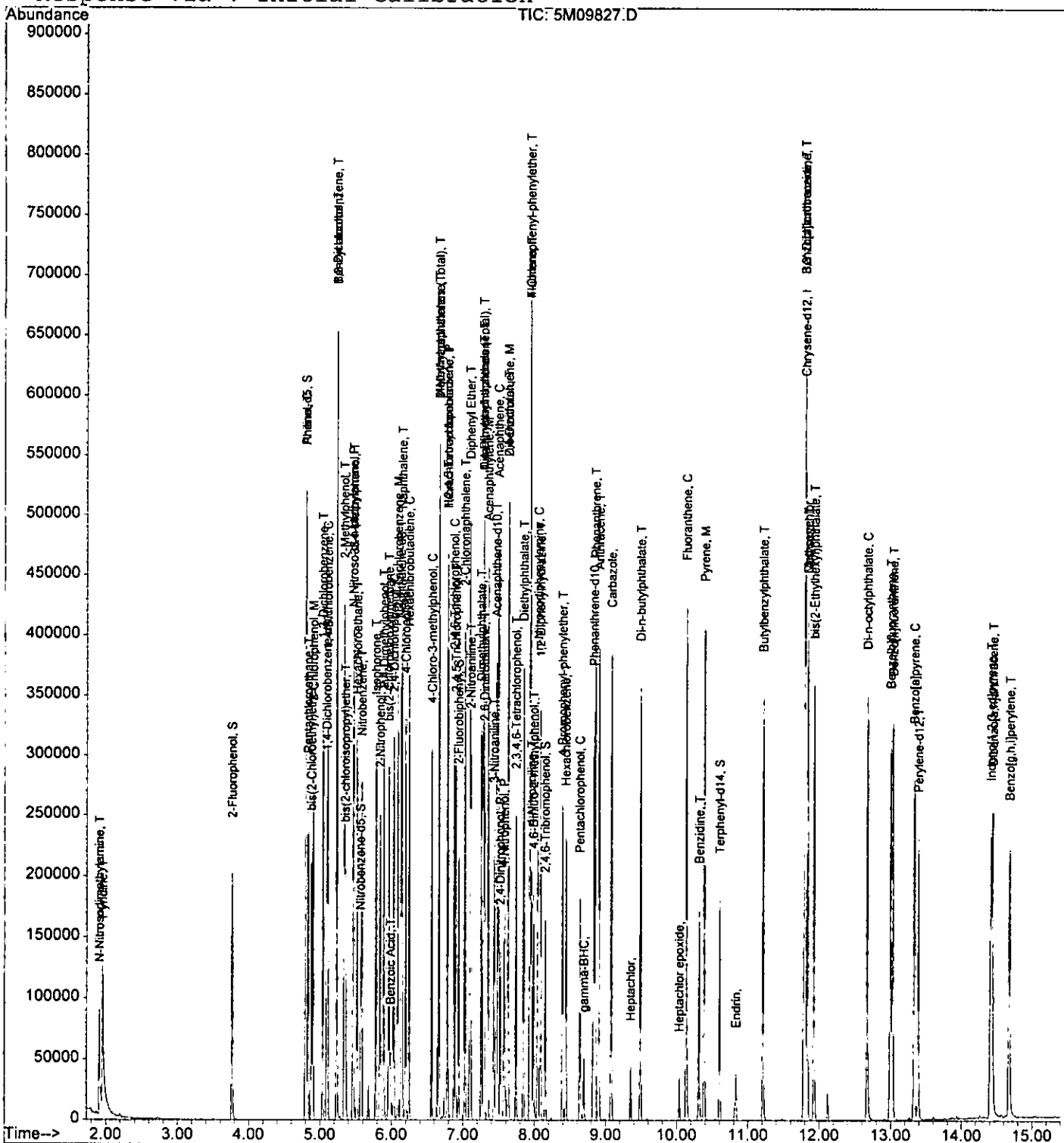
(#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09827.D Vial: 2
 Acq On : 8 Aug 2005 6:40 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 7:16 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/8/05 6:59:00 AM

Data File: 4M05426.D
Method: 8270

Instrument: GCMS_4

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
1,4-Dichlorobenzene-d4	1	0	I	4.90	40.00	40				0.000	0.00	
Pyridine	1	0		2.28	55.57	50			1.515	1.684	11.14	
N-Nitrosodimethylamine	1	0		2.22	47.33	50			0.917	0.868	5.34	
2-Fluorophenol	1	0	S	3.75	49.89	50			1.128	1.126	0.22	
Aniline	1	0		4.63	52.98	50			1.631	1.729	5.96	
bis(2-Chloroethyl)ether	1	0		4.69	54.15	50			1.226	1.328	8.30	
Phenol-d5	1	0	S	4.61	51.12	50			1.502	1.536	2.24	
Phenol	1	0	CC	4.62	53.85	50	20		1.625	1.750	7.70	
2-Chlorophenol	1	0		4.73	53.31	50			1.244	1.327	6.62	
1,3-Dichlorobenzene	1	0		4.85	54.32	50			1.335	1.451	8.64	
1,4-Dichlorobenzene	1	0	CC	4.91	55.18	50	20		1.306	1.442	10.36	
1,2-Dichlorobenzene	1	0		5.03	54.94	50			1.281	1.408	9.88	
Benzyl alcohol	1	0		5.02	49.27	50			0.775	0.764	1.46	
bis(2-chloroisopropyl)ether	1	0		5.12	46.67	50			3.011	2.810	6.66	
2-Methylphenol	1	0		5.11	50.65	50			1.038	1.051	1.30	
Hexachloroethane	1	0		5.30	54.13	50			0.610	0.660	8.26	
N-Nitroso-di-n-propylamine	1	0	CP	5.23	53.40	50	0.05		1.070	1.142	6.80	
3&4-Methylphenol	1	0		5.24	49.03	50			1.070	1.050	1.94	
Naphthalene-d8	1	0	I	5.90	40.00	40				0.000	0.00	
Nitrobenzene-d5	1	0	S	5.34	25.74	25			0.200	0.206	2.96	
Nitrobenzene	1	0		5.35	50.01	50			0.417	0.417	0.02	
Isophorone	1	0		5.54	47.40	50			0.797	0.755	5.20	
2-Nitrophenol	1	0	CC	5.60	48.63	50	20		0.216	0.210	2.74	
2,4-Dimethylphenol	1	0		5.65	45.18	50			0.398	0.360	9.64	
Benzoic Acid	1	0		5.72	48.51	50			0.069	0.067	2.98	
bis(2-Chloroethoxy)methane	1	0		5.72	48.78	50			0.476	0.464	2.44	
2,4-Dichlorophenol	1	0	CC	5.79	48.99	50	20		0.327	0.321	2.02	
1,2,4-Trichlorobenzene	1	0		5.86	51.89	50			0.351	0.364	3.78	
Naphthalene	1	0		5.92	53.01	50			0.879	0.932	6.02	
4-Chloroaniline	1	0		5.96	61.76	50			0.342	0.422	23.52	
Hexachlorobutadiene	1	0	CC	6.01	50.59	50	20		0.225	0.228	1.18	
4-Chloro-3-methylphenol	1	0	CC	6.36	46.59	50	20		0.365	0.340	6.82	
2-Methylnaphthalene	1	0		6.49	52.47	50			0.615	0.645	4.94	
Methylnaphthalene	1	0		6.49	52.47							
Acenaphthene-d10	1	0	I	7.48	40.00	40				0.000	0.00	
1,2,4,5-Tetrachlorobenzene	1	0		6.65	53.76	50			0.653	0.702	7.52	
Hexachlorocyclopentadiene	1	0	CP	6.64	50.97	50	0.05		0.459	0.468	1.94	
2,4,6-Trichlorophenol	1	0	CC	6.75	53.46	50	20		0.470	0.503	6.92	
2,4,5-Trichlorophenol	1	0		6.78	56.94	50			0.482	0.549	13.88	
2-Fluorobiphenyl	1	0	S	6.82	25.20	25			1.282	1.292	0.80	
2-Chloronaphthalene	1	0		6.93	54.37	50			1.120	1.218	8.74	
2-Nitroaniline	1	0		7.04	47.19	50			0.663	0.626	5.62	
1,4-Dimethylnaphthalene	1	0		7.25	53.61	50			0.770	0.826	7.22	
Dimethylnaphthalene	1	0		7.25	53.61							
Diphenyl Ether	1	0		7.02	52.26	50			0.977	1.021	4.52	
Acenaphthylene	1	0		7.33	53.71	50			1.719	1.847	7.42	
Dimethylphthalate	1	0		7.21	49.56	50			1.413	1.401	0.88	
2,6-Dinitrotoluene	1	0		7.27	54.96	50			0.337	0.370	9.92	
Acenaphthene	1	0	CC	7.51	53.09	50	20		1.127	1.196	6.18	
3-Nitroaniline	1	0		7.43	63.11	50			0.294	0.370	26.22	
2,4-Dinitrophenol	1	0	CP	7.55	42.85	50	0.05		0.191	0.164	14.30	
Dibenzofuran	1	0		7.68	54.74	50			1.485	1.626	9.48	
2,4-Dinitrotoluene	1	0		7.68	53.92	50			0.440	0.475	7.84	
4-Nitrophenol	1	0	CP	7.61	42.63	50	0.05		0.377	0.322	14.74	
Fluorene	1	0		8.04	55.64	50			1.092	1.216	11.28	
4-Chlorophenyl-phenylether	1	0		8.05	51.30	50			0.656	0.673	2.60	
Diethylphthalate	1	0		7.94	48.74	50			1.474	1.437	2.52	
4-Nitroaniline	1	0		8.07	47.40	50			0.378	0.358	5.20	
Phenanthrene-d10	1	0	I	9.07	40.00	40				0.000	0.00	
4,6-Dinitro-2-methylphenol	1	0		8.10	51.60	50			0.144	0.148	3.20	
n-Nitrosodiphenylamine	1	0	CC	8.17	54.54	50	20		0.507	0.553	9.08	
2,4,6-Tribromophenol	1	0	S	8.30	54.86	50			0.179	0.196	9.72	
1,2-Diphenylhydrazine	1	0		8.21	49.87	50			0.910	0.907	0.26	
4-Bromophenyl-phenylether	1	0		8.58	52.86	50			0.259	0.273	5.72	
Hexachlorobenzene	1	0		8.63	54.54	50			0.347	0.379	9.08	
Pentachlorophenol	1	0	CC	8.87	47.32	50	20		0.189	0.179	5.36	
Phenanthrene	1	0		9.10	55.04	50			0.999	1.100	10.08	
Anthracene	1	0		9.15	55.23	50			1.015	1.121	10.46	
Carbazole	1	0		9.35	56.49	50			0.931	1.052	12.98	
Di-n-butylphthalate	1	0		9.80	52.33	50			1.362	1.425	4.66	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 1 of 2

** - No limit specified in method

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Form 7

Continuing Calibration

Calibration Name: CAL BNA@50PPM
Cont Calibration Date/Time 8/8/05 6:59:00 AM

Data File: 4M05426.D
Method: 8270

Instrument: GCMS_4

TxtCompd:	Col	Multi Num	Type	RT	Conc	Conc Exp	Lo Lim	Hi Lim	Initial RF	RF	%Diff	Flag
Fluoranthene	1	0	CC	10.49	58.26	50	20		1.039	1.211	16.52	
Chrysene-d12	1	0	I	12.27	40.00	40				0.000	0.00	
Pyrene	1	0		10.75	46.49	50			1.538	1.431	7.02	
Benzidine	1	0		10.68	51.73	50			0.434	0.449	3.46	
Terphenyl-d14	1	0	S	10.97	22.47	25			1.127	1.013	10.12	
Butylbenzylphthalate	1	0		11.61	48.46	50			0.750	0.727	3.08	
3,3'-Dichlorobenzidine	1	0		12.26	84.71	50			0.303	0.513	69.42	
Benzo[a]anthracene	1	0		12.26	53.25	50			1.255	1.337	6.50	
Chrysene	1	0		12.30	52.96	50			1.121	1.188	5.92	
bis(2-Ethylhexyl)phthalate	1	0		12.39	55.37	50			0.976	1.080	10.74	
Perylene-d12	1	0	I	14.12	40.00	40				0.000	0.00	
Di-n-octylphthalate	1	0	CC	13.26	47.50	50	20		2.191	2.082	5.00	
Benzo[b]fluoranthene	1	0		13.65	47.03	50			1.664	1.565	5.94	
Benzo[k]fluoranthene	1	0		13.69	49.49	50			1.441	1.426	1.02	
Benzo[a]pyrene	1	0	CC	14.06	49.74	50	20		1.368	1.361	0.52	
Indeno[1,2,3-cd]pyrene	1	0		15.37	61.29	50			1.241	1.521	22.58	
Dibenzo[a,h]anthracene	1	0		15.39	61.30	50			1.016	1.246	22.60	
Benzo[g,h,i]perylene	1	0		15.64	63.75	50			0.990	1.263	27.50	
2,4 Diaminotoluene	1	1E		0.00	0.00	50				0.000	100.00	
Heptachlor_epoxide	1	1E		0.00	0.00	10				0.000	100.00	
Methoxychlor	1	1E		0.00	0.00	10				0.000	100.00	
Diaminotoluene Dihydrochloride	1	1E		0.00	0.00	50				0.000	100.00	
Toluene Diisocyanate	1	1E		0.00	0.00	50				0.000	100.00	
Endrin	1	1E		0.00	0.00	10				0.000	100.00	
2,3,4,6-Tetrachlorophenol	1	1E		0.00	0.00	50				0.000	100.00	
gamma-BHC	1	1E		0.00	0.00	10				0.000	100.00	
Pentachloroethane	1	1E		0.00	0.00	50				0.000	100.00	
Heptachlor	1	1E		0.00	0.00	10				0.000	100.00	

CC - Continuing Calibration Check Compound
N/O or N/Q - Not applicable for this run

CP - System Performance Check Compound I - Internal Standard
* - Failed the C or P Criteria

Page 2 of 2

Note:

8260/8270 limits are compared against the %DIFF/R.F.
624 limits are compared against the concentration found.

625 limits are compared against the %DIFF.
524.2 limits are compared against the %DIFF

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05426.D Vial: 2
 Acq On : 8 Aug 2005 6:59 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 8 7:16 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.90	152	30327	40.00	ng	-0.04
19) Naphthalene-d8	5.90	136	95647	40.00	ng	-0.04
35) Acenaphthene-d10	7.48	164	50656	40.00	ng	-0.05
59) Phenanthrene-d10	9.07	188	82469	40.00	ng	-0.06
72) Chrysene-d12	12.27	240	68852	40.00	ng	-0.06
81) Perylene-d12	14.12	264	58010	40.00	ng	-0.06

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.75	112	42680	49.89	ng	-0.04
Spiked Amount				200.000		
			Recovery	=	24.95%	
7) Phenol-d5	4.61	99	58212	51.12	ng	-0.04
Spiked Amount				200.000		
			Recovery	=	25.56%	
20) Nitrobenzene-d5	5.34	128	12324	25.74	ng	-0.04
Spiked Amount				100.000		
			Recovery	=	25.74%	
40) 2-Fluorobiphenyl	6.82	172	40907	25.20	ng	-0.05
Spiked Amount				100.000		
			Recovery	=	25.20%	
62) 2,4,6-Tribromophenol	8.30	332	20249	54.86	ng	-0.05
Spiked Amount				200.000		
			Recovery	=	27.43%	
75) Terphenyl-d14	10.97	244	43587	22.47	ng	-0.05
Spiked Amount				100.000		
			Recovery	=	22.47%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	2.28	79	63836	55.57	ng	96
3) N-Nitrosodimethylamine	2.22	74	32900	47.33	ng	99
5) Aniline	4.63	93	65532	52.98	ng	40
6) bis(2-Chloroethyl)ether	4.69	93	50356	54.15	ng	85
8) Phenol	4.62	94	66352	53.85	ng	78
9) 2-Chlorophenol	4.73	128	50287	53.31	ng	86
10) 1,3-Dichlorobenzene	4.85	146	54991	54.32	ng	99
11) 1,4-Dichlorobenzene	4.91	146	54650	55.18	ng	98
12) 1,2-Dichlorobenzene	5.03	146	53366	54.94	ng	97
13) Benzyl alcohol	5.02	108	28961	49.27	ng	94
14) bis(2-chloroisopropyl)ethe	5.12	45	106527	46.67	ng	98
15) 2-Methylphenol	5.11	108	39843	50.65	ng	100
16) Hexachloroethane	5.30	117	25036	54.13	ng	65
17) N-Nitroso-di-n-propylamine	5.23	70	43297	53.40	ng	88
18) 3&4-Methylphenol	5.24	108	39790	49.03	ng	94
21) Nitrobenzene	5.35	77	49858	50.01	ng	84
22) Isophorone	5.54	82	90295	47.40	ng	98
23) 2-Nitrophenol	5.60	139	25087	48.63	ng	73
24) 2,4-Dimethylphenol	5.65	107	43050	45.18	ng	97

(#) = qualifier out of range (m) = manual integration

Handwritten signature

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05426.D Vial: 2
 Acq On : 8 Aug 2005 6:59 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 7:16 2005 Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
25) Benzoic Acid	5.72	105	8028	48.51	ng	87
26) bis(2-Chloroethoxy)methane	5.72	93	55518	48.78	ng	98
27) 2,4-Dichlorophenol	5.79	162	38341	48.99	ng	98
28) 1,2,4-Trichlorobenzene	5.86	180	43553	51.89	ng	97
29) Naphthalene	5.92	128	111383	53.01	ng	98
30) 4-Chloroaniline	5.96	127	50447	61.76	ng	98
31) Hexachlorobutadiene	6.01	225	27247	50.59	ng	95
32) 4-Chloro-3-methylphenol	6.36	107	40654	46.59	ng	80
33) 2-Methylnaphthalene	6.49	142	77140	52.47	ng	99
34) Methylnaphthalene (Total)	6.49	142	77140	52.47	ng	99
36) 1,2,4,5-Tetrachlorobenzene	6.65	216	44457	53.76	ng	97
37) Hexachlorocyclopentadiene	6.64	237	29649	50.97	ng	98
38) 2,4,6-Trichlorophenol	6.75	196	31848	53.46	ng	98
39) 2,4,5-Trichlorophenol	6.78	196	34746	56.94	ng	99
41) 2-Chloronaphthalene	6.93	162	77118	54.37	ng	100
42) 2-Nitroaniline	7.04	65	39629	47.19	ng	64
43) 1,4-Dimethylnaphthalene	7.25	156	52295	53.61	ng	91
44) Dimethylnaphthalene (Total)	7.25	156	52295	53.61	ng	91
45) Diphenyl Ether	7.02	170	64679	52.26	ng	93
46) Acenaphthylene	7.33	152	116924	53.71	ng	98
47) Dimethylphthalate	7.21	163	88714	49.56	ng	100
48) 2,6-Dinitrotoluene	7.27	165	23420	54.96	ng	96
49) Acenaphthene	7.51	153	75753	53.09	ng	96
50) 3-Nitroaniline	7.43	138	23458	63.11	ng	99
51) 2,4-Dinitrophenol	7.55	184	10355	42.85	ng	99
52) Dibenzofuran	7.68	168	102961	54.74	ng	94
53) 2,4-Dinitrotoluene	7.68	165	30050	53.92	ng	87
54) 4-Nitrophenol	7.61	65	20360	42.63	ng	97
55) Fluorene	8.04	166	76972	55.64	ng	97
56) 4-Chlorophenyl-phenylether	8.05	204	42603	51.30	ng	95
57) Diethylphthalate	7.94	149	90970	48.74	ng	99
58) 4-Nitroaniline	8.07	138	22686	47.40	ng	95
60) 4,6-Dinitro-2-methylphenol	8.10	198	15268	51.60	ng	100
61) n-Nitrosodiphenylamine	8.17	169	57057	54.54	ng	99
63) 1,2-Diphenylhydrazine	8.21	77	93513	49.87	ng	98
64) 4-Bromophenyl-phenylether	8.58	248	28182	52.86	ng	93
65) Hexachlorobenzene	8.63	284	39056	54.54	ng	79
66) Pentachlorophenol	8.87	266	18462	47.32	ng	96
67) Phenanthrene	9.10	178	113373	55.04	ng	98
68) Anthracene	9.15	178	115531	55.23	ng	99
69) Carbazole	9.35	167	108457	56.49	ng	98

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05426.D Vial: 2
 Acq On : 8 Aug 2005 6:59 Operator: AHD
 Sample : CAL BNA@50PPM Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 8 7:16 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
70) Di-n-butylphthalate	9.80	149	146922	52.33	ng	98
71) Fluoranthene	10.49	202	124842	58.26	ng	85
73) Pyrene	10.75	202	123125	46.49	ng	99
74) Benzidine	10.68	184	38609	51.73	ng	96
76) Butylbenzylphthalate	11.61	149	62562	48.46	ng	96
77) 3,3'-Dichlorobenzidine	12.26	252	44111	84.71	ng	96
78) Benzo[a]anthracene	12.26	228	115071	53.25	ng	100
79) Chrysene	12.30	228	102231	52.96	ng	98
80) bis(2-Ethylhexyl)phthalate	12.39	149	92969	55.37	ng	97
82) Di-n-octylphthalate	13.26	149	150947	47.50	ng	99
83) Benzo[b]fluoranthene	13.65	252	113500	47.03	ng	97
84) Benzo[k]fluoranthene	13.69	252	103425	49.49	ng	96
85) Benzo[a]pyrene	14.06	252	98669	49.74	ng	94
86) Indeno[1,2,3-cd]pyrene	15.37	276	110279	61.29	ng	79
87) Dibenzo[a,h]anthracene	15.39	278	90317	61.30	ng	96
88) Benzo[g,h,i]perylene	15.64	276	91551	63.75	ng	93

 (#) = qualifier out of range (m) = manual integration

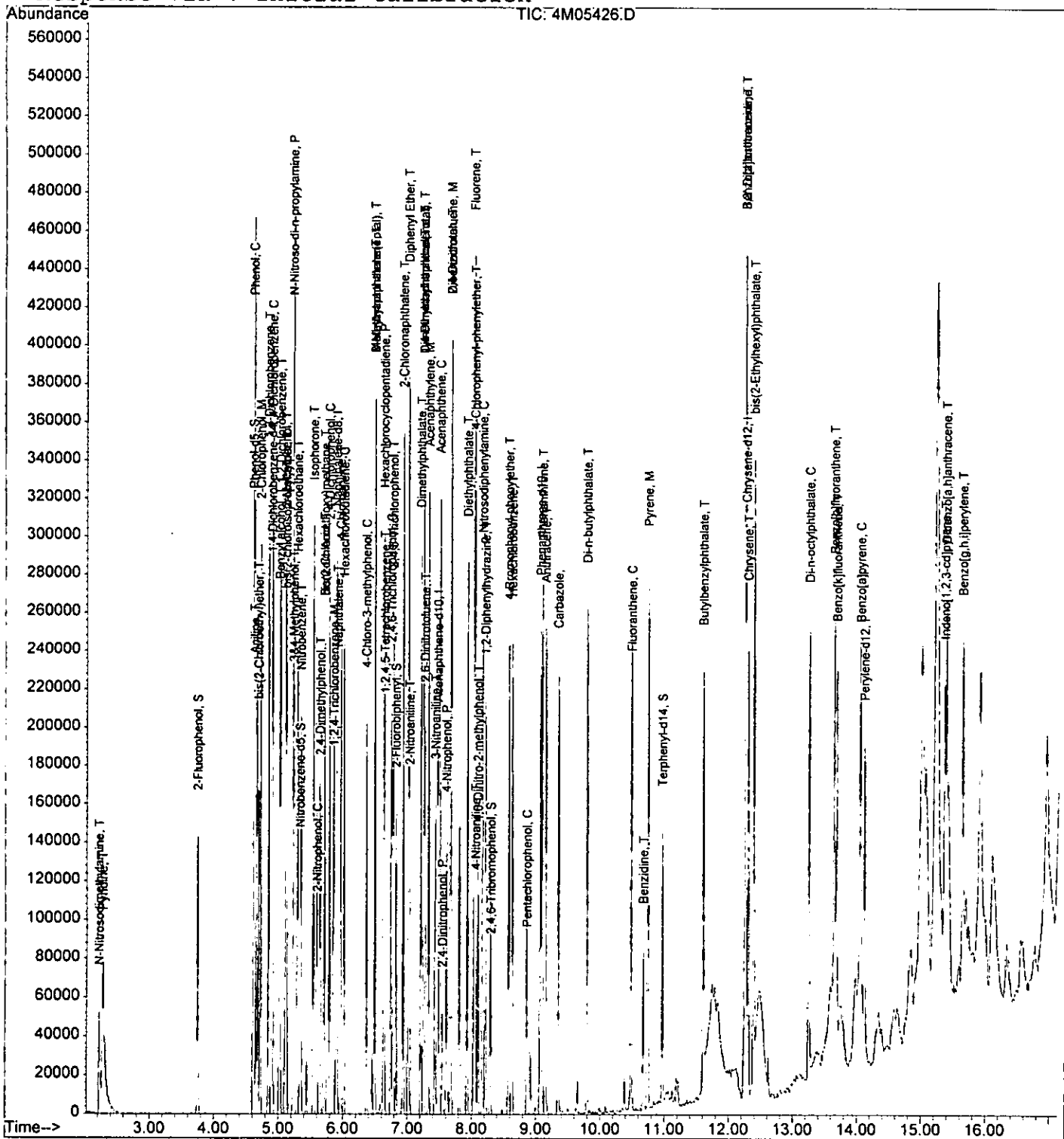
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05426.D Vial: 2
Acq On : 8 Aug 2005 6:59 Operator: AHD
Sample : CAL BNA@50PPM Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 8 7:16 2005



Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



GC/MS Semi-Volatile Data
Raw QC Data

Form 5

Tune Name: CAL DFTPP

Data File: 5M09384.D

Instrument: GCMS_5

Analysis Date: 07/22/05 08:08

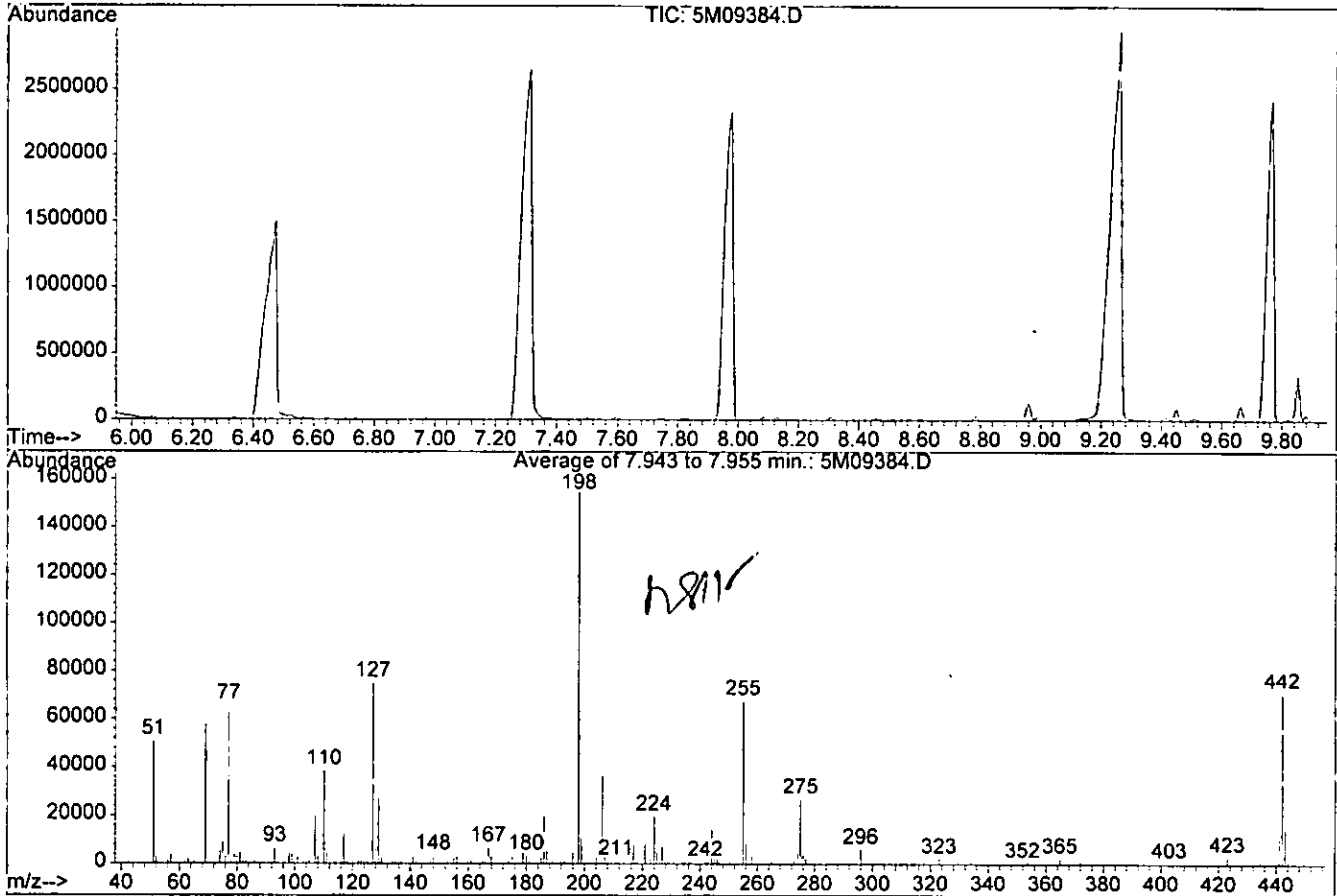
Tune Scan/Time Range: Average of 7.943 to 7.955 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/ Fail
51	198	30	60	32.8	50680	PASS
68	69	0.00	2	0.4	237	PASS
69	198	0.00	100	37.6	58061	PASS
70	69	0.00	2	0.0	0	PASS
127	198	40	60	48.5	74997	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	154539	PASS
199	198	5	9	6.9	10733	PASS
275	198	10	30	17.4	26872	PASS
365	198	1	100	1.5	2276	PASS
441	443	0.01	100	76.5	10892	PASS
442	198	40	100	46.1	71181	PASS
443	442	17	23	20.0	14247	PASS

Data File	Sample Number	Analysis Date:
5M09385.D	CAL BNA@50PPM	07/22/05 08:30
5M09386.D	CAL BNA@10PPM	07/22/05 08:53
5M09387.D	CAL BNA@25PPM	07/22/05 09:16
5M09388.D	CAL BNA@80PPM	07/22/05 09:39
5M09389.D	CAL BNA@120PP	07/22/05 10:01
5M09390.D	CAL BNA@160PP	07/22/05 10:24
5M09391.D	CAL BNA@200PP	07/22/05 10:47
5M09392.D	AC18716-003	07/22/05 11:29
5M09393.D	AC18623-013(R)	07/22/05 11:52
5M09394.D	AC18669-004(T)	07/22/05 12:15
5M09395.D	WMB2620	07/22/05 12:38
5M09396.D	AC18716-001	07/22/05 13:00
5M09397.D	AC18716-002	07/22/05 13:23
5M09398.D	WMB2620(MS)	07/22/05 13:46
5M09399.D	AC18623-007(R)	07/22/05 14:09
5M09400.D	WMB2621	07/22/05 14:32
5M09401.D	WMB2621(MS)	07/22/05 14:55
5M09402.D	AC18667-001	07/22/05 15:18
5M09403.D	AC18667-001(MS)	07/22/05 15:41
5M09404.D	AC18667-001(MS)	07/22/05 16:04
5M09405.D	SMB2594	07/22/05 16:27
5M09406.D	SMB2594(MS)	07/22/05 16:50
5M09407.D	AC18689-002	07/22/05 17:13
5M09408.D	AC18689-002(MS)	07/22/05 17:36
5M09409.D	AC18689-002(MS)	07/22/05 17:59
5M09410.D	AC18689-007	07/22/05 18:22
5M09411.D	AC18475-001(T)	07/22/05 18:46
5M09412.D	EF2V4993	07/22/05 19:09
5M09413.D	AC18681-001(5X)	07/22/05 19:31
5M09414.D	AC18657-001	07/22/05 19:54
5M09415.D	AC18666-001	07/22/05 20:17
5M09416.D	AC18691-001	07/22/05 20:40
5M09417.D	AC18698-005	07/22/05 21:03
5M09418.D	AC18661-001(R)	07/22/05 21:26
5M09419.D	AC18711-001	07/22/05 21:49

DFTPP

Data File : G:\GcMsData\2005\Gcms_5\Data\07-2205\5M09384.D Vial: 1
 Acq On : 22 Jul 2005 8:08 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0711.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 7.943 to 7.955 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	32.8	50680	PASS
68	69	0.00	2	0.4	237	PASS
69	198	0.00	100	37.6	58061	PASS
70	69	0.00	2	0.0	0	PASS
127	198	40	60	48.5	74997	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	154539	PASS
199	198	5	9	6.9	10733	PASS
275	198	10	30	17.4	26872	PASS
365	198	1	100	1.5	2276	PASS
441	443	0.01	100	76.5	10892	PASS
442	198	40	100	46.1	71181	PASS
443	442	17	23	20.0	14247	PASS

Form 5

Tune Name: CAL DFTPP

Data File: 4M05297.D

Instrument: GCMS_4

Analysis Date: 08/03/05 08:09

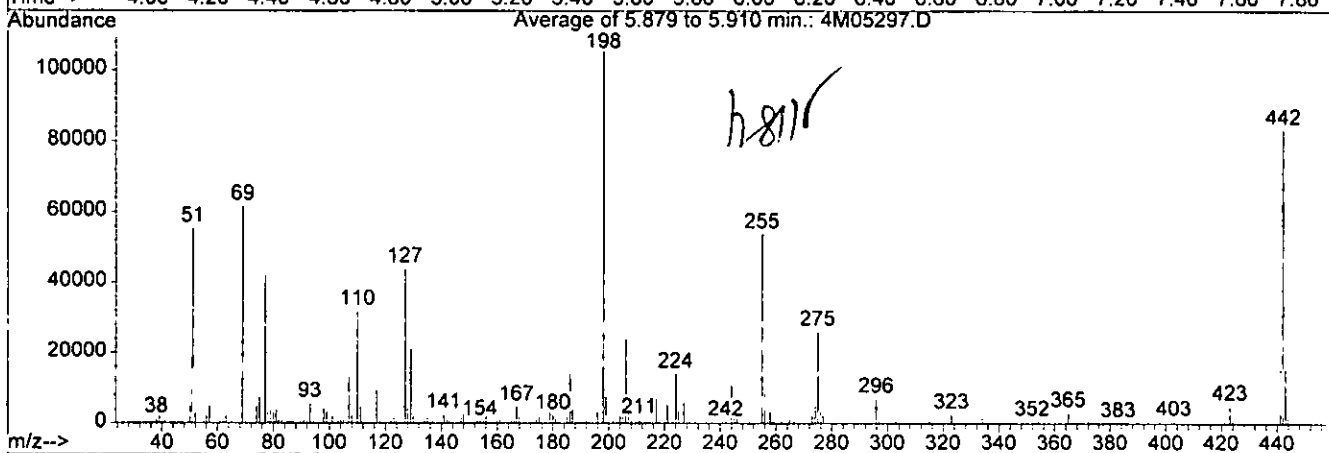
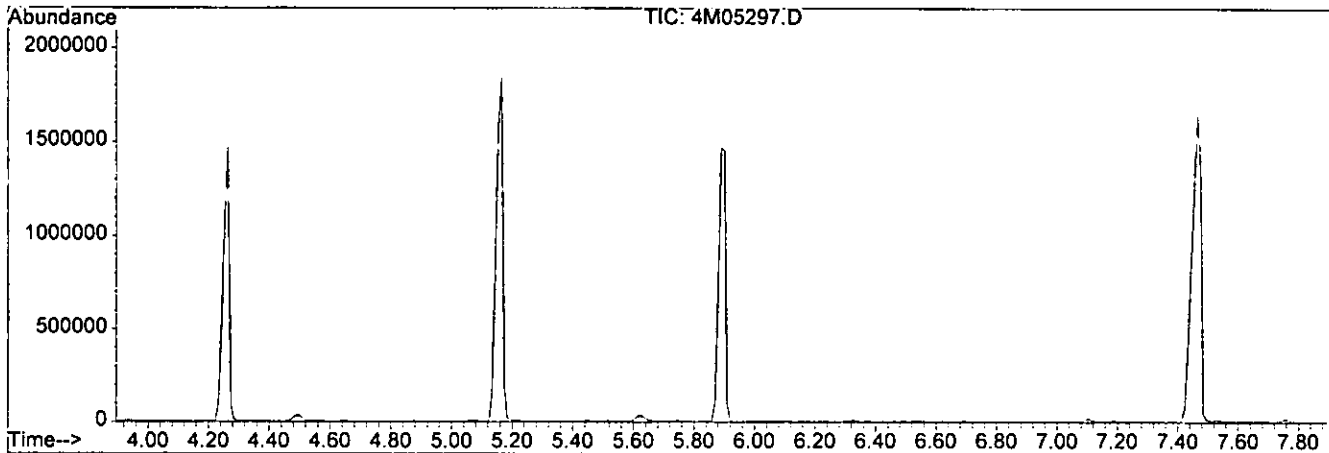
Tune Scan/Time Range: Average of 5.879 to 5.910 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/Fail
51	198	30	60	52.6	55404	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	58.5	61615	PASS
70	69	0.00	2	0.5	338	PASS
127	198	40	60	41.7	43931	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	105386	PASS
199	198	5	9	7.3	7679	PASS
275	198	10	30	24.7	26080	PASS
365	198	1	100	3.0	3133	PASS
441	443	0.01	100	91.8	15388	PASS
442	198	40	100	79.0	83290	PASS
443	442	17	23	20.1	16769	PASS

Data File	Sample Number	Analysis Date:
4M05299.D	CAL BNA@50PPM	08/03/05 08:52
4M05300.D	CAL BNA@10PPM	08/03/05 09:19
4M05301.D	CAL BNA@25PPM	08/03/05 09:43
4M05302.D	CAL BNA@80PPM	08/03/05 10:07
4M05303.D	CAL BNA@120PP	08/03/05 10:31
4M05304.D	CAL BNA@160PP	08/03/05 10:55
4M05305.D	CAL BNA@200PP	08/03/05 11:19
4M05306.D	SMB2606	08/03/05 11:43
4M05307.D	AC18819-004	08/03/05 12:06
4M05308.D	AC18819-006	08/03/05 12:30
4M05309.D	AC18819-012	08/03/05 12:54
4M05310.D	AC18819-018	08/03/05 13:18
4M05311.D	SMB2606	08/03/05 13:42
4M05312.D	SMB2605(MS)	08/03/05 14:06
4M05313.D	AC18819-008(MS)	08/03/05 14:30
4M05314.D	AC18819-008(MS)	08/03/05 14:54
4M05315.D	AC18802-004	08/03/05 15:18
4M05316.D	AC18802-006	08/03/05 15:41
4M05317.D	AC18853-002	08/03/05 16:05
4M05318.D	AC18853-003	08/03/05 16:29
4M05319.D	AC18853-004	08/03/05 16:53
4M05320.D	AC18808-001	08/03/05 17:17
4M05321.D	AC18802-002	08/03/05 17:42
4M05322.D	AC18802-005	08/03/05 18:06
4M05323.D	AC18852-001	08/03/05 18:30
4M05324.D	AC18853-001	08/03/05 18:54
4M05325.D	AC18847-001	08/03/05 19:18
4M05326.D	AC18802-001	08/03/05 19:42
4M05327.D	AC18786-013	08/03/05 20:06
4M05328.D	AC18786-014	08/03/05 20:30
4M05329.D	AC18796-007	08/03/05 20:54

DFTPP

Data File : G:\GcMsData\2005\Gcms_4\Data\08-03-05\4M05297.D Vial: 1
 Acq On : 3 Aug 2005 8:09 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_4
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 5.879 to 5.910 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	52.6	55404	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	58.5	61615	PASS
70	69	0.00	2	0.5	338	PASS
127	198	40	60	41.7	43931	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	105386	PASS
199	198	5	9	7.3	7679	PASS
275	198	10	30	24.7	26080	PASS
365	198	1	100	3.0	3133	PASS
441	443	0.01	100	91.8	15388	PASS
442	198	40	100	79.0	83290	PASS
443	442	17	23	20.1	16769	PASS

Form 5

Tune Name: CAL DFTPP
Instrument: GCMS_5

Data File: 5M09735.D
Analysis Date: 08/04/05 06:25

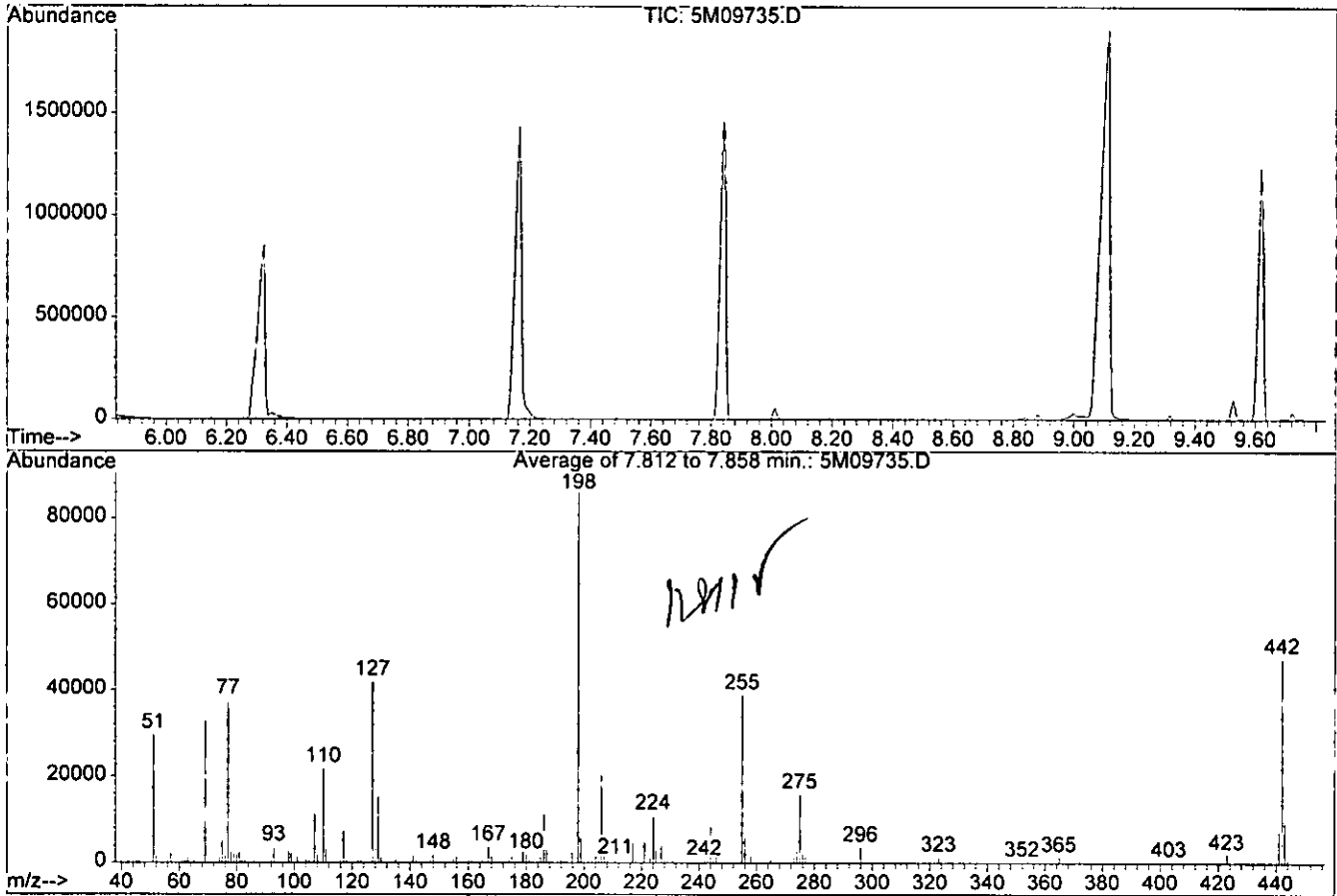
Tune Scan/Time Range: Average of 7.812 to 7.858 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/ Fail
51	198	30	60	34.5	29672	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	38.4	33009	PASS
70	69	0.00	2	0.2	53	PASS
127	198	40	60	48.9	42075	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	86042	PASS
199	198	5	9	6.7	5746	PASS
275	198	10	30	18.5	15876	PASS
365	198	1	100	1.6	1394	PASS
441	443	0.01	100	79.0	7136	PASS
442	198	40	100	55.2	47514	PASS
443	442	17	23	19.0	9034	PASS

Data File	Sample Number	Analysis Date:
5M09736.D	CAL BNA@50PPM	08/04/05 06:44
5M09737.D	WMB2631	08/04/05 07:06
5M09738.D	WMB2631(MS)	08/04/05 07:31
5M09739.D	AC18852-001(T)	08/04/05 07:53
5M09740.D	AC18832-004	08/04/05 08:15
5M09741.D	AC18832-004(MS)	08/04/05 08:37
5M09742.D	AC18832-004(MS)	08/04/05 08:59
5M09743.D	AC18897-001	08/04/05 09:20
5M09744.D	AC18897-002	08/04/05 09:42
5M09745.D	SMB2607	08/04/05 10:04
5M09746.D	SMB2608	08/04/05 10:26
5M09747.D	SMB2608(MS)	08/04/05 10:48
5M09748.D	AC18855-001	08/04/05 11:09
5M09749.D	AC18855-001(MS)	08/04/05 11:31
5M09750.D	AC18855-001(MS)	08/04/05 11:53
5M09751.D	AC18807-007	08/04/05 12:15
5M09752.D	AC18820-012	08/04/05 12:37
5M09753.D	AC18847-005	08/04/05 12:59
5M09754.D	AC18847-013	08/04/05 13:21
5M09755.D	AC18847-014	08/04/05 13:42
5M09756.D	AC18847-015	08/04/05 14:04
5M09757.D	AC18847-016	08/04/05 14:26
5M09758.D	AC18847-017	08/04/05 14:48
5M09759.D	AC18786-005	08/04/05 15:10
5M09760.D	AC18786-007	08/04/05 15:32
5M09761.D	AC18786-008	08/04/05 15:54
5M09762.D	AC18786-009	08/04/05 16:16
5M09763.D	AC18786-010	08/04/05 16:38
5M09764.D	AC18786-017	08/04/05 17:00
5M09765.D	AC18796-015	08/04/05 17:22
5M09766.D	AC18796-016	08/04/05 17:44
5M09767.D	AC18796-018	08/04/05 18:06
5M09768.D	AC18796-019	08/04/05 18:29
5M09769.D	AC18832-001	08/04/05 18:50
5M09770.D	AC18832-002	08/04/05 19:12
5M09771.D	AC18832-003	08/04/05 19:34
5M09772.D	AC18825-005	08/04/05 19:56
5M09773.D	AC18825-007	08/04/05 20:18
5M09774.D	AC18823-001	08/04/05 20:40
5M09775.D	AC18823-003	08/04/05 21:02
5M09776.D	AC18841-001	08/04/05 21:24
5M09777.D	AC18841-002	08/04/05 21:46

DFTPP

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09735.D Vial: 1
 Acq On : 4 Aug 2005 6:25 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 7.812 to 7.858 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	34.5	29672	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	38.4	33009	PASS
70	69	0.00	2	0.2	53	PASS
127	198	40	60	48.9	42075	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	86042	PASS
199	198	5	9	6.7	5746	PASS
275	198	10	30	18.5	15876	PASS
365	198	1	100	1.6	1394	PASS
441	443	0.01	100	79.0	7136	PASS
442	198	40	100	55.2	47514	PASS
443	442	17	23	19.0	9034	PASS

Form 5

Tune Name: CAL DFTPP

Data File: 5M09778.D

Instrument: GCMS_5

Analysis Date: 08/05/05 06:24

Tune Scan/Time Range: Average of 7.800 to 7.846 min

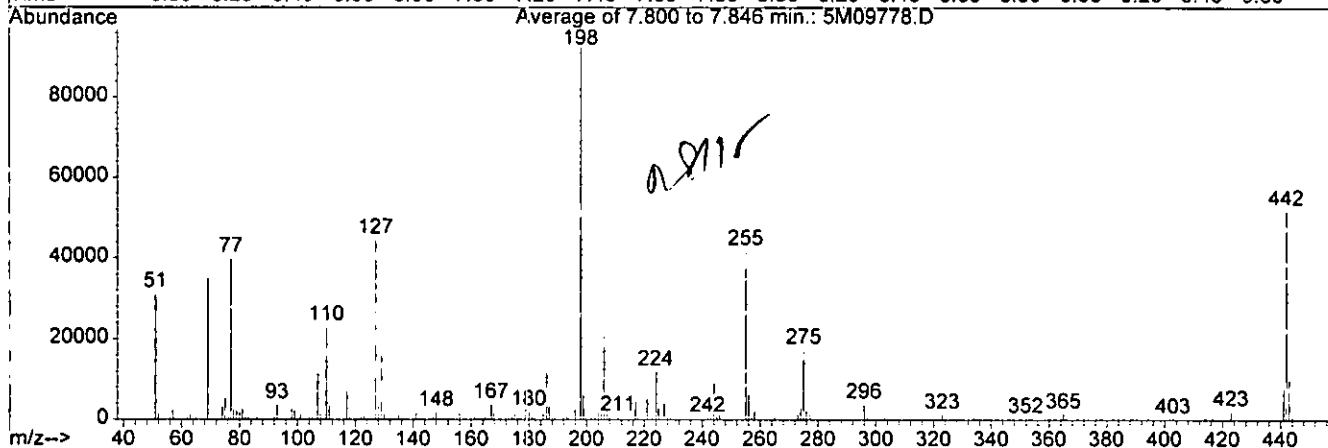
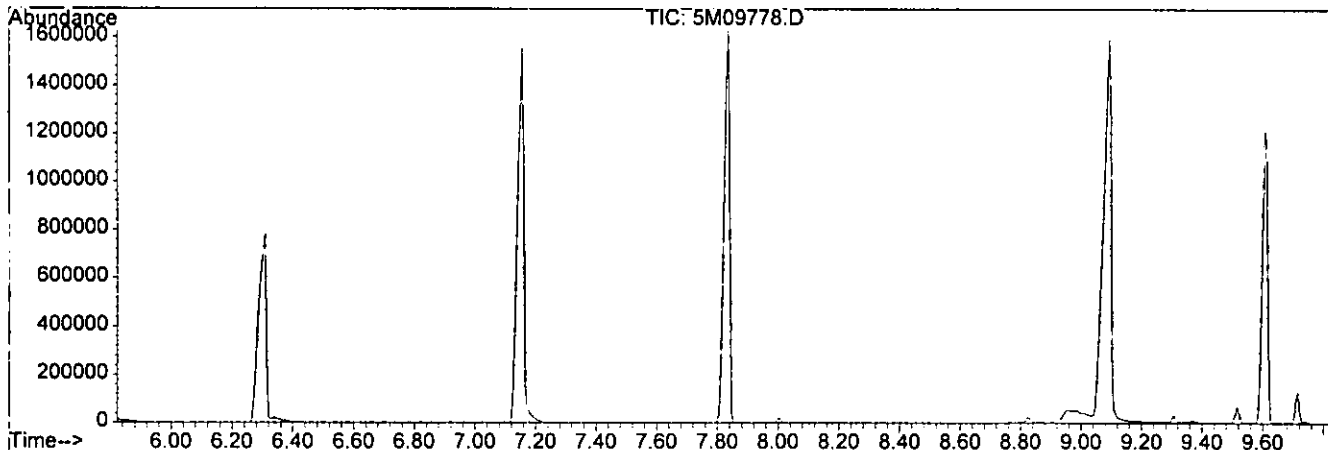
Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/Fail
51	198	30	60	33.8	31227	PASS
68	69	0.00	2	0.2	75	PASS
69	198	0.00	100	38.0	35044	PASS
70	69	0.00	2	0.1	52	PASS
127	198	40	60	48.3	44568	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	92293	PASS
199	198	5	9	6.8	6241	PASS
275	198	10	30	18.8	17366	PASS
365	198	1	100	1.5	1425	PASS
441	443	0.01	100	79.0	7831	PASS
442	198	40	100	56.2	51853	PASS
443	442	17	23	19.1	9913	PASS

Data File	Sample Number	Analysis Date:
5M09779.D	CAL BNA@50PPM	08/05/05 06:42
5M09780.D	SMB2610	08/05/05 07:09
5M09781.D	SMB2611	08/05/05 07:31
5M09782.D	SMB2611(MS)	08/05/05 07:52
5M09783.D	WMB2632	08/05/05 08:14
5M09784.D	WMB2632(MS)	08/05/05 08:36
5M09785.D	AC18883-001(T)	08/05/05 08:57
5M09786.D	AC18796-019	08/05/05 09:19
5M09787.D	AC18881-005	08/05/05 09:41
5M09788.D	AC18778-003	08/05/05 10:02
5M09789.D	AC18778-005	08/05/05 10:23
5M09790.D	AC18778-008	08/05/05 10:45
5M09791.D	AC18778-011	08/05/05 11:07
5M09792.D	AC18778-014	08/05/05 11:29
5M09793.D	AC18778-015	08/05/05 11:51
5M09794.D	WMB2633	08/05/05 12:13
5M09795.D	WMB2633(MS)	08/05/05 12:34
5M09796.D	AC18907-005(T)	08/05/05 12:56
5M09797.D	AC18807-002	08/05/05 13:18
5M09798.D	AC18807-003	08/05/05 13:40
5M09799.D	AC18807-005	08/05/05 14:02
5M09800.D	AC18807-006	08/05/05 14:24
5M09801.D	AC18807-008	08/05/05 14:46
5M09802.D	AC18807-010	08/05/05 15:08
5M09803.D	AC18807-013	08/05/05 15:30
5M09804.D	WMB2633	08/05/05 15:52
5M09805.D	AC18807-014	08/05/05 16:14
5M09806.D	AC18778-020	08/05/05 16:36
5M09807.D	AC18778-006	08/05/05 16:58
5M09808.D	AC18778-009	08/05/05 17:20
5M09809.D	AC18778-010	08/05/05 17:41
5M09810.D	AC18778-017	08/05/05 18:03
5M09811.D	AC18807-001	08/05/05 18:25
5M09812.D	AC18778-018	08/05/05 18:47
5M09813.D	AC18832-006	08/05/05 19:09
5M09814.D	AC18832-007	08/05/05 19:30
5M09815.D	AC18836-001	08/05/05 19:52
5M09816.D	AC18836-002	08/05/05 20:14
5M09817.D	AC18839-002	08/05/05 20:35
5M09818.D	AC18839-003	08/05/05 20:57
5M09819.D	AC18858-001	08/05/05 21:18
5M09820.D	AC18858-002	08/05/05 21:40
5M09821.D	AC18858-003	08/05/05 22:01
5M09822.D	AC18858-004	08/05/05 22:23
5M09823.D	AC18858-005	08/05/05 22:44
5M09824.D	AC18858-006	08/05/05 23:06
5M09825.D	AC18832-005	08/05/05 23:27

4860

DFTPP

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09778.D Vial: 1
 Acq On : 5 Aug 2005 6:24 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270



Spectrum Information: Average of 7.800 to 7.846 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	33.8	31227	PASS
68	69	0.00	2	0.2	75	PASS
69	198	0.00	100	38.0	35044	PASS
70	69	0.00	2	0.1	52	PASS
127	198	40	60	48.3	44568	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	92293	PASS
199	198	5	9	6.8	6241	PASS
275	198	10	30	18.8	17366	PASS
365	198	1	100	1.5	1425	PASS
441	443	0.01	100	79.0	7831	PASS
442	198	40	100	56.2	51853	PASS
443	442	17	23	19.1	9913	PASS

Form 5

Tune Name: CAL DFTPP Data File: 4M05383.D
 Instrument: GCMS_4 Analysis Date: 08/05/05 06:36

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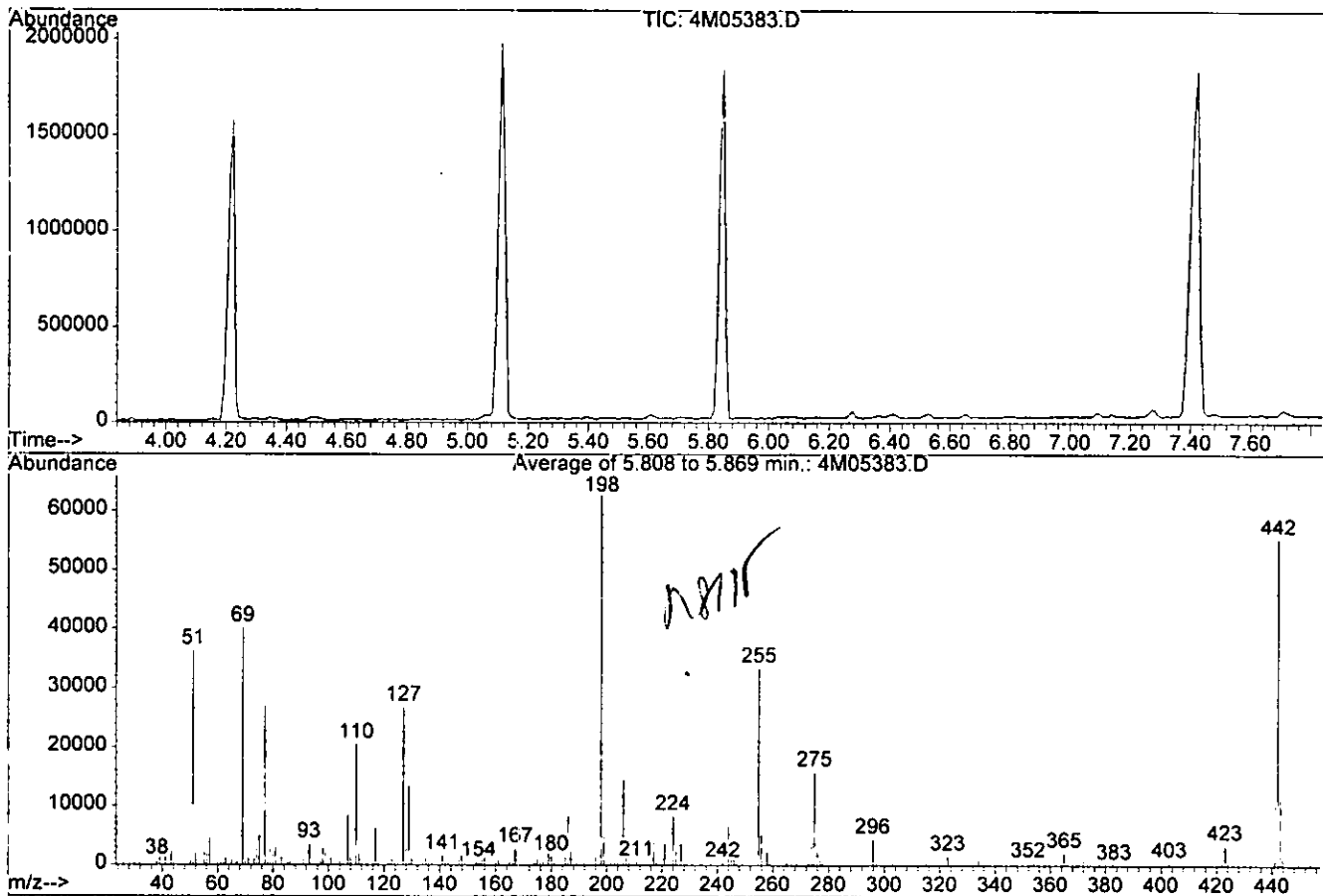
Tune Scan/Time Range: Average of 5.808 to 5.869 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/ Fail
51	198	30	60	57.8	36280	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	64.0	40160	PASS
70	69	0.00	2	1.7	678	PASS
127	198	40	60	42.7	26809	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	62744	PASS
199	198	5	9	7.5	4697	PASS
275	198	10	30	25.2	15835	PASS
365	198	1	100	3.3	2098	PASS
441	443	0.01	100	91.5	10190	PASS
442	198	40	100	88.2	55330	PASS
443	442	17	23	20.1	11137	PASS

Data File	Sample Number	Analysis Date:
4M05384.D	CAL BNA@50PPM	08/05/05 06:58
4M05385.D	CAL BNA@50PPM	08/05/05 08:18
4M05386.D	SMB2610	08/05/05 08:42
4M05387.D	SMB2610(MS)	08/05/05 09:06
4M05388.D	AC18807-009	08/05/05 09:30
4M05389.D	AC18807-011(MS:	08/05/05 09:54
4M05390.D	AC18807-012(MS	08/05/05 10:18
4M05391.D	AC18847-004(R)	08/05/05 10:42
4M05392.D	SMB2611	08/05/05 11:06
4M05393.D	AC18881-001	08/05/05 11:30
4M05394.D	AC18883-001(R)	08/05/05 11:54
4M05395.D	AC18881-001(R)	08/05/05 12:18
4M05396.D	AC18778-012	08/05/05 12:42
4M05397.D	AC18881-002(3X)	08/05/05 13:06
4M05398.D	AC18881-006(3X)	08/05/05 13:30
4M05399.D	AC18881-003	08/05/05 13:54
4M05400.D	AC18881-004	08/05/05 14:18
4M05401.D	AC18855-002	08/05/05 14:42
4M05402.D	AC18855-003	08/05/05 15:06
4M05403.D	AC18855-004	08/05/05 15:31
4M05404.D	AC18807-004	08/05/05 15:55
4M05405.D	SMB2612(MS)	08/05/05 16:19
4M05406.D	SMB2612	08/05/05 16:43
4M05407.D	AC18907-005	08/05/05 17:07
4M05408.D	AC18881-004(3X)	08/05/05 17:31
4M05409.D	AC18881-002(30X)	08/05/05 17:55
4M05410.D	AC18920-002(20X)	08/05/05 18:19
4M05411.D	AC18920-003(20X)	08/05/05 18:43
4M05412.D	AC18920-001(10X)	08/05/05 19:07
4M05413.D	AC18778-013	08/05/05 19:31
4M05414.D	AC18778-004	08/05/05 19:55
4M05415.D	AC18778-001	08/05/05 20:19
4M05416.D	AC18778-019	08/05/05 20:43
4M05417.D	AC18778-016	08/05/05 21:07
4M05418.D	AC18778-007	08/05/05 21:31
4M05419.D	TEST	08/05/05 21:55
4M05420.D	TEST	08/05/05 22:19
4M05421.D	TEST	08/05/05 22:43
4M05422.D	TEST	08/05/05 23:07
4M05423.D	TEST	08/05/05 23:31
4M05424.D	TEST	08/05/05 23:54

DFTPP

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05383.D Vial: 1
 Acq On : 5 Aug 2005 6:36 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_4
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\SM_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 5.808 to 5.869 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	57.8	36280	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	64.0	40160	PASS
70	69	0.00	2	1.7	678	PASS
127	198	40	60	42.7	26809	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	62744	PASS
199	198	5	9	7.5	4697	PASS
275	198	10	30	25.2	15835	PASS
365	198	1	100	3.3	2098	PASS
441	443	0.01	100	91.5	10190	PASS
442	198	40	100	88.2	55330	PASS
443	442	17	23	20.1	11137	PASS

Form 5

Tune Name: CAL DFTPP

Data File: 5M09826.D

Instrument: GCMS_5

Analysis Date: 08/08/05 06:23

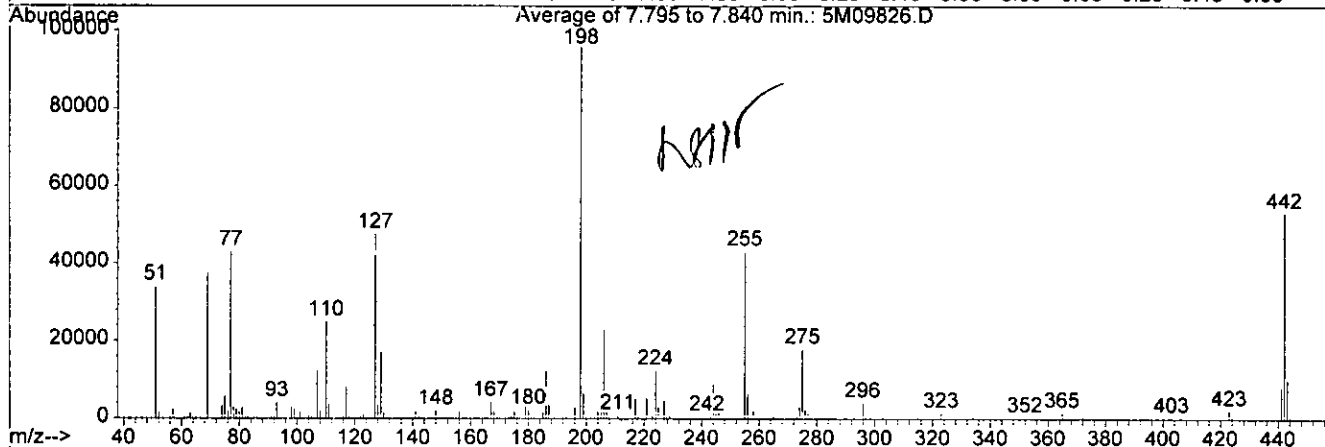
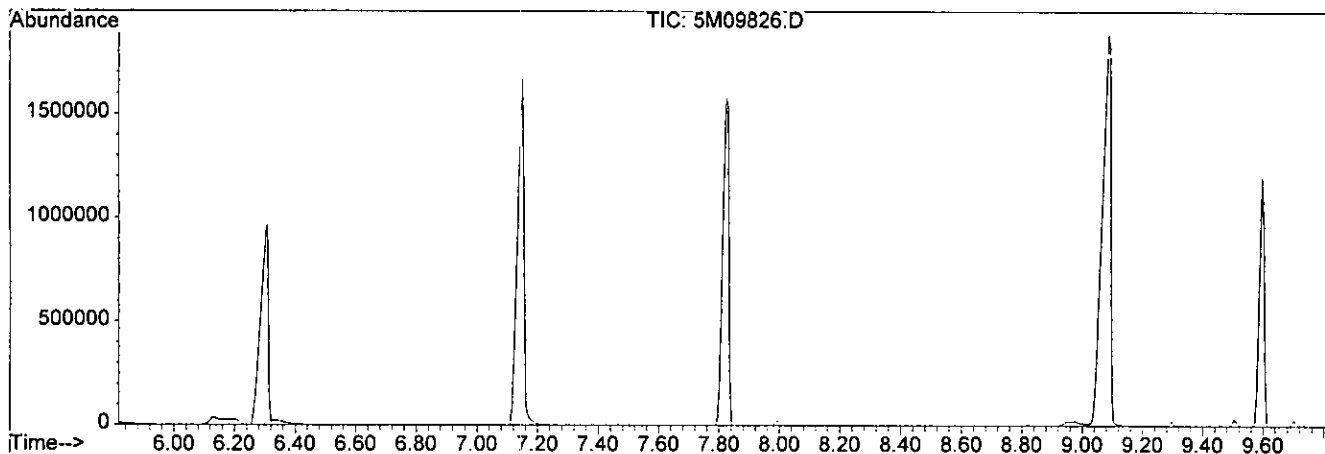
Tune Scan/Time Range: Average of 7.795 to 7.840 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/Fail
51	198	30	60	35.5	34016	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	39.2	37569	PASS
70	69	0.00	2	0.0	0	PASS
127	198	40	60	49.6	47553	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	95784	PASS
199	198	5	9	6.7	6464	PASS
275	198	10	30	18.7	17881	PASS
365	198	1	100	1.6	1555	PASS
441	443	0.01	100	78.8	8053	PASS
442	198	40	100	55.5	53126	PASS
443	442	17	23	19.2	10216	PASS

Data File	Sample Number	Analysis Date:
5M09827.D	CAL BNA@50PPM	08/08/05 06:40
5M09828.D	WMB2633	08/08/05 07:15
5M09829.D	WMB2634	08/08/05 07:36
5M09830.D	SMB2613	08/08/05 07:58
5M09831.D	SMB2613(MS)	08/08/05 08:19
5M09832.D	AC18807-021	08/08/05 08:41
5M09833.D	AC18807-021(MS)	08/08/05 09:02
5M09834.D	AC18807-021(MS)	08/08/05 09:24
5M09835.D	WMB2634(MS)	08/08/05 09:46
5M09836.D	AC18892-001	08/08/05 10:07
5M09837.D	AC18892-001(MS)	08/08/05 10:29
5M09838.D	AC18892-001(MS)	08/08/05 10:51
5M09839.D	AC18778-017	08/08/05 11:12
5M09840.D	AC18807-001	08/08/05 11:34
5M09841.D	AC18888-001	08/08/05 11:56
5M09842.D	AC18892-002	08/08/05 12:17
5M09843.D	AC18892-003	08/08/05 12:39
5M09844.D	AC18873-014	08/08/05 13:01
5M09845.D	AC18778-022	08/08/05 13:23
5M09846.D	AC18778-023	08/08/05 13:44
5M09847.D	AC18807-024	08/08/05 14:06
5M09848.D	AC18807-025	08/08/05 14:28
5M09849.D	AC18807-017	08/08/05 14:50
5M09850.D	AC18807-018	08/08/05 15:11
5M09851.D	AC18807-020	08/08/05 15:33
5M09852.D	AC18778-018	08/08/05 15:55
5M09853.D	AC18884-004	08/08/05 16:16
5M09854.D	SMB2614	08/08/05 16:38
5M09855.D	AC18873-017	08/08/05 17:00
5M09856.D	AC18830-001	08/08/05 17:21
5M09857.D	AC18845-002	08/08/05 17:43
5M09858.D	AC18939-001	08/08/05 18:04
5M09859.D	AC18845-004	08/08/05 18:26
5M09860.D	AC18882-001	08/08/05 18:47
5M09861.D	AC18882-002	08/08/05 19:09
5M09862.D	AC18884-001	08/08/05 19:30
5M09863.D	AC18884-002	08/08/05 19:51
5M09864.D	AC18884-003	08/08/05 20:13
5M09865.D	AC18866-001	08/08/05 20:34
5M09866.D	AC18866-002	08/08/05 20:55
5M09867.D	CH2CL2(#1)	08/08/05 21:17
5M09868.D	CH2CL2(#2)	08/08/05 21:38

DFTPP

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09826.D Vial: 1
 Acq On : 8 Aug 2005 6:23 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 7.795 to 7.840 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	35.5	34016	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	39.2	37569	PASS
70	69	0.00	2	0.0	0	PASS
127	198	40	60	49.6	47553	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	95784	PASS
199	198	5	9	6.7	6464	PASS
275	198	10	30	18.7	17881	PASS
365	198	1	100	1.6	1555	PASS
441	443	0.01	100	78.8	8053	PASS
442	198	40	100	55.5	53126	PASS
443	442	17	23	19.2	10216	PASS

Form 5

Tune Name: CAL DFTPP

Data File: 4M05425.D

Instrument: GCMS_4

Analysis Date: 08/08/05 06:40

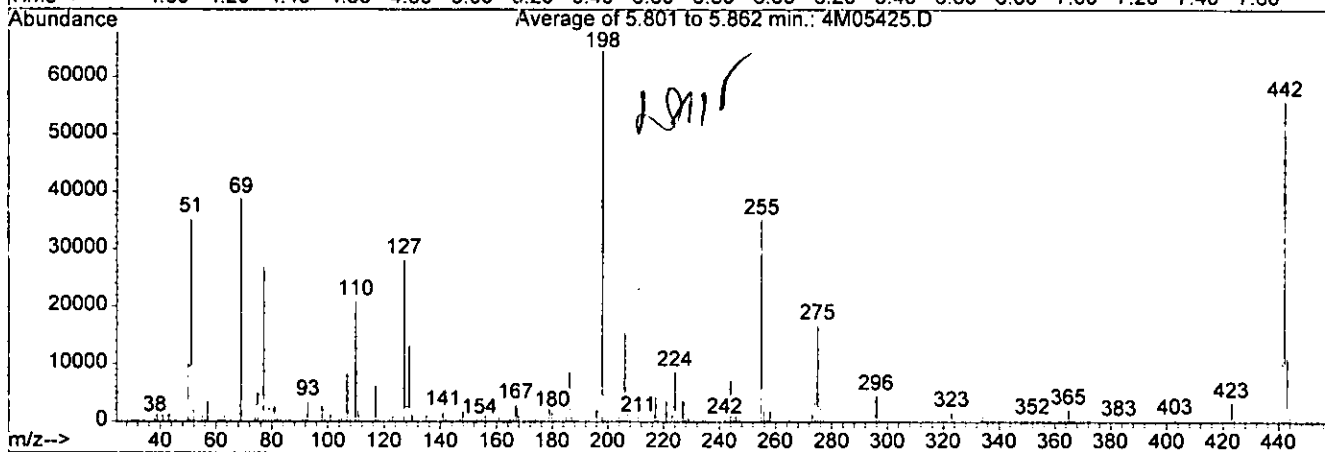
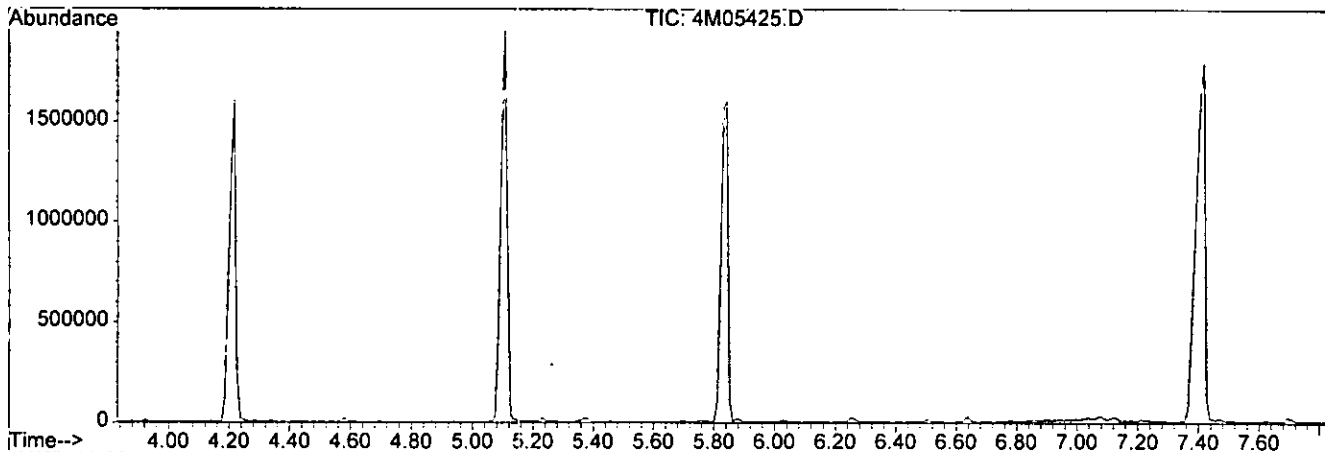
Tune Scan/Time Range: Average of 5.801 to 5.862 min

Tgt Mass	Rel Mass	Lo Lim	Hi Lim	Rel Abund	Raw Abund	Pass/Fail
51	198	30	60	54.5	35295	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	59.9	38787	PASS
70	69	0.00	2	0.3	119	PASS
127	198	40	60	43.5	28152	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	64745	PASS
199	198	5	9	7.6	4939	PASS
275	198	10	30	25.9	16788	PASS
365	198	1	100	3.2	2063	PASS
441	443	0.01	100	91.4	10260	PASS
442	198	40	100	86.4	55931	PASS
443	442	17	23	20.1	11227	PASS

Data File	Sample Number	Analysis Date:
4M05426.D	CAL BNA@50PPM	08/08/05 06:59
4M05427.D	SMB2613(MS)	08/08/05 07:22
4M05428.D	SMB2613	08/08/05 07:49
4M05429.D	AC18920-001	08/08/05 08:13
4M05430.D	AC18920-002	08/08/05 08:36
4M05431.D	AC18920-003	08/08/05 09:00
4M05432.D	AC18778-024	08/08/05 09:24
4M05433.D	AC18807-019	08/08/05 09:48
4M05434.D	AC18820-001	08/08/05 10:12
4M05435.D	AC18807-023(5X)	08/08/05 10:36
4M05436.D	AC18820-003(5X)	08/08/05 11:00
4M05437.D	AC18820-004(5X)	08/08/05 11:24
4M05438.D	AC18820-002(3X)	08/08/05 11:47
4M05439.D	AC18806-001(20X)	08/08/05 12:11
4M05440.D	AC18778-021	08/08/05 12:35
4M05441.D	AC18807-015	08/08/05 12:59
4M05442.D	AC18807-016	08/08/05 13:23
4M05443.D	AC18778-004(5X)	08/08/05 13:47
4M05444.D	AC18778-016(5X)	08/08/05 14:11
4M05445.D	AC18778-019(5X)	08/08/05 14:35
4M05446.D	AC18778-013	08/08/05 14:59
4M05447.D	AC18778-001	08/08/05 15:23
4M05448.D	AC18778-007	08/08/05 15:47
4M05449.D	AC18807-022	08/08/05 16:11
4M05450.D	AC18820-003	08/08/05 16:35
4M05451.D	AC18820-004	08/08/05 16:59
4M05452.D	AC18820-002	08/08/05 17:22
4M05453.D	AC18806-001(10X)	08/08/05 17:46
4M05454.D	AC18873-016	08/08/05 18:10
4M05455.D	TEST	08/08/05 18:34
4M05456.D	TEST	08/08/05 18:58
4M05457.D	TEST	08/08/05 19:22
4M05458.D	TEST	08/08/05 19:45
4M05459.D	TEST	08/08/05 20:09
4M05460.D	TEST	08/08/05 20:33

1005

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05425.D Vial: 1
 Acq On : 8 Aug 2005 6:40 Operator: AHD
 Sample : CAL DFTPP Inst : GCMS_4
 Misc : A,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270



Spectrum Information: Average of 5.801 to 5.862 min.

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
51	198	30	60	54.5	35295	PASS
68	69	0.00	2	0.0	0	PASS
69	198	0.00	100	59.9	38787	PASS
70	69	0.00	2	0.3	119	PASS
127	198	40	60	43.5	28152	PASS
197	198	0.00	1	0.0	0	PASS
198	198	100	100	100.0	64745	PASS
199	198	5	9	7.6	4939	PASS
275	198	10	30	25.9	16788	PASS
365	198	1	100	3.2	2063	PASS
441	443	0.01	100	91.4	10260	PASS
442	198	40	100	86.4	55931	PASS
443	442	17	23	20.1	11227	PASS

Form1

ORGANICS SEMIVOLATILE REPORT

1892

Sample Number: WMB2631
 Client Id:
 Data File: 5M09737.D
 Analysis Date: 08/04/05 07:06
 Date Rec/Extracted: NA-08/03/05

Matrix: Aqueous
 Initial Vol: 1000ml
 Final Vol: 1ml
 Dilution: 1
 Solids: 0

Units: ug/L

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
120-82-1	1,2,4-Trichlorobenzene	0.17	U	205-99-2	Benzo[b]fluoranthene	0.28	U
95-50-1	1,2-Dichlorobenzene	0.40	U	191-24-2	Benzo[g,h,i]perylene	0.14	U
122-66-7	1,2-Diphenylhydrazine	0.33	U	207-08-9	Benzo[k]fluoranthene	0.35	U
541-73-1	1,3-Dichlorobenzene	0.28	U	111-91-1	bis(2-Chloroethoxy)methan	0.23	U
106-46-7	1,4-Dichlorobenzene	0.18	U	111-44-4	bis(2-Chloroethyl)ether	0.44	U
95-95-4	2,4,5-Trichlorophenol	1.6	U	108-60-1	bis(2-chloroisopropyl)ether	0.21	U
88-06-2	2,4,6-Trichlorophenol	0.75	U	117-81-7	bis(2-Ethylhexyl)phthalate	0.63	U
120-83-2	2,4-Dichlorophenol	1.3	U	85-68-7	Butylbenzylphthalate	0.27	U
105-67-9	2,4-Dimethylphenol	0.85	U	86-74-8	Carbazole	0.19	U
51-28-5	2,4-Dinitrophenol	1.8	U	218-01-9	Chrysene	0.28	U
121-14-2	2,4-Dinitrotoluene	0.36	U	84-74-2	Di-n-butylphthalate	0.20	U
606-20-2	2,6-Dinitrotoluene	0.45	U	117-84-0	Di-n-octylphthalate	0.34	U
91-58-7	2-Chloronaphthalene	0.11	U	53-70-3	Dibenzo[a,h]anthracene	0.18	U
95-57-8	2-Chlorophenol	1.8	U	132-64-9	Dibenzofuran	1.3	U
91-57-6	2-Methylnaphthalene	1.7	U	84-66-2	Diethylphthalate	0.24	U
95-48-7	2-Methylphenol	3.7	U	131-11-3	Dimethylphthalate	0.17	U
88-74-4	2-Nitroaniline	1.3	U	206-44-0	Fluoranthene	0.16	U
88-75-5	2-Nitrophenol	1.2	U	86-73-7	Fluorene	0.24	U
106-44-5	3&4-Methylphenol	3.7	U	118-74-1	Hexachlorobenzene	0.41	U
91-94-1	3,3'-Dichlorobenzidine	1.8	U	87-68-3	Hexachlorobutadiene	0.25	U
99-09-2	3-Nitroaniline	2.5	U	77-47-4	Hexachlorocyclopentadiene	2.7	U
534-52-1	4,6-Dinitro-2-methylphenol	1.9	U	67-72-1	Hexachloroethane	0.35	U
101-55-3	4-Bromophenyl-phenylether	0.41	U	193-39-5	Indeno[1,2,3-cd]pyrene	0.17	U
59-50-7	4-Chloro-3-methylphenol	2.0	U	78-59-1	Isophorone	5.3	U
106-47-8	4-Chloroaniline	6.8	U	621-64-7	N-Nitroso-di-n-propylamine	0.32	U
7005-72-3	4-Chlorophenyl-phenylether	0.28	U	62-75-9	N-Nitrosodimethylamine	11	U
100-01-6	4-Nitroaniline	1.5	U	86-30-6	n-Nitrosodiphenylamine	0.27	U
100-02-7	4-Nitrophenol	1.4	U	91-20-3	Naphthalene	0.097	U
83-32-9	Acenaphthene	0.16	U	98-95-3	Nitrobenzene	0.28	U
208-96-8	Acenaphthylene	0.15	U	87-86-5	Pentachlorophenol	0.97	U
120-12-7	Anthracene	0.20	U	85-01-8	Phenanthrene	0.22	U
92-87-5	Benzidine	10	U	108-95-2	Phenol	1.7	U
56-55-3	Benzo[a]anthracene	0.14	U	129-00-0	Pyrene	0.23	U
50-32-8	Benzo[a]pyrene	0.17	U				

Worksheet #: 18122

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09737.D Vial: 3
 Acq On : 4 Aug 2005 7:06 Operator: AHD
 Sample : WMB2631 Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 4 11:23 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.11	152	19696	40.00	ng	-0.14
20) Naphthalene-d8	6.14	136	75749	40.00	ng	-0.14
36) Acenaphthene-d10	7.48	164	47430	40.00	ng	-0.16
61) Phenanthrene-d10	8.85	188	80535	40.00	ng	-0.18
77) Chrysene-d12	11.82	240	62453	40.00	ng	-0.21
88) Perylene-d12	13.41	264	50792	40.00	ng	-0.21
System Monitoring Compounds						
4) 2-Fluorophenol	3.79	112	75675	114.07	ng	-0.18
Spiked Amount	200.000		Recovery	=	57.03%	
8) Phenol-d5	4.81	99	78753	81.19	ng	-0.14
Spiked Amount	200.000		Recovery	=	40.60%	
21) Nitrobenzene-d5	5.58	128	28052	84.58	ng	-0.14
Spiked Amount	100.000		Recovery	=	84.58%	
41) 2-Fluorobiphenyl	6.95	172	119823	80.82	ng	-0.14
Spiked Amount	100.000		Recovery	=	80.82%	
64) 2,4,6-Tribromophenol	8.17	330	30820	178.81	ng	-0.17
Spiked Amount	200.000		Recovery	=	89.41%	
80) Terphenyl-d14	10.62	244	134174	90.94	ng	-0.19
Spiked Amount	100.000		Recovery	=	90.94%	

Target Compounds Qvalue

12815

(#) = qualifier out of range (m) = manual integration

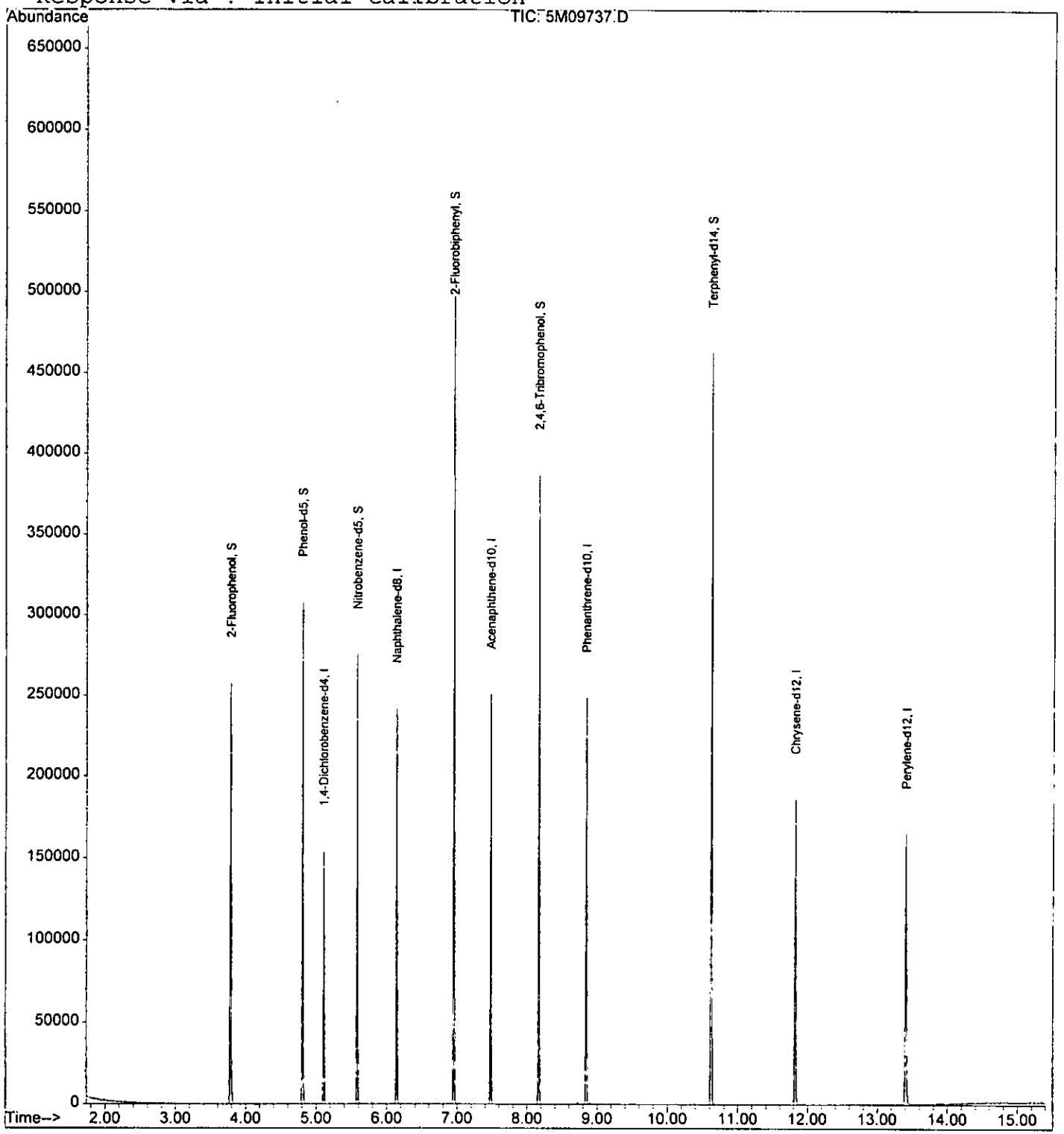
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09737.D Vial: 3
Acq On : 4 Aug 2005 7:06 Operator: AHD
Sample : WMB2631 Inst : GCMS_5
Misc : A,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 4 11:23 2005

1594

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Form1

ORGANICS SEMIVOLATILE REPORT

Sample Number: SMB2610
 Client Id:
 Data File: 5M09780.D
 Analysis Date: 08/05/05 07:09
 Date Rec/Extracted: NA-08/04/05

Matrix: Soil
 Initial Vol: 30g
 Final Vol: 1ml
 Dilution: 1
 Solids: 100

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
120-82-1	1,2,4-Trichlorobenzene	0.0058	U	205-99-2	Benzo[b]fluoranthene	0.0093	U
95-50-1	1,2-Dichlorobenzene	0.013	U	191-24-2	Benzo[g,h,i]perylene	0.0048	U
122-66-7	1,2-Diphenylhydrazine	0.011	U	207-08-9	Benzo[k]fluoranthene	0.012	U
541-73-1	1,3-Dichlorobenzene	0.0095	U	111-91-1	bis(2-Chloroethoxy)methan	0.0078	U
106-46-7	1,4-Dichlorobenzene	0.0058	U	111-44-4	bis(2-Chloroethyl)ether	0.015	U
95-95-4	2,4,5-Trichlorophenol	0.052	U	108-60-1	bis(2-chloroisopropyl)ether	0.0069	U
88-06-2	2,4,6-Trichlorophenol	0.025	U	117-81-7	bis(2-Ethylhexyl)phthalate	0.021	U
120-83-2	2,4-Dichlorophenol	0.044	U	85-68-7	Butylbenzylphthalate	0.0090	U
105-67-9	2,4-Dimethylphenol	0.028	U	86-74-8	Carbazole	0.0064	U
51-28-5	2,4-Dinitrophenol	0.061	U	218-01-9	Chrysene	0.0095	U
121-14-2	2,4-Dinitrotoluene	0.012	U	84-74-2	Di-n-butylphthalate	0.0067	U
606-20-2	2,6-Dinitrotoluene	0.015	U	117-84-0	Di-n-octylphthalate	0.011	U
91-58-7	2-Chloronaphthalene	0.0038	U	53-70-3	Dibenzo[a,h]anthracene	0.0061	U
95-57-8	2-Chlorophenol	0.061	U	132-64-9	Dibenzofuran	0.043	U
91-57-6	2-Methylnaphthalene	0.057	U	84-66-2	Diethylphthalate	0.0078	U
95-48-7	2-Methylphenol	0.12	U	131-11-3	Dimethylphthalate	0.0057	U
88-74-4	2-Nitroaniline	0.043	U	206-44-0	Fluoranthene	0.0055	U
88-75-5	2-Nitrophenol	0.041	U	86-73-7	Fluorene	0.0080	U
106-44-5	3&4-Methylphenol	0.12	U	118-74-1	Hexachlorobenzene	0.014	U
91-94-1	3,3'-Dichlorobenzidine	0.058	U	87-68-3	Hexachlorobutadiene	0.0082	U
99-09-2	3-Nitroaniline	0.084	U	77-47-4	Hexachlorocyclopentadiene	0.090	U
534-52-1	4,6-Dinitro-2-methylphenol	0.063	U	67-72-1	Hexachloroethane	0.012	U
101-55-3	4-Bromophenyl-phenylether	0.014	U	193-39-5	Indeno[1,2,3-cd]pyrene	0.0057	U
59-50-7	4-Chloro-3-methylphenol	0.067	U	78-59-1	Isophorone	0.18	U
106-47-8	4-Chloroaniline	0.23	U	621-64-7	N-Nitroso-di-n-propylamine	0.011	U
7005-72-3	4-Chlorophenyl-phenylether	0.0094	U	62-75-9	N-Nitrosodimethylamine	0.37	U
100-01-6	4-Nitroaniline	0.050	U	86-30-6	n-Nitrosodiphenylamine	0.0091	U
100-02-7	4-Nitrophenol	0.047	U	91-20-3	Naphthalene	0.0032	U
83-32-9	Acenaphthene	0.0055	U	98-95-3	Nitrobenzene	0.0094	U
208-96-8	Acenaphthylene	0.0050	U	87-86-5	Pentachlorophenol	0.032	U
120-12-7	Anthracene	0.0066	U	85-01-8	Phenanthrene	0.0074	U
92-87-5	Benzidine	0.34	U	108-95-2	Phenol	0.055	U
56-55-3	Benzo[a]anthracene	0.0046	U	129-00-0	Pyrene	0.0076	U
50-32-8	Benzo[a]pyrene	0.0055	U				

Worksheet #: 18122

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09780.D Vial: 3
 Acq On : 5 Aug 2005 7:09 Operator: AHD
 Sample : SMB2610 Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 9 18:32 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	23226	40.00	ng	-0.15
20) Naphthalene-d8	6.14	136	88373	40.00	ng	-0.14
36) Acenaphthene-d10	7.48	164	51486	40.00	ng	-0.16
61) Phenanthrene-d10	8.84	188	90184	40.00	ng	-0.19
77) Chrysene-d12	11.82	240	65517	40.00	ng	-0.22
88) Perylene-d12	13.40	264	47027	40.00	ng	-0.21

System Monitoring Compounds

4) 2-Fluorophenol	3.78	112	124697	159.40	ng	-0.19
Spiked Amount	200.000		Recovery	=	79.70%	
8) Phenol-d5	4.81	99	169212	147.93	ng	-0.14
Spiked Amount	200.000		Recovery	=	73.97%	
21) Nitrobenzene-d5	5.58	128	31044	80.23	ng	-0.14
Spiked Amount	100.000		Recovery	=	80.23%	
41) 2-Fluorobiphenyl	6.95	172	129930	80.73	ng	-0.14
Spiked Amount	100.000		Recovery	=	80.73%	
64) 2,4,6-Tribromophenol	8.17	330	32558	168.69	ng	-0.18
Spiked Amount	200.000		Recovery	=	84.35%	
80) Terphenyl-d14	10.62	244	151100	97.62	ng	-0.19
Spiked Amount	100.000		Recovery	=	97.62%	

Target Compounds

Qvalue

12811

 (#) = qualifier out of range (m) = manual integration

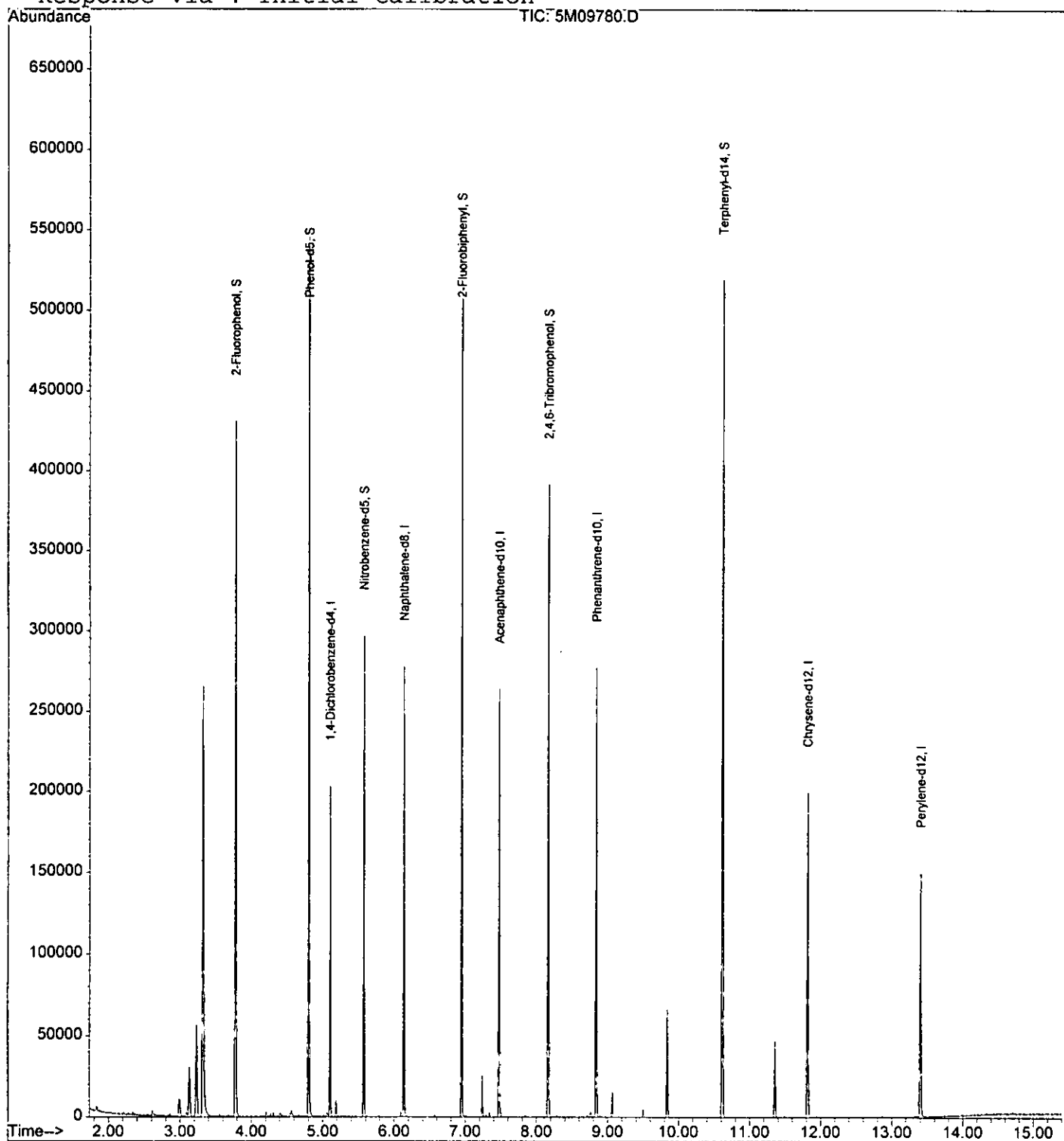
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09780.D Vial: 3
 Acq On : 5 Aug 2005 7:09 Operator: AHD
 Sample : SMB2610 Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 9 18:32 2005

1585

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Form1

ORGANICS SEMIVOLATILE REPORT

1558

Sample Number: SMB2611
 Client Id:
 Data File: 5M09781.D
 Analysis Date: 08/05/05 07:31
 Date Rec/Extracted: NA-08/04/05

Matrix: Soil
 Initial Vol: 30g
 Final Vol: 1ml
 Dilution: 1
 Solids: 100

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
120-82-1	1,2,4-Trichlorobenzene	0.0058	U	205-99-2	Benzo[b]fluoranthene	0.0093	U
95-50-1	1,2-Dichlorobenzene	0.013	U	191-24-2	Benzo[g,h,i]perylene	0.0048	U
122-66-7	1,2-Diphenylhydrazine	0.011	U	207-08-9	Benzo[k]fluoranthene	0.012	U
541-73-1	1,3-Dichlorobenzene	0.0095	U	111-91-1	bis(2-Chloroethoxy)methan	0.0078	U
106-46-7	1,4-Dichlorobenzene	0.0058	U	111-44-4	bis(2-Chloroethyl)ether	0.015	U
95-95-4	2,4,5-Trichlorophenol	0.052	U	108-60-1	bis(2-chloroisopropyl)ether	0.0069	U
88-06-2	2,4,6-Trichlorophenol	0.025	U	117-81-7	bis(2-Ethylhexyl)phthalate	0.021	U
120-83-2	2,4-Dichlorophenol	0.044	U	85-68-7	Butylbenzylphthalate	0.0090	U
105-67-9	2,4-Dimethylphenol	0.028	U	86-74-8	Carbazole	0.0064	U
51-28-5	2,4-Dinitrophenol	0.061	U	218-01-9	Chrysene	0.0095	U
121-14-2	2,4-Dinitrotoluene	0.012	U	84-74-2	Di-n-butylphthalate	0.0067	U
606-20-2	2,6-Dinitrotoluene	0.015	U	117-84-0	Di-n-octylphthalate	0.011	U
91-58-7	2-Chloronaphthalene	0.0038	U	53-70-3	Dibenzo[a,h]anthracene	0.0061	U
95-57-8	2-Chlorophenol	0.061	U	132-64-9	Dibenzofuran	0.043	U
91-57-6	2-Methylnaphthalene	0.057	U	84-66-2	Diethylphthalate	0.0078	U
95-48-7	2-Methylphenol	0.12	U	131-11-3	Dimethylphthalate	0.0057	U
88-74-4	2-Nitroaniline	0.043	U	206-44-0	Fluoranthene	0.0055	U
88-75-5	2-Nitrophenol	0.041	U	86-73-7	Fluorene	0.0080	U
106-44-5	3&4-Methylphenol	0.12	U	118-74-1	Hexachlorobenzene	0.014	U
91-94-1	3,3'-Dichlorobenzidine	0.058	U	87-68-3	Hexachlorobutadiene	0.0082	U
99-09-2	3-Nitroaniline	0.084	U	77-47-4	Hexachlorocyclopentadiene	0.090	U
534-52-1	4,6-Dinitro-2-methylphenol	0.063	U	67-72-1	Hexachloroethane	0.012	U
101-55-3	4-Bromophenyl-phenylether	0.014	U	193-39-5	Indeno[1,2,3-cd]pyrene	0.0057	U
59-50-7	4-Chloro-3-methylphenol	0.067	U	78-59-1	Isophorone	0.18	U
106-47-8	4-Chloroaniline	0.23	U	621-64-7	N-Nitroso-di-n-propylamine	0.011	U
7005-72-3	4-Chlorophenyl-phenylether	0.0094	U	62-75-9	N-Nitrosodimethylamine	0.37	U
100-01-6	4-Nitroaniline	0.050	U	86-30-6	n-Nitrosodiphenylamine	0.0091	U
100-02-7	4-Nitrophenol	0.047	U	91-20-3	Naphthalene	0.0032	U
83-32-9	Acenaphthene	0.0055	U	98-95-3	Nitrobenzene	0.0094	U
208-96-8	Acenaphthylene	0.0050	U	87-86-5	Pentachlorophenol	0.032	U
120-12-7	Anthracene	0.0066	U	85-01-8	Phenanthrene	0.0074	U
92-87-5	Benzidine	0.34	U	108-95-2	Phenol	0.055	U
56-55-3	Benzo[a]anthracene	0.0046	U	129-00-0	Pyrene	0.0076	U
50-32-8	Benzo[a]pyrene	0.0055	U				

Worksheet #: 18122

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09781.D Vial: 4
 Acq On : 5 Aug 2005 7:31 Operator: AHD
 Sample : SMB2611 Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 11 11:36 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	25216	40.00	ng	-0.15
20) Naphthalene-d8	6.14	136	97755	40.00	ng	-0.14
36) Acenaphthene-d10	7.47	164	57927	40.00	ng	-0.16
61) Phenanthrene-d10	8.84	188	99553	40.00	ng	-0.19
77) Chrysene-d12	11.82	240	80231	40.00	ng	-0.22
88) Perylene-d12	13.40	264	57685	40.00	ng	-0.22

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.78	112	79406	93.50	ng	-0.19
Spiked Amount				200.000		
				Recovery =		46.75%
8) Phenol-d5	4.80	99	110124	88.68	ng	-0.15
Spiked Amount				200.000		
				Recovery =		44.34%
21) Nitrobenzene-d5	5.58	128	20932	48.91	ng	-0.14
Spiked Amount				100.000		
				Recovery =		48.91%
41) 2-Fluorobiphenyl	6.95	172	90119	49.77	ng	-0.14
Spiked Amount				100.000		
				Recovery =		49.77%
64) 2,4,6-Tribromophenol	8.16	330	20952	98.34	ng	-0.18
Spiked Amount				200.000		
				Recovery =		49.17%
80) Terphenyl-d14	10.61	244	98907	52.18	ng	-0.20
Spiked Amount				100.000		
				Recovery =		52.18%

Target Compounds Qvalue

12.8115

(#) = qualifier out of range (m) = manual integration

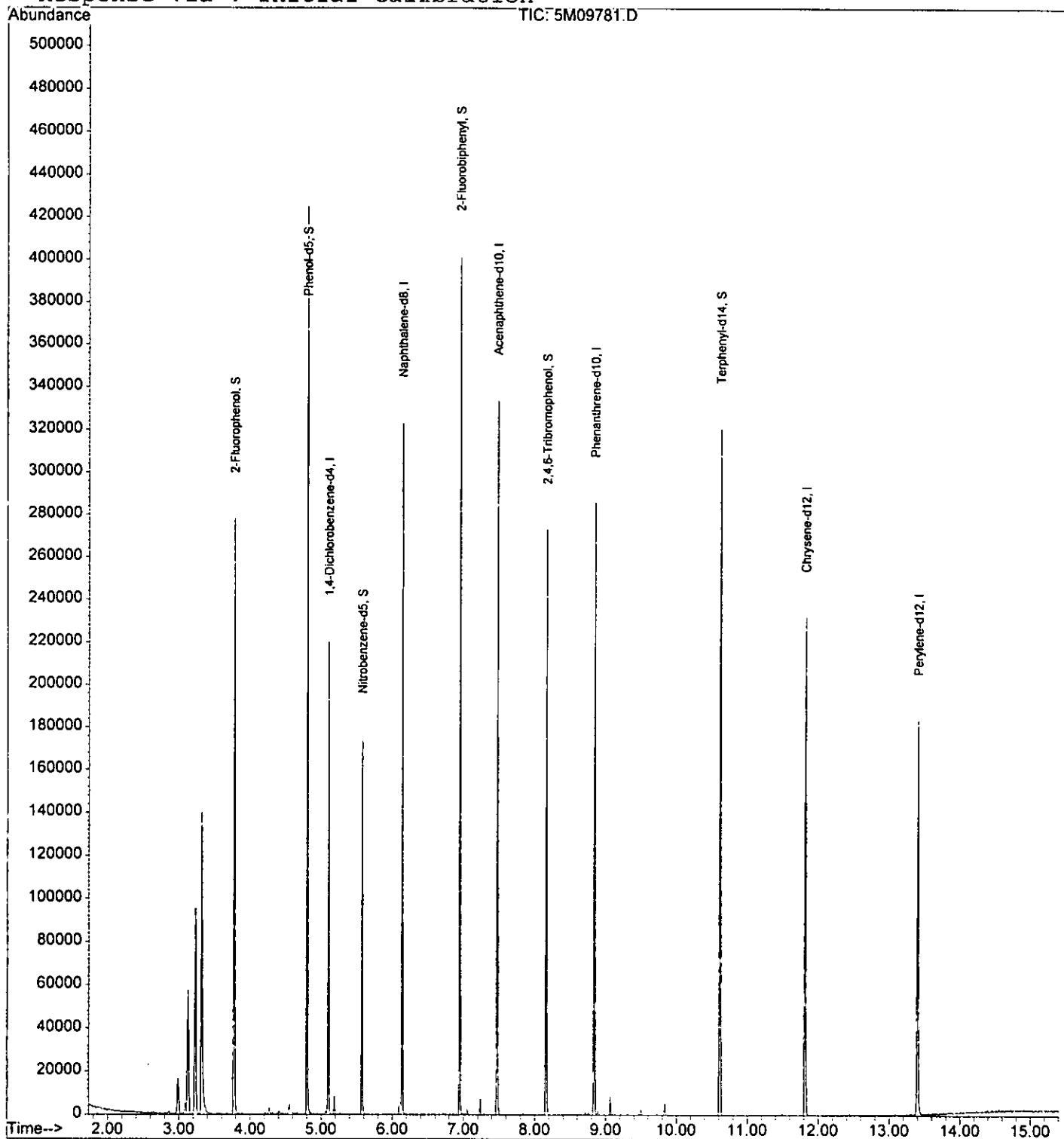
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09781.D Vial: 4
Acq On : 5 Aug 2005 7:31 Operator: AHD
Sample : SMB2611 Inst : GCMS_5
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 11 11:36 2005

1919

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Form1

ORGANICS SEMIVOLATILE REPORT

1812

Sample Number: SMB2613
 Client Id:
 Data File: 4M05428.D
 Analysis Date: 08/08/05 07:49
 Date Rec/Extracted: NA-08/07/05

Matrix: Soil
 Initial Vol: 30g
 Final Vol: 1ml
 Dilution: 1
 Solids: 100

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
120-82-1	1,2,4-Trichlorobenzene	0.0090	U	205-99-2	Benzo[b]fluoranthene	0.010	U
95-50-1	1,2-Dichlorobenzene	0.015	U	191-24-2	Benzo[g,h,i]perylene	0.0063	U
122-66-7	1,2-Diphenylhydrazine	0.0096	U	207-08-9	Benzo[k]fluoranthene	0.011	U
541-73-1	1,3-Dichlorobenzene	0.014	U	111-91-1	bis(2-Chloroethoxy)methan	0.0076	U
106-46-7	1,4-Dichlorobenzene	0.017	U	111-44-4	bis(2-Chloroethyl)ether	0.018	U
95-95-4	2,4,5-Trichlorophenol	0.45	U	108-60-1	bis(2-chloroisopropyl)ether	0.011	U
88-06-2	2,4,6-Trichlorophenol	0.81	U	117-81-7	bis(2-Ethylhexyl)phthalate	0.030	U
120-83-2	2,4-Dichlorophenol	0.054	U	85-68-7	Butylbenzylphthalate	0.013	U
105-67-9	2,4-Dimethylphenol	0.046	U	86-74-8	Carbazole	0.0099	U
51-28-5	2,4-Dinitrophenol	0.23	U	218-01-9	Chrysene	0.0069	U
121-14-2	2,4-Dinitrotoluene	0.012	U	84-74-2	Di-n-butylphthalate	0.0075	U
606-20-2	2,6-Dinitrotoluene	0.014	U	117-84-0	Di-n-octylphthalate	0.0079	U
91-58-7	2-Chloronaphthalene	0.0092	U	53-70-3	Dibenzo[a,h]anthracene	0.012	U
95-57-8	2-Chlorophenol	0.068	U	132-64-9	Dibenzofuran	0.042	U
91-57-6	2-Methylnaphthalene	0.043	U	84-66-2	Diethylphthalate	0.0092	U
95-48-7	2-Methylphenol	0.16	U	131-11-3	Dimethylphthalate	0.0075	U
88-74-4	2-Nitroaniline	0.023	U	206-44-0	Fluoranthene	0.0096	U
88-75-5	2-Nitrophenol	0.039	U	86-73-7	Fluorene	0.0084	U
106-44-5	3&4-Methylphenol	0.18	U	118-74-1	Hexachlorobenzene	0.015	U
91-94-1	3,3'-Dichlorobenzidine	0.073	U	87-68-3	Hexachlorobutadiene	0.014	U
99-09-2	3-Nitroaniline	0.14	U	77-47-4	Hexachlorocyclopentadiene	0.089	U
534-52-1	4,6-Dinitro-2-methylphenol	0.063	U	67-72-1	Hexachloroethane	0.025	U
101-55-3	4-Bromophenyl-phenylether	0.013	U	193-39-5	Indeno[1,2,3-cd]pyrene	0.0046	U
59-50-7	4-Chloro-3-methylphenol	0.085	U	78-59-1	Isophorone	0.010	U
106-47-8	4-Chloroaniline	0.26	U	621-64-7	N-Nitroso-di-n-propylamine	0.016	U
7005-72-3	4-Chlorophenyl-phenylether	0.015	U	62-75-9	N-Nitrosodimethylamine	0.39	U
100-01-6	4-Nitroaniline	0.082	U	86-30-6	n-Nitrosodiphenylamine	0.016	U
100-02-7	4-Nitrophenol	0.059	U	91-20-3	Naphthalene	0.0078	U
83-32-9	Acenaphthene	0.014	U	98-95-3	Nitrobenzene	0.013	U
208-96-8	Acenaphthylene	0.0077	U	87-86-5	Pentachlorophenol	0.041	U
120-12-7	Anthracene	0.0087	U	85-01-8	Phenanthrene	0.0077	U
92-87-5	Benzidine	0.076	U	108-95-2	Phenol	0.051	U
56-55-3	Benzo[a]anthracene	0.0058	U	129-00-0	Pyrene	0.0078	U
50-32-8	Benzo[a]pyrene	0.0077	U				

Worksheet #: 18122

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05428.D Vial: 4
 Acq On : 8 Aug 2005 7:49 Operator: AHD
 Sample : SMB2613 Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 9 18:33 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.90	152	39428	40.00	ng	-0.04
19) Naphthalene-d8	5.90	136	119863	40.00	ng	-0.04
35) Acenaphthene-d10	7.46	164	65103	40.00	ng	-0.06
59) Phenanthrene-d10	9.07	188	102453	40.00	ng	-0.06
72) Chrysene-d12	12.27	240	94992	40.00	ng	-0.06
81) Perylene-d12	14.12	264	83360	40.00	ng	-0.06
System Monitoring Compounds						
4) 2-Fluorophenol	3.74	112	119701	107.63	ng	-0.05
Spiked Amount	200.000		Recovery	=	53.82%	
7) Phenol-d5	4.61	99	170363	115.07	ng	-0.04
Spiked Amount	200.000		Recovery	=	57.53%	
20) Nitrobenzene-d5	5.34	128	34300	57.16	ng	-0.04
Spiked Amount	100.000		Recovery	=	57.16%	
40) 2-Fluorobiphenyl	6.82	172	118128	56.62	ng	-0.05
Spiked Amount	100.000		Recovery	=	56.62%	
62) 2,4,6-Tribromophenol	8.29	332	52531	114.55	ng	-0.06
Spiked Amount	200.000		Recovery	=	57.28%	
75) Terphenyl-d14	10.97	244	124188	46.40	ng	-0.05
Spiked Amount	100.000		Recovery	=	46.40%	

Target Compounds

Qvalue

Handwritten signature

(#) = qualifier out of range (m) = manual integration

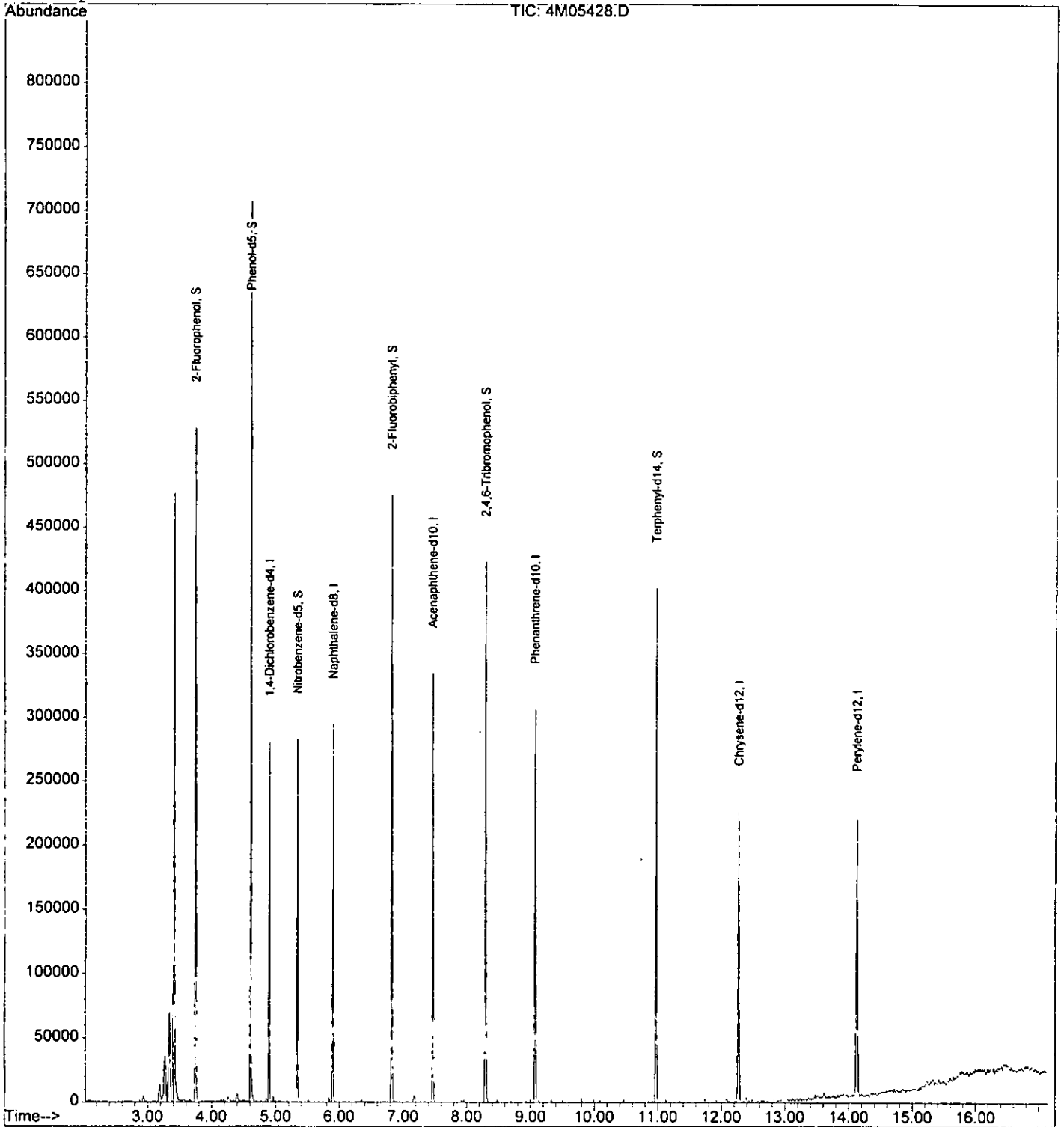
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-08-05\4M05428.D Vial: 4
Acq On : 8 Aug 2005 7:49 Operator: AHD
Sample : SMB2613 Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 9 18:33 2005

1913

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



1514

Data File:====>
Data/Batch/Sample ID:====>
Date/Time:====>

Compound	Limit(s)		Col	Mr	5M09782.D			5M09738.D											
	Soil	Aq			Conc	Exp	%	Conc	Exp	%	Conc	Exp	%	Conc	Exp	%	Conc	Exp	%
					Conc	Exp	Rec	Conc	Exp	Rec	Conc	Exp	Rec	Conc	Exp	Rec	Conc	Exp	Rec
1,2,4-Trichlorobenz	38-107	39-98	1	0	82.52	100	83	90	100	90									
1,4-Dichlorobenzen	28-104	36-97	1	0	84.74	100	85	85.45	100	85									
2,4-Dinitrotoluene	28-89	24-96	1	0	84.14	100	84	88.88	100	89									
2-Chlorophenol	25-102	27-123	1	0	147.9	200	74	78.02	100	78									
4-Chloro-3-methylp	26-103	23-97	1	0	161.1	200	81	82.28	100	82									
4-Nitrophenol	11-114	10-80	1	0	153.8	200	77	50.6	100	51									
Acenaphthene	31-137	46-118	1	0	84.6	100	85	89.38	100	89									
N-Nitroso-di-n-propy	41-126	41-116	1	0	77.85	100	78	79.47	100	79									
Pentachlorophenol	17-109	9-103	1	0	162.8	200	81	97.1	100	97									
Phenol	26-90	12-89	1	0	148.4	200	74	43.41	100	43									
Pyrene	35-142	26-127	1	0	86.64	100	87	91.64	100	92									

1815

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09782.D Vial: 5
 Acq On : 5 Aug 2005 7:52 Operator: AHD
 Sample : SMB2611 (MS) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 5 8:57 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	19213	40.00	ng	-0.15
20) Naphthalene-d8	6.14	136	74090	40.00	ng	-0.14
36) Acenaphthene-d10	7.47	164	44434	40.00	ng	-0.16
61) Phenanthrene-d10	8.84	188	74172	40.00	ng	-0.19
77) Chrysene-d12	11.82	240	58240	40.00	ng	-0.22
88) Perylene-d12	13.40	264	43488	40.00	ng	-0.22

System Monitoring Compounds

4) 2-Fluorophenol	3.78	112	106168	164.06	ng	-0.19	
Spiked Amount				200.000			
				Recovery	=		82.03%
8) Phenol-d5	4.81	99	134454	142.09	ng	-0.14	
Spiked Amount				200.000			
				Recovery	=		71.05%
21) Nitrobenzene-d5	5.58	128	25578	78.85	ng	-0.14	
Spiked Amount				100.000			
				Recovery	=		78.85%
41) 2-Fluorobiphenyl	6.95	172	110902	79.85	ng	-0.14	
Spiked Amount				100.000			
				Recovery	=		79.85%
64) 2,4,6-Tribromophenol	8.16	330	26953	169.79	ng	-0.18	
Spiked Amount				200.000			
				Recovery	=		84.90%
80) Terphenyl-d14	10.61	244	107449	78.09	ng	-0.20	
Spiked Amount				100.000			
				Recovery	=		78.09%

Target Compounds

						Qvalue
9) Phenol	4.82	94	148799	148.36	ng	98
10) 2-Chlorophenol	4.91	128	112600	147.92	ng	97
11) 1,3-Dichlorobenzene	5.12	146	61079	86.72	ng	98
12) 1,4-Dichlorobenzene	5.12	146	61079	84.74	ng	99
13) 1,2-Dichlorobenzene	5.12	146	61079	88.89	ng	97
18) N-Nitroso-di-n-propylamine	5.46	70	42656	77.85	ng	99
23) Isophorone	5.58	82	58024	42.84	ng	60
29) 1,2,4-Trichlorobenzene	6.09	180	50108	82.52	ng	98
33) 4-Chloro-3-methylphenol	6.57	107	105362	161.09	ng	97
45) Diphenyl Ether	6.95	170	24965	30.08	ng	27
49) 2,6-Dinitrotoluene	7.47	165	5628	16.85	ng	29
50) Acenaphthene	7.50	153	103822	84.60	ng	98
54) 2,4-Dinitrotoluene	7.64	165	38813	84.14	ng	97
55) 4-Nitrophenol	7.58	65	44640	153.82	ng	93
57) Fluorene	8.16	166	2011	1.40	ng	87
69) Pentachlorophenol	8.65	266	38325	162.82	ng	96
74) Di-n-butylphthalate	9.50	149	2430	1.01	ng	92
78) Pyrene	10.40	202	202097	86.64	ng	99

(#) = qualifier out of range (m) = manual integration

h.811

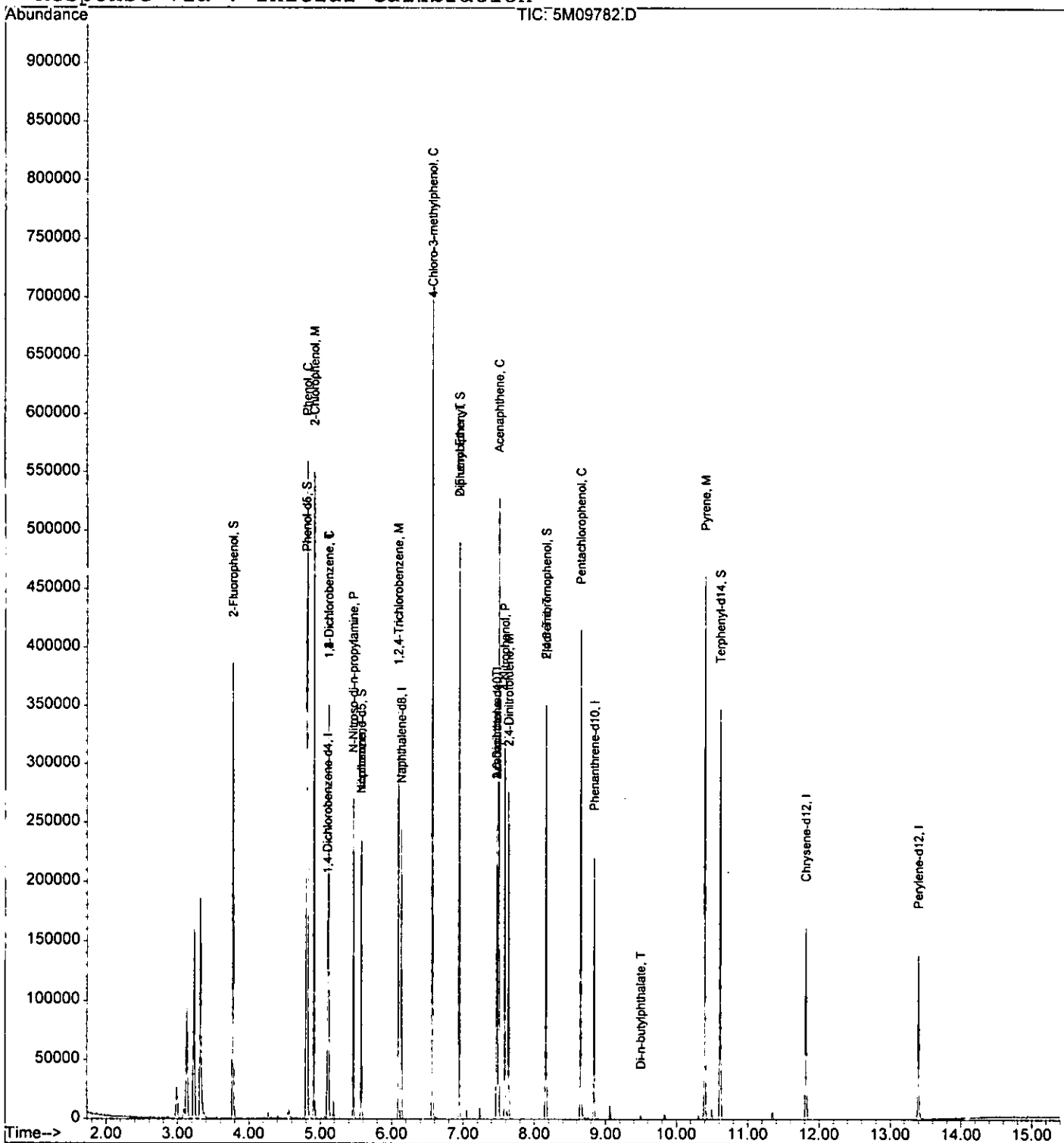
Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-05-05\5M09782.D Vial: 5
 Acq On : 5 Aug 2005 7:52 Operator: AHD
 Sample : SMB2611 (MS) Inst : GCMS_5
 Misc : S, BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 5 8:57 2005

3101

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09738.D Vial: 4
 Acq On : 4 Aug 2005 7:31 Operator: AHD
 Sample : WMB2631 (MS) Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 4 8:55 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.11	152	21674	40.00	ng	-0.14
20) Naphthalene-d8	6.15	136	80909	40.00	ng	-0.13
36) Acenaphthene-d10	7.48	164	48168	40.00	ng	-0.15
61) Phenanthrene-d10	8.85	188	81912	40.00	ng	-0.18
77) Chrysene-d12	11.83	240	68362	40.00	ng	-0.20
88) Perylene-d12	13.41	264	51235	40.00	ng	-0.20

System Monitoring Compounds

4) 2-Fluorophenol	3.79	112	88145	120.75	ng	-0.18
Spiked Amount	200.000		Recovery	=	60.38%	
8) Phenol-d5	4.81	99	92178	86.35	ng	-0.14
Spiked Amount	200.000		Recovery	=	43.18%	
21) Nitrobenzene-d5	5.59	128	32880	92.82	ng	-0.13
Spiked Amount	100.000		Recovery	=	92.82%	
41) 2-Fluorobiphenyl	6.96	172	132107	87.74	ng	-0.13
Spiked Amount	100.000		Recovery	=	87.74%	
64) 2,4,6-Tribromophenol	8.18	330	30999	176.83	ng	-0.16
Spiked Amount	200.000		Recovery	=	88.42%	
80) Terphenyl-d14	10.62	244	148058	91.68	ng	-0.19
Spiked Amount	100.000		Recovery	=	91.68%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Pyridine	1.99	79	31097	34.41	ng	93
3) N-Nitrosodimethylamine	1.93	74	31788	57.83	ng	91
7) bis(2-Chloroethyl)ether	4.90	93	64505	84.11	ng	96
9) Phenol	4.82	94	49109	43.41	ng	95
10) 2-Chlorophenol	4.92	128	66998	78.02	ng	96
11) 1,3-Dichlorobenzene	5.05	146	70722	89.01	ng	97
12) 1,4-Dichlorobenzene	5.12	146	69478	85.45	ng	99
13) 1,2-Dichlorobenzene	5.25	146	64902	83.72	ng	98
15) bis(2-chloroisopropyl)ethe	5.37	45	101968	87.53	ng	94
16) 2-Methylphenol	5.35	108	55950	71.36	ng	100
17) Hexachloroethane	5.53	117	27469	81.45	ng	95
18) N-Nitroso-di-n-propylamine	5.47	70	49122	79.47	ng	98
19) 3&4-Methylphenol	5.49	108	53577	64.27	ng	98
22) Nitrobenzene	5.60	77	75559	95.09	ng	95
23) Isophorone	5.80	82	125986	85.18	ng	99
24) 2-Nitrophenol	5.85	139	36087	88.50	ng	92
25) 2,4-Dimethylphenol	5.91	107	65636	84.62	ng	99
26) Benzoic Acid	5.99	105	15661	35.17	ng	93
27) bis(2-Chloroethoxy)methane	5.98	93	79204	93.59	ng	100

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09738.D Vial: 4
 Acq On : 4 Aug 2005 7:31 Operator: AHD
 Sample : WMB2631 (MS) Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 4 8:55 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
28) 2,4-Dichlorophenol	6.05	162	55215	85.37	ng	98
29) 1,2,4-Trichlorobenzene	6.11	180	59682	90.00	ng	99
30) Naphthalene	6.16	128	185279	87.39	ng	100
32) Hexachlorobutadiene	6.26	225	33729	92.24	ng	99
33) 4-Chloro-3-methylphenol	6.58	107	58770	82.28	ng	98
38) Hexachlorocyclopentadiene	6.79	237	31529	78.70	ng	96
39) 2,4,6-Trichlorophenol	6.89	196	41103	87.67	ng	100
40) 2,4,5-Trichlorophenol	6.92	196	41936	82.24	ng	97
42) 2-Chloronaphthalene	7.05	162	122222	89.76	ng	98
45) Diphenyl Ether	6.95	170	29958	33.30	ng	27
47) Acenaphthylene	7.37	152	189908	88.29	ng	99
48) Dimethylphthalate	7.28	163	138917	88.39	ng	100
49) 2,6-Dinitrotoluene	7.32	165	32571	89.98	ng	91
50) Acenaphthene	7.51	153	118904	89.38	ng	98
52) 2,4-Dinitrophenol	7.54	184	20279	91.05	ng	77
54) 2,4-Dinitrotoluene	7.65	165	44444	88.88	ng	97
55) 4-Nitrophenol	7.59	65	15919	50.60	ng	93
57) Fluorene	7.95	166	136871	87.72	ng	99
58) 4-Chlorophenyl-phenylether	7.96	204	65333	86.14	ng	96
59) Diethylphthalate	7.87	149	140379	87.16	ng	98
60) 4-Nitroaniline	7.95	138	1820	3.97	ng	29
62) 4,6-Dinitro-2-methylphenol	8.01	198	29710	94.34	ng	100
63) n-Nitrosodiphenylamine	8.07	169	101227	89.64	ng	97
65) 1,2-Diphenylhydrazine	8.10	77	152613	94.71	ng	97
66) 4-Bromophenyl-phenylether	8.41	248	39556	94.18	ng	90
67) Hexachlorobenzene	8.47	284	36448	91.92	ng	87
69) Pentachlorophenol	8.66	266	25241	97.10	ng	92
70) Phenanthrene	8.88	178	219835	93.06	ng	99
71) Anthracene	8.93	178	217105	90.42	ng	99
74) Di-n-butylphthalate	9.51	149	251844	94.88	ng	99
76) Fluoranthene	10.16	202	244841	95.07	ng	99
78) Pyrene	10.41	202	250892	91.64	ng	99
79) Benzidine	10.34	184	28901	28.59	ng	97
82) Butylbenzylphthalate	11.23	149	118391	98.25	ng	92
83) Methoxychlor	11.86	227	21854	17.39	ng	98
84) 3,3'-Dichlorobenzidine	11.82	252	69282	88.03	ng	99
85) Benzo[a]anthracene	11.82	228	217094	86.37	ng	99
86) Chrysene	11.86	228	196650	85.30	ng	100
87) bis(2-Ethylhexyl)phthalate	11.95	149	150533	90.41	ng	96
89) Di-n-octylphthalate	12.70	149	261481	93.20	ng	100
90) Benzo[b]fluoranthene	13.02	252	188100	92.98	ng	98

(#) = qualifier out of range (m) = manual integration

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09738.D Vial: 4 .
 Acq On : 4 Aug 2005 7:31 Operator: AHD
 Sample : WMB2631 (MS) Inst : GCMS_5
 Misc : A,BNA Multiplr: 1.00

1919

MS Integration Params: RTEINT.P

Quant Time: Aug 4 8:55 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:58:10 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
91) Benzo[k]fluoranthene	13.06	252	167986	82.04	ng	96
92) Benzo[a]pyrene	13.35	252	173373	91.07	ng	100
93) Indeno[1,2,3-cd]pyrene	14.44	276	182021	88.25	ng	89
94) Dibenzo[a,h]anthracene	14.47	278	146409	85.53	ng	98
95) Benzo[g,h,i]perylene	14.72	276	152079	88.16	ng	93

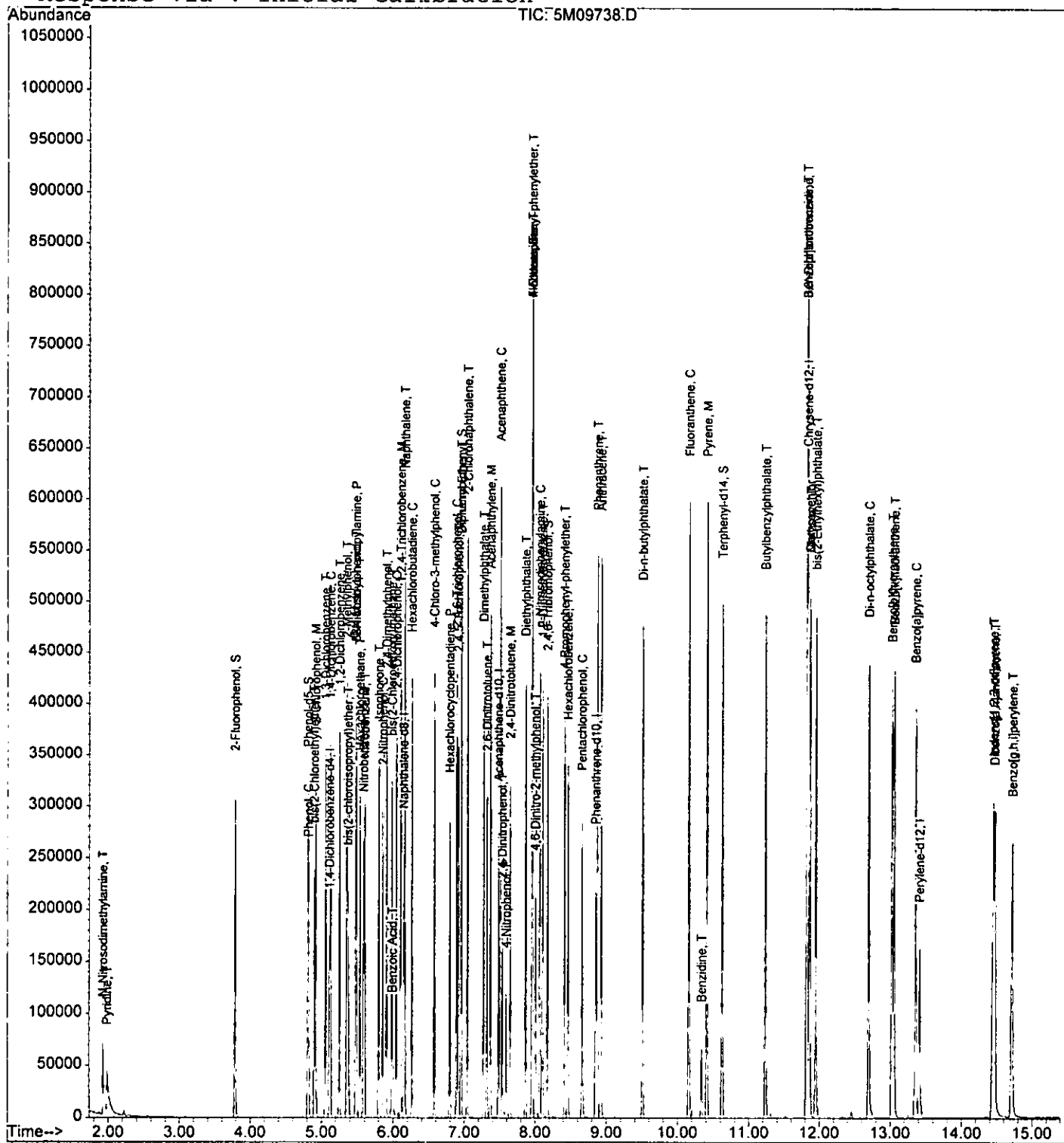
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-04-05\5M09738.D
Acq On : 4 Aug 2005 7:31
Sample : WMB2631 (MS)
Misc : A, BNA
MS Integration Params: RTE.INT.P
Quant Time: Aug 4 8:55 2005

Vial: 4
Operator: AHD
Inst : GCMS_5
Multiplr: 1.00
Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



FORM 3
Spike Recovery

Batch Number: SMB2610

Mbs File: 4M05387.D

Mbs Name: SMB2610(MS)

Non Spk'd File: 4M05388.D

Ns Name: AC18807-009

Spike File: 4M05389.D

Ms Name: AC18807-011(MS)

Spike Dup File: 4M05390.D

Msd Name: AC18807-012(MS)

Matrix: Soil

Method: 8270

1991

Compound	Col	Mr	Conc	Lo	Hi	Rpd	Mbs	Sample	Spike	Spike	Mbs	MS	Msd	Rpd
			Exp	Llm	Lim	Llm				Dup				
Phenol	1	0	200	26	90	35	133.22	0.00	172.15	140.74	67	86	70	20
2-Chlorophenol	1	0	200	25	102	50	135.23	0.00	168.67	145.80	68	84	73	15
1,4-Dichlorobenzene	1	0	100	28	104	27	78.99	0.00	94.13	85.67	79	94	86	9.4
N-Nitroso-di-n-propyla	1	0	100	41	126	38	67.99	0.00	79.67	79.91	68	80	80	0.3
1,2,4-Trichlorobenzene	1	0	100	38	107	23	82.70	0.00	94.33	80.48	83	94	80	16
4-Chloro-3-methylphen	1	0	200	26	103	33	142.76	0.00	181.87	142.32	71	91	71	24
Acenaphthene	1	0	100	31	137	19	86.63	6.60	94.85	92.25	87	88	86	2.8
2,4-Dinitrotoluene	1	0	100	28	89	47	94.37	0.00	94.95	96.04	94 Mo	95 Mo	96 Mo	1.1
4-Nitrophenol	1	0	200	11	114	50	128.68	0.00	163.81	138.02	64	82	69	17
Pentachlorophenol	1	0	200	17	109	47	164.18	0.00	182.90	161.55	82	91	81	12
Pyrene	1	0	100	35	142	36	77.76	131.50	183.75	181.30	78	52	50	1.3

Note:

Rp = Failed Rpd Criteria

Mo = Failed Recovery Criteria

^ - Both Ms and Msd Recoveries = 0 ... no valid information can be calculated

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05387.D Vial: 4
 Acq On : 5 Aug 2005 9:06 Operator: AHD
 Sample : SMB2610 (MS) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 5 9:23 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.90	152	32436	40.00	ng	-0.04
19) Naphthalene-d8	5.91	136	98141	40.00	ng	-0.04
35) Acenaphthene-d10	7.48	164	50928	40.00	ng	-0.05
59) Phenanthrene-d10	9.08	188	83681	40.00	ng	-0.06
72) Chrysene-d12	12.28	240	72967	40.00	ng	-0.06
81) Perylene-d12	14.13	264	62170	40.00	ng	-0.06
System Monitoring Compounds						
4) 2-Fluorophenol	3.76	112	127380	139.22	ng	-0.04
Spiked Amount	200.000		Recovery	=	69.61%	
7) Phenol-d5	4.62	99	162573	133.48	ng	-0.04
Spiked Amount	200.000		Recovery	=	66.74%	
20) Nitrobenzene-d5	5.34	128	34838	70.91	ng	-0.04
Spiked Amount	100.000		Recovery	=	70.91%	
40) 2-Fluorobiphenyl	6.84	172	120982	74.13	ng	-0.04
Spiked Amount	100.000		Recovery	=	74.13%	
62) 2,4,6-Tribromophenol	8.31	332	66760	178.24	ng	-0.05
Spiked Amount	200.000		Recovery	=	89.12%	
75) Terphenyl-d14	10.99	244	138599	67.42	ng	-0.04
Spiked Amount	100.000		Recovery	=	67.42%	
Target Compounds						Qvalue
8) Phenol	4.64	94	175567	133.22	ng	47
9) 2-Chlorophenol	4.74	128	136432	135.23	ng	98
10) 1,3-Dichlorobenzene	4.92	146	83673	77.28	ng	96
11) 1,4-Dichlorobenzene	4.92	146	83673	78.99	ng	96
12) 1,2-Dichlorobenzene	4.92	146	83673	80.54	ng	98
17) N-Nitroso-di-n-propylamine	5.23	70	58968	67.99	ng	96
22) Isophorone	5.34	82	81698	41.79	ng	61
28) 1,2,4-Trichlorobenzene	5.87	180	71219	82.70	ng	98
32) 4-Chloro-3-methylphenol	6.38	107	127820	142.76	ng	91
48) 2,6-Dinitrotoluene	7.48	165	7515	17.54	ng	40
49) Acenaphthene	7.51	153	124277	86.63	ng	98
53) 2,4-Dinitrotoluene	7.69	165	52873	94.37	ng	76
54) 4-Nitrophenol	7.61	65	61784	128.68	ng	97
66) Pentachlorophenol	8.87	266	64991	164.18	ng	97
70) Di-n-butylphthalate	9.81	149	8413	2.95	ng	94
73) Pyrene	10.76	202	218233	77.76	ng	88

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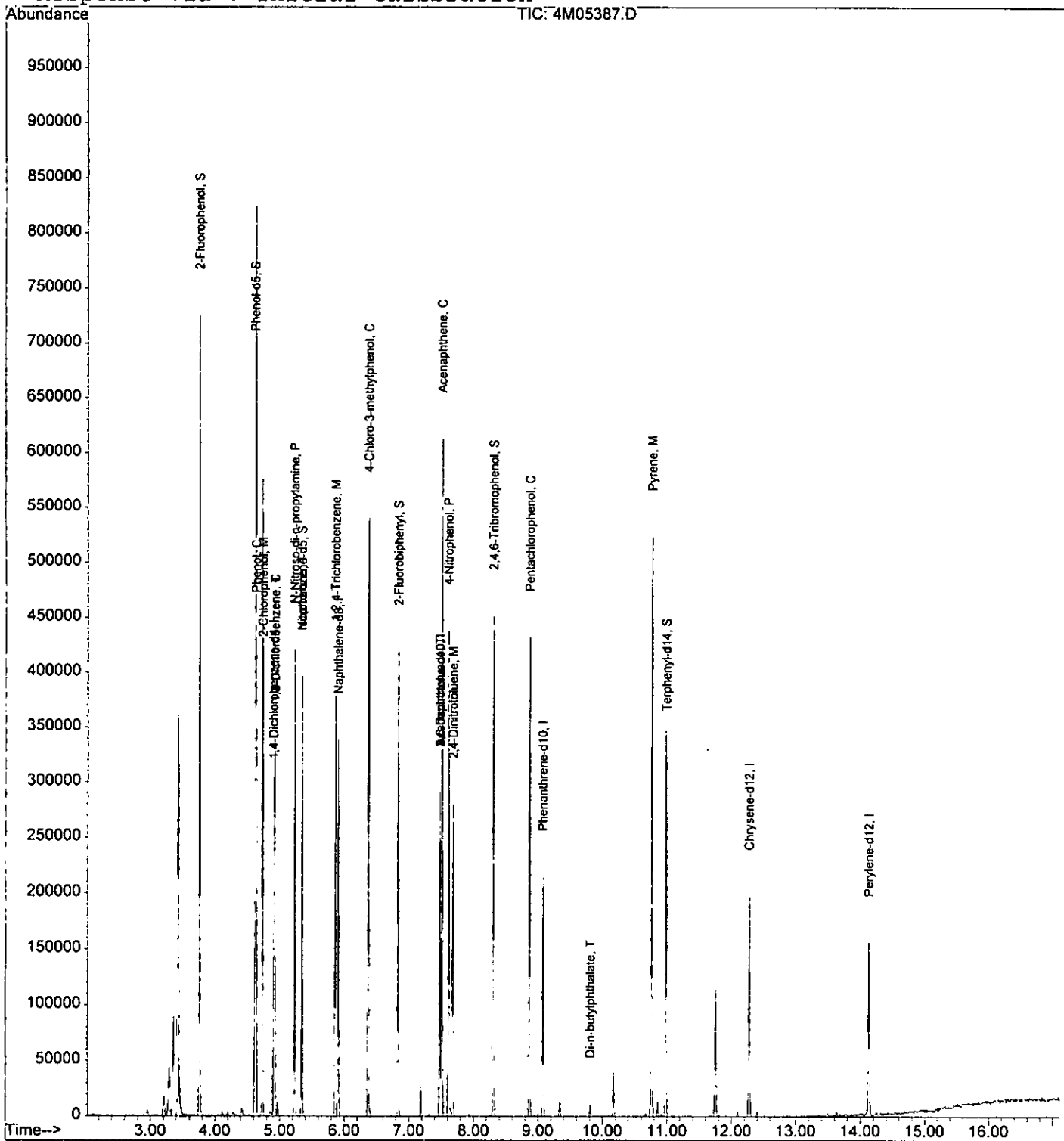
(#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05387.D Vial: 4
Acq On : 5 Aug 2005 9:06 Operator: AHD
Sample : SMB2610 (MS) Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 5 9:23 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05389.D Vial: 6
 Acq On : 5 Aug 2005 9:54 Operator: AHD
 Sample : AC18807-011 (MS:AC18807-009) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 11 12:09 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.91	152	27971	40.00	ng	-0.03
19) Naphthalene-d8	5.90	136	84816	40.00	ng	-0.04
35) Acenaphthene-d10	7.48	164	47798	40.00	ng	-0.05
59) Phenanthrene-d10	9.08	188	76921	40.00	ng	-0.05
72) Chrysene-d12	12.28	240	52651	40.00	ng	-0.05
81) Perylene-d12	14.13	264	47092	40.00	ng	-0.05

System Monitoring Compounds

4) 2-Fluorophenol	3.76	112	124266	157.50	ng	-0.04
Spiked Amount	200.000		Recovery	=	78.75%	
7) Phenol-d5	4.62	99	167658	159.63	ng	-0.03
Spiked Amount	200.000		Recovery	=	79.82%	
20) Nitrobenzene-d5	5.34	128	30551	71.95	ng	-0.04
Spiked Amount	100.000		Recovery	=	71.95%	
40) 2-Fluorobiphenyl	6.83	172	124752	81.45	ng	-0.04
Spiked Amount	100.000		Recovery	=	81.45%	
62) 2,4,6-Tribromophenol	8.31	332	63515	184.48	ng	-0.05
Spiked Amount	200.000		Recovery	=	92.24%	
75) Terphenyl-d14	10.98	244	115331	77.75	ng	-0.04
Spiked Amount	100.000		Recovery	=	77.75%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
8) Phenol	4.64	94	195642	172.15	ng	51
9) 2-Chlorophenol	4.74	128	146741	168.67	ng	91
11) 1,4-Dichlorobenzene	4.92	146	85979	94.13	ng	99
17) N-Nitroso-di-n-propylamine	5.24	70	59582	79.67	ng	72
28) 1,2,4-Trichlorobenzene	5.86	180	70209	94.33	ng	97
29) Naphthalene	5.92	128	24276	13.03	ng	96
32) 4-Chloro-3-methylphenol	6.37	107	140724	181.87	ng	95
33) 2-Methylnaphthalene	6.51	142	9037	6.93	ng	94
46) Acenaphthylene	7.33	152	2233	1.09	ng	57
49) Acenaphthene	7.51	153	127706	94.85	ng	99
53) 2,4-Dinitrotoluene	7.68	165	49929	94.95	ng	94
54) 4-Nitrophenol	7.62	65	73818	163.81	ng	80
55) Fluorene	8.05	166	4247	3.25	ng	97
66) Pentachlorophenol	8.87	266	66552	182.90	ng	94
67) Phenanthrene	9.10	178	90914	47.32	ng	98
68) Anthracene	9.16	178	23166	11.87	ng	98
69) Carbazole	9.36	167	15826	8.84	ng	95
70) Di-n-butylphthalate	9.81	149	6493	2.48	ng	85
71) Fluoranthene	10.50	202	220156	110.15	ng	93

(#) = qualifier out of range (m) = manual integration

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Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05389.D Vial: 6
 Acq On : 5 Aug 2005 9:54 Operator: AHD
 Sample : AC18807-011 (MS:AC18807-009) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 11 12:09 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
73) Pyrene	10.77	202	372112	183.75	ng	81
78) Benzo[a]anthracene	12.27	228	160853	97.34	ng	97
79) Chrysene	12.31	228	142427	96.48	ng	98
80) bis(2-Ethylhexyl)phthalate	12.41	149	7639	5.95	ng	97
83) Benzo[b]fluoranthene	13.66	252	218703m	111.64	ng	
84) Benzo[k]fluoranthene	13.69	252	81436m	48.00	ng	
85) Benzo[a]pyrene	14.06	252	166595	103.46	ng	97
86) Indeno[1,2,3-cd]pyrene	15.37	276	121517	83.19	ng	89
87) Dibenzo[a,h]anthracene	15.40	278	40587	33.93	ng	93
88) Benzo[g,h,i]perylene	15.66	276	115215	98.82	ng	91

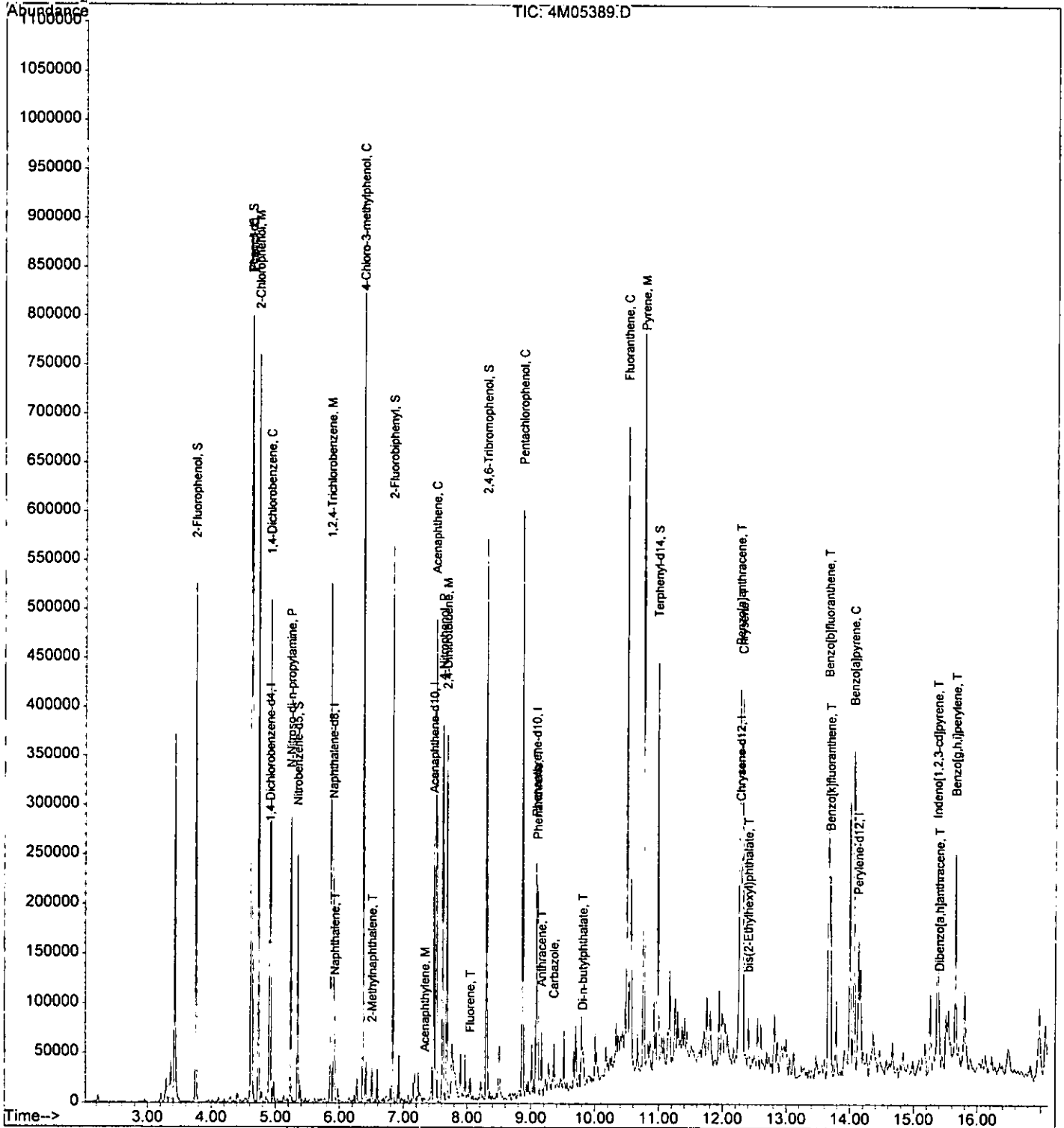
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05389.D Vial: 6
 Acq On : 5 Aug 2005 9:54 Operator: AHD
 Sample : AC18807-011 (MS:AC18807-009) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 11 12:09 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05390.D Vial: 7
 Acq On : 5 Aug 2005 10:18 Operator: AHD
 Sample : AC18807-012 (MSD:AC18807-009) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 11 12:10 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	4.90	152	29435	40.00	ng	-0.04
19) Naphthalene-d8	5.91	136	96821	40.00	ng	-0.04
35) Acenaphthene-d10	7.48	164	48896	40.00	ng	-0.05
59) Phenanthrene-d10	9.08	188	77744	40.00	ng	-0.06
72) Chrysene-d12	12.29	240	54276	40.00	ng	-0.05
81) Perylene-d12	14.14	264	46828	40.00	ng	-0.05

System Monitoring Compounds

4) 2-Fluorophenol	3.76	112	127414	153.46	ng	-0.04
Spiked Amount	200.000		Recovery	=	76.73%	
7) Phenol-d5	4.62	99	164560	148.89	ng	-0.04
Spiked Amount	200.000		Recovery	=	74.44%	
20) Nitrobenzene-d5	5.34	128	35338	72.91	ng	-0.04
Spiked Amount	100.000		Recovery	=	72.91%	
40) 2-Fluorobiphenyl	6.84	172	118876	75.87	ng	-0.04
Spiked Amount	100.000		Recovery	=	75.87%	
62) 2,4,6-Tribromophenol	8.31	332	64407	185.09	ng	-0.05
Spiked Amount	200.000		Recovery	=	92.55%	
75) Terphenyl-d14	10.99	244	114975	75.19	ng	-0.04
Spiked Amount	100.000		Recovery	=	75.19%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
8) Phenol	4.63	94	168322	140.74	ng	61
9) 2-Chlorophenol	4.73	128	133490	145.80	ng	69
11) 1,4-Dichlorobenzene	4.93	146	82346	85.67	ng	98
17) N-Nitroso-di-n-propylamine	5.23	70	62888	79.91	ng	91
28) 1,2,4-Trichlorobenzene	5.87	180	68375	80.48	ng	95
29) Naphthalene	5.93	128	18779	8.83	ng	98
32) 4-Chloro-3-methylphenol	6.37	107	125710	142.32	ng	82
33) 2-Methylnaphthalene	6.50	142	8799	5.91	ng	99
49) Acenaphthene	7.51	153	127060	92.25	ng	100
53) 2,4-Dinitrotoluene	7.69	165	51662	96.04	ng	72
54) 4-Nitrophenol	7.61	65	63627	138.02	ng	96
55) Fluorene	8.04	166	4268	3.20	ng	89
66) Pentachlorophenol	8.87	266	59411	161.55	ng	95
67) Phenanthrene	9.11	178	94537	48.68	ng	98
68) Anthracene	9.17	178	25730	13.05	ng	97
69) Carbazole	9.36	167	17205	9.51	ng	98
70) Di-n-butylphthalate	9.81	149	5880	2.22	ng	88
71) Fluoranthene	10.51	202	234596	116.13	ng	93
73) Pyrene	10.76	202	378479	181.30	ng	95

(#) = qualifier out of range (m) = manual integration

hmm

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05390.D Vial: 7
 Acq On : 5 Aug 2005 10:18 Operator: AHD
 Sample : AC18807-012 (MSD:AC18807-009) Inst : GCMS_4
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 11 12:10 2005

Quant Results File: 4M_0803.RES

Quant Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
 Title : @GCMS_4,mg,625,8270
 Last Update : Wed Aug 03 12:10:40 2005
 Response via : Initial Calibration
 DataAcq Meth : 4M_0803

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
78) Benzo[a]anthracene	12.27	228	168738	99.05	ng	98
79) Chrysene	12.32	228	181344	119.17	ng	97
80) bis(2-Ethylhexyl)phthalate	12.41	149	8017	6.06	ng	95
83) Benzo[b]fluoranthene	13.67	252	270301m	138.76	ng	
84) Benzo[k]fluoranthene	13.70	252	86354m	51.19	ng	
85) Benzo[a]pyrene	14.07	252	190926	119.24	ng	98
86) Indeno[1,2,3-cd]pyrene	15.37	276	132135	90.97	ng	86
87) Dibenzo[a,h]anthracene	15.40	278	48832	41.06	ng	94
88) Benzo[g,h,i]perylene	15.66	276	119489	103.06	ng	93

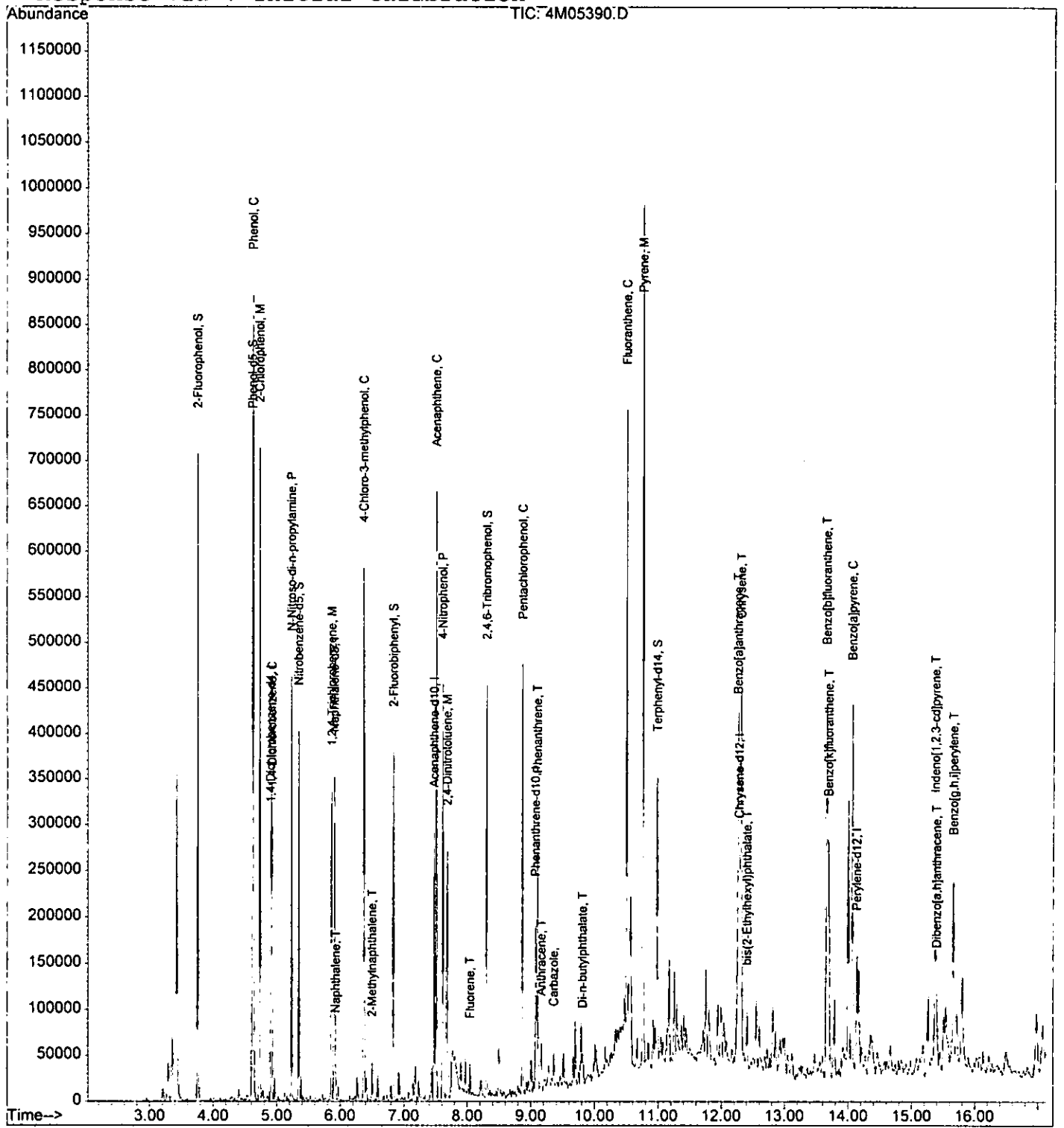
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_4\Data\08-05-05\4M05390.D Vial: 7
Acq On : 5 Aug 2005 10:18 Operator: AHD
Sample : AC18807-012 (MSD:AC18807-009) Inst : GCMS_4
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 11 12:10 2005

Quant Results File: 4M_0803.RES

Method : G:\GCMSDATA\2005\GCMS_4\METHODS\4M_0803.M (RTE Integrator)
Title : @GCMS_4,mg,625,8270
Last Update : Wed Aug 03 12:10:40 2005
Response via : Initial Calibration



FORM 3
Spike Recovery

Batch Number: SMB2613

Mbs File: 5M09831.D

Mbs Name: SMB2613(MS)

Non Spk'd File: 5M09832.D

Ns Name: AC18807-021

Spike File: 5M09833.D

Ms Name: AC18807-021(MS)

Spike Dup File: 5M09834.D

Msd Name: AC18807-021(MS)

Matrix: Soil

Method: 8270

Compound	Col	Mr	Conc Exp	Lo Llm	Hi Lim	Rpd Llm	Mbs Conc	Sample Conc	Spike Conc	Spike Dup Conc	Mbs Rec	MS Rec	Msd Rec	Rpd
Phenol	1	0	200	26	90	35	147.86	0.00	143.63	153.49	74	72	77	6.6
2-Chlorophenol	1	0	200	25	102	50	146.61	0.00	133.72	147.48	73	67	74	9.8
1,4-Dichlorobenzene	1	0	100	28	104	27	80.84	0.00	69.72	85.31	81	70	85	20
N-Nitroso-di-n-propyla	1	0	100	41	126	38	78.31	0.00	76.52	80.88	78	77	81	5.5
1,2,4-Trichlorobenzene	1	0	100	38	107	23	76.91	0.00	71.96	78.73	77	72	79	9
4-Chloro-3-methylphen	1	0	200	26	103	33	145.20	0.00	151.98	151.51	73	76	76	0.31
Acenaphthene	1	0	100	31	137	19	82.57	0.00	78.51	86.92	83	79	87	10
2,4-Dinitrotoluene	1	0	100	28	89	47	84.43	0.00	76.86	88.56	84	77	89	14
4-Nitrophenol	1	0	200	11	114	50	162.73	0.00	183.15	182.85	81	92	91	0.16
Pentachlorophenol	1	0	200	17	109	47	156.30	0.00	165.58	174.20	78	83	87	5.1
Pyrene	1	0	100	35	142	36	87.65	2.46	97.34	93.60	88	95	91	3.9

Note:

Rp = Failed Rpd Criteria

Mo = Failed Recovery Criteria

^ - Both Ms and Msd Recoveries = 0 ... no valid information can be calculated

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09831.D Vial: 6
 Acq On : 8 Aug 2005 3:19 Operator: AHD
 Sample : SMB2613 (MS) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P
 Quant Time: Aug 8 9:57 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	29466	40.00	ng	-0.15
20) Naphthalene-d8	6.13	136	120361	40.00	ng	-0.15
36) Acenaphthene-d10	7.46	164	65295	40.00	ng	-0.18
61) Phenanthrene-d10	8.83	188	108689	40.00	ng	-0.20
77) Chrysene-d12	11.80	240	85966	40.00	ng	-0.23
88) Perylene-d12	13.38	264	64418	40.00	ng	-0.23

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev (Min)
4) 2-Fluorophenol	3.77	112	148470	149.60	ng	-0.19
Spiked Amount				200.000		
			Recovery	=	74.80%	
8) Phenol-d5	4.80	99	198423	136.73	ng	-0.15
Spiked Amount				200.000		
			Recovery	=	68.36%	
21) Nitrobenzene-d5	5.57	128	36827	69.88	ng	-0.15
Spiked Amount				100.000		
			Recovery	=	69.88%	
41) 2-Fluorobiphenyl	6.94	172	147091	72.07	ng	-0.15
Spiked Amount				100.000		
			Recovery	=	72.07%	
64) 2,4,6-Tribromophenol	8.15	330	37367	160.64	ng	-0.19
Spiked Amount				200.000		
			Recovery	=	80.32%	
80) Terphenyl-d14	10.60	244	155513	76.57	ng	-0.21
Spiked Amount				100.000		
			Recovery	=	76.57%	

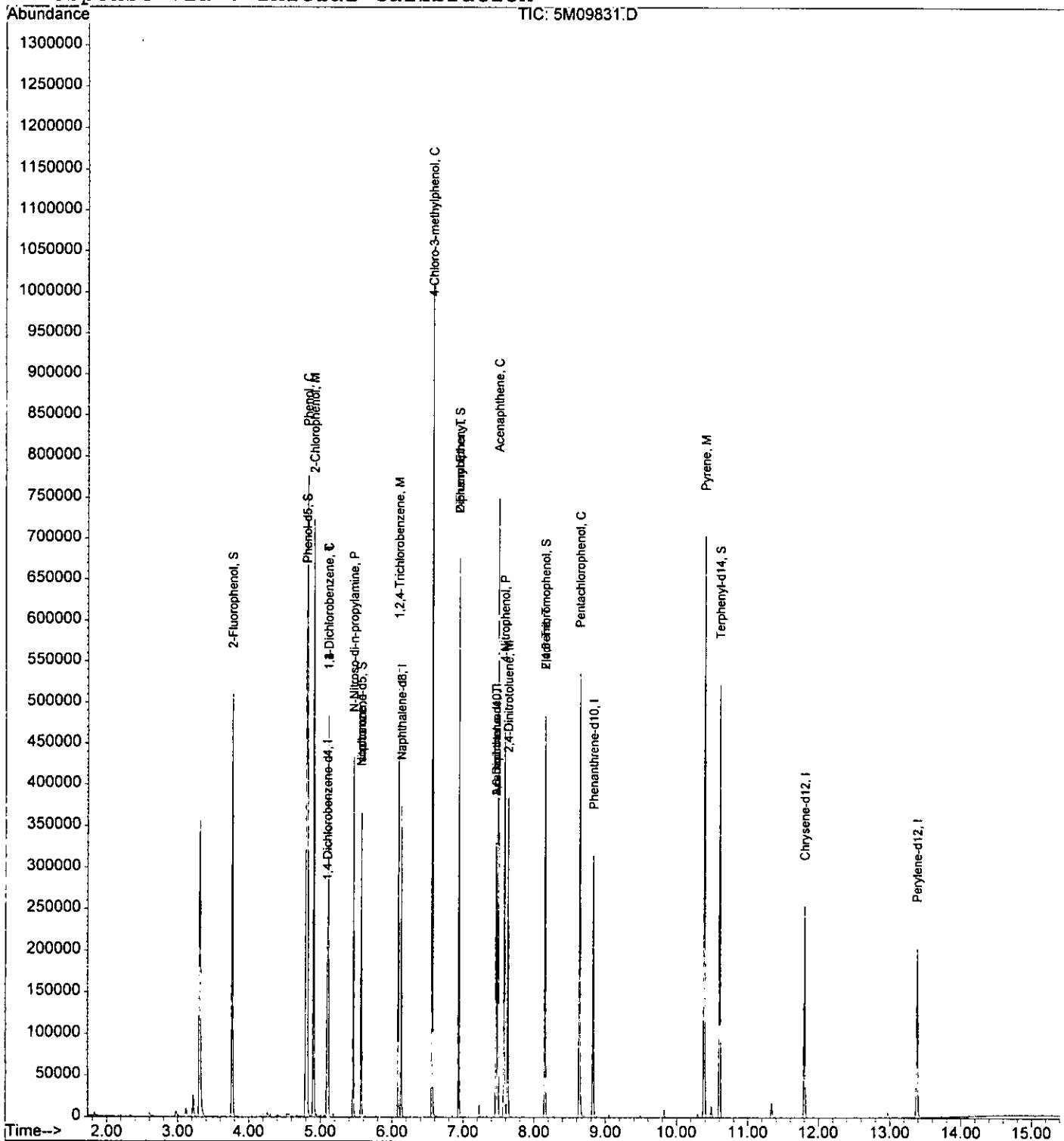
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
9) Phenol	4.82	94	227431	147.86	ng	95
10) 2-Chlorophenol	4.91	128	171161	146.61	ng	99
11) 1,3-Dichlorobenzene	5.11	146	89363	82.73	ng	98
12) 1,4-Dichlorobenzene	5.11	146	89363	80.84	ng	100
13) 1,2-Dichlorobenzene	5.11	146	89363	84.79	ng	97
18) N-Nitroso-di-n-propylamine	5.46	70	65810	78.31	ng	98
23) Isophorone	5.57	82	85024	38.64	ng	60
29) 1,2,4-Trichlorobenzene	6.09	180	75868	76.91	ng	99
33) 4-Chloro-3-methylphenol	6.56	107	154272	145.20	ng	94
45) Diphenyl Ether	6.94	170	33229	27.25	ng	27
49) 2,6-Dinitrotoluene	7.46	165	8422	17.16	ng	29
50) Acenaphthene	7.49	153	148903	82.57	ng	100
54) 2,4-Dinitrotoluene	7.63	165	57233	84.43	ng	91
55) 4-Nitrophenol	7.58	65	69395	162.73	ng	92
57) Fluorene	8.15	166	2867	1.36	ng	74
69) Pentachlorophenol	8.64	266	53909	156.30	ng	93
78) Pyrene	10.39	202	301771	87.65	ng	98

12811

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09831.D Vial: 6
Acq On : 8 Aug 2005 3:19 Operator: AHD
Sample : SMB2613 (MS) Inst : GCMS_5
Misc : S,BNA Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Aug 8 9:57 2005 Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
Title : @GCMS_5,mg,625,8270
Last Update : Fri Jul 22 11:19:45 2005
Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09833.D Vial: 8
 Acq On : 8 Aug 2005 9:02 Operator: AHD
 Sample : AC18807-021(MS) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 9:57 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	24317	40.00	ng	-0.15
20) Naphthalene-d8	6.13	136	92443	40.00	ng	-0.15
36) Acenaphthene-d10	7.46	164	52500	40.00	ng	-0.18
61) Phenanthrene-d10	8.83	188	80267	40.00	ng	-0.20
77) Chrysene-d12	11.80	240	55829	40.00	ng	-0.23
88) Perylene-d12	13.39	264	39737	40.00	ng	-0.23

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
4) 2-Fluorophenol	3.77	112	121793	148.71	ng	-0.20
Spiked Amount	200.000		Recovery	=	74.36%	
8) Phenol-d5	4.80	99	168974	141.09	ng	-0.15
Spiked Amount	200.000		Recovery	=	70.55%	
21) Nitrobenzene-d5	5.57	128	29809	73.65	ng	-0.15
Spiked Amount	100.000		Recovery	=	73.65%	
41) 2-Fluorobiphenyl	6.94	172	121111	73.80	ng	-0.15
Spiked Amount	100.000		Recovery	=	73.80%	
64) 2,4,6-Tribromophenol	8.15	330	27195	158.31	ng	-0.19
Spiked Amount	200.000		Recovery	=	79.16%	
80) Terphenyl-d14	10.60	244	109999	83.40	ng	-0.21
Spiked Amount	100.000		Recovery	=	83.40%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
9) Phenol	4.82	94	182317	143.63	ng	99
10) 2-Chlorophenol	4.91	128	128835	133.72	ng	95
11) 1,3-Dichlorobenzene	5.11	146	63602	71.35	ng	99
12) 1,4-Dichlorobenzene	5.11	146	63602	69.72	ng	99
13) 1,2-Dichlorobenzene	5.11	146	63602	73.13	ng	98
18) N-Nitroso-di-n-propylamine	5.46	70	53069	76.52	ng	95
23) Isophorone	5.57	82	73286	43.37	ng	60
29) 1,2,4-Trichlorobenzene	6.09	180	54524	71.96	ng	98
33) 4-Chloro-3-methylphenol	6.56	107	124025	151.98	ng	98
45) Diphenyl Ether	6.94	170	27386	27.93	ng	27
49) 2,6-Dinitrotoluene	7.46	165	6779	17.18	ng	29
50) Acenaphthene	7.49	153	113841	78.51	ng	99
54) 2,4-Dinitrotoluene	7.63	165	41893	76.86	ng	98
55) 4-Nitrophenol	7.58	65	62800	183.15	ng	87
57) Fluorene	8.15	166	2330	1.37	ng	83
69) Pentachlorophenol	8.64	266	42175	165.58	ng	94
70) Phenanthrene	8.85	178	3814	1.65	ng	99
78) Pyrene	10.39	202	217657	97.34	ng	98
85) Benzo[a]anthracene	11.79	228	2212	1.08	ng	93

(#) = qualifier out of range (m) = manual integration

hsh

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09833.D Vial: 8
 Acq On : 8 Aug 2005 9:02 Operator: AHD
 Sample : AC18807-021(MS) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00

1834

MS Integration Params: RTEINT.P

Quant Time: Aug 8 9:57 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
86) Chrysene	11.83	228	2406	1.28	ng	94
87) bis(2-Ethylhexyl)phthalate	11.93	149	4245	3.12	ng	77
90) Benzo[b]fluoranthene	12.99	252	2771	1.77	ng	96
91) Benzo[k]fluoranthene	12.99	252	2771	1.74	ng	96

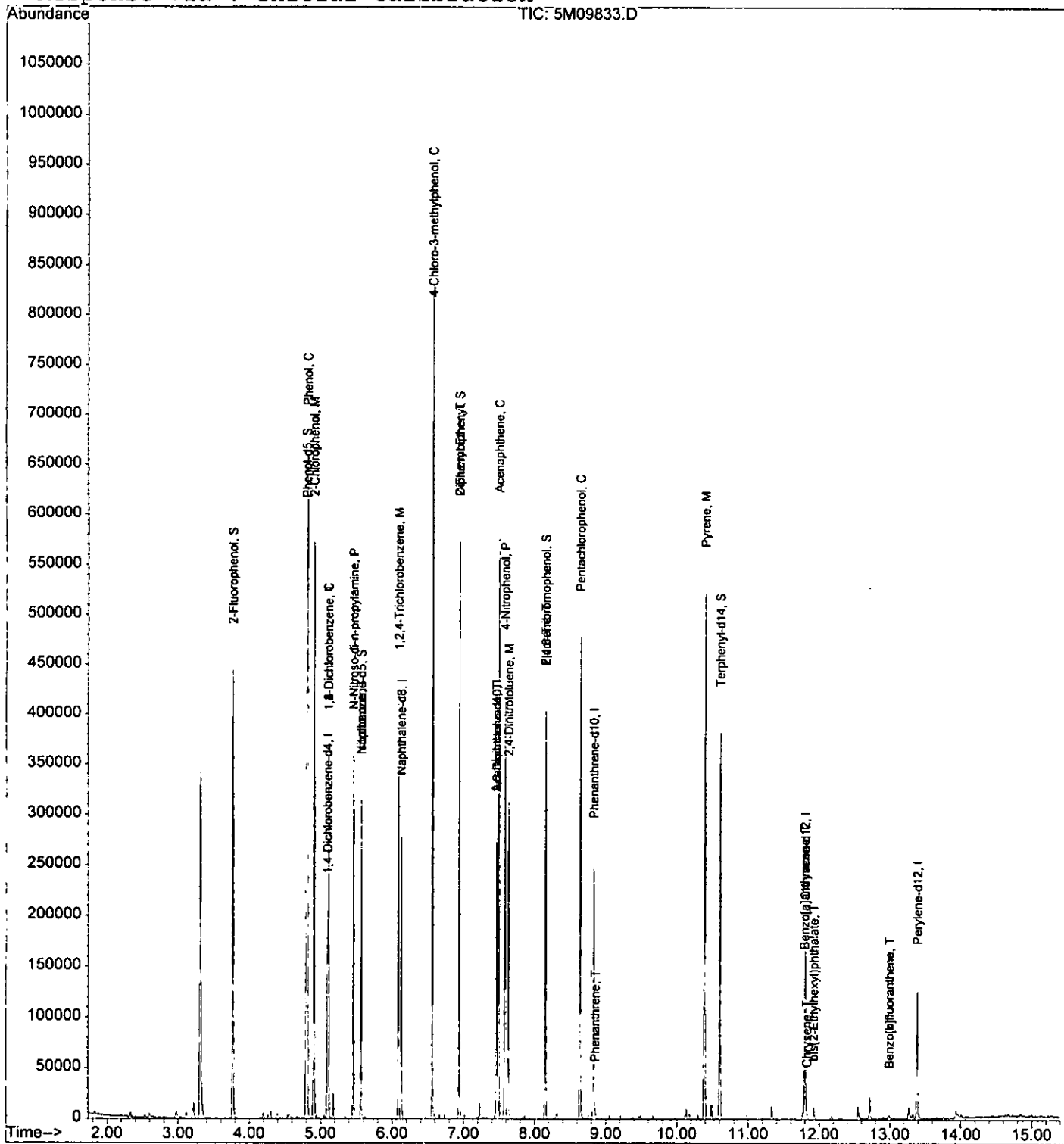
 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09833.D Vial: 8
 Acq On : 8 Aug 2005 9:02 Operator: AHD
 Sample : AC18807-021 (MS) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 9:57 2005

Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09834.D Vial: 9
 Acq On : 8 Aug 2005 9:24 Operator: AHD
 Sample : AC18807-021(MSD) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Aug 8 9:58 2005

Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	5.10	152	25141	40.00	ng	-0.15
20) Naphthalene-d8	6.13	136	101209	40.00	ng	-0.15
36) Acenaphthene-d10	7.47	164	53756	40.00	ng	-0.17
61) Phenanthrene-d10	8.83	188	88987	40.00	ng	-0.20
77) Chrysene-d12	11.80	240	64455	40.00	ng	-0.23
88) Perylene-d12	13.39	264	48834	40.00	ng	-0.23

System Monitoring Compounds

4) 2-Fluorophenol	3.77	112	133492	157.65	ng	-0.20
Spiked Amount	200.000		Recovery	=	78.83%	
8) Phenol-d5	4.80	99	180944	146.14	ng	-0.15
Spiked Amount	200.000		Recovery	=	73.07%	
21) Nitrobenzene-d5	5.57	128	33936	76.58	ng	-0.15
Spiked Amount	100.000		Recovery	=	76.58%	
41) 2-Fluorobiphenyl	6.94	172	130100	77.43	ng	-0.15
Spiked Amount	100.000		Recovery	=	77.43%	
64) 2,4,6-Tribromophenol	8.16	330	30298	159.09	ng	-0.19
Spiked Amount	200.000		Recovery	=	79.55%	
80) Terphenyl-d14	10.60	244	126614	83.15	ng	-0.21
Spiked Amount	100.000		Recovery	=	83.15%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
9) Phenol	4.82	94	201437	153.49	ng	97
10) 2-Chlorophenol	4.91	128	146904	147.48	ng	96
11) 1,3-Dichlorobenzene	5.11	146	80464	87.30	ng	98
12) 1,4-Dichlorobenzene	5.11	146	80464	85.31	ng	98
13) 1,2-Dichlorobenzene	5.11	146	80464	89.49	ng	96
18) N-Nitroso-di-n-propylamine	5.46	70	57992	80.88	ng	96
23) Isophorone	5.57	82	79064	42.74	ng	60
29) 1,2,4-Trichlorobenzene	6.09	180	65304	78.73	ng	98
33) 4-Chloro-3-methylphenol	6.56	107	135369	151.51	ng	96
45) Diphenyl Ether	6.94	170	29215	29.10	ng	27
49) 2,6-Dinitrotoluene	7.47	165	6942	17.18	ng	29
50) Acenaphthene	7.49	153	129058	86.92	ng	98
54) 2,4-Dinitrotoluene	7.63	165	49423	88.56	ng	96
55) 4-Nitrophenol	7.58	65	64197	182.85	ng	90
57) Fluorene	8.15	166	2408	1.38	ng	71
69) Pentachlorophenol	8.64	266	49191	174.20	ng	93
70) Phenanthrene	8.85	178	4433	1.73	ng	96
78) Pyrene	10.39	202	241633	93.60	ng	99
85) Benzo[a]anthracene	11.79	228	3095	1.31	ng	93

(#) = qualifier out of range (m) = manual integration

Handwritten signature

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09834.D Vial: 9
 Acq On : 8 Aug 2005 9:24 Operator: AHD
 Sample : AC18807-021(MSD) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 9:58 2005 Quant Results File: 5M_0722.RES

Quant Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration
 DataAcq Meth : 5M_RUN5

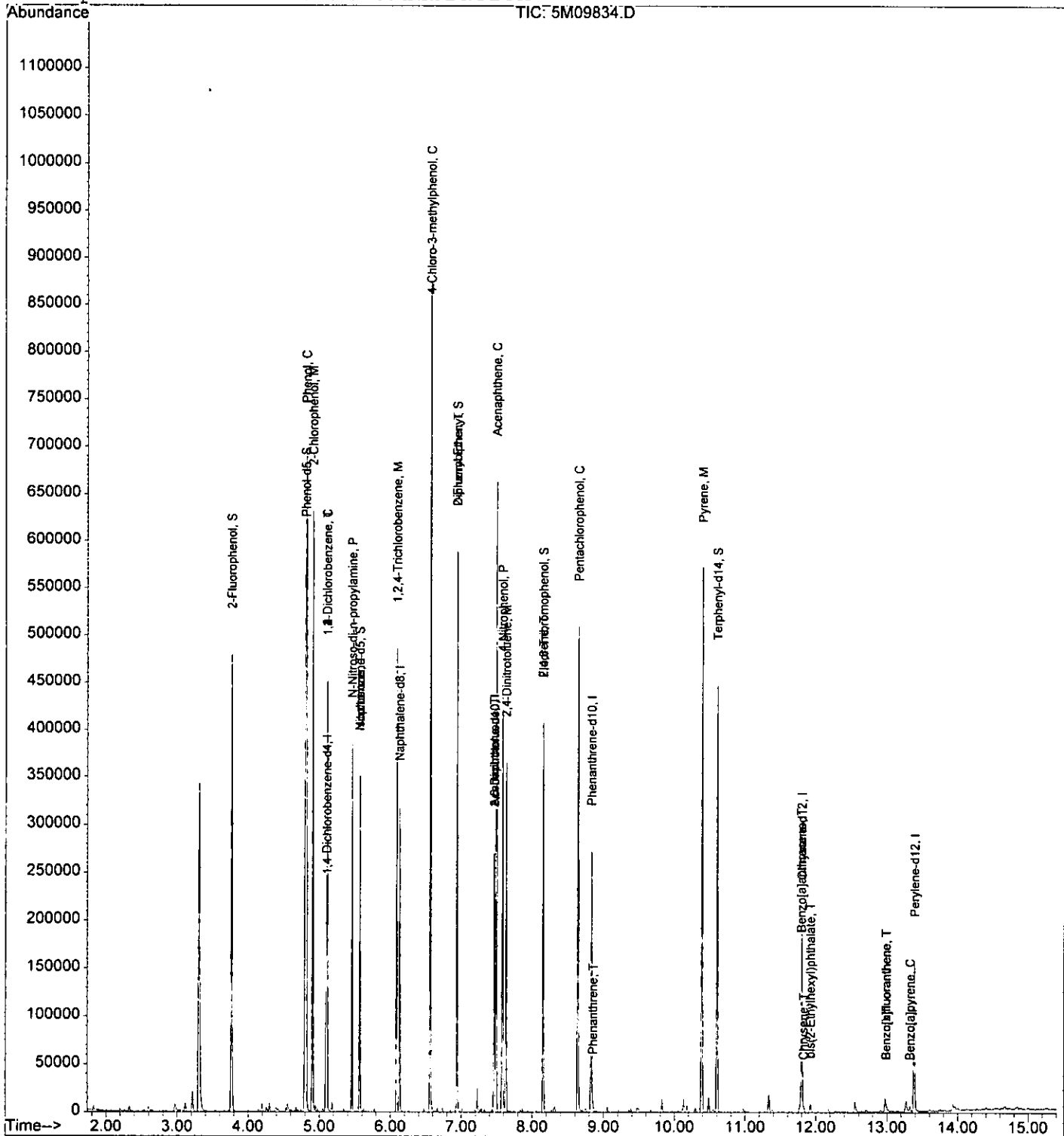
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
86) Chrysene	11.83	228	3164	1.46	ng	92
87) bis(2-Ethylhexyl)phthalate	11.93	149	3584	2.28	ng	83
90) Benzo[b]fluoranthene	12.99	252	3977	2.06	ng	89
91) Benzo[k]fluoranthene	12.99	252	3977	2.04	ng	90
92) Benzo[a]pyrene	13.32	252	2146	1.18	ng	100

 (#) = qualifier out of range (m) = manual integration

Quantitation Report

Data File : G:\GcMsData\2005\Gcms_5\Data\08-08-05\5M09834.D Vial: 9
 Acq On : 8 Aug 2005 9:24 Operator: AHD
 Sample : AC18807-021 (MSD) Inst : GCMS_5
 Misc : S,BNA Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Aug 8 9:58 2005 Quant Results File: 5M_0722.RES

Method : G:\GCMSDATA\2005\GCMS_5\METHODS\5M_0722.M (RTE Integrator)
 Title : @GCMS_5,mg,625,8270
 Last Update : Fri Jul 22 11:19:45 2005
 Response via : Initial Calibration



**GC/MS Semi-Volatile Data
Extraction/Logbook Data**

RUN LOG

Instrument: GCMS_5 Year: 2005
Analyst: AHD

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	8000			BIKFile	
											Cal 600	Beg Cal	End Cal		
5M09384	CAL DFTPP								07/22 08:08						
5M09385	CAL BNA@50PPM				Aqueou	1	1	625 8270	07/22 08:30	5M09385					
5M09386	CAL BNA@10PPM				Aqueou	1	1	625 8270	07/22 08:53	5M09385					
5M09387	CAL BNA@25PPM				Aqueou	1	1	625 8270	07/22 09:16	5M09385					
5M09388	CAL BNA@80PPM				Aqueou	1	1	625 8270	07/22 09:39	5M09385					
5M09389	CAL BNA@120PPM				Aqueou	1	1	625 8270	07/22 10:01	5M09385					
5M09390	CAL BNA@160PPM	Oc			Aqueou	1	1	625 8270	07/22 10:24	5M09385					
5M09391	CAL BNA@200PPM	Oc			Aqueou	1	1	625 8270	07/22 10:47	5M09385					
5M09392	AC18716-003				BNPAH-8270	Aqueou	1	1	8270	07/22 11:29	5M09385		5M09385		
5M09393	AC18623-013(R)				BN15-625	Aqueou	1	1	625	07/22 11:52	5M09385	5M09385	5M09385		
5M09394	AC18669-004(T)				BNATCLP-82	Aqueou	1	1	8270	07/22 12:15	5M09385		5M09385		
5M09395	WMB2620				Aqueou	1	1	625 8270	07/22 12:38	5M09385	5M09385	5M09385			
5M09396	AC18716-001				BNPAH-8270	Soil	1	1	8270	07/22 13:00	5M09385		5M09385		
5M09397	AC18716-002				BNPAH-8270	Soil	1	1	8270	07/22 13:23	5M09385		5M09385		
5M09398	WMB2620(MS)	M18b	WMB2620		Aqueou	1	1	625 8270	07/22 13:46	5M09385	5M09385	5M09385			
5M09399	AC18623-007(R)	Sb6			BN15-625	Aqueou	1	1	625	07/22 14:09	5M09385	5M09385	5M09385		
5M09400	WMB2621				Aqueou	1	1	625 8270	07/22 14:32	5M09385	5M09385	5M09385			
5M09401	WMB2621(MS)	M18a	WMB2621		Aqueou	1	1	625 8270	07/22 14:55	5M09385	5M09385	5M09385			
5M09402	AC18667-001		WMB2621		BNA-625	Aqueou	1	1	625	07/22 15:18	5M09385	5M09385	5M09385		
5M09403	AC18667-001(MS)	M16bM18a	WMB2621		BNA-625	Aqueou	1	1	625 8270	07/22 15:41	5M09385	5M09385	5M09385		
5M09404	AC18667-001(MSD)	M18aM18b	WMB2621		BNA-625	Aqueou	1	1	625 8270	07/22 16:04	5M09385	5M09385	5M09385		
5M09405	SMB2594				Soil	1	1	8270	07/22 16:27	5M09385		5M09385			
5M09406	SMB2594(MS)	OcM18aM18b	SMB2594		Soil	1	1	8270	07/22 16:50	5M09385		5M09385			
5M09407	AC18689-002		SMB2594		BNPAH-8270	Soil	1	1	8270	07/22 17:13	5M09385		5M09385		
5M09408	AC18689-002(MS)	OcM18aM18b	SMB2594		BNPAH-8270	Soil	1	1	8270	07/22 17:36	5M09385		5M09385		
5M09409	AC18689-002(MSD)	OcM18b	SMB2594		BNPAH-8270	Soil	1	1	8270	07/22 17:59	5M09385		5M09385		
5M09410	AC18689-007				BNA25-8270	Soil	1	1	8270	07/22 18:22	5M09385		5M09385		
5M09411	AC18475-001(T)				BNATCLP-82	Aqueou	1	1	8270	07/22 18:46	5M09385		5M09385		
5M09412	EF2V4993				Aqueou	1	1	8270	07/22 19:09	5M09385		5M09385			
5M09413	AC18681-001(5X)				BNSTAR2-82	Aqueou	5	5	8270	07/22 19:31	5M09385		5M09385		
5M09414	AC18657-001				BN15-625	Aqueou	1	1	625	07/22 19:54	5M09385	5M09385	5M09385		
5M09415	AC18666-001				BNA-625	Aqueou	1	1	625	07/22 20:17	5M09385	5M09385	5M09385		
5M09416	AC18691-001				BN15-625	Aqueou	1	1	625	07/22 20:40	5M09385	5M09385	5M09385		
5M09417	AC18698-005				BN15-625	Aqueou	1	1	625	07/22 21:03	5M09385	5M09385	5M09385		
5M09418	AC18661-001(R)				BN15-625	Aqueou	1	1	625	07/22 21:26	5M09385	5M09385	5M09385		
5M09419	AC18711-001				BN15-625	Aqueou	1	1	625	07/22 21:49	5M09385	5M09385	5M09385		

Anc	Area Not Checked	Er	Extraction Performed Past Hold	Co	Warning Possible Carry Over
Ao	Area Out	EsM	Solvent Extraction Date Missing/Not check'd	R16,R26	Rpd Out on MstMsd (col1 and or col2) 8000 series
B6m	Blank 600 series missing	Ein	Top/Solvent Extraction Date Missing/Not check'd	R18,R28	Rpd Out on MstMsd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eto	Top Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C16	Calibration Column 1 Out (8000 Series)	Hb	Analysis Before Collection Date	S6	8000 series surrogate out
C18	Calibration Column 2 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C26	Calibration Column 1 Out (8000 Series)	I16,I26	Initial cal 6000 series failed Column 1 and or 2	Sa6,Sb6	Acid and or BN Surrogate Out (600 series)
C28	Calibration Column 2 Out (8000 Series)	I18,I28	Initial cal 8000 series failed Column 1 and or 2	Sa8,Sb8	Acid and or BN Surrogate Out (8000 series)
C6f	600 series sample/blank did not have passing cal	Ia	Initial Cal Not Checked	Sd	Surrogate Diluted Out
C6t	6000 series sample/blank did not have passing cal	Iv	Prob with calprt csv for int calibration check rts	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial cal warning. Ini cal file <- method.	T6	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a sample	T8	Outside of 8000 series Tune time/Cal Time
D1o,D2o	Drift Out Column 1 or Column 2 Cals or Init Cals	M16,M26	Spike Out Col 1 and or Col 2 600 series	T8	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M16a,M16b	Spike Out Col 1 600 series Acid and or BN	Tm	Too Many Samples/ for beginning Calibration
Do	Drift Out	M18,M28	Spike Out Col 1 and or Col 2 8000 series	Trw	If for 600 ser Too many samples begin Calibration
E6e	An Extraction Before Collection Date	M18a,M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune Not Checked
Emp	Problem Checking Prep/updates modcheck/preprundates	Mnc	Spike Not Checked for this mst/msd	To	Tune File Failed
En	Eval Time Not Checked	Oc	Warning Compnd(s) Over Calibration	W6	Warning Instrument Id not in TxtLoc field

RUN LOG

Instrument: GCMS_4 Year: 2005
Analyst: AHD

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	Cal 600	8000 Beg Cal	End Cal	BlkFile
4M05295	CAL DFTPP								08/03 06:21					
4M05296		TnlsCnSnc	_Not_Quant'd_											
4M05297	CAL DFTPP								08/03 08:09					
4M05298		TnlsCnSnc	_Not_Quant'd_											
4M05299	CAL BNA@50PPM				Soil	1	1	625 8270	08/03 08:52	4M05299				
4M05300	CAL BNA@10PPM				Soil	1	1	625 8270	08/03 09:19	4M05299				
4M05301	CAL BNA@25PPM				Soil	1	1	625 8270	08/03 09:43	4M05299				
4M05302	CAL BNA@80PPM				Soil	1	1	625 8270	08/03 10:07	4M05299				
4M05303	CAL BNA@120PPM				Soil	1	1	625 8270	08/03 10:31	4M05299				
4M05304	CAL BNA@160PPM	Oc			Soil	1	1	625 8270	08/03 10:55	4M05299				
4M05305	CAL BNA@200PPM	Oc			Soil	1	1	625 8270	08/03 11:19	4M05299				
4M05306	SMB2606	Ao			Soil	1	1	8270	08/03 11:43	4M05299		4M05299		
4M05307	AC18819-004		BNA25-8270		Soil	1	1	8270	08/03 12:06	4M05299		4M05299		
4M05308	AC18819-006		BNA25-8270		Soil	1	1	8270	08/03 12:30	4M05299		4M05299		
4M05309	AC18819-012		BNA25-8270		Soil	1	1	8270	08/03 12:54	4M05299		4M05299		
4M05310	AC18819-018		BNA25-8270		Soil	1	1	8270	08/03 13:18	4M05299		4M05299		
4M05311	SMB2606				Soil	1	1	8270	08/03 13:42	4M05299		4M05299		
4M05312	SMB2605(MS)	M18b	SMB2605		Soil	1	1	8270	08/03 14:06	4M05299		4M05299		
4M05313	AC18819-008(MS)	AoMnc	SMB2605	BNA25-8270	Soil	1	1	8270	08/03 14:30	4M05299		4M05299		
4M05314	AC18819-008(MSD)	Mnc	SMB2605	BNA25-8270	Soil	1	1	8270	08/03 14:54	4M05299		4M05299		
4M05315	AC18802-004		BNA25-8270		Soil	1	1	8270	08/03 15:18	4M05299		4M05299		
4M05316	AC18802-006		BNA25-8270		Soil	1	1	8270	08/03 15:41	4M05299		4M05299		
4M05317	AC18853-002		BNA25-8270		Soil	1	1	8270	08/03 16:05	4M05299		4M05299		
4M05318	AC18853-003		BNA25-8270		Soil	1	1	8270	08/03 16:29	4M05299		4M05299		
4M05319	AC18853-004		BNA25-8270		Soil	1	1	8270	08/03 16:53	4M05299		4M05299		
4M05320	AC18808-001	Ao	BNA-8270		Soil	1	1	8270	08/03 17:17	4M05299		4M05299		
4M05321	AC18802-002	Ao	BNA25-8270		Soil	1	1	8270	08/03 17:42	4M05299		4M05299		
4M05322	AC18802-005		BNA25-8270		Soil	1	1	8270	08/03 18:06	4M05299		4M05299		
4M05323	AC18852-001	Ao	BNPAH-8270		Soil	1	1	8270	08/03 18:30	4M05299		4M05299		
4M05324	AC18853-001	Ao	BNA25-8270		Soil	1	1	8270	08/03 18:54	4M05299		4M05299		
4M05325	AC18847-001		BN-8270		Soil	1	1	8270	08/03 19:18	4M05299		4M05299		
4M05326	AC18802-001	Ao	BNA25-8270		Soil	1	1	8270	08/03 19:42	4M05299		4M05299		
4M05327	AC18786-013	Ao	BNA25-8270		Soil	1	1	8270	08/03 20:06	4M05299		4M05299		
4M05328	AC18786-014	Ti8Ao	BNA25-8270		Soil	1	1	8270	08/03 20:30	4M05299		4M05299		
4M05329	AC18796-007	Ti8	BNA-8270		Soil	1	1	8270	08/03 20:54	4M05299		4M05299		

Anc	Area Not Checked	Ex	Extraction Performed Past Hold	Co	Warning Possible Carry Over
Ao	Area Out	Exm	Solvent Extraction Date Missing/Not check'd	R18, R28	Rpd Out on MsMsd (col1 and or col2) 600 series
B6m	Blank 600 series missing	Ein	Top/Solvent Extraction Date Missing/Not check'd	R18, R28	Rpd Out on MsMsd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eto	Top Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C16	Calibration Column 1 Out (800 Series)	Hb	Analysis Before Collection Date	S6	600 series surrogate out
C18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C26	Calibration Column 2 Out (600 Series)	I16, I28	Initial cal 600 series failed Column 1 and or 2	Se6, Sb6	Acid and or BN Surrogate Out (600 series)
C28	Calibration Column 2 Out (8000 Series)	I18, I28	Initial cal 8000 series failed Column 1 and or 2	Se8, Sb8	Acid and or BN Surrogate Out (8000 series)
C6f	600 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Deleted Out
C8f	8000 series sample/blank did not have passing cal	Iv	Prob with calrpt.csv for int calibration check rfs	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial cal warning; Ini cal file <- method..	T15	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a sample	T6	Outside of 600 series Tune time/Cal Time
D1o, D2o	Drift Out Column 1 or Column 2 Cals or Int Cals	M16, M28	Spike Out Col 1 and or Col 2 600 series	T8	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M16a, M18b	Spike Out Col 1 600 series Acid and or BN	Tm	Too Many Samples/ for beginning Calibration
Do	Drift Out	M18, M28	Spike Out Col 1 and or Col 2 8000 series	Tmw	If for 600 ser Too many samples begin Calibration
Eba	An Extraction Before Collection Date	M18a, M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune Not Checked
Emp	Problem Checking Prep/updates mod/check/prep/updates	Mnc	Spike Not Check'ed for this ms/msd	To	Tune File Failed
En	Eval Time Not Checked	Oc	Warning Compound(s) Over Calibration	W6	Warning... Instrument Id not in TxtLoc field

RUN LOG

Instrument: GCMS_5 Year: 2005
Analyst: AHD

1845

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis		Cal 600	8000		BlkFile
									Date	IniCal		Beg Cal	End Cal	
5M09735	CAL DFTPP								08/04 06:25					
5M09736	CAL BNA@50PPM				Aqueou	1	1	625 8270	08/04 06:44	5M09385				
5M09737	WMB2631				Aqueou	1	1	625 8270	08/04 07:06	5M09385	5M09736	5M09736		
5M09738	WMB2631(MS)		WMB2631		Aqueou	1	1	625 8270	08/04 07:31	5M09385	5M09736	5M09736		
5M09739	AC18852-001(T)			BNATCLP-82	Aqueou	1	1	8270	08/04 07:53	5M09385		5M09736		
5M09740	AC18832-004		WMB2631	BN15-625	Aqueou	1	1	625	08/04 08:15	5M09385	5M09736	5M09736		
5M09741	AC18832-004(MS)		WMB2631	BN15-625	Aqueou	1	1	625 8270	08/04 08:37	5M09385	5M09736	5M09736		
5M09742	AC18832-004(MSD)	M18a	WMB2631	BN15-625	Aqueou	1	1	625 8270	08/04 08:59	5M09385	5M09736	5M09736		
5M09743	AC18897-001			BN15-625	Aqueou	1	1	625	08/04 09:20	5M09385	5M09736	5M09736		
5M09744	AC18897-002			BN15-625	Aqueou	1	1	625	08/04 09:42	5M09385	5M09736	5M09736		
5M09745	SMB2607				Soil	1	1	8270	08/04 10:04	5M09385		5M09736		
5M09746	SMB2608				Soil	1	1	8270	08/04 10:26	5M09385		5M09736		
5M09747	SMB2608(MS)		SMB2608		Soil	1	1	8270	08/04 10:48	5M09385		5M09736		
5M09748	AC18855-001		SMB2608	BNA25-8270	Soil	1	1	8270	08/04 11:09	5M09385		5M09736		
5M09749	AC18855-001(MS)		SMB2608	BNA25-8270	Soil	1	1	8270	08/04 11:31	5M09385		5M09736		
5M09750	AC18855-001(MSD)		SMB2608	BNA25-8270	Soil	1	1	8270	08/04 11:53	5M09385		5M09736		
5M09751	AC18807-007			BNA-8270	Aqueou	1	1	8270	08/04 12:15	5M09385		5M09736		
5M09752	AC18820-012			BN-8270	Aqueou	1	1	8270	08/04 12:37	5M09385		5M09736		
5M09753	AC18847-005			BN-8270	Soil	1	1	8270	08/04 12:59	5M09385		5M09736		
5M09754	AC18847-013			BN-8270	Soil	1	1	8270	08/04 13:21	5M09385		5M09736		
5M09755	AC18847-014			BN-8270	Soil	1	1	8270	08/04 13:42	5M09385		5M09736		
5M09756	AC18847-015			BN-8270	Soil	1	1	8270	08/04 14:04	5M09385		5M09736		
5M09757	AC18847-016			BN-8270	Soil	1	1	8270	08/04 14:26	5M09385		5M09736		
5M09758	AC18847-017			BN-8270	Soil	1	1	8270	08/04 14:48	5M09385		5M09736		
5M09759	AC18786-005			BNA25-8270	Soil	1	1	8270	08/04 15:10	5M09385		5M09736		
5M09760	AC18786-007			BNA25-8270	Soil	1	1	8270	08/04 15:32	5M09385		5M09736		
5M09761	AC18786-008			BNA25-8270	Soil	1	1	8270	08/04 15:54	5M09385		5M09736		
5M09762	AC18786-009			BNA25-8270	Soil	1	1	8270	08/04 16:16	5M09385		5M09736		
5M09763	AC18786-010			BNA25-8270	Soil	1	1	8270	08/04 16:38	5M09385		5M09736		
5M09764	AC18786-017			BNA25-8270	Soil	1	1	8270	08/04 17:00	5M09385		5M09736		
5M09765	AC18796-015			BNA-8270	Soil	1	1	8270	08/04 17:22	5M09385		5M09736		
5M09766	AC18796-016			BNA-8270	Soil	1	1	8270	08/04 17:44	5M09385		5M09736		
5M09767	AC18796-018			BNA-8270	Soil	1	1	8270	08/04 18:06	5M09385		5M09736		
5M09768	AC18796-019	Ti8		BNA-8270	Soil	1	1	8270	08/04 18:29	5M09385		5M09736		
5M09769	AC18832-001			BN15-625	Aqueou	1	1	625	08/04 18:50	5M09385	5M09736	5M09736		
5M09770	AC18832-002			BN15-625	Aqueou	1	1	625	08/04 19:12	5M09385	5M09736	5M09736		
5M09771	AC18832-003			BN15-625	Aqueou	1	1	625	08/04 19:34	5M09385	5M09736	5M09736		
5M09772	AC18825-005			BN15-625	Aqueou	1	1	625	08/04 19:56	5M09385	5M09736	5M09736		
5M09773	AC18825-007			BN15-625	Aqueou	1	1	625	08/04 20:18	5M09385	5M09736	5M09736		
5M09774	AC18823-001			BN15-625	Aqueou	1	1	625	08/04 20:40	5M09385	5M09736	5M09736		
5M09775	AC18823-003			BN15-625	Aqueou	1	1	625	08/04 21:02	5M09385	5M09736	5M09736		
5M09776	AC18841-001			BN15-625	Aqueou	1	1	625	08/04 21:24	5M09385	5M09736	5M09736		
5M09777	AC18841-002			BN15-625	Aqueou	1	1	625	08/04 21:46	5M09385	5M09736	5M09736		

Anc	Area Not Checked	Ed	Extraction Performed Past Hold	Co	Warning Possible Carry Over
As	Area Out	EsM	Solvent Extraction Date Missing/Not check'd	R18,R26	Rpd Out on MsMsd (col1 and or col2) 600 series
B8m	Blank 600 series missing	Ein	Tcp/Solvent Extraction Date Missing/Not check'd	R18,R28	Rpd Out on MsMsd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eto	Tcp Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Blf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C18	Calibration Column 1 Out (800 Series)	Hb	Analysis Before Collection Date	S6	600 series surrogate out
C18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C28	Calibration Column 2 Out (800 Series)	I18,I26	Initial cal 600 series failed Column 1 and or 2	Sa6,Sb6	Acid and or BN Surrogate Out (600 series)
C28	Calibration Column 2 Out (8000 Series)	I18,I28	Initial cal 8000 series failed Column 1 and or 2	Sa8,Sb8	Acid and or BN Surrogate Out (8000 series)
C8f	600 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Diluted Out
C8f	8000 series sample/blank did not have passing cal	Iv	Prob with calprt csv for int calibration check its	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial cal warning. Ini cal file <- method.	T15	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a sampl	T16	Outside of 600 series Tune time/Cal Time
D1o,D2o	Drift Out Column 1 or Column 2 Cals or Ini Cals	M16,M26	Spike Out Col 1 and or Col 2 600 series	T18	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M18a,M18b	Spike Out Col 1 600 series Acid and or BN	Trn	Too Many Samples/ for beginning Calibration
Do	Drift Out	M16,M28	Spike Out Col 1 and or Col 2 8000 series	Trmw	If for 600 ser Too many samples begin Calibration
Eba	An Extraction Before Collection Date	M18a,M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune Not Checked
Emp	Problem Checking Prep/rundates modcheck/preprund	Mnc	Spike Not Checked for this ms/msd	To	Tune File Failed
En	Eval Time Not Checked	Ioc	Warning Compund(s) Over Calibration	Wle	Warning... Instrument Id not in TxLoc field

RUN LOG

Instrument: GCMS_4 Year: 2005
Analyst: AHD

10/11/05

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis		Cal 600	8000		BlkFile
									Date	IniCal		Beg Cal	End Cal	
4M05383.	CAL DFTPP								08/05 06:36					
4M05384.	CAL BNA@50PPM	AoC16			Soil	1	1	625 8270	08/05 06:58	4M05299				
4M05385.	CAL BNA@50PPM	C16			Soil	1	1	625 8270	08/05 08:18	4M05299				
4M05386.	SMB2610				Soil	1	1	8270	08/05 08:42	4M05299			4M05385	
4M05387.	SMB2610(MS)	M18b	SMB2610		Soil	1	1	8270	08/05 09:06	4M05299			4M05385	
4M05388.	AC18807-009		SMB2610	BNA-8270	Soil	1	1	8270	08/05 09:30	4M05299			4M05385	
4M05389.	AC18807-011(MS:AC18807-011)	M18b	SMB2610	BNA-8270	Soil	1	1	8270	08/05 09:54	4M05299			4M05385	
4M05390.	AC18807-012(MSD:AC18807-012)	M18b	SMB2610	BNA-8270	Soil	1	1	8270	08/05 10:18	4M05299			4M05385	
4M05391.	AC18847-004(R)			BN-8270	Soil	1	1	8270	08/05 10:42	4M05299			4M05385	
4M05392.	SMB2611				Soil	1	1	8270	08/05 11:06	4M05299			4M05385	
4M05393.	AC18881-001			BNA25-8270	Soil	1	1	8270	08/05 11:30	4M05299			4M05385	
4M05394.	AC18883-001(R)			BNA25-8270	Soil	1	1	8270	08/05 11:54	4M05299			4M05385	
4M05395.	AC18881-001(R)			BNA25-8270	Soil	1	1	8270	08/05 12:18	4M05299			4M05385	
4M05396.	AC18778-012			BNA-8270	Soil	1	1	8270	08/05 12:42	4M05299			4M05385	
4M05397.	AC18881-002(3X)	Oc		BNA25-8270	Soil	3	3	8270	08/05 13:06	4M05299			4M05385	
4M05398.	AC18881-006(3X)			BNA25-8270	Soil	3	3	8270	08/05 13:30	4M05299			4M05385	
4M05399.	AC18881-003			BNA25-8270	Soil	1	1	8270	08/05 13:54	4M05299			4M05385	
4M05400.	AC18881-004	Oc		BNA25-8270	Soil	1	1	8270	08/05 14:18	4M05299			4M05385	
4M05401.	AC18855-002			BNA25-8270	Soil	1	1	8270	08/05 14:42	4M05299			4M05385	
4M05402.	AC18855-003			BNA25-8270	Soil	1	1	8270	08/05 15:06	4M05299			4M05385	
4M05403.	AC18855-004			BNA25-8270	Soil	1	1	8270	08/05 15:31	4M05299			4M05385	
4M05404.	AC18807-004			BNA-8270	Soil	1	1	8270	08/05 15:55	4M05299			4M05385	
4M05405.	SMB2612(MS)		SMB2612		Soil	1	1	8270	08/05 16:19	4M05299			4M05385	
4M05406.	SMB2612				Soil	1	1	8270	08/05 16:43	4M05299			4M05385	
4M05407.	AC18907-005			BNPAH-8270	Soil	1	1	8270	08/05 17:07	4M05299			4M05385	
4M05408.	AC18881-004(3X)			BNA25-8270	Soil	3	3	8270	08/05 17:31	4M05299			4M05385	
4M05409.	AC18881-002(30X)			BNA25-8270	Soil	30	30	8270	08/05 17:55	4M05299			4M05385	
4M05410.	AC18920-002(20X)	Sd		BNPAH-8270	Soil	20	20	8270	08/05 18:19	4M05299			4M05385	
4M05411.	AC18920-003(20X)	Ti8		BNPAH-8270	Soil	20	20	8270	08/05 18:43	4M05299			4M05385	
4M05412.	AC18920-001(10X)	SdTi8		BNA25-8270	Soil	10	10	8270	08/05 19:07	4M05299			4M05385	
4M05413.	AC18778-013	Ti8		BNA-8270	Soil	1	1	8270	08/05 19:31	4M05299			4M05385	
4M05414.	AC18778-004	Ti8Oc		BNA-8270	Soil	1	1	8270	08/05 19:55	4M05299			4M05385	
4M05415.	AC18778-001	Ti8		BNA-8270	Soil	1	1	8270	08/05 20:19	4M05299			4M05385	
4M05416.	AC18778-019	Ti8Oc		BNA-8270	Soil	1	1	8270	08/05 20:43	4M05299			4M05385	
4M05417.	AC18778-016	Ti8Oc		BNA-8270	Soil	1	1	8270	08/05 21:07	4M05299			4M05385	
4M05418.	AC18778-007	Ti8		BNA-8270	Soil	1	1	8270	08/05 21:31	4M05299			4M05385	
4M05419.	TEST	Ti8			Soil	1	1	8270	08/05 21:55	4M05299			4M05385	
4M05420.	TEST	Ti8			Soil	1	1	8270	08/05 22:19	4M05299			4M05385	
4M05421.	TEST	Ti8			Soil	1	1	8270	08/05 22:43	4M05299			4M05385	
4M05422.	TEST	Ti8			Soil	1	1	8270	08/05 23:07	4M05299			4M05385	
4M05423.	TEST	Ti8			Soil	1	1	8270	08/05 23:31	4M05299			4M05385	
4M05424.	TEST	Ti8			Soil	1	1	8270	08/05 23:54	4M05299			4M05385	

Anc	Area Not Checked	Er	Extraction Performed Past Hold	Co	Warning Possible Carry Over
As	Area Out	EsM	Solvent Extraction Date Missing/Not check'd	R18, R28	Rpd Out on MsMsd (col1 and or col2) 8000 series
B4m	Blank 8000 series missing	Ein	Top/Solvent Extraction Date Missing/Not check'd	R18, R28	Rpd Out on MsMsd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eio	Top Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C16	Calibration Column 1 Out (8000 Series)	Hb	Analysis Before Collection Date	S6	8000 series surrogate out
C18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C26	Calibration Column 2 Out (8000 Series)	I18, I28	Initial cal 8000 series failed Column 1 and or 2	Sa6, Sb6	Acid and or BN Surrogate Out (800 series)
C28	Calibration Column 2 Out (8000 Series)	I8	Initial Cal Not Checked	Sa8, Sb8	Acid and or BN Surrogate Out (8000 series)
C4f	8000 series sample/blank did not have passing cal	Iv	Prob with calprt.csv for init calibration check rts	Sd	Surrogate Diluted Out
C4f	8000 series sample/blank did not have passing cal	Iw	Initial cal warnin g. Ini cal file <- method.	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Ix	Initial Cal Files Not Updated Properly for a sample	T15	Outside of 8000 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	M16, M28	Spike Out Col 1 and or Col 2 8000 series	T6	Outside of 8000 series Tune time/Cal Time
D1o, D2o	Drift Out Column 1 or Column 2 Cals or Init Cals	M16a, M18b	Spike Out Col 1 8000 series Acid and or BN	T8	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M18a, M18b	Spike Out Col 1 and or Col 2 8000 series	Tm	Too Many Samples/ for beginning Calibration
Do	Drift Out	M18a, M18b	Spike Out Col 1 8000 series Acid and or BN	Tmw	If for 800 ser Too many samples begin Calibration
Eba	An Extraction Before Collection Date	Mnc	Spike Not Checked for this ms/msd	Tn	Tune Not Checked
Emp	Problem Checking Prep/updates modcheck/prep/updates	Oc	Warning Compound(s) Over Calibration	To	Tune File Failed
En	Eval Time Not Checked			We	Warning ... Instrument Id not in TxtLoc field

RUN LOG

Instrument: GCMS_5 Year: 2005
Analyst: AHD

Bioscience Resource Project

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	Cal 600	8000	End Cal	BikFile
												Beg Cal		
5M09826	CAL DFTPP								08/08 06:23					
5M09827	CAL BNA@50PPM				Aqueou	1	1	625 8270	08/08 06:40	5M09385				
5M09828	WMB2633				Aqueou	1	1	625 8270	08/08 07:15	5M09385	5M09827	5M09827		
5M09829	WMB2634				Aqueou	1	1	625 8270	08/08 07:36	5M09385	5M09827	5M09827		
5M09830	SMB2613				Soil	1	1	8270	08/08 07:58	5M09385		5M09827		
5M09831	SMB2613(MS)		SMB2613		Soil	1	1	8270	08/08 08:19	5M09385		5M09827		
5M09832	AC18807-021		SMB2613	BNA-8270	Soil	1	1	8270	08/08 08:41	5M09385		5M09827		
5M09833	AC18807-021(MS)		SMB2613	BNA-8270	Soil	1	1	8270	08/08 09:02	5M09385		5M09827		
5M09834	AC18807-021(MSD)		SMB2613	BNA-8270	Soil	1	1	8270	08/08 09:24	5M09385		5M09827		
5M09835	WMB2634(MS)		WMB2634		Aqueou	1	1	625 8270	08/08 09:46	5M09385	5M09827	5M09827		
5M09836	AC18892-001		WMB2634	BN-8270	Aqueou	1	1	8270	08/08 10:07	5M09385		5M09827		
5M09837	AC18892-001(MS)		WMB2634	BN-8270	Aqueou	1	1	625 8270	08/08 10:29	5M09385	5M09827	5M09827		
5M09838	AC18892-001(MSD)		WMB2634	BN-8270	Aqueou	1	1	625 8270	08/08 10:51	5M09385	5M09827	5M09827		
5M09839	AC18778-017			BNA-8270	Soil	1	1	8270	08/08 11:12	5M09385		5M09827		
5M09840	AC18807-001			BNA-8270	Soil	1	1	8270	08/08 11:34	5M09385		5M09827		
5M09841	AC18888-001			BNA25-8270	Aqueou	1	1	8270	08/08 11:56	5M09385		5M09827		
5M09842	AC18892-002			BN-8270	Aqueou	1	1	8270	08/08 12:17	5M09385		5M09827		
5M09843	AC18892-003			BN-8270	Aqueou	1	1	8270	08/08 12:39	5M09385		5M09827		
5M09844	AC18873-014			BNA-8270	Aqueou	1	1	8270	08/08 13:01	5M09385		5M09827		
5M09845	AC18778-022			BNA-8270	Soil	1	1	8270	08/08 13:23	5M09385		5M09827		
5M09846	AC18778-023			BNA-8270	Soil	1	1	8270	08/08 13:44	5M09385		5M09827		
5M09847	AC18807-024			BNA-8270	Soil	1	1	8270	08/08 14:06	5M09385		5M09827		
5M09848	AC18807-025			BNA-8270	Soil	1	1	8270	08/08 14:28	5M09385		5M09827		
5M09849	AC18807-017			BNA-8270	Soil	1	1	8270	08/08 14:50	5M09385		5M09827		
5M09850	AC18807-018			BNA-8270	Soil	1	1	8270	08/08 15:11	5M09385		5M09827		
5M09851	AC18807-020			BNA-8270	Soil	1	1	8270	08/08 15:33	5M09385		5M09827		
5M09852	AC18778-018			BNA-8270	Soil	1	1	8270	08/08 15:55	5M09385		5M09827		
5M09853	AC18884-004	Sa6Sa8		ERROR	Aqueou	1	1	625 8270	08/08 16:16	5M09385	5M09827	5M09827		
5M09854	SMB2614				Soil	1	1	8270	08/08 16:38	5M09385		5M09827		
5M09855	AC18873-017				Soil	1	1	8270	08/08 17:00	5M09385		5M09827		
5M09856	AC18830-001				Soil	1	1	8270	08/08 17:21	5M09385		5M09827		
5M09857	AC18845-002				Soil	1	1	8270	08/08 17:43	5M09385		5M09827		
5M09858	AC18939-001				Soil	1	1	8270	08/08 18:04	5M09385		5M09827		
5M09859	AC18845-004	Ti8			Soil	1	1	8270	08/08 18:26	5M09385		5M09827		
5M09860	AC18882-001				Aqueou	1	1	625	08/08 18:47	5M09385	5M09827	5M09827		
5M09861	AC18882-002				Aqueou	1	1	625	08/08 19:09	5M09385	5M09827	5M09827		
5M09862	AC18884-001				Aqueou	1	1	625	08/08 19:30	5M09385	5M09827	5M09827		
5M09863	AC18884-002				Aqueou	1	1	625	08/08 19:51	5M09385	5M09827	5M09827		
5M09864	AC18884-003				Aqueou	1	1	625	08/08 20:13	5M09385	5M09827	5M09827		
5M09865	AC18866-001				Aqueou	1	1	625	08/08 20:34	5M09385	5M09827	5M09827		
5M09866	AC18866-002				Aqueou	1	1	625	08/08 20:55	5M09385	5M09827	5M09827		
5M09867	CH2CL2(#1)	Ti8			Aqueou	1	1	625 8270	08/08 21:17	5M09385	5M09827	5M09827		
5M09868	CH2CL2(#2)	Ti8			Aqueou	1	1	625 8270	08/08 21:38	5M09385	5M09827	5M09827		

Anc	Area Not Checked	Ex	Extraction Performed Past Hold	Co	Warning Possible Carry Over
As	Area Out	Esm	Solvent Extraction Date Missing/Not check'd	R18,R26	Rpd Out on MsMsd (col1 and or col2) 8000 series
B6m	Blank 8000 series missing	Ein	Top/Solvent Extraction Date Missing/Not check'd	R18,R26	Rpd Out on MsMsd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eio	Top Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Off Out
Bnl	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C16	Calibration Column 1 Out (8000 Series)	Hb	Analysis Before Collection Date	S6	600 series surrogate out
C18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C26	Calibration Column 2 Out (8000 Series)	I18,I26	Initial cal 800 series failed Column 1 and or 2	Sa6,Sb6	Acid and or BN Surrogate Out (800 series)
C28	Calibration Column 2 Out (8000 Series)	I18,I26	Initial cal 8000 series failed Column 1 and or 2	Sa6,Sb6	Acid and or BN Surrogate Out (8000 series)
C6f	800 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Diluted Out
C6f	8000 series sample/blank did not have passing cal	Iv	Prob with calrpt csv for init calibration chck rts	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial cal warning, ini cal file <- method..	Ti5	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a samp	Ti6	Outside of 800 series Tune time/Cal Time
D1o,D2o	Drift Out Column 1 or Column 2 Cals or Int Cals	M16,M26	Spike Out Col 1 and or Col 2 8000 series	Ti8	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M16a,M18b	Spike Out Col 1 800 series Acid and or BN	Tm	Too Many Samples/ for beginning Calibration
Do	Drift Out	M18,M26	Spike Out Col 1 and or Col 2 8000 series	Trmw	If for 600 ser Too many samples begin Calibration
Eba	An Extraction Before Collection Date	M18a,M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune Not Checked
Emp	Problem Checking Prep/updates modcheck/prep/nd	Mnc	Spike Not Checked for this ms/msd	To	Tune File Failed
En	Eval Time Not Checked	IOc	Warning Compund(s) Over Calibration	Wte	Warning... Instrument Id not in TxLoc field

RUN LOG

Instrument: GCMS_4 Year: 2005
Analyst: AHD

1945

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr	Sam	Method(s)		Analysis Date	Cal	Beg	End	BlkFile
						Dil	Dil							
4M05425	CAL DFTPP									08/08 06:40				
4M05426	CAL BNA@50PPM	C16			Soil	1	1	625	8270	08/08 06:59	4M05299			
4M05427	SMB2613(MS)	M18b	SMB2613		Soil	1	1		8270	08/08 07:22	4M05299		4M05426	
4M05428	SMB2613				Soil	1	1		8270	08/08 07:49	4M05299		4M05426	
4M05429	AC18920-001			BNA25-8270	Soil	1	1		8270	08/08 08:13	4M05299		4M05426	
4M05430	AC18920-002			BNPAH-8270	Soil	1	1		8270	08/08 08:36	4M05299		4M05426	
4M05431	AC18920-003	Ao		BNPAH-8270	Soil	1	1		8270	08/08 09:00	4M05299		4M05426	
4M05432	AC18778-024			BNA-8270	Soil	1	1		8270	08/08 09:24	4M05299		4M05426	
4M05433	AC18807-019			BNA-8270	Soil	1	1		8270	08/08 09:48	4M05299		4M05426	
4M05434	AC18820-001			BN-8270	Soil	1	1		8270	08/08 10:12	4M05299		4M05426	
4M05435	AC18807-023(5X)			BNA-8270	Soil	5	5		8270	08/08 10:36	4M05299		4M05426	
4M05436	AC18820-003(5X)			BN-8270	Soil	5	5		8270	08/08 11:00	4M05299		4M05426	
4M05437	AC18820-004(5X)			BN-8270	Soil	5	5		8270	08/08 11:24	4M05299		4M05426	
4M05438	AC18820-002(3X)			BN-8270	Soil	3	3		8270	08/08 11:47	4M05299		4M05426	
4M05439	AC18806-001(20X)	Sd		BNPAH-8270	Soil	60	20		8270	08/08 12:11	4M05299		4M05426	
4M05440	AC18778-021	Ao		BNA-8270	Soil	1	1		8270	08/08 12:35	4M05299		4M05426	
4M05441	AC18807-015			BNA-8270	Soil	1	1		8270	08/08 12:59	4M05299		4M05426	
4M05442	AC18807-016			BNA-8270	Soil	1	1		8270	08/08 13:23	4M05299		4M05426	
4M05443	AC18778-004(5X)			BNA-8270	Soil	5	5		8270	08/08 13:47	4M05299		4M05426	
4M05444	AC18778-016(5X)			BNA-8270	Soil	5	5		8270	08/08 14:11	4M05299		4M05426	
4M05445	AC18778-019(5X)			BNA-8270	Soil	5	5		8270	08/08 14:35	4M05299		4M05426	
4M05446	AC18778-013			BNA-8270	Soil	1	1		8270	08/08 14:59	4M05299		4M05426	
4M05447	AC18778-001			BNA-8270	Soil	1	1		8270	08/08 15:23	4M05299		4M05426	
4M05448	AC18778-007			BNA-8270	Soil	1	1		8270	08/08 15:47	4M05299		4M05426	
4M05449	AC18807-022			BNA-8270	Soil	1	1		8270	08/08 16:11	4M05299		4M05426	
4M05450	AC18820-003			BN-8270	Soil	1	1		8270	08/08 16:35	4M05299		4M05426	
4M05451	AC18820-004	Ao		BN-8270	Soil	1	1		8270	08/08 16:59	4M05299		4M05426	
4M05452	AC18820-002			BN-8270	Soil	1	1		8270	08/08 17:22	4M05299		4M05426	
4M05453	AC18806-001(10X)	Sd		BNPAH-8270	Soil	30	10		8270	08/08 17:46	4M05299		4M05426	
4M05454	AC18873-016	Ao		BNA-8270	Soil	1	1		8270	08/08 18:10	4M05299		4M05426	
4M05455	TEST	Ao			Soil	1	1		8270	08/08 18:34	4M05299		4M05426	
4M05456	TEST	Ti8Ao			Soil	1	1		8270	08/08 18:58	4M05299		4M05426	
4M05457	TEST	Ti8Ao			Soil	1	1		8270	08/08 19:22	4M05299		4M05426	
4M05458	TEST	Ti8Ao			Soil	1	1		8270	08/08 19:45	4M05299		4M05426	
4M05459	TEST	Ti8Ao			Soil	1	1		8270	08/08 20:09	4M05299		4M05426	
4M05460	TEST	Ti8Ao			Soil	1	1		8270	08/08 20:33	4M05299		4M05426	

Anc	Area Not Checked	Er	Extraction Performed Past Hold	Co	Warning Possible Carry Over
Ao	Area Out	Esm	Solvent Extraction Date Missing/Not check'd	R16,R26	Rpd Out on Ms/Msd (col1 and or col2) 600 series
B6m	Blank 600 series missing	Etn	Top/Solvent Extraction Date Missing/Not check'd	R18,R28	Rpd Out on Ms/Msd (col1 and or col2) 8000 series
B8m	Blank 8000 series missing	Eto	Top Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Extended	Rin	Can't Calculate Drift
C16	Calibration Column 1 Out (600 Series)	Hb	Analysis Before Collection Date	S6	600 series surrogate out
C18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
C26	Calibration Column 2 Out (600 Series)	I16,I26	Initial cal 600 series failed Column 1 and or 2	Sa8,Sb8	Acid and or BN Surrogate Out (600 series)
C28	Calibration Column 2 Out (8000 Series)	I18,I28	Initial cal 8000 series failed Column 1 and or 2	Sa8,Sb8	Acid and or BN Surrogate Out (8000 series)
C6f	600 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Diluted Out
C8f	8000 series sample/blank did not have passing cal	Iv	Prob with calrpt.csv for init calibration check rfs	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial cal warning..ini cal file <- method..	T15	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a sample	T6	Outside of 600 series Tune time/Cal Time
D1o,D2o	Drift Out Column 1 or Column 2 Cals or Int Cals	M16,M26	Spike Out Col 1 and or Col 2 600 series	T8	Outside of 8000 series Tune time/Cal Time
Dnc	Drift Not Checked	M16a,M18b	Spike Out Col 1 600 series Acid and or BN	Trn	Too Many Samples/ for beginning Calibration
Do	Drift Out	M18,M28	Spike Out Col 1 and or Col 2 8000 series	Trmw	If for 600 ser Too many samples begin Calibration
Eba	An Extraction Before Collection Date	M18a,M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune Not Checked
Emp	Problem Checking Prep/rundates modcheck/prepund	Mnc	Spike Not Checked for this ms/msd	To	Tune File Failed
En	Eval Time Not Checked	Loc	Warning Compound(s) Over Calibration	Wie	Warning Instrument Id not in TxLoc field

Veritech Internally Prepared Standard Log

1347

Veritech Lot Number: V-204

Prepared By: Akmal		Department: Organics		
Description: BNA Surrog St		BatchNumber:		
Prep Date: 9/10/2004		Concentration: 1000-2000 ppm		
Expiration Date: 9/10/2005		Final Volume: 1000 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
605	2,4,6-Tribromophenol	2 g	Neat g	2000 ppm
582	Nitrobenzene-d5	800 ul	Neat	1000 ppm
583	Phenol-d6	2 g	neat	2000 ppm
586	p-Terphenyl-d14	1 g	neat	1000 ppm
606	2-Fluorobiphenyl	1 g	Neat g	1000 ppm
772	Acetone	1000 ml	Neat ml	neat
584	2-Fluorophenol	1.6 ml	neat	2000 ppm

Veritech Lot Number: V-295

Prepared By: Akmal		Department: Organics		
Description: 1,4-Dimethylnaphthalene		BatchNumber:		
Prep Date: 11/18/2004		Concentration: 10,000ppm		
Expiration Date: 11/17/2005		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
866	1,4-Dimethylnaphthalene	10 ul	neat	10,000 ppm
854	Methylene Chloride	990 ul	Neat	

Veritech Lot Number: V-498

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA Spike For Soil		BatchNumber:		
Prep Date: 2/24/2005		Concentration: 1000-2000PPM		
Expiration Date: 2/23/2006		Final Volume: 100 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
768	2,4-Dinitrotoluene	.1 g	Neat g	1000 ppm
764	4-Nitrophenol	.2 g	Neat g	2000 ppm
761	2-Chlorophenol	.2 g	Neat g	2000 ppm
767	1,4-Dichlorobenzene	.1 g	Neat g	1000 ppm
946	Phenol	.2 g	neat g	2000 ppm
769	N-Nitrosodi-n-propylamine	.1 g	Neat g	1000 ppm
762	Pentachlorophenol	.2 g	Neat g	2000 ppm
770	Pyrene	.1 g	Neat g	1000 ppm
771	1,2,4-Trichlorobenzene	.1 g	Neat g	1000 ppm
948	Acenaphthene	.1 g	neat g	1000 ppm
947	4-Chloro-3-methylphenol	.2 g	neat g	2000 ppm
950	Acetone	100 ml	Neat ml	

Veritech Lot Number: V-4045

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA Internal Std.		BatchNumber:		
Prep Date: 6/13/2005		Concentration: 2000 ppm		
Expiration Date: 6/12/2006		Final Volume: 250 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
571	Acenaphthene-D10	.5 g	Neat g	2000
570	Chrysene-d12	.5 g	Neat g	2000
567	Perylene-d12	.5 g	Neat g	2000
565	Phenanthrene-d10	.5 g	Neat g	2000
564	Naphthlene-d8	.5 g	Neat g	2000
1218	Methylene Chloride	250 ml	Neat l	
563	1,4 Dichlorobenzene-d4	.5 g	Neat g	2000

Veritech Internally Prepared Standard Log

1548

Veritech Lot Number: V-4046

Prepared By: Hamid, Akmal		Department: Organics		
Description: Pyridine Stock Std.		BatchNumber:		
Prep Date: 6/13/2005		Concentration: 10,000 ppm		
Expiration Date: 6/12/2006		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
1218	Methylene Chloride	990 ul	Neat l	
1225	Pyridine	10 ul	Neat ml	10000 ppm

Veritech Lot Number: V-4604

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA STOCK Std.		BatchNumber:		
Prep Date: 7/1/2005		Concentration: 200 ppm		
Expiration Date: 9/10/2005		Final Volume: 1.5 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
1091	EPA TCLP Pesticides Mix	60 ul	1000 ppm	40 ppm
1090	1,2,4,5-Tetrachlorobenzene	300 ul	1000 ppm	200 ppm
1089	Diphenyl Ether	150 ul	2000 ppm	200 ppm
1218	Methylene Chloride	60 ul	Neat	
1087	TCL Base-Neutrals Mix	150 ul	2000 ppm	200 ppm
1086	TCL Polynuclear Aromatic Hydrocarbons mix	150 ul	2000 ppm	200 ppm
1085	TCL Phenols/benzidine Mix	150 ul	2000 ppm	200 ppm
V-4046	Pyridine Stock Std.	30 ul	10,000 ppm	200 ppm
V-295	1,4-Dimethylnaphthalene	30 ul	10,000ppm	200 ppm
V-204	BNA Surrog St	150 ul	1000-2000 pp	100-200 ppm
1235	Pentachloroethane	60 ul	5000 ppm	200 ppm
1234	2,3,4,6-Tetrachlorophenol	60 ul	5000 ppm	200 ppm
1088	TCL Hazardous substances Mix	150 ul	2000 ppm	200 ppm

Veritech Lot Number: V-5045

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 10 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 10 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	5 ul	200 ppm	10 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	95	Neat	

Veritech Lot Number: V-5046

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 25 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 25 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	12.5 ul	200 ppm	25 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	87.5	Neat	

Veritech Internally Prepared Standard Log

1849

Veritech Lot Number: V-5047

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 50 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 50 ppm		
Expiration Date: 9/10/2005		Final Volume: 200 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	50 ul	200 ppm	50 ppm
V-4045	BNA Internal Std.	4 ul	2000 ppm	40 ppm
1218	Methylene Chloride	150	Neat	

Veritech Lot Number: V-5048

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 80 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 80 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	40 ul	200 ppm	80 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	60	Neat	

Veritech Lot Number: V-5049

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 120 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 120 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	60 ul	200 ppm	120 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	40	Neat	

Veritech Lot Number: V-5050

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 160 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 160 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	80 ul	200 ppm	160 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	20	Neat	

Veritech Lot Number: V-5051

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: BNA 200 ppm curve		BatchNumber: B-532		
Prep Date: 7/21/2005		Concentration: 200 ppm		
Expiration Date: 9/10/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4604	BNA STOCK Std.	100 ul	200 ppm	200 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	0	Neat	

Veritech Internally Prepared Standard Log

Veritech Lot Number: V-5264

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA Surrog.Std.		BatchNumber:		
Prep Date: 8/1/2005		Concentration: 1000-2000 ppm		
Expiration Date: 7/31/2006		Final Volume: 1000 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
585	2-fluorobiphenyl	1 g	neat	1000
788	p-Terphenyl-D14	1 g	Neat	1000
789	Phenol-d6	2 g	Neat	2000
790	2-Fluorophenol	1.6 ml	Neat	2000
582	Nitrobenzene-d5	800 ul	Neat	1000
605	2,4,6-Tribromophenol	2 g	Neat	2000
853	Acetone	1000 ml	Neat	neat

Veritech Lot Number: V-5267

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA STOCK Std.		BatchNumber:		
Prep Date: 8/2/2005		Concentration: 200 ppm		
Expiration Date: 11/17/2005		Final Volume: 1.5 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
1091	EPA TCLP Pesticides Mix	60 ul	1000 ppm	40 ppm
1090	1,2,4,5-Tetrachlorobenzene	300 ul	1000 ppm	200 ppm
1089	Diphenyl Ether	150 ul	2000 ppm	200 ppm
1218	Methylene Chloride	60 ul	Neat	
1087	TCL Base-Neutrals Mix	150 ul	2000 ppm	200 ppm
1086	TCL Polynuclear Aromatic Hydrocarbons mix	150 ul	2000 ppm	200 ppm
1085	TCLPhenols/benzidine Mix	150 ul	2000 ppm	200 ppm
V-4046	Pyridine Stock Std.	30 ul	10,000 ppm	200 ppm
V-295	1,4-Dimethylnaphthalene	30 ul	10,000ppm	200 ppm
1235	Pentachloroethane	60 ul	5000 ppm	200 ppm
1234	2,3,4,6-Tetrachlorophenol	60 ul	5000 ppm	200 ppm
V-5264	BNA Surrog.Std.	150	1000-2000 pp	200 ppm
1088	TCL Hazardous substances Mix	150 ul	2000 ppm	200 ppm

Veritech Lot Number: V-5269

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 10 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 10 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	5 ul	200 ppm	10 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	95	Neat	

Veritech Lot Number: V-5270

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 25 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 25 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	12.5 ul	200 ppm	25 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	87.5	Neat	

Veritech Internally Prepared Standard Log

1851

Veritech Lot Number: V-5271

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 50 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 50 ppm		
Expiration Date: 11/17/2005		Final Volume: 200 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	50 ul	200 ppm	50 ppm
V-4045	BNA Internal Std.	4 ul	2000 ppm	40 ppm
1218	Methylene Chloride	150	Neat	

Veritech Lot Number: V-5272

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 80 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 80 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	40 ul	200 ppm	80 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	60	Neat	

Veritech Lot Number: V-5273

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 120 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 120 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	60 ul	200 ppm	120 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	40	Neat	

Veritech Lot Number: V-5274

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 160 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 160 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	80 ul	200 ppm	160 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	20	Neat	

Veritech Lot Number: V-5275

Prepared By: Hamid, Akmal		Department: Organics		
Description: BNA 200 ppm curve		BatchNumber: B-551		
Prep Date: 8/2/2005		Concentration: 200 ppm		
Expiration Date: 11/17/2005		Final Volume: 100 ul		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-5267	BNA STOCK Std.	100 ul	200 ppm	200 ppm
V-4045	BNA Internal Std.	2 ul	2000 ppm	40 ppm
1218	Methylene Chloride	0	Neat	

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 563

Description
1,4 Dichlorobenzene-d4

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC.	DLM-268	PR-11537	04/04/01	04/04/10	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 564

Description
Naphthlene-d8

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC.	DLM-365	P-9785	04/04/01	04/04/10	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 565

Description
Phenanthrene-d10

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC	DLM-371	PR-11746	02/07/02	02/07/10	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 567

Description
Perylene-d12

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC	DLM-366	PR-10838	02/07/02	02/07/10	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 570

Description
Chrysene-d12

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC	DLM-261	PR-13120	06/17/03	06/17/10	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 571

Description
Acenaphthene-D10

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
CIL INC	DLM108-1	PR-11604	06/17/03	06/17/10	Akmal	2	2g	Neat	

Veritech Control/Receipt Number: 582

Description
Nitrobenzene-d5

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
ALDRICH	15195-5	09405MO	10/01/02	10/01/10	Akmal	1	5g	Neat	

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 583

Description

Phenol-d6

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Isotech	176060	07752cb	11/19/03	11/19/10	Akmal	1	1g	neat	

Veritech Control/Receipt Number: 584

Description

2-Fluorophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Aldrich	F1280-4	04515bu	02/07/02	02/07/10	Akmal	1	10g	neat	

Veritech Control/Receipt Number: 585

Description

2-fluorobiphenyl

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Aldrich	102741	06511cb	11/19/03	11/19/10	Akmal	1	2.5g	neat	

Veritech Control/Receipt Number: 586

Description

p-Terphenyl-d14

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Isotech	364630	00551kb	11/19/03	11/19/10	Akmal	3	1.5g	neat	

Veritech Control/Receipt Number: 605

Description

2,4,6-Tribromophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Aldrich	13771-5	18324MR	11/23/03	03/10/10	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 606

Description

2-Fluorobiphenyl

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
Aldrich	10274-1	02520TK	11/23/03	03/10/10	Akmal	1	2.5g	Neat	

Veritech Control/Receipt Number: 761

Description

2-Chlorophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F24	274-13A	10/08/02	10/01/05	Akmal	1	5g	Neat	

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 762

Description
Pentachlorophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F64	293-1A	10/08/02	09/01/07	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 764

Description
4-Nitrophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F58	281-142A	10/08/02	05/01/06	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 767

Description
1,4-Dichlorobenzene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F27	282-14B	10/08/02	03/01/07	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 768

Description
2,4-Dinitrotoluene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F35	270-148A	10/08/02	10/01/06	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 769

Description
N-Nitrosodi-n-propylamine

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F63	290-2B	10/08/02	08/01/06	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 770

Description
Pyrene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F84	266-23B	10/08/02	06/01/06	Akmal	1	1g	Neat	

Veritech Control/Receipt Number: 771

Description
1,2,4-Trichlorobenzene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
CHEM SERVICE	F8	274-89B	10/08/02	01/01/07	Akmal	1	1g	Neat	

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 772

Description

Acetone

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Pharmaco	329000DIS	PL000071ACE	06/11/04	06/09/09	Akmal	1	4000	Neat	

Veritech Control/Receipt Number: 788

Description

p-Terphenyl-D14

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Isotech	364630	10278AE	09/15/04	06/22/10	Akmal	5	2.5g	Neat	

Veritech Control/Receipt Number: 789

Description

Phenol-d6

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Aldrich	176060	02422JC	09/15/04	06/22/10	Akmal	1	5g	Neat	

Veritech Control/Receipt Number: 790

Description

2-Fluorophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Aldrich	F-12804	09006DO	09/15/04	06/22/10	Akmal	1	10g	Neat	

Veritech Control/Receipt Number: 853

Description

Acetone

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Fisher	A40-4	038622	08/24/04	08/18/10	Akmal	1	4000	Neat	

Veritech Control/Receipt Number: 854

Description

Methylene Chloride

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Fisher	D142-4	043063	11/02/04	08/18/10	Akmal	1	4000	Neat	

Veritech Control/Receipt Number: 866

Description

1,4-Dimethlnaphthlene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Aldrich	D17,030-5	14523CZ	11/18/97	11/25/10	Akmal	1	1ml	neat	

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 946

Description

Phenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Chem Service	F65	328-88B	02/10/05	09/30/10	Akmal	1	5g	neat	

Veritech Control/Receipt Number: 947

Description

4-Chloro-3-methylphenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Chem Service	F22	326-123B	02/10/05	08/30/07	Akmal	1	5g	neat	

Veritech Control/Receipt Number: 948

Description

Acenaphthene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Chem Service	0-782	306-17B	02/10/05	06/30/09	Akmal	1	5g	neat	

Veritech Control/Receipt Number: 950

Description

Acetone

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Fisher Scientific	A40-4	043780	12/13/04	11/17/10	Akmal	1	4L	Neat	

Veritech Control/Receipt Number: 1085

Description

TCLPhenols/benzidine Mix

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	47992-U	LB27910	04/07/05	03/31/08	Hamid, Akmal	1	1ml	2000	ppm

Veritech Control/Receipt Number: 1086

Description

TCL Polynuclear Aromatic Hydrocarbons mix

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	48905-U	LB24244	04/07/05	12/31/07	Hamid, Akmal	1	1ml	2000	ppm

Veritech Control/Receipt Number: 1087

Description

TCL Base-Neutrals Mix

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	47991-U	LB15949	04/07/05	11/30/06	Hamid, Akmal	1	1ml	2000	ppm

Veritech Standard Receipt Log

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Veritech Control/Receipt Number: 1088

Description
TCL Hazardous substances Mix

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	47990-U	LB10279	04/07/05	02/28/06	Hamid, Akmal	1	1ml	2000	ppm

Veritech Control/Receipt Number: 1089

Description
Diphenyl Ether

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	4-8155	LB23430	04/07/05	09/30/07	Hamid, Akmal	1	1ml	2000	ppm

Veritech Control/Receipt Number: 1090

Description
1,2,4,5-Tetrachlorobenzene

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	4-0177	LB25240	04/07/05	11/30/07	Hamid, Akmal	1	1ml	1000	ppm

Veritech Control/Receipt Number: 1091

Description
EPA TCLP Pesticides Mix

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Supelco	4-8139	LB09919	04/07/05	02/28/06	Hamid, Akmal	1	1ml	1000	ppm

Veritech Control/Receipt Number: 1218

Description
Methylene Chloride

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Fisher	D151-4	050241	05/20/05	05/19/10	Hamid, Akmal	1	4L	Neat	

Veritech Control/Receipt Number: 1225

Description
Pyridine

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
Fisher	P368-500	043764	12/16/04	12/16/08	Hamid, Akmal	1	500ml	Neat	

Veritech Control/Receipt Number: 1234

Description
2,3,4,6-Tetrachlorophenol

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
supelco	4-8264	lb26120	06/29/05	12/30/07	Hamid, Akmal	1	1ml	5000	ppm

Veritech Standard Receipt Log

1858

Veritech Control/Receipt Number: 1235

Description
Pentachloroethane

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/ Cont	Conc:	Units:
supelco	40300-u	1b13126	06/29/05	07/30/06	Hamid, Akmal	1	1ml	5000	ppm

Veritech Standard Receipt Log

1859

Veritech Control/Receipt Number: 1243

Description
PHENOL MIX

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
ACCUSTANDAR	Z-014H-PAK	B5050097	07/06/05	06/05/08	Wickliffe, David	5	1 ML	2000	PPM

Veritech Control/Receipt Number: 1245

Description
B/N COMPOSITE MIX

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
AccuStandard	CLP-HC-PAK	B5050070	07/06/05	05/11/06	Wickliffe, David	5	1mL	2000	PPM

Veritech Control/Receipt Number: 1246

Description
TOXIC SUBSTANCES MIX 1

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume/Cont	Conc:	Units:
AccuStandard	Z-014D-PAK	B5020076	07/06/05	07/05/08	Wickliffe, David	5	1mL	2000	PPM

Method Blank No. WMB- 2630
 Blank Spike (MBS): 2628; 2630

Date: 8/2/05
 Matrix Spike: 14417-003; 18819-004

Sample Number	Number in Batch	Initial Volume	Final Volume	Fraction			Comments	TCLP QC	Extract Fluid
				BN	BNA	AE			
MB 2630	X	1000 ml	1 ml		X		18817-003	V5084	
MBS 2630	X	↓	↓		X				
18766-001	15	250 ml	↓		↓		2	4	
18766-002	16	↓	↓		↓		3	5	
14808-001	17	↓	↓		↓		4	V5263(1)	
18840-012	18	950 ml	↓	X					
18840-013	19	↓	↓	X					
EFL V5263	X	250 ml	↓		X				
18819-002	20	250 ml	1 ml				5	2	
MS 18819-004	X	↓	↓						
MSD 18819-004	X	↓	↓				6	3	
18819-004	1	↓	↓				7	4	
18819-006	2	↓	↓				8	5	
18819-008	3	↓	↓				9	6	
18819-010	4	↓	↓				10	7	
18819-012	5	↓	↓				11	8	
18819-014	6	↓	↓				12	9	
18819-016	7	↓	↓				13	10	
18819-018	8	↓	↓						

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	2000	1245	BN Spike
↓	↓	1243	Ae Spike
↓	↓	1246	Ae box
		V320	Pyridine

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	V204	BN+ Succ

Reagent Lots: MeCl₂ 051907 Acetone _____ Hexane _____ Na₂SO₄ 052002

Other _____

Relinquished By: alv sy / AB
 Received By: _____

Date: 8/2/05
 Date: 02/02/05

Method Blank No. WMB-2631
 Blank Spike (MBS): 2630, 2631

Date: 8/3/05
 Matrix Spike: 18817-004, 18832-004

Sample Number	Number in Batch	Initial Volume	Final Volume	Fraction			Comments	TCLP QC	Extract Fluid
				BN	BNA	AE			
MBS <u>2631</u>	X	1000ml	1ml		X				
MBS 2631		↓			↓				
<u>18852-001</u>	9	250ml			X			14	11
<u>18801-007</u>	10	1000ml			X		Hsl		
<u>18820-012</u>	11	730ml		X					
<u>18823-001</u>	12	1000ml							
<u>18823-003</u>	13	↓							
<u>18841-001</u>	14	1000ml							
<u>18841-002</u>	15	900ml							
<u>18825-005</u>	16	1000ml							
<u>18825-007</u>	17	900ml							
<u>18832-001</u>	18	900ml							
<u>18832-002</u>	19	↓							
<u>18832-003</u>	20	820ml							
<u>18832-004</u>	X	50ml			X				
MBS <u>18832-004</u>	X	↓	↓		↓				
<u>18832-004</u>	1	↓	↓		↓				
<u>18897-001</u>	2	820ml	1ml						
<u>18897-002</u>	3	900ml	↓						

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
50	2000	V.1245	Bd spike
↓	↓	V.1243	AE spike
↓	↓	V.1046	AE Fox
↓	↓	V.320	Pyridine

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000ppb	V.204	BdA-Surr.

Reagent Lots: MeCl₂ 21407 Acetone _____ Hexane _____ Na₂SO₄ 152002

Other _____
 Relinquished By: PM / HSL
 Received By: _____

Date: 8/3/05
 Date: 08/04/05

Method Blank No. SMB-2610
 Blank Spike (SMBS): 2609, 2610
 Blank Spike (SMBS): _____

Date: 8/4/05
 Matrix Spike: 1883-001, 18807-011-18807-012
 Matrix Spike: _____

Analysis: BN/BNA/AE

Sample Number	# in Batch	Initial Volume	Final Volume	Fraction			Comments
				BN	BNA	AE	
MB-2610	X	30g	1ml		X		
MBS 26101	X						
18778-008	8						
18778-009	9						
18778-010	10						
18778-011	11						
18778-012	12						
18778-013	13						
18778-014	14						
18778-015	15						
18778-016	16						
18778-017	17						
18778-018	18						
18778-019	19						
18778-020	20						
MS 18807-011	X						
MSA 18807-012	X						
18807-009	1						
18807-010	2						
18807-001	3						
18807-002	4						
18807-003	5						
18807-004	6						
18807-005	7	↓	↓		↓		

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	1498	BdA spike

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	Y204	BdA Surr

Reagent Lots: MeCL₂ 051907 Acetone 043285 Hexane _____ Na₂SO₄ 052002 Ether _____
 MTBE _____ Other _____

Relinquished By: MSL
 Received By: _____

Date: 8/4/05
 Date: 6/6/05

Method Blank No. SMB- 2611
 Blank Spike (SMBS): 2610
 Blank Spike (SMBS): _____

Date: 08/04/05
 Matrix Spike: 18807-011, 18807-012
 Matrix Spike: _____

Analysis: BN/ BNA / AE

Sample Number	# in Batch	Initial Volume	Final Volume	Fraction			Comments
				BN	BNA	AE	
MB 2611	X	305	1ml		X		
MB 2611	X	↓	↓		↓		
18807-006	8						
18807-008	9						
18807-013	10						
18807-014	11						
18847-004	R					X	

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	V498	BNA SPIKE

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	V204	BNA SURR

Reagent Lots: MeCL2 051907 Acetone 050776 Hexane _____ Na2SO4 052002 Ether _____
 MTBE _____ Other _____ AB

Relinquished By: AB
 Received By: R

Date: 08/04/05
 Date: 08/07/05

Method Blank No. SMB- 2613
 Blank Spike (SMBS): 2610; 2613
 Blank Spike (SMBS): _____

Date: 08/07/05
 Matrix Spike: 18807-011; 18807-012; 18807-021
 Matrix Spike: _____

Analysis: BN BNA / AE

Sample Number	# in Batch	Initial Volume	Final Volume	Fraction			Comments
				BN	BNA	AE	
MB 2613							
MBS 2613	x	30g	1ml		x		
18778-021	16						
18778-022	17						
18778-023	18						
18778-024	19						
18807-015	20						
18807-021 MS	x						
18807-021 MSD	x						
18807-021	1						
18807-016	2						
18807-017	3						
18807-018	4						
18807-019	5						
18807-020	6						
18807-022	7						
18807-023	8						
18807-024	9						
18807-025	10						
18806-001	11	30g	3ml				
18820-001	12		3ml				
18820-002	13		1.0ml				
18820-003	14						
18820-004	15						

MSL NAS / Y Samples
 MS / Samples

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	V-498	BNA SPIKE

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	1000/2000	V-204	BNA SURROGATE

Reagent Lots: MeCL2 051907 Acetone 043785 Hexane _____ Na2SO4 052002 Ether _____
 MTBE _____ Other _____

Relinquished By: AB / MSL
 Received By: _____

Date: 08/07/05
 Date: 08/08/05

GC PCB Data

**GC PCB Data
QC Summary**

1585

FORM2

Surrogate Recovery

Dfile	Sample#	Matrix	Surr Dil	Dilute Out Flag	Column1	Column2	Column1	Column2	Column0	Column0
					S1 Recov	S2 Recov	S3 Recov	S4 Recov	S5 Recov	S6 Recov
2G10427.	WMB2305	Aqueous	1		103	95	114	113		
2G10513.	SMB727B	Soil	1		80	76	96	93		
2G10591.	SMB729B	Soil	1		98	84	111	90		
2G10598.	AC18807-001	Soil	1		99	88	90	70		
2G10599.	AC18807-004	Soil	1		100	89	104	80		
2G10431.	AC18807-007	Aqueous	1		117	108	156 *	103		
2G10611.	AC18807-008	Soil	1		113	101	103	83		
2G10608.	AC18807-014	Soil	1		103	93	98	79		
2G10609.	AC18807-017	Soil	1		110	99	94	78		
2G10610.	AC18807-020	Soil	1		93	84	81	63		
2G10607.	AC18807-023	Soil	1		98	89	103	75		
2G10433.	WMB2305(MS)	Aqueous	1		98	93	103	97		
2G10514.	SMB727B(MS)	Soil	1		75	72	100	95		
2G10525.	AC18778-020(MS)	Soil	1		102	95	114	94		
2G10526.	AC18778-020(MSD)	Soil	1		54 *	52 *	56	51		
2G10592.	SMB729B(MS)	Soil	1		98	85	113	90		

Flags: SD=Surrogate diluted out
 *=Surrogate out

Method: 8082

Soil Limits

Compound	Spike Amt	Limits
S1=TCMX-Surrogate	100	60-150
S2=TCMX-Surrogate	100	60-150
S3=DCB-Surrogate	100	20-150
S4=DCB-Surrogate	100	20-150

Aqueous Limits

Compound	Spike Amt	Limits
S1=TCMX-Surrogate	100	60-150
S2=TCMX-Surrogate	100	60-150
S3=DCB-Surrogate	100	20-150
S4=DCB-Surrogate	100	20-150

Form3
MBS Data
Method: 8082

15298

Data File:====>
Data/Batch/Sample ID:====>
Date/Time:====>

2G10592.D	2G10433.D			
SMB729B(MS)	WMB2305(MS)			
08/08/05 11:20	08/03/05 13:14			

Compound	Limit(s)				Conc			Conc			Conc			Conc		
	Soil	Aq	Col	Mr	Conc	Exp	% Rec	Conc	Exp	% Rec	Conc	Exp	% Rec	Conc	Exp	% Rec
Aroclor-1016	29-131	29-131	1	0	1039	1000	104	982.1	1000	98						
Aroclor-1260	29-131	29-131	1	0	1096	1000	110	957.8	1000	96						

FORM 3
Spike Recovery

6551

Batch Number: SMB727B	Mbs File: 2G10514.D
Mbs Name: SMB727B(MS)	Non Spk'd File: 2G10527.D
Ns Name: AC18778-020	Spike File: 2G10525.D
Ms Name: AC18778-020(MS)	Spike Dup File: 2G10526.D
Msd Name: AC18778-020(MSD)	Matrix: Soil
	Method: 8082

Compound	Col	Mr	Conc	Lo	Hi	Rpd	Mbs	Sample	Spike	Spike	Mbs	MS	Msd	Rpd
			Exp	Llm	Lim	Llm	Conc	Conc	Conc	Dup	Rec	Rec	Rec	
Aroclor-1016	1	0	1000	29	131	40	915.31	0.00	1045.59	559.62	92	105	56	61Rp
Aroclor-1260	1	0	1000	29	131	40	987.76	0.00	1106.18	594.80	99	111	59	60Rp

Note:

Rp = Failed Rpd Criteria

Mo = Failed Recovery Criteria

^ - Both Ms and Msd Recoveries = 0 ... no valid information can be calculated

FORM 4
Blank Summary

Blank Number: WMB2305
Blank Data File: 2G10427.D
Matrix: Aqueous

Blank Analysis Date: 08/03/05 11:33
Blank Extraction Date: 08/02/05
(If Applicable)

Sample Number	Data File	Analysis Date
AC18807-007	2G10431.D	08/03/05 12:45
WMB2305(MS)	2G10433.D	08/03/05 13:14

FORM 4
Blank Summary

Blank Number: SMB727B
Blank Data File: 2G10513.D
Matrix: Soil

Blank Analysis Date: 08/05/05 06:11
Blank Extraction Date: 08/04/05
(If Applicable)

Sample Number	Data File	Analysis Date
AC18778-020(MSD)	2G10526.D	08/05/05 09:18
AC18778-020(MS)	2G10525.D	08/05/05 09:04
SMB727B(MS)	2G10514.D	08/05/05 06:25

FORM 4
Blank Summary

Blank Number: SMB729B
Blank Data File: 2G10591.D
Matrix: Soil

Blank Analysis Date: 08/08/05 11:06
Blank Extraction Date: 08/07/05
(If Applicable)

Sample Number	Data File	Analysis Date
AC18807-001	2G10598.D	08/08/05 12:47
AC18807-004	2G10599.D	08/08/05 13:10
AC18807-008	2G10611.D	08/08/05 16:03
AC18807-014	2G10608.D	08/08/05 15:20
AC18807-017	2G10609.D	08/08/05 15:34
AC18807-020	2G10610.D	08/08/05 15:49
AC18807-023	2G10607.D	08/08/05 15:06
SMB729B(MS)	2G10592.D	08/08/05 11:20

Form 5

Data File	Sample#	Analysis Date/Time	Matrix	Reference File	Column 1 RT	Column 1 % Drift	Column 2 RT	Column 2 % Drift
2G10408.	CAL 1660@500PPB	08/03/05 06:10	Aqueous	2G10408.	8.9934	0	9.3172	0
2G10409.	WMB2305	08/03/05 06:30	Aqueous	2G10408.	8.9991	0.0634	9.3213	0.044
2G10410.	WMB2305(MS)	08/03/05 06:44	Aqueous	2G10408.	8.9951	0.0189	9.3197	0.0268
2G10411.	AC18807-007	08/03/05 06:59	Aqueous	2G10408.	8.9949	0.0167	9.3207	0.0376
2G10412.	AC18820-012	08/03/05 07:13	Aqueous	2G10408.	8.9966	0.0356	9.3211	0.0419
2G10413.	AC18863-001	08/03/05 07:28	Aqueous	2G10408.	8.9968	0.0378	9.3217	0.0483
2G10414.	AC18819-008	08/03/05 07:42	Aqueous	2G10408.	8.9960	0.0289	9.3219	0.0504
2G10415.	AC18819-018	08/03/05 07:57	Aqueous	2G10408.	8.9952	0.02	9.3204	0.0343
2G10416.	CAL 1660@500PPB	08/03/05 08:26	Soil	2G10408.	8.9946	0.0511	9.3194	0.0236
2G10417.	CAL 1660@50PPB	08/03/05 08:47	Soil	2G10417.	8.9992	0	9.3203	0
2G10418.	CAL 1660@200PPB	08/03/05 09:04	Soil	2G10417.	8.9967	0.0278	9.3194	0.0097
2G10419.	CAL 1660@1000PPB	08/03/05 09:19	Soil	2G10417.	8.9932	0.0667	9.3178	0.0268
2G10420.	CAL 1660@2000PPB	08/03/05 09:33	Soil	2G10417.	8.9927	0.0723	9.3170	0.0354
2G10421.	CAL 1660@4000PPB	08/03/05 09:47	Soil	2G10417.	8.9924	0.0756	9.3162	0.044
2G10422.	CAL 2154@500PPB	08/03/05 10:20	Soil	2G10417.	8.9961	0.0344	9.3190	0.0139
2G10423.	CAL 1248@500PPB	08/03/05 10:35	Soil	2G10417.	8.9946	0.0511	9.3183	0.0215
2G10424.	CAL 1242@500PPB	08/03/05 10:50	Soil	2G10417.	8.9920	0.08	9.3168	0.0376
2G10425.	CAL 1232@500PPB	08/03/05 11:04	Soil	2G10417.	8.9939	0.0589	9.3183	0.0215
2G10426.	AC18863-001	08/03/05 11:18	Aqueous	2G10417.	8.9914	0.0867	9.3156	0.0504
2G10427.	WMB2305	08/03/05 11:33	Aqueous	2G10417.	8.9925	0.0745	9.3174	0.0311
2g10428.	AC18819-008	08/03/05 12:02	Soil	2G10417.	8.9927	0.0723	9.3178	0.0268
2g10429.	AC18819-018	08/03/05 12:16	Soil	2G10417.	8.9928	0.0711	9.3176	0.029
2g10430.	AC18802-005	08/03/05 12:31	Soil	2G10417.	8.9923	0.0767	9.3180	0.0247
2G10431.	AC18807-007	08/03/05 12:45	Aqueous	2G10417.	8.9926	0.0734	9.3158	0.0483
2G10432.	SMB725B(MS)	08/03/05 12:59	Soil	2G10417.	8.9929	0.07	9.3185	0.0193
2G10433.	WMB2305(MS)	08/03/05 13:14	Aqueous	2G10417.	8.9932	0.0667	9.3179	0.0258
2G10434.	AC18694-001(20X)(R)	08/03/05 13:28	Soil	2G10417.	8.9936	0.0622	9.3198	0.0054
2G10435.	SMB725B	08/03/05 13:42	Soil	2G10417.	8.9939	0.0589	9.3170	0.0354
2G10436.	AC18820-012	08/03/05 13:57	Aqueous	2G10417.	8.9916	0.0845	9.3162	0.044
2G10437.	AC18786-012	08/03/05 14:11	Soil	2G10417.	8.9928	0.0711	9.3167	0.0386
2G10438.	AC18786-012(MS)	08/03/05 14:26	Soil	2G10417.	8.9935	0.0634	9.3189	0.015
2G10439.	AC18786-012(MSD)	08/03/05 14:40	Soil	2G10417.	8.9929	0.07	9.3179	0.0258
2G10440.	SMB671	08/03/05 14:55	Soil	2G10417.	8.9930	0.0689	9.3179	0.0258
2G10441.	SMB671(MS)	08/03/05 15:09	Soil	2G10417.	8.9931	0.0678	9.3173	0.0322
2G10442.	AC18772-001	08/03/05 15:23	Soil	2G10417.	8.9568	0.4723	9.3317	0.1222
2G10443.	AC18772-001(MS)(200X	08/03/05 15:58	Soil	2G10417.	8.9655	0.3752	9.3413	0.2251
2G10444.	AC18772-001(MSD)(200	08/03/05 16:12	Soil	2G10417.	8.9639	0.393	9.3388	0.1983
2G10445.	AC18786-001	08/03/05 16:27	Soil	2G10417.	8.9935	0.0634	9.3174	0.0311
2G10446.	CAL 1660@1000PPB	08/03/05 16:41	Soil	2G10417.	8.9930	0.0689	9.3178	0.0268
2G10447.	1000PPB	08/03/05 16:56	Soil	2G10446.	8.9938	0.0089	9.3193	0.0161
2G10448.	2000PPB	08/03/05 17:10	Soil	2G10446.	8.9927	0.0033	9.3191	0.014
2G10449.	AC18786-002	08/03/05 17:25	Soil	2G10446.	8.9934	0.0044	9.3192	0.015
2G10450.	AC18786-003	08/03/05 17:39	Soil	2G10446.	8.9944	0.0156	9.3194	0.0172
2G10451.	AC18786-004	08/03/05 17:54	Soil	2G10446.	8.9936	0.0067	9.3192	0.015
2G10452.	AC18786-005	08/03/05 18:08	Soil	2G10446.	8.9948	0.02	9.3192	0.015
2G10453.	AC18786-006	08/03/05 18:23	Soil	2G10446.	8.9932	0.0022	9.3186	0.0086
2G10454.	AC18786-007	08/03/05 18:37	Soil	2G10446.	8.9951	0.0233	9.3189	0.0118
2G10455.	AC18786-008	08/03/05 18:51	Soil	2G10446.	8.9925	0.0056	9.3185	0.0075
2G10456.	AC18786-009	08/03/05 19:06	Soil	2G10446.	8.9929	0.0011	9.3186	0.0086
2G10457.	AC18786-010	08/03/05 19:20	Soil	2G10446.	8.9937	0.0078	9.3186	0.0086
2G10458.	AC18786-011	08/03/05 19:35	Soil	2G10446.	8.9921	0.01	9.3171	0.0075
2G10458.	AC18772-001(200X)	08/03/05 19:49	Soil	2G10446.	0.0000	200 *	0.0000	200 *
2G10459.	AC18786-013	08/03/05 20:04	Soil	2G10446.	8.9926	0.0044	9.3170	0.0086
2G10460.	AC18786-014	08/03/05 20:18	Soil	2G10446.	8.9913	0.0189	9.3160	0.0193
2G10461.	AC18786-015	08/03/05 20:33	Soil	2G10446.	8.9895	0.0389	9.3157	0.0225
2G10462.	AC18786-016	08/03/05 20:47	Soil	2G10446.	8.9916	0.0156	9.3162	0.0172
2G10463.	AC18786-017	08/03/05 21:01	Soil	2G10446.	8.9915	0.0167	9.3172	0.0064
2G10464.	AC18855-001	08/03/05 21:16	Soil	2G10446.	8.9912	0.02	9.3165	0.014
2G10465.	500PPB	08/03/05 21:30	Soil	2G10446.	8.9919	0.0122	9.3173	0.0054
2G10466.	500PPB	08/03/05 21:45	Soil	2G10446.	8.9917	0.0145	9.3166	0.0129
2G10467.	1000PPB	08/03/05 21:59	Soil	2G10446.	8.9914	0.0178	9.3168	0.0107
2G10468.	1000PPB	08/03/05 22:13	Soil	2G10446.	8.9909	0.0234	9.3167	0.0118

Drift Compound: DCB-Surrogate

Drift Limit(s): 0.5 (Pest/Pcb) 1.5(Herb/Tph)

* - Values outside of limits for this column/run

Form 5

Data File	Sample#	Analysis Date/Time	Matrix	Reference File	Column 1 RT	Column 1 % Drift	Column 2 RT	Column 2 % Drift
2G10469.	CAL 1660@2000PPB	08/03/05 22:28	Soil	2G10446.	8.9898	0.0356	9.3149	0.0311
2G10470.	2000PPB	08/03/05 22:42	Soil	2G10469.	8.9902	0.0044	9.3151	0.0021
2G10471.	AC18772-001(4000X)	08/04/05 05:19	Soil	2G10469.	0.0000	200 *	0.0000	200 *
2G10472.	CAL 1660@1000PPB	08/04/05 05:53	Soil	2G10469.	8.9911	0.0145	9.3140	0.0097

Drift Compound: DCB-Surrogate

Drift Limit(s): 0.5 (Pest/Pcb) 1.5(Herb/Tph)

* - Values outside of limits for this column/run

Form 5

Data File	Sample#	Analysis Date/Time	Matrix	Reference File	Column 1 RT	Column 1 % Drift	Column 2 RT	Column 2 % Drift
2G10502.	CAL 1660@50PPB	08/05/05 02:34	Soil		8.9710	0	9.2828	0
2G10503.	CAL 1660@50PPB	08/05/05 02:48	Soil		8.9584	0	9.2783	0
2G10504.	CAL 1660@200PPB	08/05/05 03:02	Soil	2G10503.	8.9572	0.0134	9.2794	0.0119
2G10505.	CAL 1660@500PPB	08/05/05 03:17	Soil	2G10503.	8.9575	0.01	9.2804	0.0226
2G10506.	CAL 1660@1000PPB	08/05/05 03:31	Soil	2G10503.	8.9577	0.0078	9.2819	0.0388
2G10507.	CAL 1660@2000PPB	08/05/05 03:46	Soil	2G10503.	8.9587	0.0034	9.2823	0.0431
2G10508.	CAL 1660@4000PPB	08/05/05 04:00	Soil	2G10503.	8.9589	0.0056	9.2830	0.0506
2G10509.	CAL 2154@500PPB	08/05/05 04:15	Soil	2G10503.	8.9595	0.0123	9.2837	0.0582
2G10510.	CAL 1248@500PPB	08/05/05 04:29	Soil	2G10503.	8.9596	0.0134	9.2843	0.0646
2G10511.	CAL 1242@500PPB	08/05/05 04:43	Soil	2G10503.	8.9603	0.0212	9.2848	0.07
2G10512.	CAL 1232@500PPB	08/05/05 04:58	Soil	2G10503.	8.9597	0.0145	9.2845	0.0668
2G10513.	SMB727B	08/05/05 06:11	Soil	2G10503.	8.9709	0.1394	9.2851	0.0733
2G10514.	SMB727B(MS)	08/05/05 06:25	Soil	2G10503.	8.9576	0.0089	9.2797	0.0151
2G10515.	AC18737-033	08/05/05 06:40	Soil	2G10503.	8.9556	0.0313	9.2795	0.0129
2G10516.	AC18919-001	08/05/05 06:54	Soil	2G10503.	8.9556	0.0313	9.2805	0.0237
2G10517.	AC18919-002	08/05/05 07:09	Soil	2G10503.	8.9571	0.0145	9.2812	0.0313
2G10518.	AC18919-003	08/05/05 07:23	Soil	2G10503.	8.9578	0.0067	9.2821	0.0409
2G10519.	SMB2405	08/05/05 07:37	Soil	2G10503.	8.9586	0.0022	9.2833	0.0539
2G10520.	SMB2405(MS)	08/05/05 07:52	Soil	2G10503.	8.9586	0.0022	9.2839	0.0603
2G10521.	AC18876-002(MS)	08/05/05 08:06	Soil	2G10503.	8.9581	0.0033	9.2834	0.0549
2G10522.	AC18876-002(MSD)	08/05/05 08:21	Soil	2G10503.	8.9590	0.0067	9.2836	0.0571
2G10523.	AC18876-002	08/05/05 08:35	Soil	2G10503.	8.9592	0.0089	9.2844	0.0657
2G10524.	AC18876-001	08/05/05 08:50	Soil	2G10503.	8.9591	0.0078	9.2859	0.0819
2G10525.	AC18778-020(MS)	08/05/05 09:04	Soil	2G10503.	8.9599	0.0167	9.2852	0.0743
2G10526.	AC18778-020(MSD)	08/05/05 09:18	Soil	2G10503.	8.9609	0.0279	9.2865	0.0883
2G10527.	AC18778-020	08/05/05 09:33	Soil	2G10503.	8.9614	0.0335	9.2866	0.0894
2G10528.	AC18737-033(10X)	08/05/05 09:47	Soil	2G10503.	8.9625	0.0458	9.2870	0.0937
2G10529.	AC18778-010	08/05/05 10:02	Soil	2G10503.	8.9604	0.0223	9.2870	0.0937
2G10530.	AC18778-011	08/05/05 10:16	Soil	2G10503.	8.9620	0.0402	9.2872	0.0959
2G10531.	AC18778-012	08/05/05 10:30	Soil	2G10503.	8.9621	0.0413	9.2877	0.1013
2G10532.	AC18778-013	08/05/05 10:45	Soil	2G10503.	8.9618	0.0379	9.2876	0.1002
2G10533.	CAL1660@1000PPB	08/05/05 10:59	Soil	2G10503.	8.9622	0.0424	9.2878	0.1023
2G10534.	AC18778-014	08/05/05 11:17	Soil	2G10533.	8.9686	0.0714	9.2911	0.0355
2G10535.	AC18778-003(R)	08/05/05 11:32	Soil	2G10533.	8.9631	0.01	9.2887	0.0097
2G10536.	TEST0805	08/05/05 11:46	Soil	2G10533.	8.9613	0.01	9.2863	0.0162
2G10537.	AC18778-024	08/05/05 12:01	Soil	2G10533.	8.9626	0.0045	9.2885	0.0075
2G10538.	AC18778-016	08/05/05 12:15	Soil	2G10533.	8.9618	0.0045	9.2873	0.0054
2G10539.	18786-009	08/05/05 12:30	Soil	2G10533.	8.9626	0.0045	9.2879	0.0011
2G10540.	AC18778-018	08/05/05 13:17	Soil	2G10533.	8.9628	0.0067	9.2880	0.0022
2G10541.	AC18778-019	08/05/05 13:46	Soil	2G10533.	8.9613	0.01	9.2870	0.0086
2G10541.	AC18919-001	08/05/05 14:00	Soil	2G10533.	8.9619	0.0034	9.2878	0
2G10541.	AC18919-002	08/05/05 14:14	Soil	2G10533.	8.9620	0.0022	9.2886	0.0086
2G10541.	AC18919-003	08/05/05 14:29	Soil	2G10533.	8.9624	0.0022	9.2884	0.0065
2G10542.	AC18778-021	08/05/05 14:43	Soil	2G10533.	8.9643	0.0234	9.2894	0.0172
2G10543.	AC18778-023	08/05/05 14:58	Soil	2G10533.	8.9640	0.0201	9.2895	0.0183
2G10544.	AC18778-022	08/05/05 15:12	Soil	2G10533.	8.9635	0.0145	9.2895	0.0183
2G10545.	AC18778-015	08/05/05 15:29	Soil	2G10533.	8.9677	0.0613	9.2906	0.0301
2G10546.	AC18778-017	08/05/05 15:43	Soil	2G10533.	8.9636	0.0156	9.2889	0.0118
2G10547.	CAL 1660@2000PPB	08/05/05 15:58	Soil	2G10533.	8.9640	0.0201	9.2896	0.0194
2G10548.	2000PPB	08/05/05 16:12	Soil	2G10547.	8.9643	0.0034	9.2897	0.0011
2G10549.	2000PPB	08/05/05 16:26	Soil	2G10547.	8.9635	0.0056	9.2893	0.0032
2G10550.	AC18778-014(R)	08/05/05 16:41	Soil	2G10547.	8.9631	0.01	9.2891	0.0054
2G10551.	AC18778-024(R)	08/05/05 16:55	Soil	2G10547.	8.9637	0.0033	9.2889	0.0075
2G10552.	test0805	08/05/05 17:10	Soil	2G10547.	8.9648	0.0089	9.2899	0.0032
2G10553.	AC18778-003(R)	08/05/05 17:24	Soil	2G10547.	8.9643	0.0034	9.2896	0
2G10554.	AC18778-010	08/05/05 17:39	Soil	2G10547.	8.9640	0	9.2899	0.0032
2G10555.	AC18778-011	08/05/05 17:53	Soil	2G10547.	8.9629	0.0123	9.2889	0.0075
2G10556.	AC18778-012	08/05/05 18:07	Soil	2G10547.	8.9634	0.0067	9.2891	0.0054
2G10557.	AC18778-013	08/05/05 18:22	Soil	2G10547.	8.9623	0.019	9.2882	0.0151
2G10558.	AC18778-014	08/05/05 18:36	Soil	2G10547.	8.9631	0.01	9.2886	0.0108
2G10559.	AC18778-015	08/05/05 18:51	Soil	2G10547.	8.9626	0.0156	9.2881	0.0161
2G10560.	AC18778-016	08/05/05 19:05	Soil	2G10547.	8.9606	0.0379	9.2868	0.0301

Drift Compound: DCB-Surrogate

Drift Limit(s): 0.5 (Pest/Pcb) 1.5(Herb/Tph)

* - Values outside of limits for this column/run

Form 5

Data File	Sample#	Analysis Date/Time	Matrix	Reference File	Column 1 RT	Column 1 % Drift	Column 2 RT	Column 2 % Drift
2G10561.	AC18778-017	08/05/05 19:19	Soil	2G10547.	8.9614	0.029	9.2875	0.0226
2G10562.	AC18778-018	08/05/05 19:34	Soil	2G10547.	8.9629	0.0123	9.2884	0.0129
2G10563.	AC18778-019	08/05/05 19:48	Soil	2G10547.	8.9596	0.0491	9.2853	0.0463
2G10564.	AC18778-020	08/05/05 20:03	Soil	2G10547.	8.9603	0.0413	9.2859	0.0398
2G10565.	AC18778-021	08/05/05 20:17	Soil	2G10547.	8.9609	0.0346	9.2860	0.0388
2G10566.	AC18778-022	08/05/05 20:32	Soil	2G10547.	8.9598	0.0469	9.2854	0.0452
2G10567.	AC18778-023	08/05/05 20:46	Soil	2G10547.	8.9610	0.0335	9.2870	0.028
2G10568.	AC18778-024	08/05/05 21:01	Soil	2G10547.	8.9610	0.0335	9.2858	0.0409
2G10569.	CAL 1660@1000PPB	08/05/05 21:15	Soil	2G10547.	8.9610	0.0335	9.2861	0.0377
2G10570.	_1000PPB	08/05/05 21:29	Soil	2G10569.	8.9596	0.0156	9.2855	0.0065
2G10571.	2000PPB	08/05/05 21:44	Soil	2G10569.	8.9592	0.0201	9.2842	0.0205
2G10572.	2000PPB	08/05/05 21:58	Soil	2G10569.	8.9601	0.01	9.2861	0

Drift Compound: DCB-Surrogate

Drift Limit(s): 0.5 (Pest/Pcb) 1.5(Herb/Tph)

* - Values outside of limits for this column/run

10/25/05

Form 5

Data File	Sample#	Analysis Date/Time	Matrix	Reference File	Column 1 RT	Column 1 % Drift	Column 2 RT	Column 2 % Drift
2G10580.	CAL 1660@500PPB	08/08/05 08:12	Soil	2G10580.	8.9572	0	9.2805	0
2G10581.	AC18920-001	08/08/05 08:27	Soil	2G10580.	8.9522	0.0558	9.2782	0.0248
2G10582.	AC18907-005	08/08/05 08:41	Soil	2G10580.	8.9539	0.0368	9.2798	0.0075
2G10583.	WMB2310	08/08/05 08:56	Aqueous	2G10580.	8.9538	0.038	9.2791	0.0151
2G10584.	WMB2310(MS)	08/08/05 09:10	Aqueous	2G10580.	8.9550	0.0246	9.2806	0.0011
2G10585.	AC18873-014	08/08/05 09:25	Aqueous	2G10580.	8.9558	0.0156	9.2818	0.014
2G10586.	AC18886-009	08/08/05 09:39	Aqueous	2G10580.	8.9561	0.0123	9.2820	0.0162
2G10587.	AC18888-001	08/08/05 09:53	Aqueous	2G10580.	8.9564	0.0089	9.2826	0.0226
2G10588.	AC18916-025	08/08/05 10:08	Aqueous	2G10580.	8.9569	0.0034	9.2822	0.0183
2G10588.	test	08/08/05 10:22	Aqueous	2G10580.	8.9569	0.0034	9.2825	0.0215
2G10589.	SMB728B	08/08/05 10:37	Soil	2G10580.	8.9568	0.0045	9.2834	0.0312
2G10590.	SMB728B(MS)	08/08/05 10:51	Soil	2G10580.	8.9563	0.0101	9.2828	0.0248
2G10591.	SMB729B	08/08/05 11:06	Soil	2G10580.	8.9574	0.0022	9.2839	0.0366
2G10592.	SMB729B(MS)	08/08/05 11:20	Soil	2G10580.	8.9574	0.0022	9.2841	0.0388
2G10593.	AC18820-005	08/08/05 11:34	Soil	2G10580.	8.9567	0.0056	9.2835	0.0323
2G10594.	AC18820-005(MS)	08/08/05 11:49	Soil	2G10580.	8.9592	0.0223	9.2863	0.0625
2G10595.	AC18820-005(MSD)	08/08/05 12:03	Soil	2G10580.	8.9609	0.0413	9.2881	0.0819
2G10596.	AC18939-001	08/08/05 12:18	Soil	2G10580.	8.9615	0.048	9.2888	0.0894
2G10597.	AC18774-029	08/08/05 12:32	Soil	2G10580.	8.9630	0.0647	9.2891	0.0926
2G10598.	AC18807-001	08/08/05 12:47	Soil	2G10580.	8.9620	0.0536	9.2890	0.0915
2G10599.	AC18807-004	08/08/05 13:10	Soil	2G10580.	8.9669	0.1082	9.2897	0.0991
2G10600.	CAL 1660@1000PPB	08/08/05 13:25	Soil	2G10580.	8.9622	0.0558	9.2882	0.0829
2G10601.	SMB730B(MS)	08/08/05 13:39	Soil	2G10600.	8.9616	0.0067	9.2878	0.0043
2G10602.	SMB730B	08/08/05 13:54	Soil	2G10600.	8.9617	0.0056	9.2880	0.0022
2G10603.	AC18820-001	08/08/05 14:08	Soil	2G10600.	8.9620	0.0022	9.2890	0.0086
2G10604.	AC18820-002	08/08/05 14:22	Soil	2G10600.	8.9625	0.0033	9.2892	0.0108
2G10605.	AC18820-003	08/08/05 14:37	Soil	2G10600.	8.9644	0.0245	9.2913	0.0334
2G10606.	AC18820-004	08/08/05 14:51	Soil	2G10600.	8.9658	0.0402	9.2929	0.0506
2G10607.	AC18807-023	08/08/05 15:06	Soil	2G10600.	8.9653	0.0346	9.2923	0.0441
2G10608.	AC18807-014	08/08/05 15:20	Soil	2G10600.	8.9656	0.0379	9.2911	0.0312
2G10609.	AC18807-017	08/08/05 15:34	Soil	2G10600.	8.9647	0.0279	9.2911	0.0312
2G10610.	AC18807-020	08/08/05 15:49	Soil	2G10600.	8.9648	0.029	9.2914	0.0344
2G10611.	AC18807-008	08/08/05 16:03	Soil	2G10600.	8.9643	0.0234	9.2911	0.0312
2G10612.	AC18848-006	08/08/05 16:18	Soil	2G10600.	8.9633	0.0123	9.2898	0.0172
2G10613.	AC18848-007	08/08/05 16:32	Soil	2G10600.	8.9636	0.0156	9.2897	0.0161
2G10614.	AC18848-008	08/08/05 16:47	Soil	2G10600.	8.9623	0.0011	9.2892	0.0108
2G10615.	AC18845-002	08/08/05 17:01	Soil	2G10600.	8.9629	0.0078	9.2891	0.0097
2G10616.	AC18845-004	08/08/05 17:16	Soil	2G10600.	8.9628	0.0067	9.2890	0.0086
2G10617.	AC18845-007	08/08/05 17:30	Soil	2G10600.	8.9620	0.0022	9.2893	0.0118
2G10618.	AC18845-010	08/08/05 17:44	Soil	2G10600.	8.9614	0.0089	9.2875	0.0075
2G10619.	AC18845-012	08/08/05 17:59	Soil	2G10600.	8.9609	0.0145	9.2870	0.0129
2G10620.	500PPB	08/08/05 18:13	Soil	2G10600.	8.9611	0.0123	9.2872	0.0108
2G10621.	500PPB	08/08/05 18:28	Soil	2G10600.	8.9611	0.0123	9.2869	0.014
2G10622.	CAL 1660@2000PPB	08/08/05 18:42	Soil	2G10600.	8.9612	0.0112	9.2872	0.0108
2G10623.	2000PPB	08/08/05 18:56	Soil	2G10622.	8.9611	0.0011	9.2868	0.0043

Drift Compound: DCB-Surrogate

Drift Limit(s): 0.5 (Pest/Pcb) 1.5(Herb/Tph)

* - Values outside of limits for this column/run

GC PCB Data
Sample Data

18089

Form1

ORGANICS PCB REPORT

Sample Number: AC18807-001	Matrix: Soil
Client Id: PCSB-39(0.5)	Initial Vol: 20g
Data File: 2G10598.D	Final Vol: 10ml
Analysis Date: 08/08/05 12:47	Dilution: 1
Date Rec/Extracted: 07/28/05-08/07/05	Solids: 77

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.032	U	12672-29-6	Aroclor-1248	0.032	U
11104-28-2	Aroclor-1221	0.032	U	11097-69-1	Aroclor-1254	0.032	U
11141-16-5	Aroclor-1232	0.032	U	11096-82-5	Aroclor-1260	0.032	U
53469-21-9	Aroclor-1242	0.032	U				

Worksheet #: 18089

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10598.D\ECD1A.CH Vial: 19
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10598.D\ECD2B.CH
 Acq On : 8 Aug 2005 12:47 Operator: JK
 Sample : AC18807-001 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 12:53 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1932618	1280200	98.945	87.690
35) DCB-Surrogate	8.96	9.29	1969904	1055365	90.493	69.890

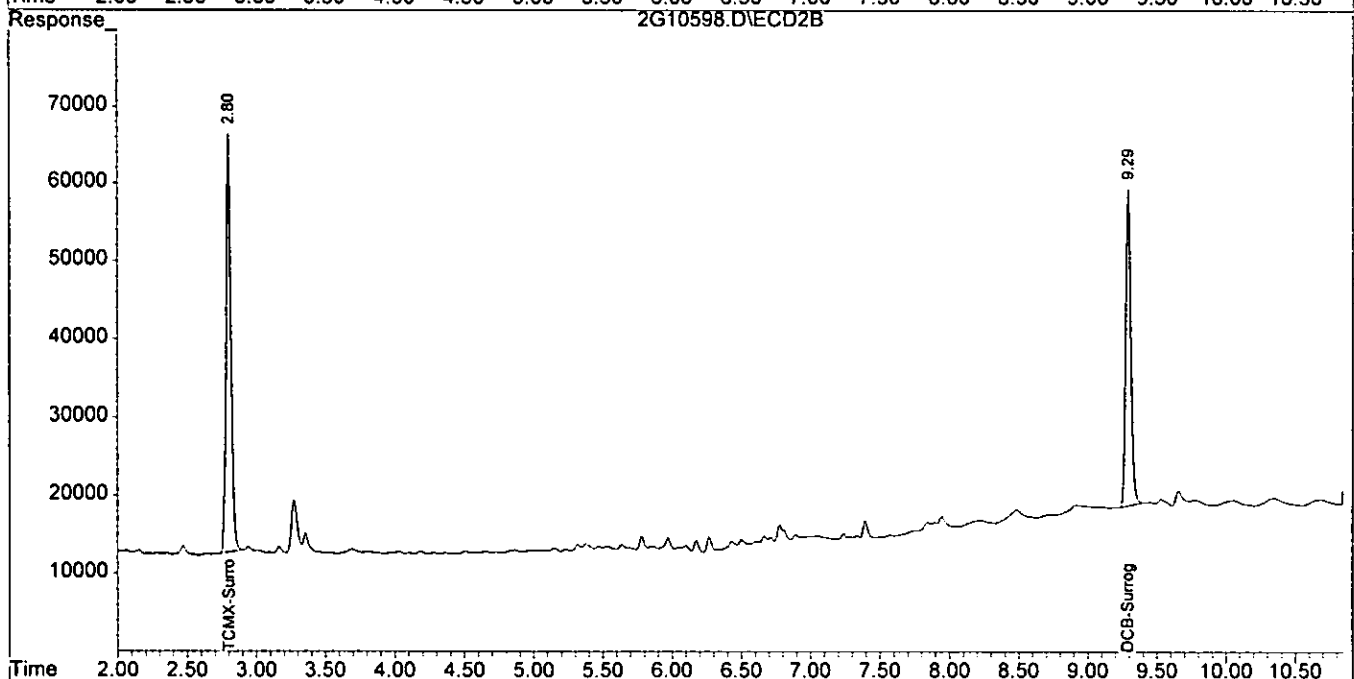
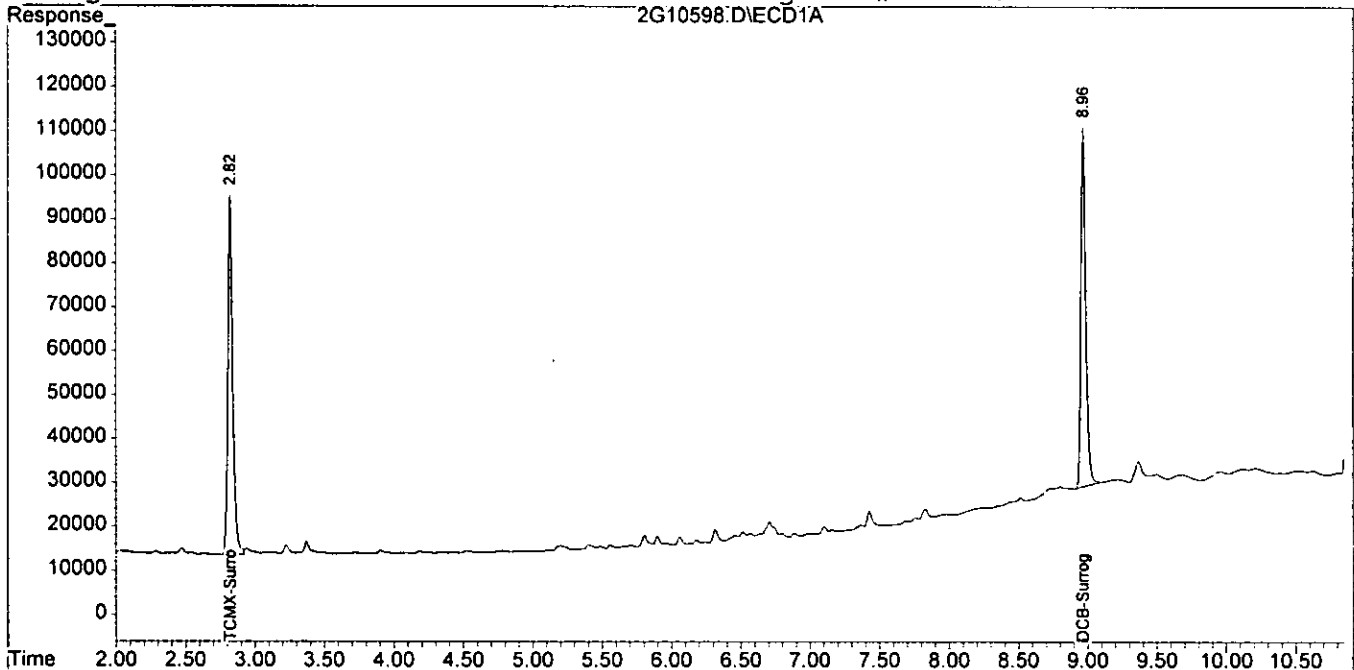
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10598.D\ECD1A.CH Vial: 19
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10598.D\ECD2B.CH
Acq On : 8 Aug 2005 12:47 Operator: JK
Sample : AC18807-001 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 12:53 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-004
 Client Id: PCSB-46(0.5)
 Data File: 2G10599.D
 Analysis Date: 08/08/05 13:10
 Date Rec/Extracted: 07/28/05-08/07/05

Matrix: Soil
 Initial Vol: 20g
 Final Vol: 10ml
 Dilution: 1
 Solids: 88

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.028	U	12672-29-6	Aroclor-1248	0.028	U
11104-28-2	Aroclor-1221	0.028	U	11097-69-1	Aroclor-1254	0.028	U
11141-16-5	Aroclor-1232	0.028	U	11096-82-5	Aroclor-1260	0.028	0.046
53469-21-9	Aroclor-1242	0.028	U				

Worksheet #: 18089

Total Target Concentration 0.046

*U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10599.D\ECD1A.CH Vial: 120
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10599.D\ECD2B.CH
 Acq On : 8 Aug 2005 13:10 Operator: JK
 Sample : AC18807-004 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 13:18 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.83	2.80	1949676	1297400	99.818	88.868
7) Aroclor-1260 {1}	6.06	6.17	78600	61199	77.997	78.973
10) Aroclor-1260 {4}	7.43	7.94	167115	70390	75.819	84.030
11) Aroclor-1260 {5}	7.83	8.48	115823	43397	71.021	77.726m
35) DCB-Surrogate	8.97	9.29	2253337	1208362	104.202	80.022

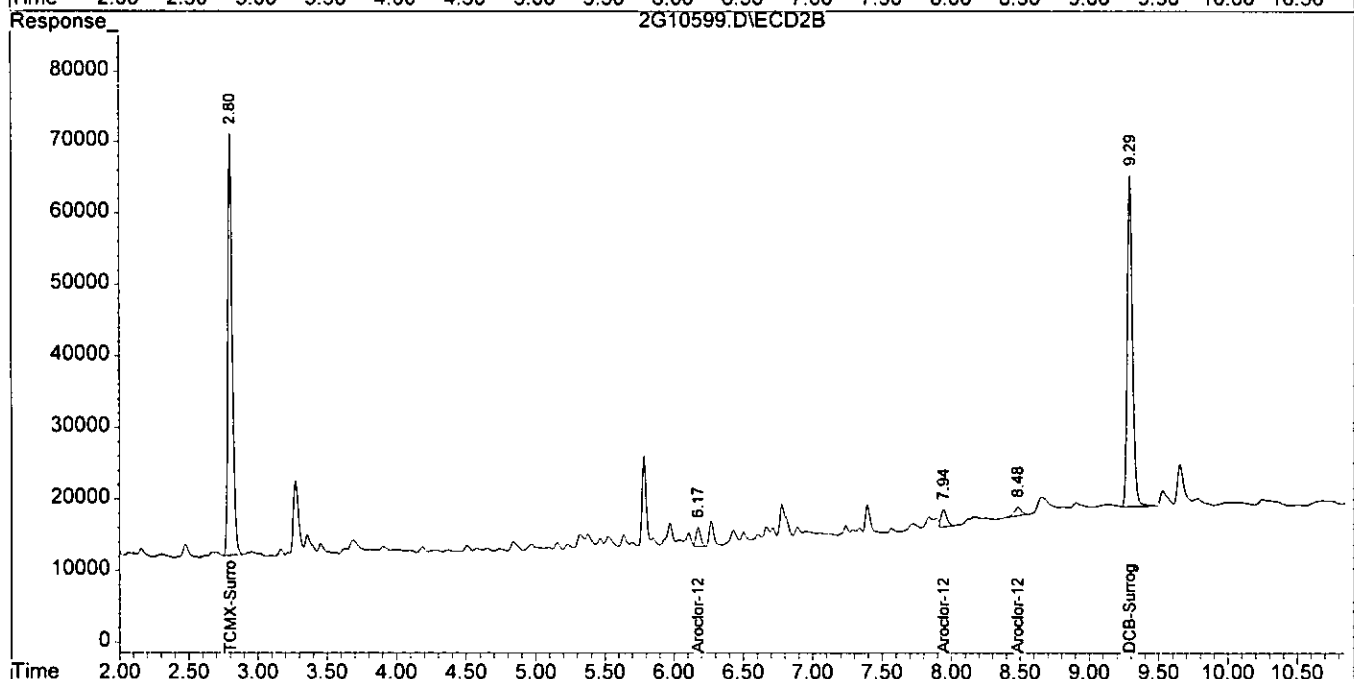
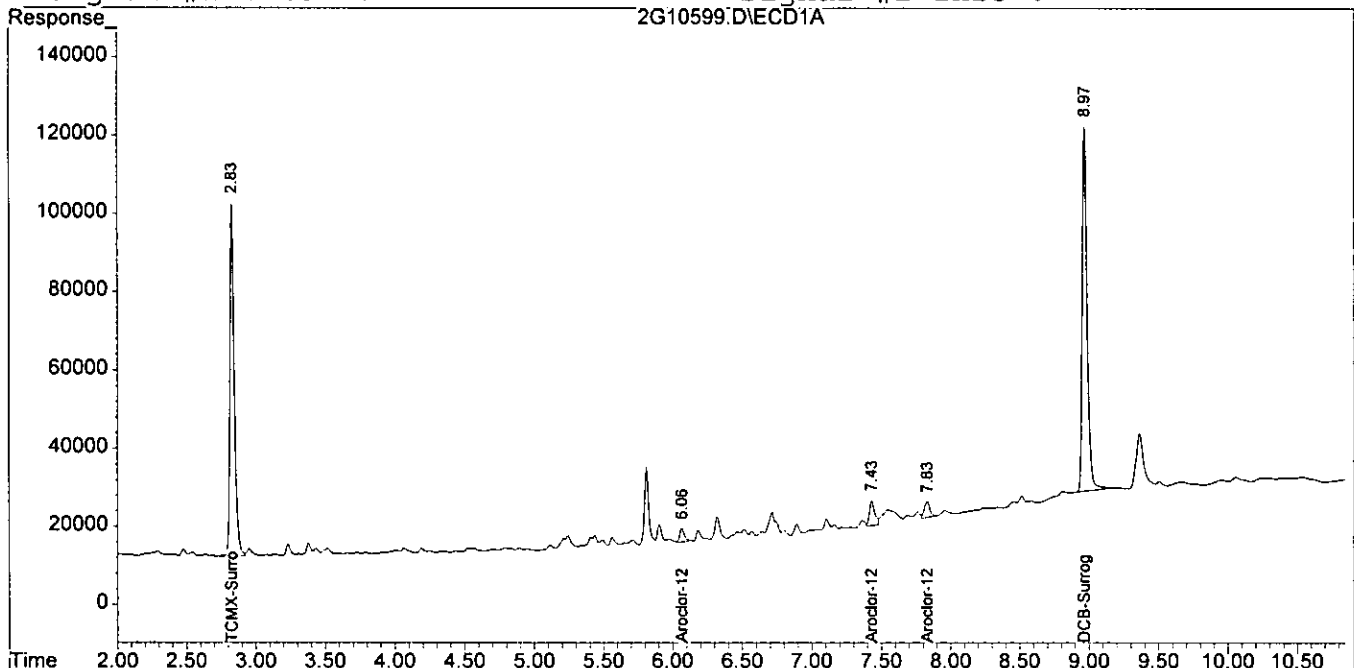
8/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10599.D\ECD1A.CH Vial: 20
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10599.D\ECD2B.CH
Acq On : 8 Aug 2005 13:10 Operator: JK
Sample : AC18807-004 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 13:18 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase :
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-007
 Client Id: FB072805
 Data File: 2G10431.D
 Analysis Date: 08/03/05 12:45
 Date Rec/Extracted: 07/28/05-08/02/05

Matrix: Aqueous
 Initial Vol: 1000ml
 Final Vol: 5ml
 Dilution: 1
 Solids: 0

Units: ug/L

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.25	U	12672-29-6	Aroclor-1248	0.25	U
11104-28-2	Aroclor-1221	0.25	U	11097-69-1	Aroclor-1254	0.25	U
11141-16-5	Aroclor-1232	0.25	U	11096-82-5	Aroclor-1260	0.25	U
53469-21-9	Aroclor-1242	0.25	U				

Worksheet #: 18089

Total Target Concentration 0

*U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10431.D\ECD1A.CH Vial: 6
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10431.D\ECD2B.CH
 Acq On : 3 Aug 2005 12:45 Operator: JK
 Sample : AC18807-007 Inst : gc_2
 Misc : A,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 13:08 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 11:18:05 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	2998979	1920640	116.559	108.323
35) DCB-Surrogate	8.99	9.32	3894180	1452250	155.631	102.739m#

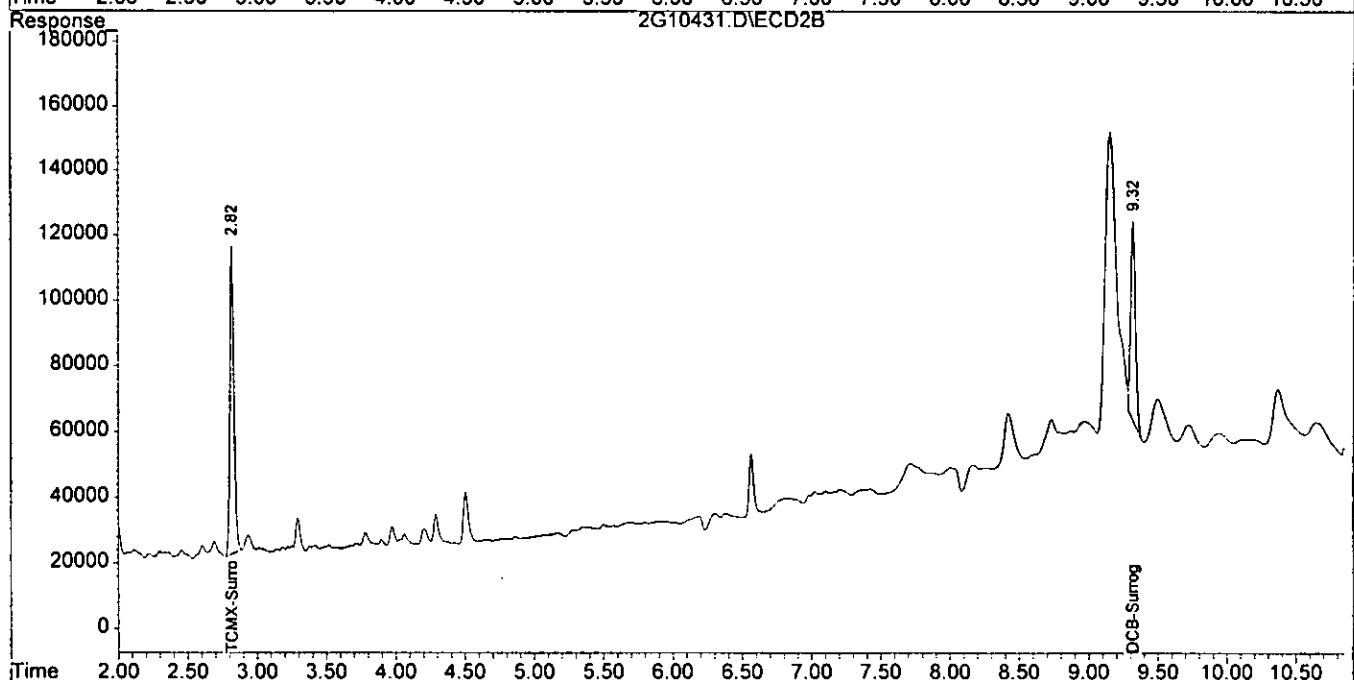
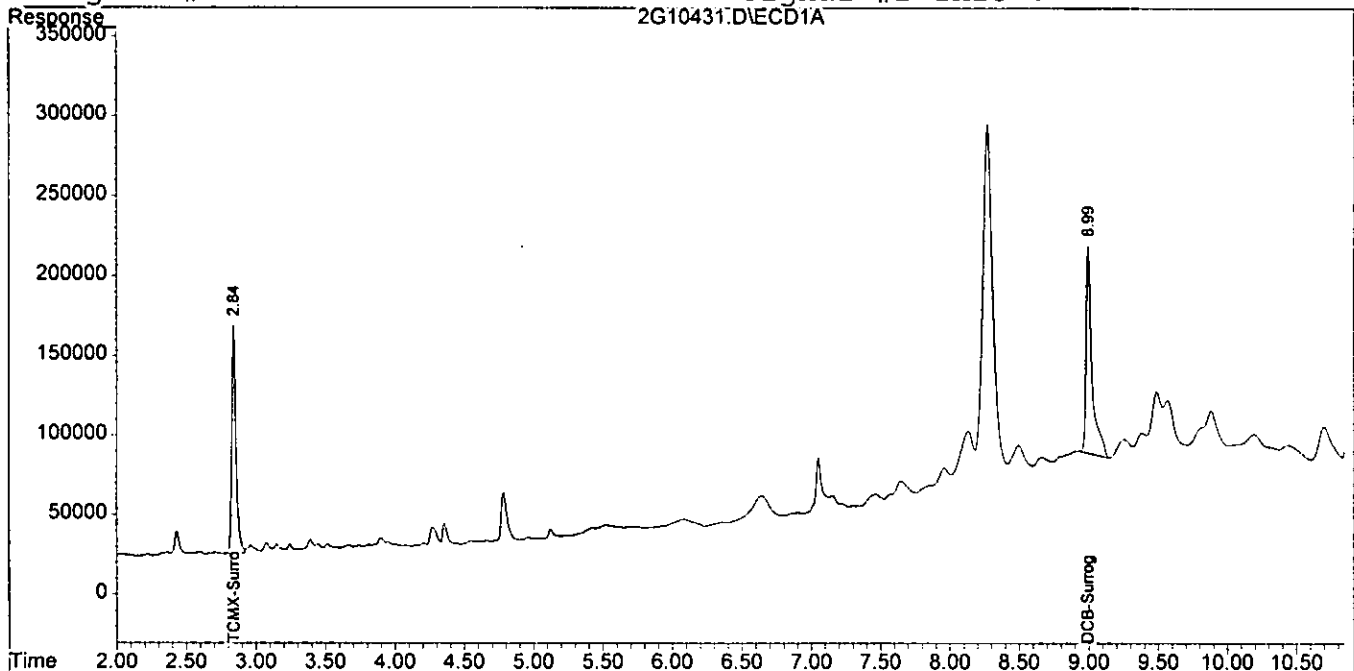
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10431.D\ECD1A.CH Vial: 16
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10431.D\ECD2B.CH
Acq On : 3 Aug 2005 12:45 Operator: JK
Sample : AC18807-007 Inst : gc_2
Misc : A,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 13:08 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 11:18:05 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-008
Client Id: PCSB-40(0.5)
Data File: 2G10611.D
Analysis Date: 08/08/05 16:03
Date Rec/Extracted: 07/28/05-08/07/05

Matrix: Soil
Initial Vol: 20g
Final Vol: 10ml
Dilution: 1
Solids: 75

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.033	U	12672-29-6	Aroclor-1248	0.033	U
11104-28-2	Aroclor-1221	0.033	U	11097-69-1	Aroclor-1254	0.033	U
11141-16-5	Aroclor-1232	0.033	U	11096-82-5	Aroclor-1260	0.033	0.053
53469-21-9	Aroclor-1242	0.033	U				

Worksheet #: 18089

Total Target Concentration 0.053

*U - Indicates the compound was analyzed but not detected.
B - Indicates the analyte was found in the blank as well as in the sample.
E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

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Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10611.D\ECD1A.CH Vial 11
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10611.D\ECD2B.CH
 Acq On : 8 Aug 2005 16:03 Operator: JK
 Sample : AC18807-008 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:48 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	2213650	1472658	113.333	100.873
7) Aroclor-1260 {1}	6.06	6.18	72291	59166	71.737	76.349
8) Aroclor-1260 {2}	6.32	6.27	114347	75234	94.609	86.733
9) Aroclor-1260 {3}	7.10	7.40	67687	112507	75.683m	66.456
10) Aroclor-1260 {4}	7.43	7.94	167279	74404	75.893	88.821
35) DCB-Surrogate	8.96	9.29	2227234	1252433	102.940	82.941

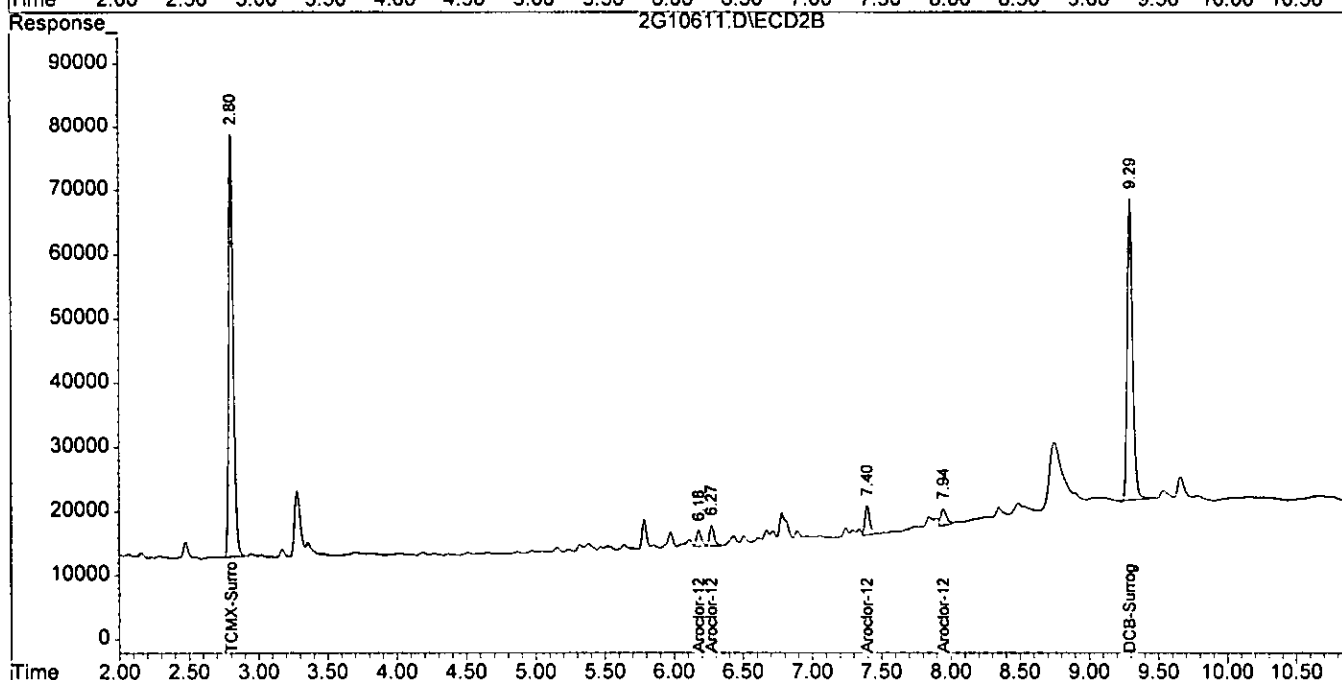
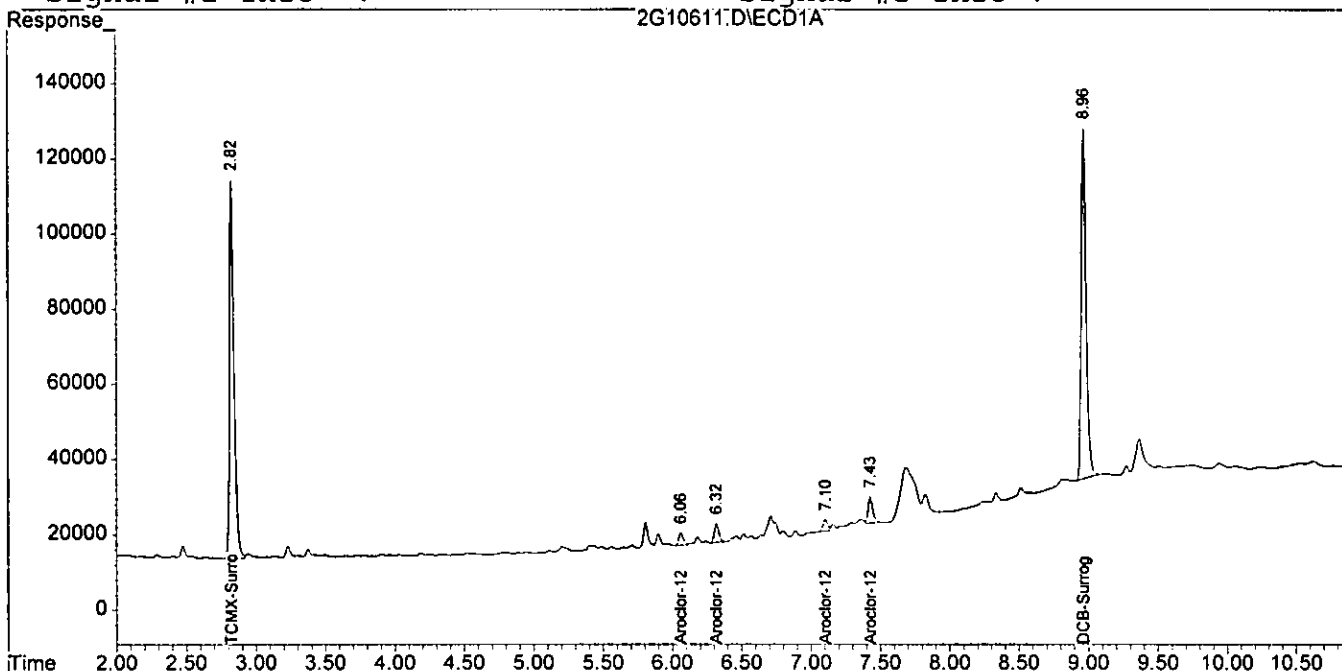
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10611.D\ECD1A.CH Vial: 1
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10611.D\ECD2B.CH
Acq On : 8 Aug 2005 16:03 Operator: JK
Sample : AC18807-008 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 9 6:48 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1
ORGANICS PCB REPORT

Sample Number: AC18807-014	Matrix: Soil
Client Id: PCSB-31(0.5)	Initial Vol: 20g
Data File: 2G10608.D	Final Vol: 10ml
Analysis Date: 08/08/05 15:20	Dilution: 1
Date Rec/Extracted: 07/28/05-08/07/05	Solids: 72

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.035	U	12672-29-6	Aroclor-1248	0.035	U
11104-28-2	Aroclor-1221	0.035	U	11097-69-1	Aroclor-1254	0.035	U
11141-16-5	Aroclor-1232	0.035	U	11096-82-5	Aroclor-1260	0.035	U
53469-21-9	Aroclor-1242	0.035	U				

Worksheet #: 18089

Total Target Concentration 0

*U - Indicates the compound was analyzed but not detected.
B - Indicates the analyte was found in the blank as well as in the sample.
E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10608.D\ECD1A.CH Vial: 8
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10608.D\ECD2B.CH
 Acq On : 8 Aug 2005 15:20 Operator: JK
 Sample : AC18807-014 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.83	2.80	2006632	1353292	102.734	92.697
35) DCB-Surrogate	8.97	9.29	2116402	1198575	97.579	79.374m

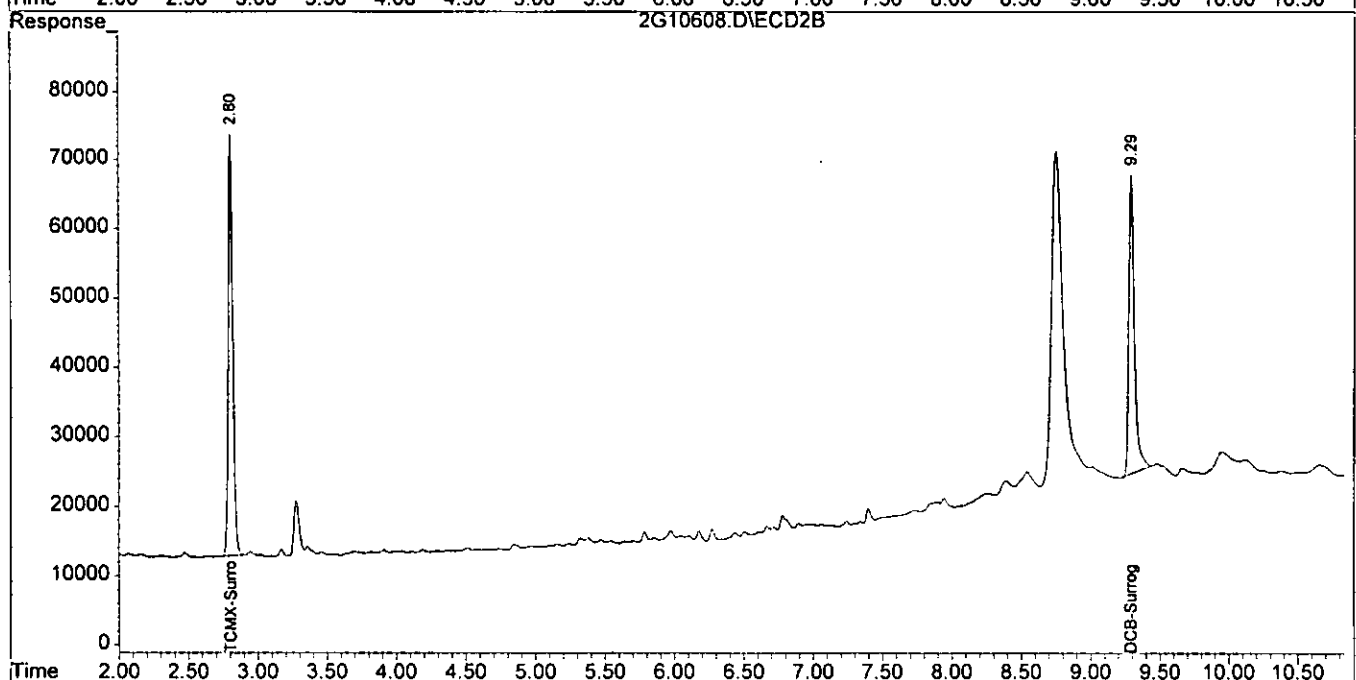
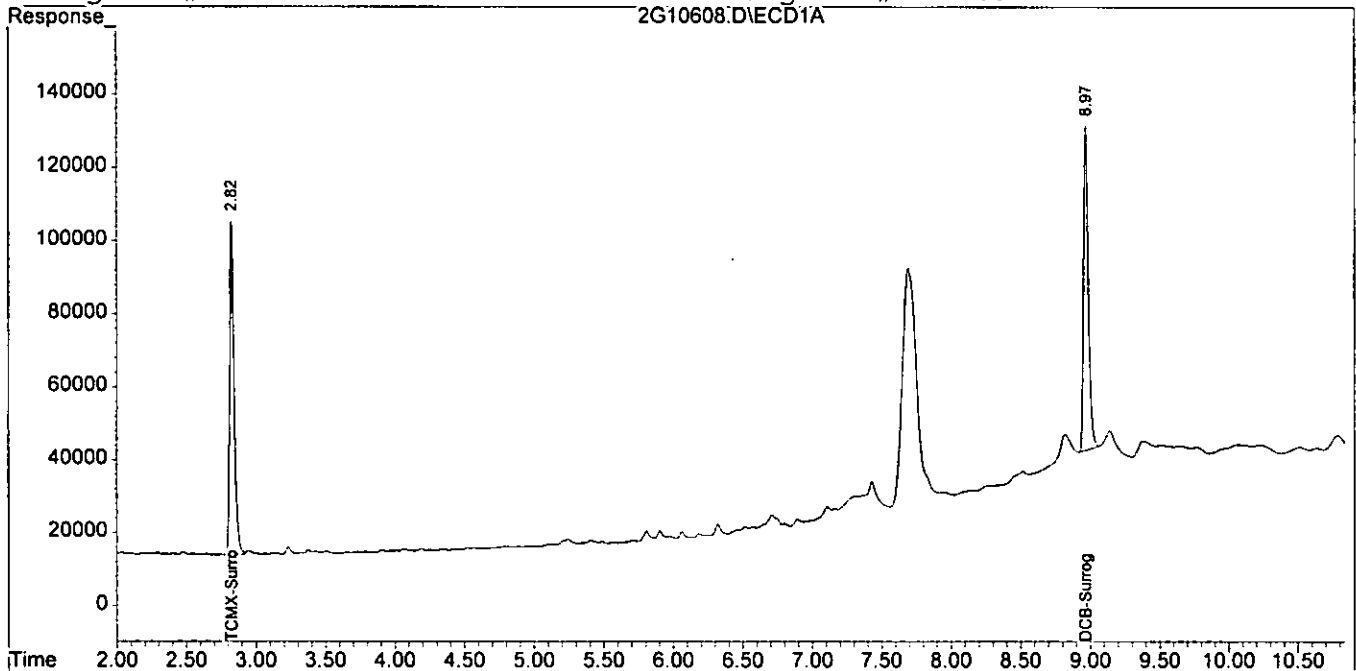
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10608.D\ECD1A.CH Vial: 151
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10608.D\ECD2B.CH
Acq On : 8 Aug 2005 15:20 Operator: JK
Sample : AC18807-014 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 9 6:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-017	Matrix: Soil
Client Id: PCSB-32(0.5)	Initial Vol: 20g
Data File: 2G10609.D	Final Vol: 10ml
Analysis Date: 08/08/05 15:34	Dilution: 1
Date Rec/Extracted: 07/28/05-08/07/05	Solids: 75

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.033	U	12672-29-6	Aroclor-1248	0.033	U
11104-28-2	Aroclor-1221	0.033	U	11097-69-1	Aroclor-1254	0.033	U
11141-16-5	Aroclor-1232	0.033	U	11096-82-5	Aroclor-1260	0.033	U
53469-21-9	Aroclor-1242	0.033	U				

Worksheet #: 18089

Total Target Concentration 0

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

1007

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10609.D\ECD1A.CH Vial 9
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10609.D\ECD2B.CH
 Acq On : 8 Aug 2005 15:34 Operator: JK
 Sample : AC18807-017 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.83	2.80	2155241	1445012	110.342	98.979
35) DCB-Surrogate	8.96	9.29	2050565	1179280	94.395m	78.096m

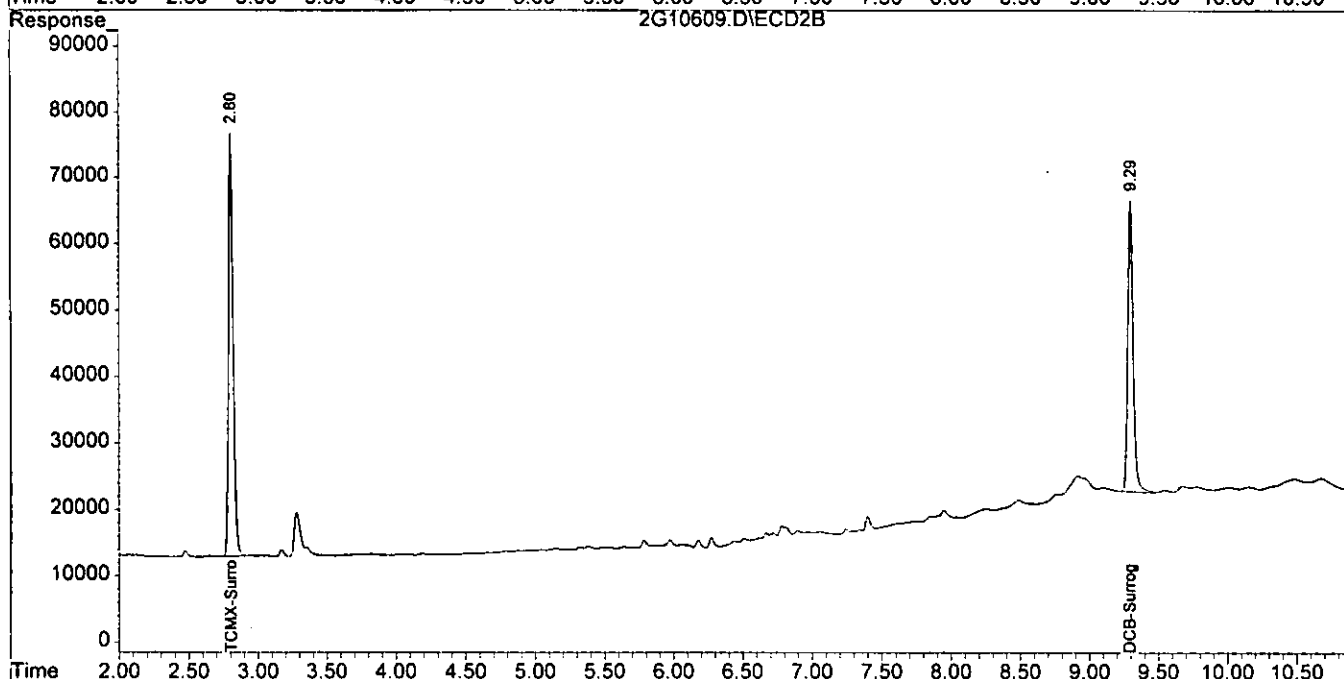
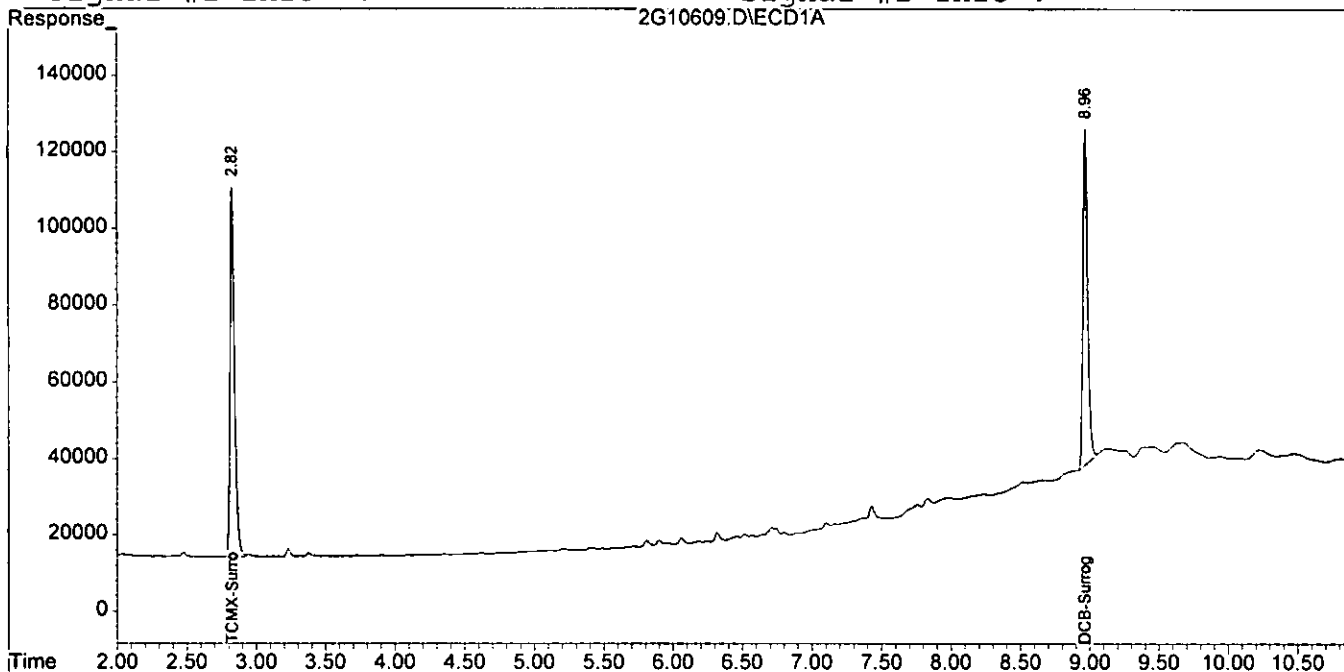
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10609.D\ECD1A.CH Vial: 20
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10609.D\ECD2B.CH
Acq On : 8 Aug 2005 15:34 Operator: JK
Sample : AC18807-017 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 9 6:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-020
 Client Id: PCSB-33(0.5)
 Data File: 2G10610.D
 Analysis Date: 08/08/05 15:49
 Date Rec/Extracted: 07/28/05-08/07/05

Matrix: Soil
 Initial Vol: 20g
 Final Vol: 10ml
 Dilution: 1
 Solids: 80

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.031	U	12672-29-6	Aroclor-1248	0.031	U
11104-28-2	Aroclor-1221	0.031	U	11097-69-1	Aroclor-1254	0.031	U
11141-16-5	Aroclor-1232	0.031	U	11096-82-5	Aroclor-1260	0.031	U
53469-21-9	Aroclor-1242	0.031	U				

Worksheet #: 18089

Total Target Concentration 0

*U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10610.D\ECD1A.CH Vial: 100
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10610.D\ECD2B.CH
 Acq On : 8 Aug 2005 15:49 Operator: JK
 Sample : AC18807-020 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:46 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1824328	1224986	93.401	83.908
35) DCB-Surrogate	8.96	9.29	1773159	958003	80.977	63.442

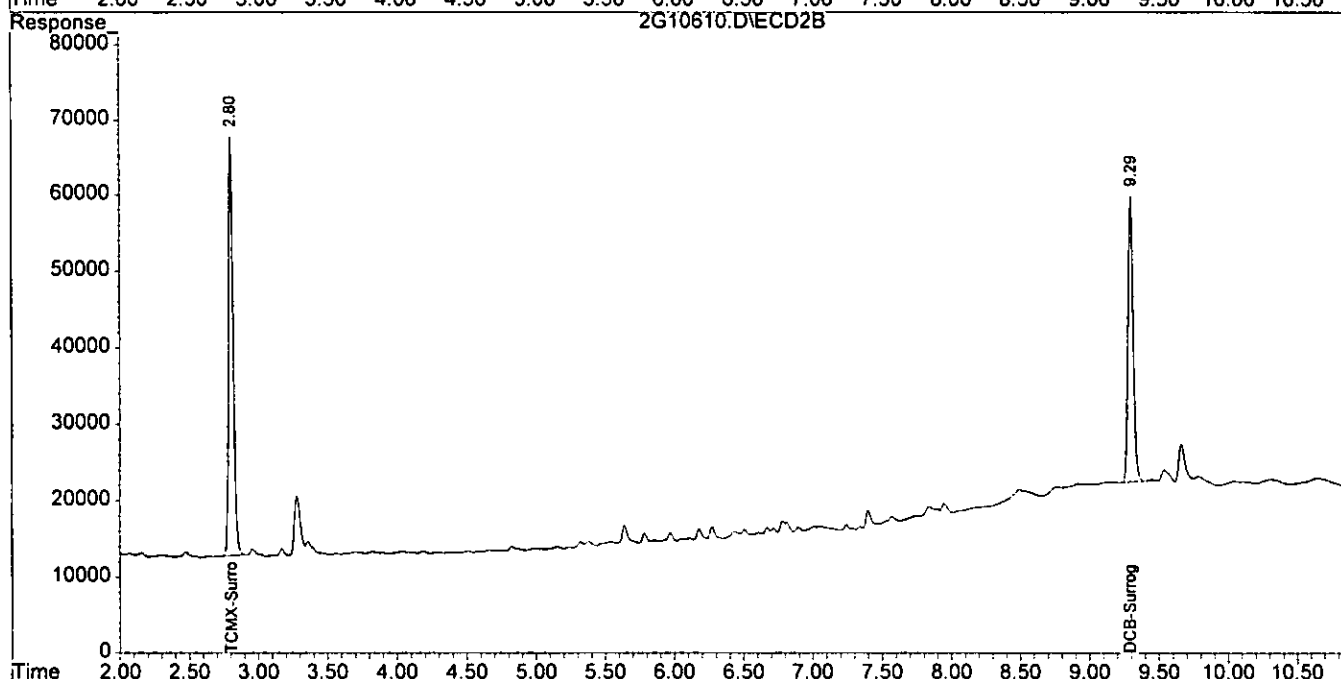
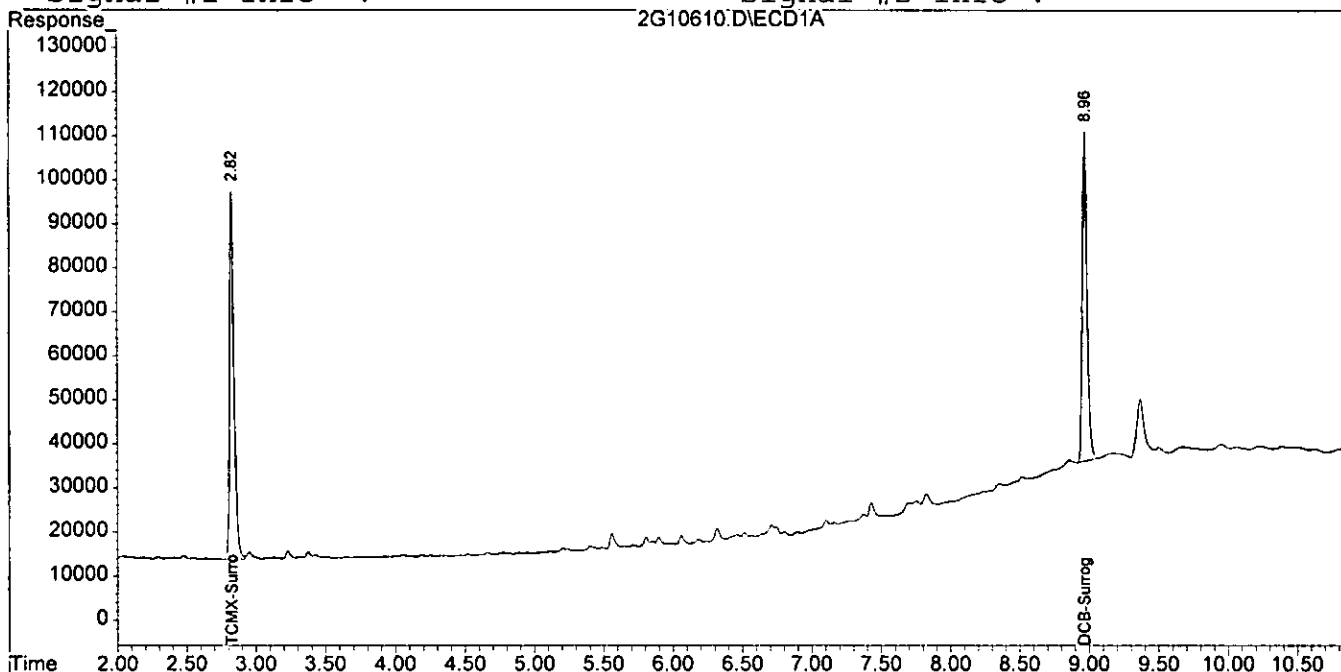
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10610.D\ECD1A.CH Vial: 0
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10610.D\ECD2B.CH
Acq On : 8 Aug 2005 15:49 Operator: JK
Sample : AC18807-020 Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 9 6:46 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: AC18807-023	Matrix: Soil
Client Id: PCSB-41(0.5)	Initial Vol: 20g
Data File: 2G10607.D	Final Vol: 10ml
Analysis Date: 08/08/05 15:06	Dilution: 1
Date Rec/Extracted: 07/28/05-08/07/05	Solids: 92

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.027	U	12672-29-6	Aroclor-1248	0.027	0.10
11104-28-2	Aroclor-1221	0.027	U	11097-69-1	Aroclor-1254	0.027	U
11141-16-5	Aroclor-1232	0.027	U	11096-82-5	Aroclor-1260	0.027	0.33
53469-21-9	Aroclor-1242	0.027	U				

Worksheet #: 18089

Total Target Concentration 0.43

U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.

R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10607.D\ECD1A.CH Vial 7
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10607.D\ECD2B.CH
 Acq On : 8 Aug 2005 15:06 Operator: JK
 Sample : AC18807-023 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:42 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.83	2.80	1916588	1299638	98.124	89.021
7) Aroclor-1260 {1}	6.06	6.18	584071	441435	579.591	569.635
9) Aroclor-1260 {3}	7.10	7.40	537839	813649	601.380m	480.607
10) Aroclor-1260 {4}	7.43	7.95	1432098	577308	649.732	689.178
11) Aroclor-1260 {5}	7.83	8.49	991348	334220	607.884	598.613m
25) Aroclor-1248 {1}	3.72	3.81	56064	51342	143.670	178.503m
26) Aroclor-1248 {2}	4.19	4.19	227663	172365	176.437	181.987
28) Aroclor-1248 {4}	4.79	4.87	198886	111959	174.813	211.856m
35) DCB-Surrogate	8.97	9.29	2227661	1138446	102.960	75.392 #

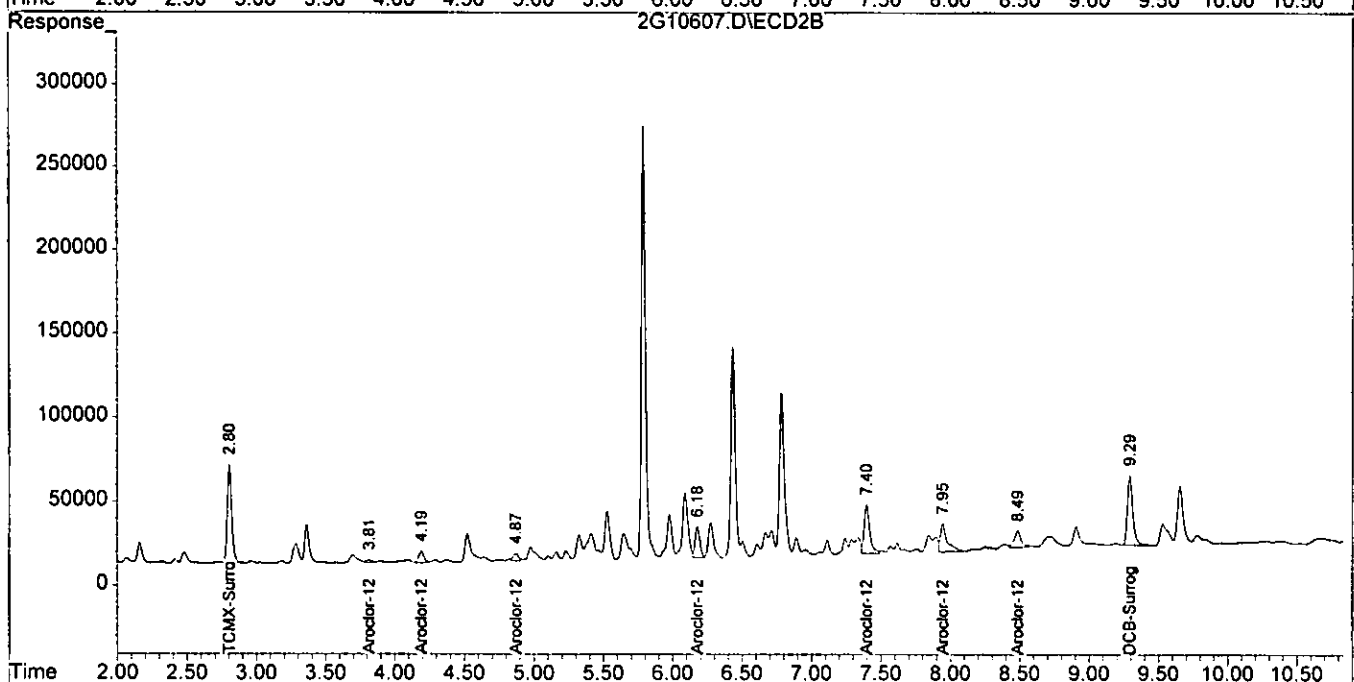
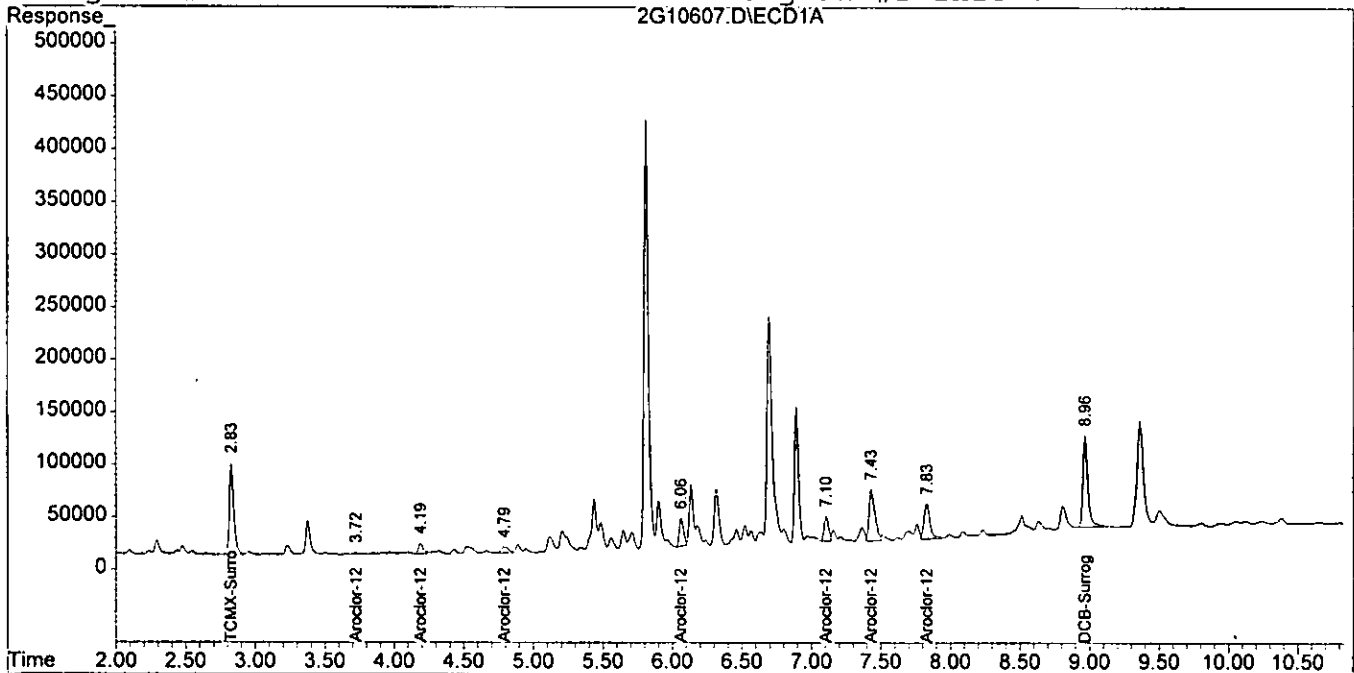
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10607.D\ECD1A.CH Vial: 7
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10607.D\ECD2B.CH
 Acq On : 8 Aug 2005 15:06 Operator: JK
 Sample : AC18807-023 Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 6:42 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Multiple Level Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :



**GC PCB Data
Standards Data**

Form 6
Initial Calibration

Instrument: GC_2

Level #	Data File:	Cal Identifier:	Analysis Date/Time	Level #:	Data File:	Cal Identifier:	Analysis Date/Time	Calibration Level Concentrations									
								Lvl1	Lvl2	Lvl3	Lvl4	Lvl5	Lvl6	Lvl7	Lvl8		
1	2G10417	CAL 1660@500PPB	08/03/05 08:47	2	2G10418	CAL 1660@200PPB	08/03/05 09:04	5.00	20.00	50.00	100.0	200.0	400.0				
3	2G10416	CAL 1660@500PPB	08/03/05 08:26	4	2G10419	CAL 1660@1000PPB	08/03/05 09:19	50.00	200.0	500.0	1000.0	2000.0	4000.0				
5	2G10420	CAL 1660@2000PPB	08/03/05 09:33	6	2G10421	CAL 1660@4000PPB	08/03/05 09:47	50.00	200.0	500.0	1000.0	2000.0	4000.0				
7	2G10425	CAL 1232@500PPB	08/03/05 11:04	8	2G10424	CAL 1242@500PPB	08/03/05 10:50	50.00	200.0	500.0	1000.0	2000.0	4000.0				
9	2G10423	CAL 1248@500PPB	08/03/05 10:35	10	2G10422	CAL 2154@500PPB	08/03/05 10:20	50.00	200.0	500.0	1000.0	2000.0	4000.0				
Compound	Col	Mr	Fit	RF1	RF2	RF3	RF4	RF5	RF6	RF7	RF8	AvgRf	RT	Corr1	Corr2	%Rsd	
TCMX-Surrogate	1	0	Avg	2.7521	2.6878	2.7601	2.5151	2.4524	2.2698	---	---	2.57	2.84	0.998	1.00	7.6	
Aroclor-1016	1	1	Qea	0.0614	0.0618	0.0611	0.0544	0.0506	0.0451	---	---	0.0558	3.37	0.995	1.00	12	
Aroclor-1016	1	2	Qea	0.1264	0.1182	0.1139	0.0998	0.0929	0.0825	---	---	0.106	3.74	0.995	1.00	16	
Aroclor-1016	1	3	Qea	0.2762	0.2401	0.2420	0.2091	0.1968	0.1743	---	---	0.223	4.20	0.995	1.00	16	
Aroclor-1016	1	4	Qea	0.1676	0.1580	0.1565	0.1382	0.1294	0.1143	---	---	0.144	4.57	0.995	1.00	14	
Aroclor-1016	1	5	Qea	0.1951	0.1328	0.1177	0.0985	0.0882	0.0765	---	---	0.118	4.81	0.994	0.999	36	
Aroclor-1260	1	1	Qea	0.1391	0.1367	0.1299	0.1161	0.1091	0.0963	---	---	0.121	6.08	0.995	1.00	14	
Aroclor-1260	1	2	Qea	0.1700	0.1613	0.1566	0.1390	0.1321	0.1170	---	---	0.146	6.34	0.995	1.00	14	
Aroclor-1260	1	3	Qea	0.1050	0.1129	0.1170	0.1000	0.0974	0.0871	---	---	0.103	7.13	0.996	1.00	11	
Aroclor-1260	1	4	Avg	0.2484	0.2620	0.2755	0.2509	0.2517	0.2328	---	---	0.254	7.46	0.998	1.00	5.6	
Aroclor-1260	1	5	Avg	0.1735	0.1808	0.2039	0.1789	0.1804	0.1667	---	---	0.181	7.86	0.998	1.00	7.0	
Aroclor-1221	1	1	Avg	---	---	---	---	---	---	---	---	0.0349	3.18	-1	-1	Lvl=10	
Aroclor-1221	1	2	Avg	---	---	---	---	---	---	---	---	0.0233	3.33	-1	-1	Lvl=10	
Aroclor-1221	1	3	Avg	---	---	---	---	---	---	---	---	0.0890	3.39	-1	-1	Lvl=10	
Aroclor-1221	1	4	Avg	---	---	---	---	---	---	---	---	0.0628	3.38	-1	-1	Lvl=7	
Aroclor-1232	1	1	Avg	---	---	---	---	---	---	---	---	0.0533	3.75	-1	-1	Lvl=7	
Aroclor-1232	1	2	Avg	---	---	---	---	---	---	---	---	0.110	4.21	-1	-1	Lvl=7	
Aroclor-1232	1	3	Avg	---	---	---	---	---	---	---	---	0.0756	4.56	-1	-1	Lvl=7	
Aroclor-1232	1	4	Avg	---	---	---	---	---	---	---	---	0.0680	4.81	-1	-1	Lvl=7	
Aroclor-1232	1	5	Avg	---	---	---	---	---	---	---	---	0.0537	3.37	-1	-1	Lvl=8	
Aroclor-1242	1	1	Avg	---	---	---	---	---	---	---	---	0.0917	3.74	-1	-1	Lvl=8	
Aroclor-1242	1	2	Avg	---	---	---	---	---	---	---	---	0.195	4.20	-1	-1	Lvl=8	
Aroclor-1242	1	3	Avg	---	---	---	---	---	---	---	---	0.127	4.56	-1	-1	Lvl=8	
Aroclor-1242	1	4	Avg	---	---	---	---	---	---	---	---	0.0948	4.81	-1	-1	Lvl=8	
Aroclor-1242	1	5	Avg	---	---	---	---	---	---	---	---	0.0459	3.75	-1	-1	Lvl=9	
Aroclor-1248	1	1	Avg	---	---	---	---	---	---	---	---	0.153	4.21	-1	-1	Lvl=9	
Aroclor-1248	1	2	Avg	---	---	---	---	---	---	---	---	0.181	4.57	-1	-1	Lvl=9	
Aroclor-1248	1	3	Avg	---	---	---	---	---	---	---	---	0.128	4.81	-1	-1	Lvl=9	
Aroclor-1248	1	4	Avg	---	---	---	---	---	---	---	---	0.209	5.26	-1	-1	Lvl=9	
Aroclor-1248	1	5	Avg	---	---	---	---	---	---	---	---	0.173	5.23	-1	-1	Lvl=10	
Aroclor-1254	1	1	Avg	---	---	---	---	---	---	---	---	0.110	5.81	-1	-1	Lvl=10	
Aroclor-1254	1	2	Avg	---	---	---	---	---	---	---	---	0.191	5.92	-1	-1	Lvl=10	
Aroclor-1254	1	3	Avg	---	---	---	---	---	---	---	---	0.127	6.20	-1	-1	Lvl=10	
Aroclor-1254	1	4	Avg	---	---	---	---	---	---	---	---	0.187	6.74	-1	-1	Lvl=10	
Aroclor-1254	1	5	Avg	---	---	---	---	---	---	---	---	2.50	8.99	0.998	1.00	6.7	
DCB-Surrogate	1	0	Avg	2.6019	2.5758	2.7116	2.4625	2.4318	2.2294	---	---	1.77	2.82	0.999	1.00	12	
TCMX-Surrogate	2	0	Avg	2.0944	1.9467	1.8077	1.6626	1.5955	1.5313	---	---	0.0342	3.42	0.999	1.00	22	
Aroclor-1016	2	1	Qea	0.0443	0.0417	0.0333	0.0320	0.0290	0.0243	---	---	0.0342	3.42	0.999	1.00	22	
Avg Rsd Col 1: 13.3 Avg Rsd Col 2: 17																	

Flags
c - failed the initial calibration criteria (if applicable)

Note:
Col = Column Number
Mr = MultiPeak Analyte 0=single peak analyte, >0=multi peak analyte (i.e. pcb/chlordane etc.)
Fit = Indicates whether Avg Rf, Linear, or Quadratic Curve was used for compound.
Corr 1 = Correlation Coefficient for linear Eq.
Corr 2 = Correlation Coefficient for quad Eq.
*Lvl: These compounds use a single pt calibration as specified by the method. The file used to update this calibration point is listed in the header under level #

All Response Factors = Response Factors / 10000
Initial Calibration Criteria: either %RSD <=20 or Corr >= .995
Columns: Signal #1 db-1701 ; Signal #2 db-608

Form 6
Initial Calibration

Level #	Data File:	Cal Identifier:	Analysis Date/Time	Level #	Data File:	Cal Identifier:	Calibration Level Concentrations										
							Lvl1	Lvl2	Lvl3	Lvl4	Lvl5	Lvl6	Lvl7	Lvl8			
1	2G10417	CAL 1660@500PPB	08/03/05 08:47	2	2G10418	CAL 1660@200PPB	50.00	200.0	500.0	1000.	2000.	4000.					
3	2G10416	CAL 1660@500PPB	08/03/05 08:26	4	2G10419	CAL 1660@1000PPB	50.00	200.0	500.0	1000.	2000.	4000.					
5	2G10420	CAL 1660@2000PPB	08/03/05 09:33	6	2G10421	CAL 1660@4000PPB	50.00	200.0	500.0	1000.	2000.	4000.					
7	2G10425	CAL 1232@500PPB	08/03/05 11:04	8	2G10424	CAL 1242@500PPB	50.00	200.0	500.0	1000.	2000.	4000.					
9	2G10423	CAL 1248@500PPB	08/03/05 10:35	10	2G10422	CAL 2154@500PPB	50.00	200.0	500.0	1000.	2000.	4000.					
	Col	Mr	Fit	RF1	RF2	RF3	RF4	RF5	RF6	RF7	RF8	AvgRf	RT	Corr1	Corr2	%Rsd	
Aroclor-1016	2	2	Qua	0.0985	0.0871	0.0769	0.0682	0.0610	0.0531	---	---	0.0742	3.83	0.993	1.00	23	
Aroclor-1016	2	3	Qua	0.2132	0.1810	0.1589	0.1404	0.1272	0.1131	---	---	0.156	4.21	0.995	1.00	24	
Aroclor-1016	2	4	Qua	0.1628	0.1057	0.0869	0.0736	0.0631	0.0535	---	---	0.0910	4.53	0.991	0.999	44	
Aroclor-1016	2	5	Avg	0.0626	0.0584	0.0532	0.0476	0.0432	0.0369	---	---	0.0504	4.89	0.992	1.00	19	
Aroclor-1260	2	1	Avg	0.1021	0.0959	0.0868	0.0807	0.0744	0.0648	---	---	0.0843	6.20	0.994	1.00	17	
Aroclor-1260	2	2	Avg	0.1138	0.1107	0.0972	0.0898	0.0830	0.0724	---	---	0.0945	6.29	0.994	1.00	17	
Aroclor-1260	2	3	Avg	0.1641	0.1891	0.1789	0.1720	0.1710	0.1651	---	---	0.173	7.42	1.00	1.00	5.4	
Aroclor-1260	2	4	Avg	0.0927	0.0840	0.0880	0.0790	0.0788	0.0731	---	---	0.0826	7.97	0.998	1.00	8.6	
Aroclor-1260	2	5	Avg	0.0540	0.0561	0.0530	0.0556	0.0541	0.0500	---	---	0.0538	8.51	0.998	1.00	4.1	
Aroclor-1221	2	1	Avg	---	---	---	---	---	---	---	---	0.0241	3.22	-1	-1	Lvl=10	
Aroclor-1221	2	2	Avg	---	---	---	---	---	---	---	---	0.0139	3.36	-1	-1	Lvl=10	
Aroclor-1221	2	3	Avg	---	---	---	---	---	---	---	---	0.0434	3.43	-1	-1	Lvl=10	
Aroclor-1232	2	1	Avg	---	---	---	---	---	---	---	---	0.0327	3.43	-1	-1	Lvl=7	
Aroclor-1232	2	2	Avg	---	---	---	---	---	---	---	---	0.0420	3.84	-1	-1	Lvl=7	
Aroclor-1232	2	3	Avg	---	---	---	---	---	---	---	---	0.0805	4.21	-1	-1	Lvl=7	
Aroclor-1232	2	4	Avg	---	---	---	---	---	---	---	---	0.0522	4.53	-1	-1	Lvl=7	
Aroclor-1232	2	5	Avg	---	---	---	---	---	---	---	---	0.0323	4.89	-1	-1	Lvl=7	
Aroclor-1242	2	1	Avg	---	---	---	---	---	---	---	---	0.0343	3.42	-1	-1	Lvl=8	
Aroclor-1242	2	2	Avg	---	---	---	---	---	---	---	---	0.0672	3.83	-1	-1	Lvl=8	
Aroclor-1242	2	3	Avg	---	---	---	---	---	---	---	---	0.138	4.21	-1	-1	Lvl=8	
Aroclor-1242	2	4	Avg	---	---	---	---	---	---	---	---	0.0719	4.53	-1	-1	Lvl=8	
Aroclor-1242	2	5	Avg	---	---	---	---	---	---	---	---	0.0498	5.17	-1	-1	Lvl=8	
Aroclor-1248	2	1	Avg	---	---	---	---	---	---	---	---	0.0333	3.84	-1	-1	Lvl=9	
Aroclor-1248	2	2	Avg	---	---	---	---	---	---	---	---	0.110	4.21	-1	-1	Lvl=9	
Aroclor-1248	2	3	Avg	---	---	---	---	---	---	---	---	0.0740	4.54	-1	-1	Lvl=9	
Aroclor-1248	2	4	Avg	---	---	---	---	---	---	---	---	0.0627	4.89	-1	-1	Lvl=9	
Aroclor-1248	2	5	Avg	---	---	---	---	---	---	---	---	0.0777	5.56	-1	-1	Lvl=9	
Aroclor-1254	2	1	Avg	---	---	---	---	---	---	---	---	0.0797	5.34	-1	-1	Lvl=10	
Aroclor-1254	2	2	Avg	---	---	---	---	---	---	---	---	0.127	5.99	-1	-1	Lvl=10	
Aroclor-1254	2	3	Avg	---	---	---	---	---	---	---	---	0.0614	6.28	-1	-1	Lvl=10	
Aroclor-1254	2	4	Avg	---	---	---	---	---	---	---	---	0.104	6.80	-1	-1	Lvl=10	
Aroclor-1254	2	5	Avg	---	---	---	---	---	---	---	---	0.0444	7.31	-1	-1	Lvl=10	
DCB-Surrogate	2	0	Avg	1.4892	1.5661	1.5090	1.3375	1.3193	1.2598	---	---	1.41	9.32	0.999	1.00	8.7	

Avg Rsd Col 1: 13.3 Avg Rsd Col 2: 17

Flags
c - failed the initial calibration criteria(if applicable)

Note:
Col = Column Number
Mir = MultiPeak Analyte 0=single peak analyte, >0=multi peak analyte (i.e. pcb/chlordane etc.)
Fit = Indicates whether Avg Rf, Linear, or Quadratic Curve was used for compound.
Corr 1 = Correlation Coefficient for linear Eq.
Corr 2 = Correlation Coefficient for quad Eq.
All Response Factors = Response Factors / 10000
Initial Calibration Criteria: either %RSD <=20 or Corr >= .995
Columns: Signal #1 db-1701 : Signal #2 db-608

*Lvl: These compounds use a single pt calibration as specified by the method. The file used to update this calibration point is listed in the header under level #

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10417.D\ECD1A.CH Vial: 112
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10417.D\ECD2B.CH
 Acq On : 3 Aug 2005 8:47 Operator: JK
 Sample : CAL 1660@50PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 8:59 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

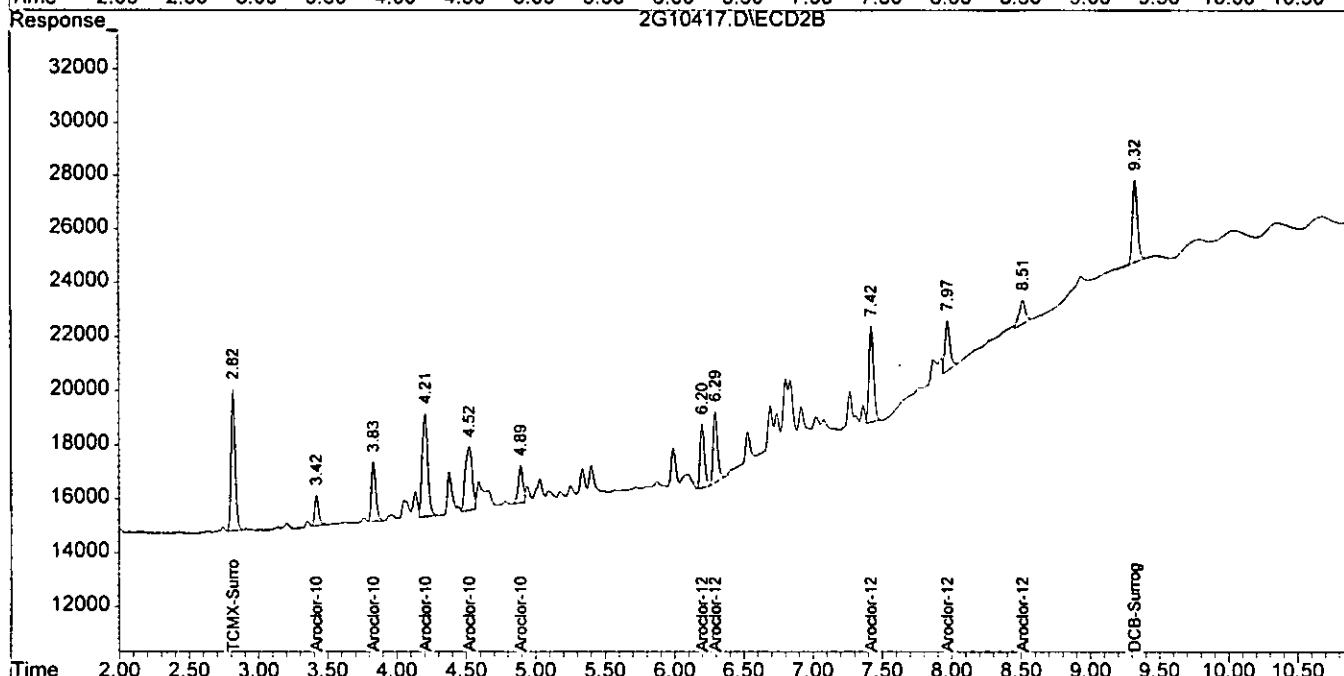
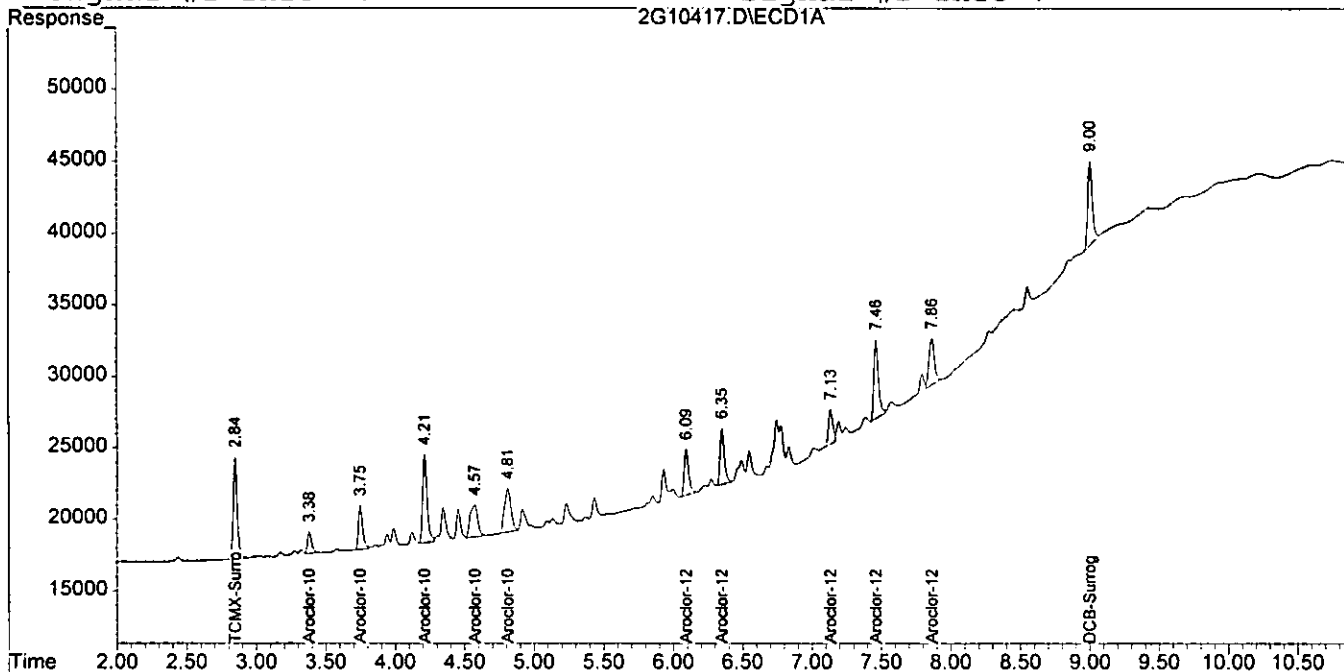
Target Compounds						
1) TCMX-Surrogate	2.85	2.82	137605	104722	7.614	7.811
2) Aroclor-1016 {1}	3.38	3.42	30735	22167	76.093	59.453
3) Aroclor-1016 {2}	3.75	3.83	63199	49263	81.395	49.106 #
4) Aroclor-1016 {3}	4.21	4.21	138146	106626	78.634	52.836 #
5) Aroclor-1016 {4}	4.57	4.52	83831	81400	69.437	42.849 #
6) Aroclor-1016 {5}	4.81	4.89	97551	31346	45.105	80.683 #
7) Aroclor-1260 {1}	6.09	6.20	69572	51085	75.769	75.081
8) Aroclor-1260 {2}	6.35	6.29	85028	56903	74.999	74.858
9) Aroclor-1260 {3}	7.13	7.42	52503	82091	68.929	57.071
10) Aroclor-1260 {4}	7.46	7.97	124210	46351	62.656	66.108m
11) Aroclor-1260 {5}	7.86	8.51	86765	27044	62.809	57.750m
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	9.00	9.32	130096	74465	6.462m	5.744

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10417.D\ECD1A.CH Vial: 142
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10417.D\ECD2B.CH
Acq On : 3 Aug 2005 8:47 Operator: JK
Sample : CAL 1660@50PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 8:59 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10418.D\ECD1A.CH Vial: 113
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10418.D\ECD2B.CH
 Acq On : 3 Aug 2005 9:04 Operator: JK
 Sample : CAL 1660@200PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 10:37 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

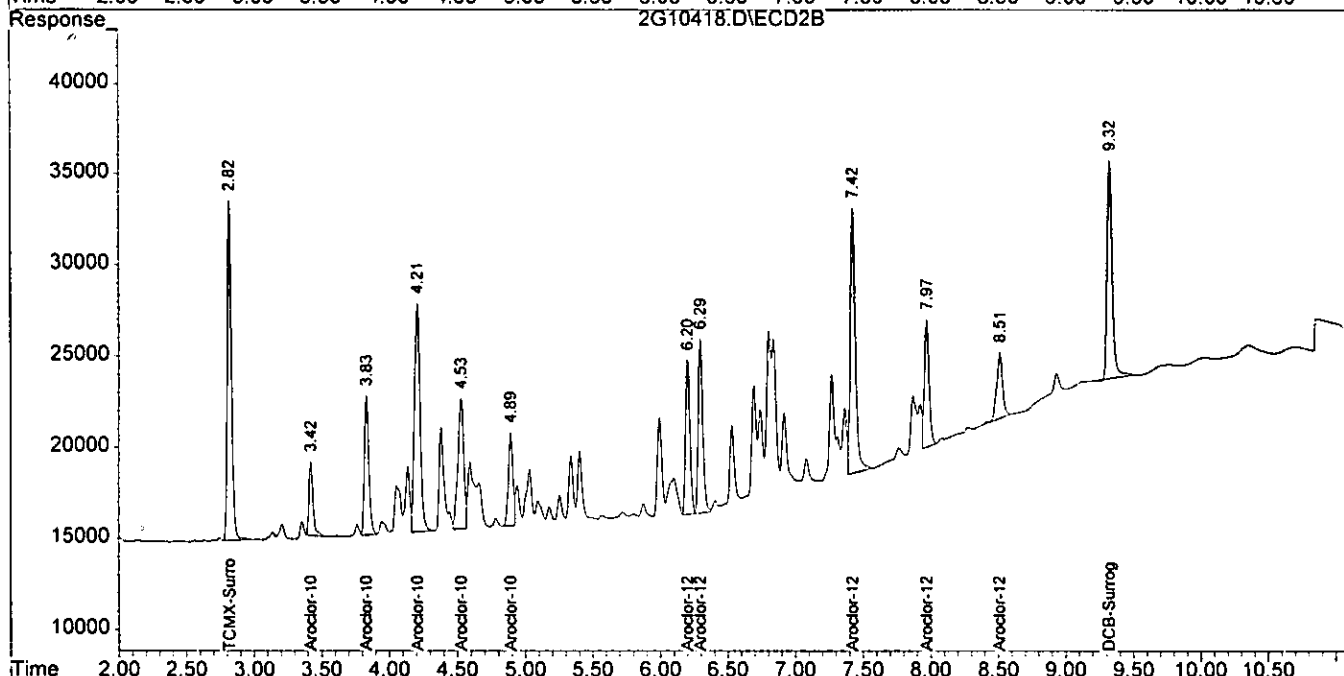
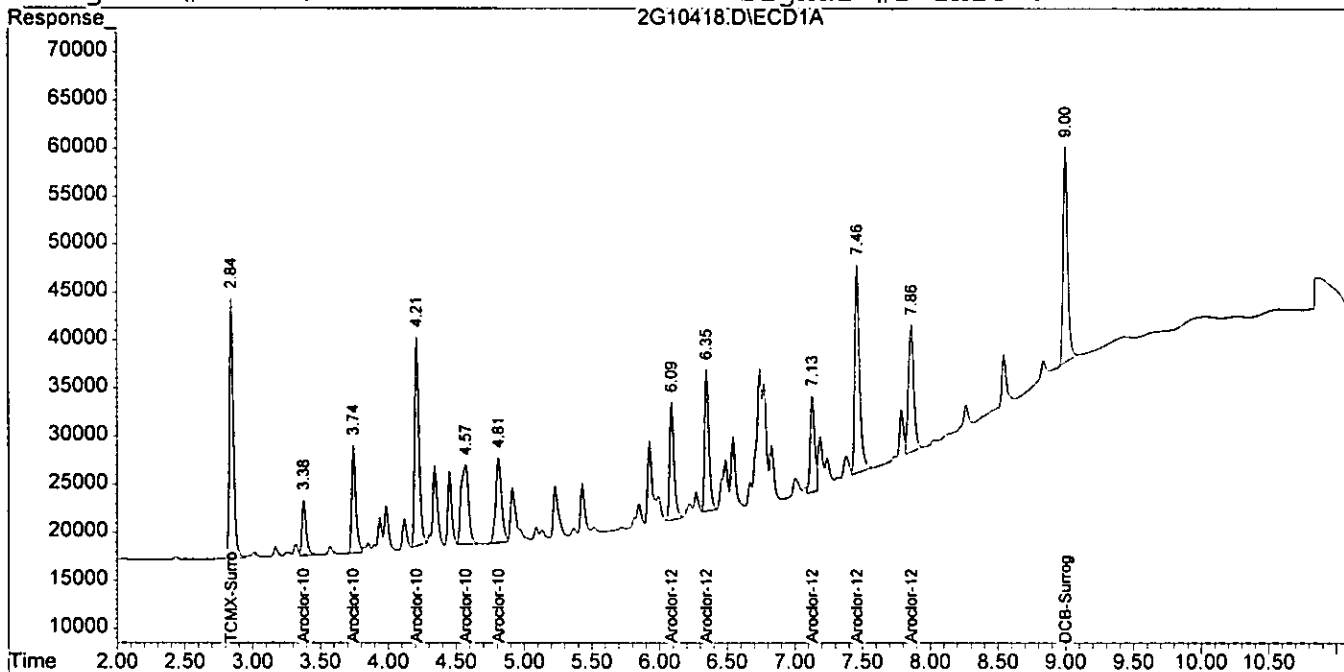
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	537576	389340	29.746	29.041
2) Aroclor-1016 {1}	3.38	3.42	123729	83411	306.324	282.803
3) Aroclor-1016 {2}	3.74	3.83	236510	174330	304.604	272.884
4) Aroclor-1016 {3}	4.21	4.21	480306	362147	278.539m	276.332
5) Aroclor-1016 {4}	4.57	4.53	316177	211512	274.064m	272.136
6) Aroclor-1016 {5}	4.81	4.89	265707	116871	270.448	300.815
7) Aroclor-1260 {1}	6.09	6.20	273566	193876	297.933	284.945
8) Aroclor-1260 {2}	6.35	6.29	322701	221397	284.637	291.254
9) Aroclor-1260 {3}	7.13	7.42	225899	378330	296.573	263.020
10) Aroclor-1260 {4}	7.46	7.97	524146	168121	264.399	239.780
11) Aroclor-1260 {5}	7.86	8.51	361621	112339	261.775	239.890
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	9.00	9.32	515176	313226	25.589	24.161

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10418.D\ECD1A.CH Vial 803
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10418.D\ECD2B.CH
Acq On : 3 Aug 2005 9:04 Operator: JK
Sample : CAL 1660@200PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 10:37 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10416.D\ECD1A.CH Vial 301
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10416.D\ECD2B.CH
 Acq On : 3 Aug 2005 8:26 Operator: JK
 Sample : CAL 1660@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 8:57 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

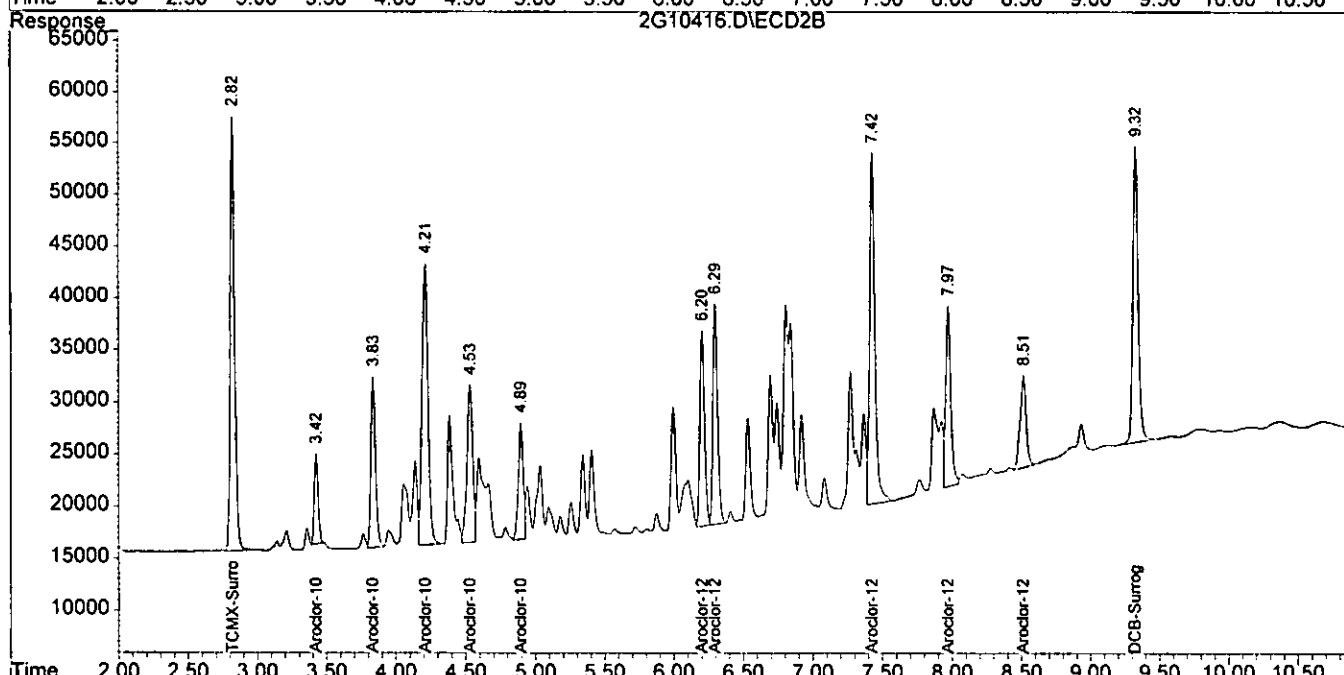
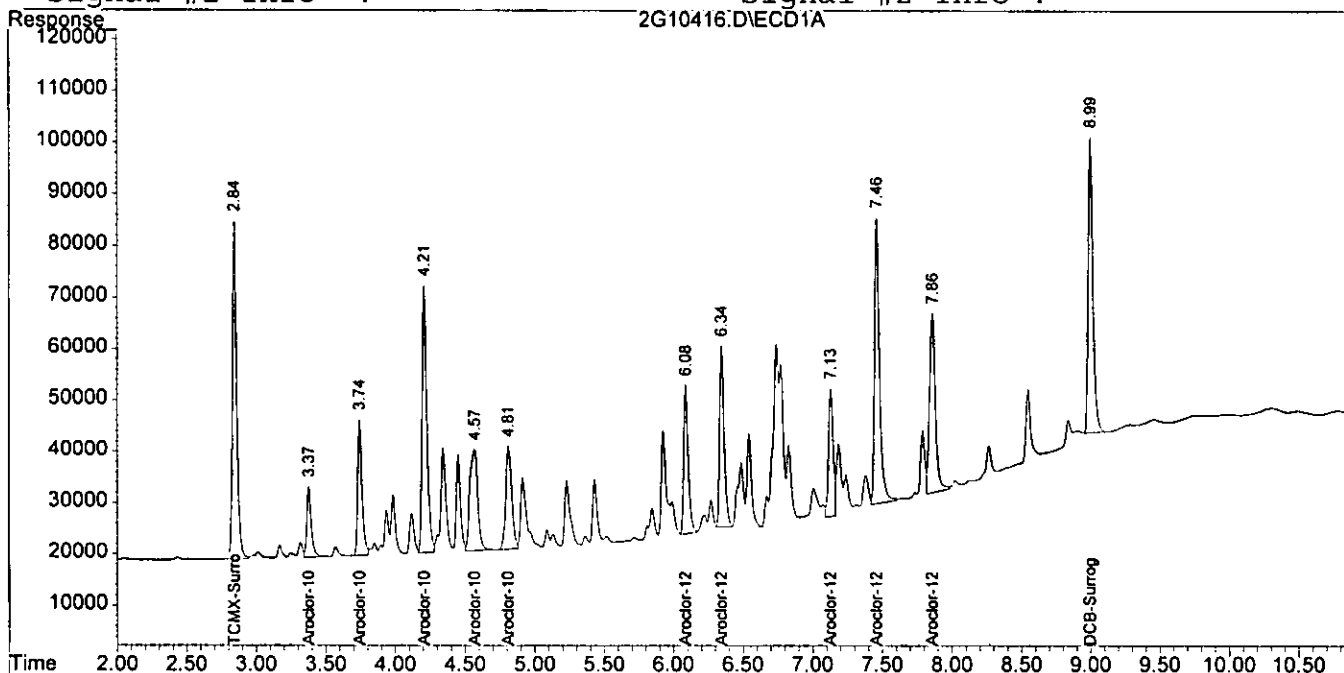
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	1380059	903887	76.365	67.421
2) Aroclor-1016 {1}	3.38	3.42	305904	166893	757.347	603.963
3) Aroclor-1016 {2}	3.74	3.83	569576	384775	733.565	667.333
4) Aroclor-1016 {3}	4.21	4.21	1210114	794926	725.013	672.158
5) Aroclor-1016 {4}	4.57	4.53	782689	434950	703.429	687.861
6) Aroclor-1016 {5}	4.81	4.89	588972	266189	725.391	685.147
7) Aroclor-1260 {1}	6.09	6.20	649718	434260	707.591	638.242
8) Aroclor-1260 {2}	6.34	6.29	783186	486263	690.806	639.693
9) Aroclor-1260 {3}	7.13	7.42	585384	894594	768.527	621.934
10) Aroclor-1260 {4}	7.46	7.97	1377810	440146	695.019	627.755
11) Aroclor-1260 {5}	7.86	8.51	1019825	265305	738.245	566.534
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	1355810	754540	67.343	58.201

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10416.D\ECD1A.CH Vial 111
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10416.D\ECD2B.CH
Acq On : 3 Aug 2005 8:26 Operator: JK
Sample : CAL 1660@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 8:57 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10419.D\ECD1A.CH Vial: 34
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10419.D\ECD2B.CH
 Acq On : 3 Aug 2005 9:19 Operator: JK
 Sample : CAL 1660@1000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 9:29 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

06/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

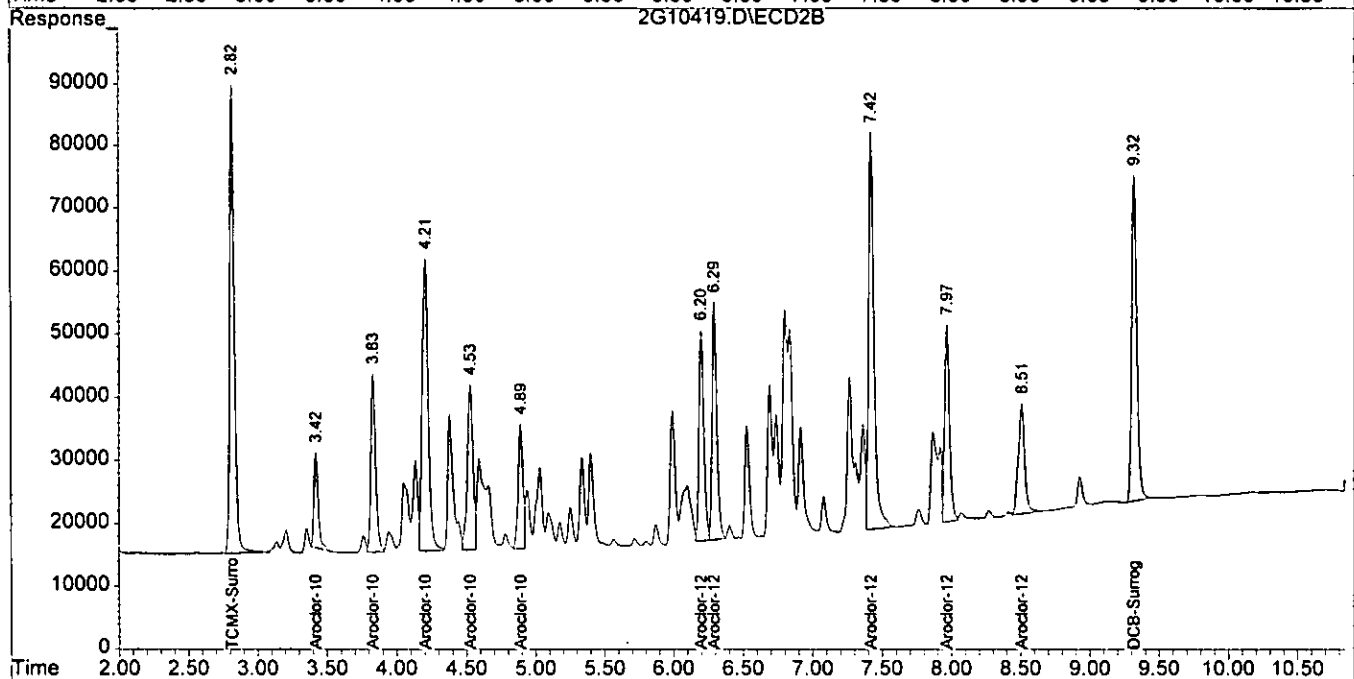
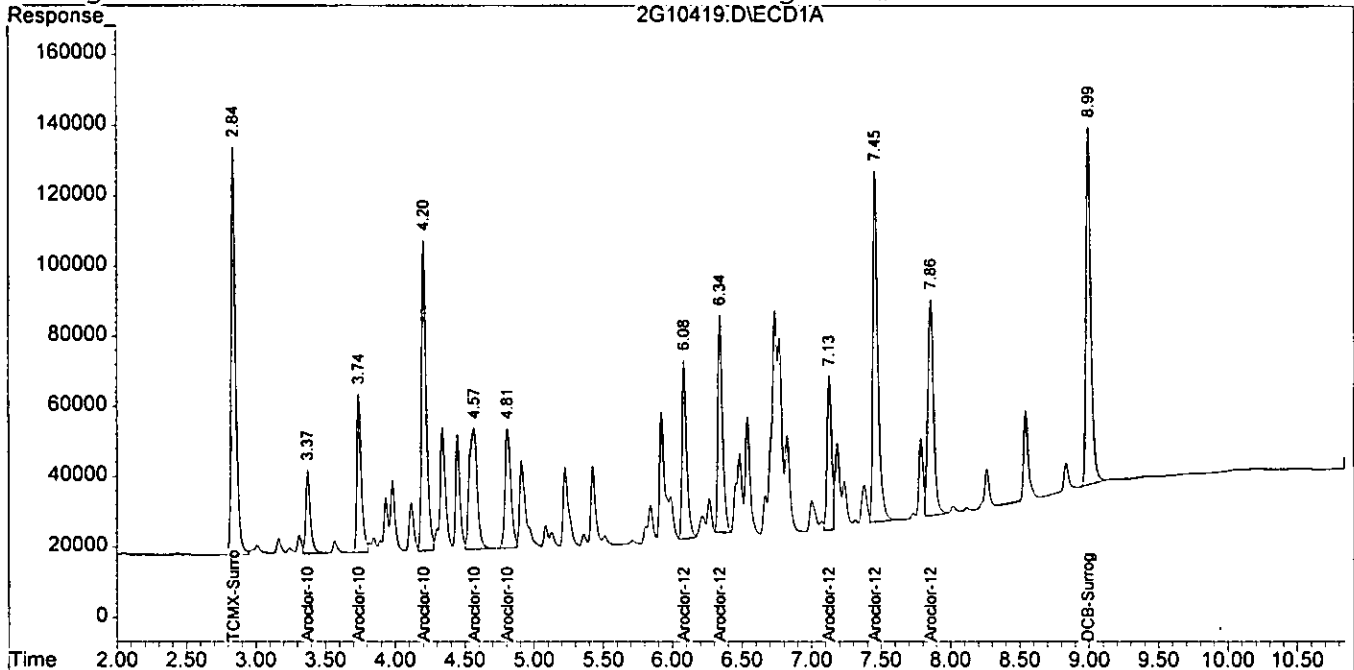
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	2515120	1662672	139.173	124.019
2) Aroclor-1016	3.37	3.42	544132	320805	1347.143	1260.817
3) Aroclor-1016	3.74	3.83	998171	682294	1285.559	1271.382
4) Aroclor-1016	4.20	4.21	2091412	1404106	1307.895	1273.544
5) Aroclor-1016	4.57	4.53	1382670	736881	1299.014	1304.746
6) Aroclor-1016	4.81	4.89	985607	476841	1330.959	1227.347
7) Aroclor-1260	6.08	6.20	1161940	807381	1265.439	1186.626
8) Aroclor-1260	6.34	6.29	1389979	898443	1226.026	1181.928
9) Aroclor-1260	7.13	7.42	1000167	1720174	1313.077	1195.889
10) Aroclor-1260	7.46	7.97	2509685	790730	1265.980	1127.771
11) Aroclor-1260	7.86	8.51	1789044	556465	1295.079	1188.279
12) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	2462508	1337558	122.313	103.172

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10419.D\ECD1A.CH Vial 34
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10419.D\ECD2B.CH
Acq On : 3 Aug 2005 9:19 Operator: JK
Sample : CAL 1660@1000PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 9:29 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10420.D\ECD1A.CH Vial: 115
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10420.D\ECD2B.CH
 Acq On : 3 Aug 2005 9:33 Operator: JK
 Sample : CAL 1660@2000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 9:45 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 09:30:47 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

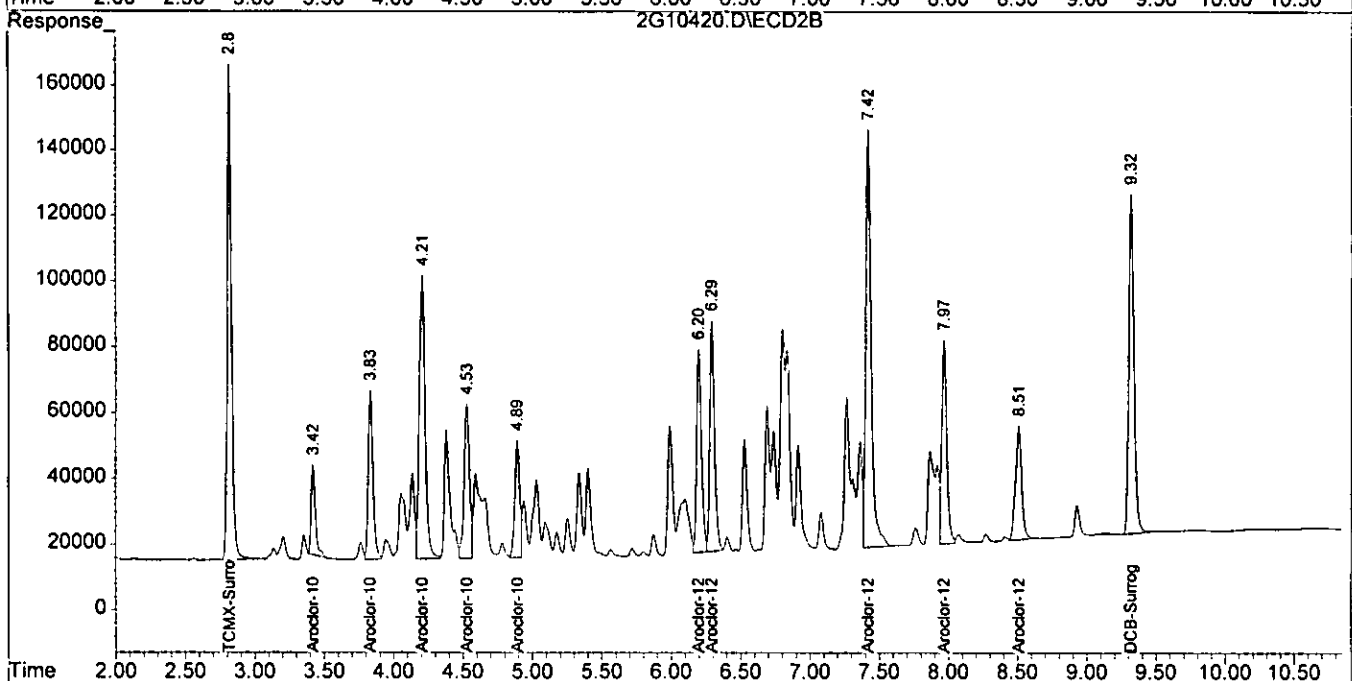
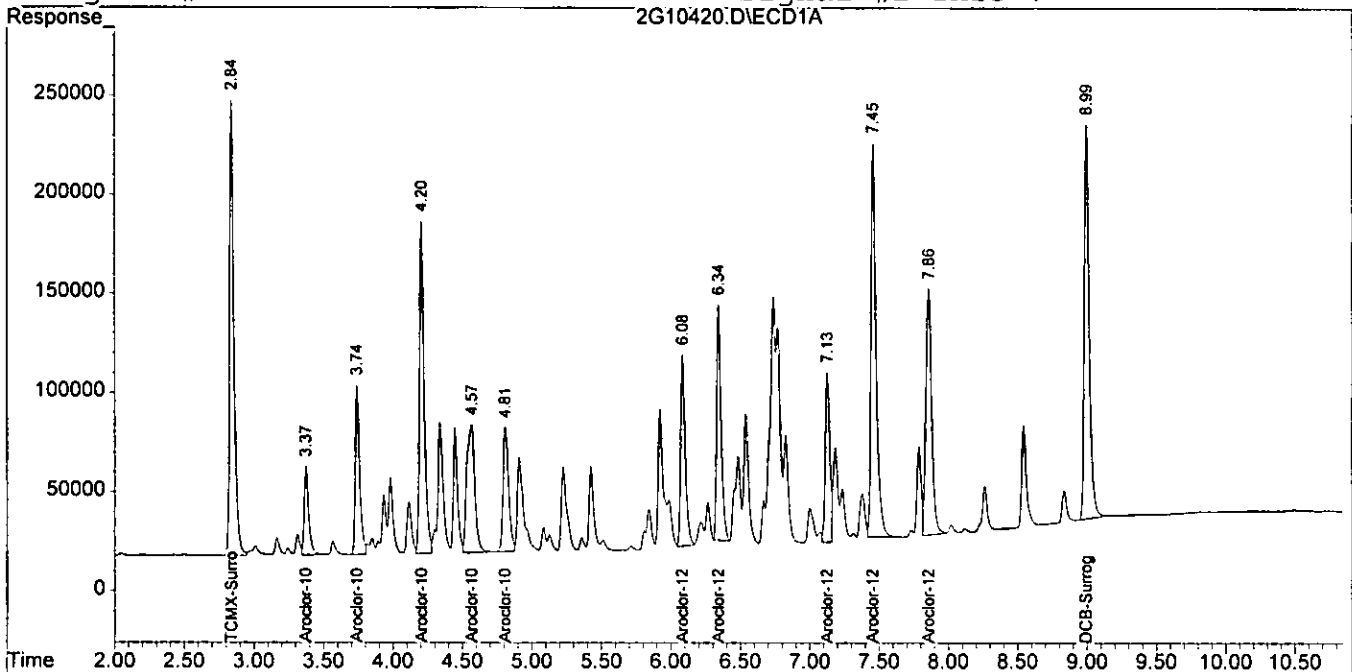
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	4904868	3190992	186.247	175.193
2) Aroclor-1016 {1}	3.37	3.42	1012689	581017	1748.648	1999.965
3) Aroclor-1016 {2}	3.74	3.83	1858956	1220640	1685.875	2003.536
4) Aroclor-1016 {3}	4.20	4.21	3936954	2545585	2005.183	2003.692
5) Aroclor-1016 {4}	4.57	4.53	2588761	1263705	2004.217	2004.283
6) Aroclor-1016 {5}	4.81	4.89	1764540	864162	2005.582	1628.910
7) Aroclor-1260 {1}	6.08	6.20	2182030	1489561	1728.572	1688.170
8) Aroclor-1260 {2}	6.34	6.29	2643029	1660729	1740.682	1678.733
9) Aroclor-1260 {3}	7.13	7.42	1948388	3420744	1829.580	1953.996
10) Aroclor-1260 {4}	7.45	7.97	5035126	1576031	1953.448	1864.392
11) Aroclor-1260 {5}	7.86	8.51	3608977	1082591	1966.405	1982.079
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	4863623	2638664	190.227	182.698

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10420.D\ECD1A.CH Vial 55
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10420.D\ECD2B.CH
Acq On : 3 Aug 2005 9:33 Operator: JK
Sample : CAL 1660@2000PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 9:45 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 09:30:47 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10421.D\ECD1A.CH Vial 116
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10421.D\ECD2B.CH
 Acq On : 3 Aug 2005 9:47 Operator: JK
 Sample : CAL 1660@4000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 10:01 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 09:47:54 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

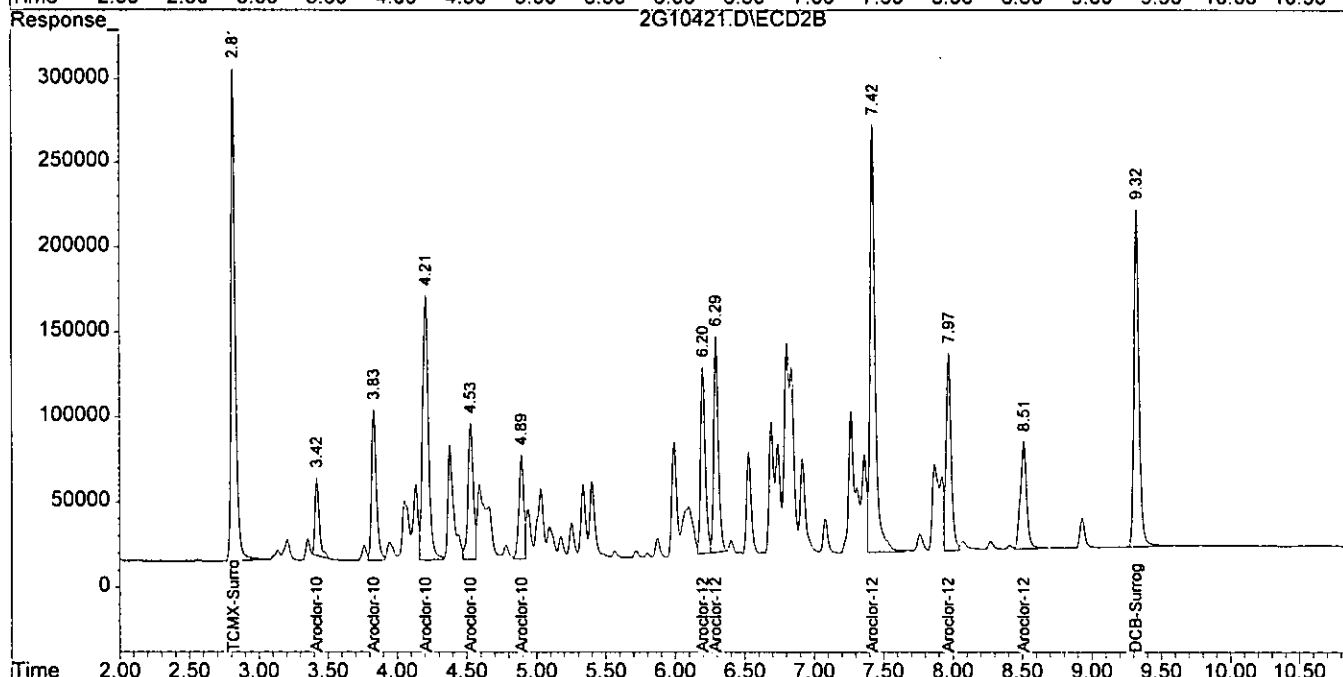
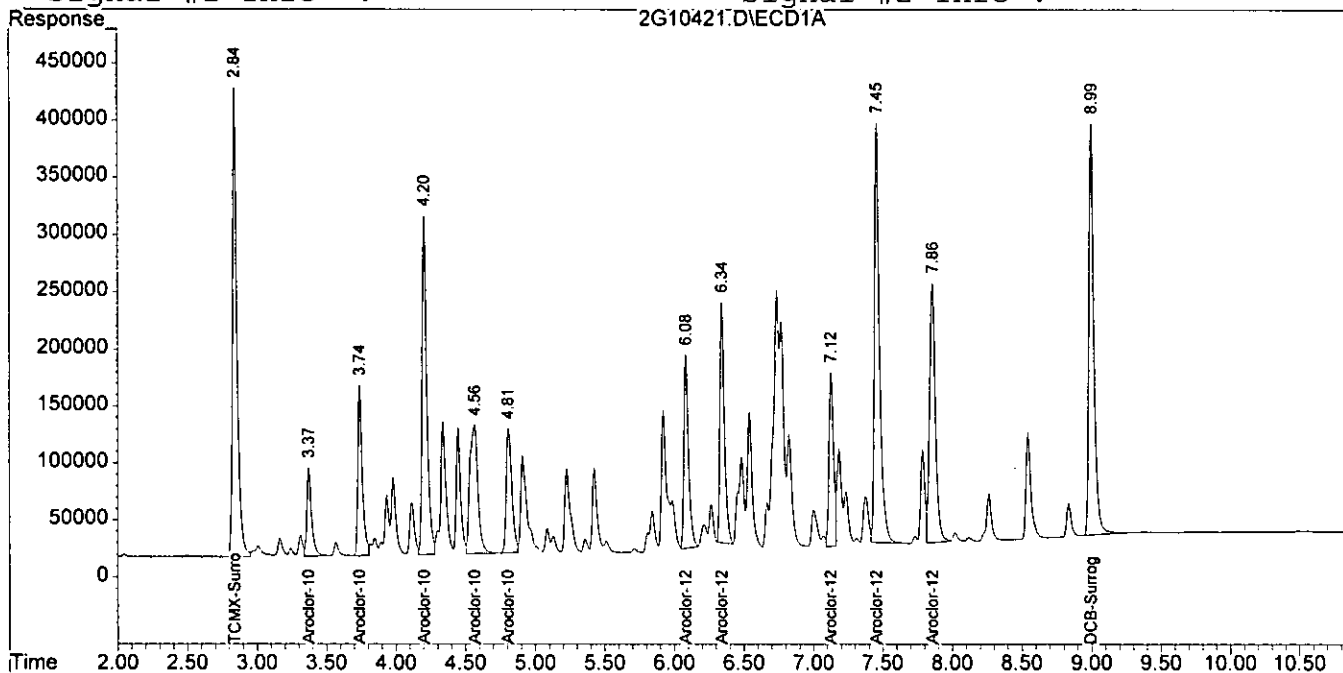
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	9079572	6125454	352.889	345.471
2) Aroclor-1016 {1}	3.37	3.42	1804347	975465	3234.834	4001.341
3) Aroclor-1016 {2}	3.74	3.83	3300014	2126653	3123.865	4007.645 #
4) Aroclor-1016 {3}	4.20	4.21	6971949	4524894	4002.896	4006.197
5) Aroclor-1016 {4}	4.56	4.53	4572495	2142050	4003.234	4011.785
6) Aroclor-1016 {5}	4.81	4.89	3062695	1477217	4007.281	2933.041 #
7) Aroclor-1260 {1}	6.08	6.20	3852270	2592961	3177.252	3074.656
8) Aroclor-1260 {2}	6.34	6.29	4680472	2898134	3205.052	3066.315
9) Aroclor-1260 {3}	7.13	7.42	3486914	6604985	3376.385	3808.940
10) Aroclor-1260 {4}	7.45	7.97	9315337	2925454	3673.089	3540.266
11) Aroclor-1260 {5}	7.86	8.51	6669996	2000849	3690.489	3715.415
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	8917603	5039323	356.392	356.505

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10421.D\ECD1A.CH Vial: 6
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10421.D\ECD2B.CH
Acq On : 3 Aug 2005 9:47 Operator: JK
Sample : CAL 1660@4000PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 10:01 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 09:47:54 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10425.D\ECD1A.CH Vial: 110
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10425.D\ECD2B.CH
 Acq On : 3 Aug 2005 11:04 Operator: JK
 Sample : CAL 1232@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 11:11 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 11:07:35 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/0

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

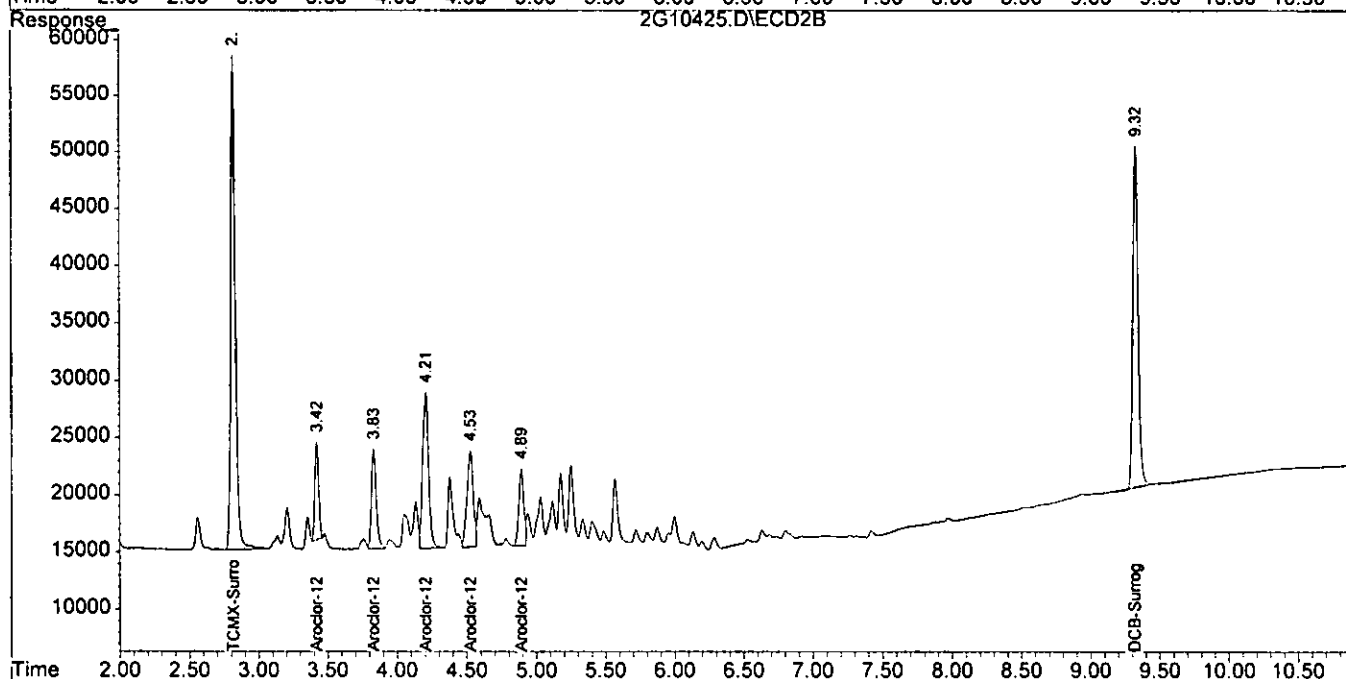
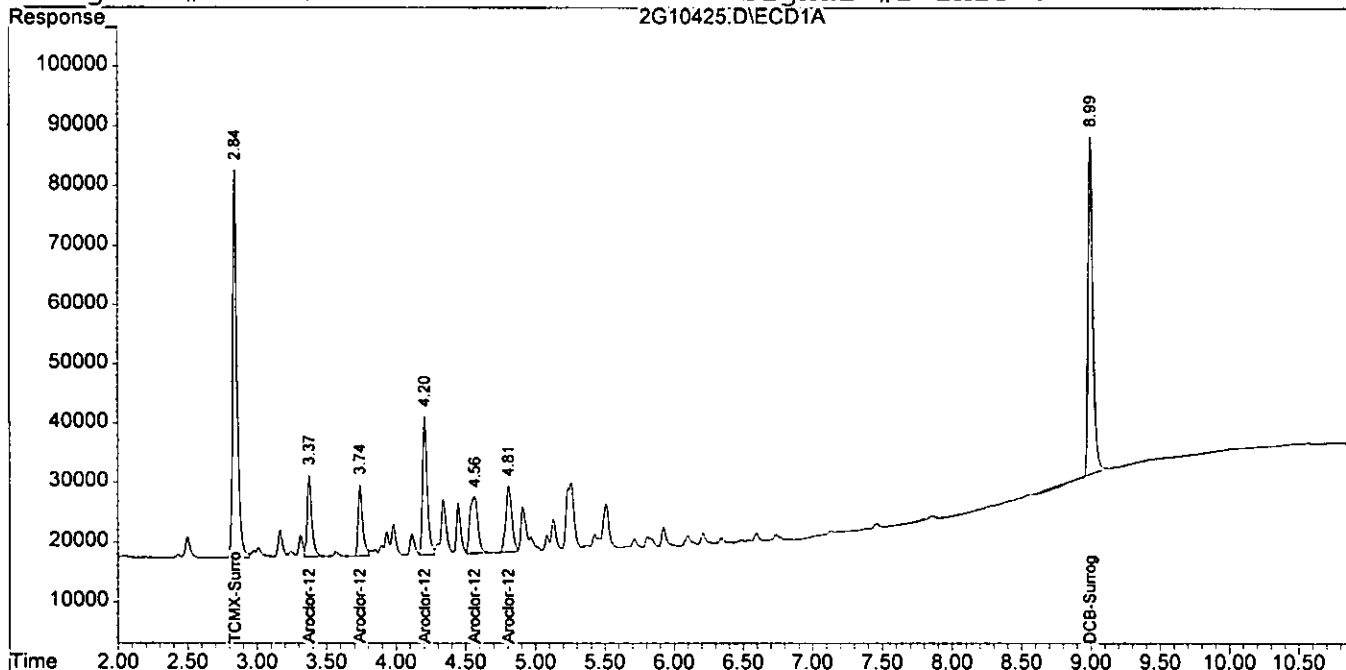
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	1424633	993078	78.831	74.074
2) Aroclor-1016 1	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 2	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 3	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 4	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 5	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 1	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 2	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 3	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 4	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 5	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 1	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 2	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 3	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 1	3.37	3.42	314128	163616	635.245	676.874
16) Aroclor-1232 2	3.74	3.83	266463	209831	616.094	647.477
17) Aroclor-1232 3	4.21	4.21	548326	402513	671.687	668.997
18) Aroclor-1232 4	4.56	4.53	378210	260887	646.242	637.641
19) Aroclor-1232 5	4.81	4.89	339761	161604	628.693	651.000
20) Aroclor-1242 1	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 2	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 3	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 4	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 5	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 1	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 2	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 3	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 4	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 5	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 1	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 2	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 3	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 4	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 5	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	1311329	771047	65.134	59.475

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10425.D\ECD1A.CH Vial 1010
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10425.D\ECD2B.CH
Acq On : 3 Aug 2005 11:04 Operator: JK
Sample : CAL 1232@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 11:11 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 11:07:35 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10424.D\ECD1A.CH Vial: 39
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10424.D\ECD2B.CH
 Acq On : 3 Aug 2005 10:50 Operator: JK
 Sample : CAL 1242@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 11:07 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

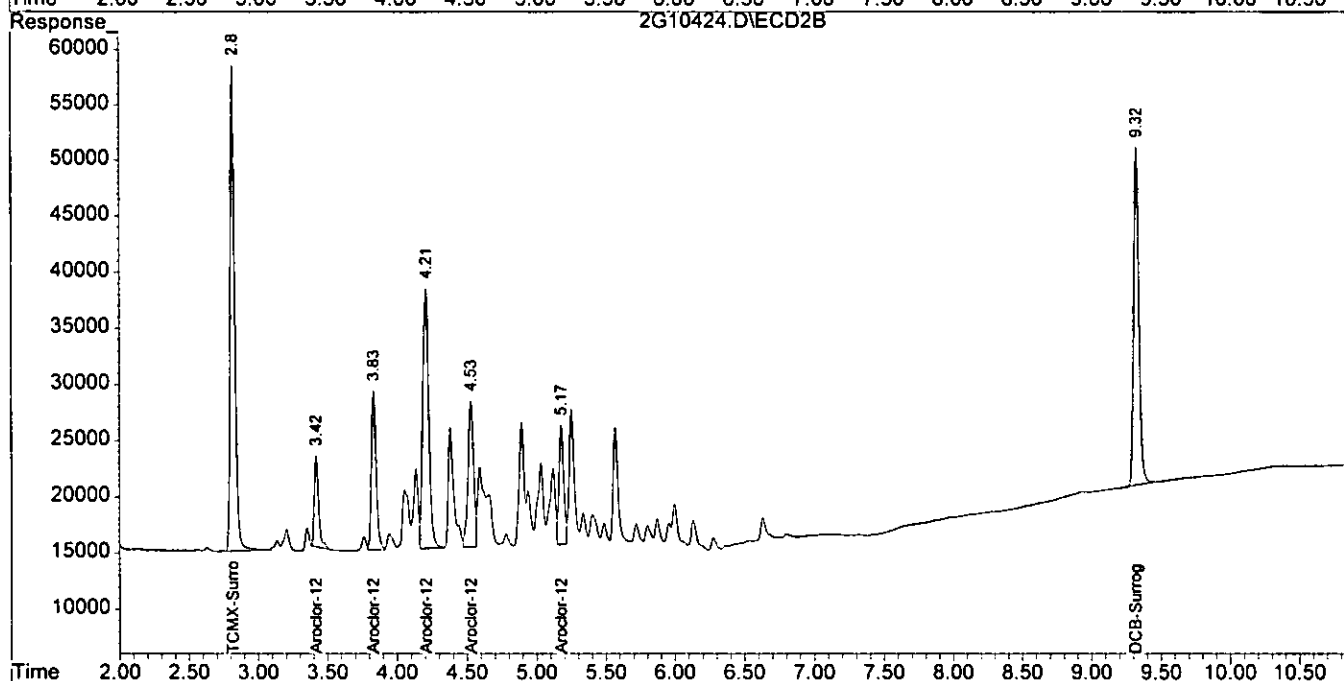
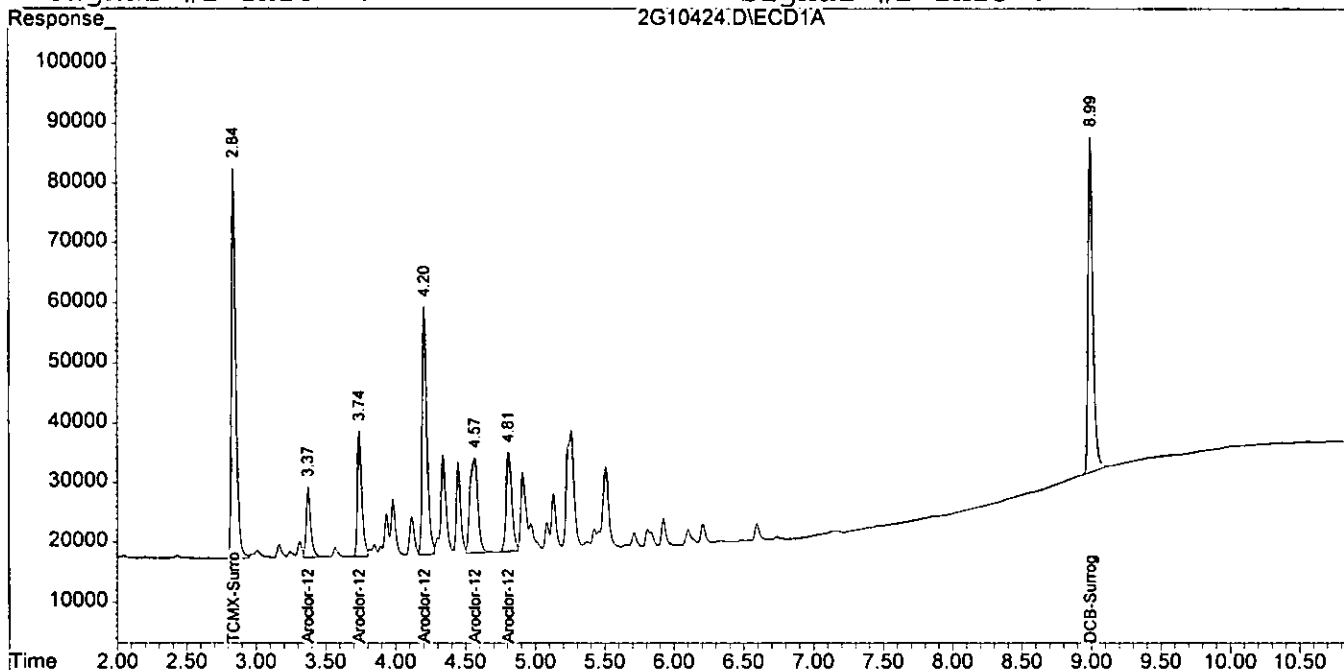
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	1378736	963401	76.292	71.861
2) Aroclor-1016 (1)	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 (2)	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 (3)	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 (4)	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 (5)	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 (1)	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 (2)	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 (3)	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 (4)	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 (5)	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 (1)	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 (2)	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 (3)	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	3.37	3.42	268715	171317	635.842	693.849
21) Aroclor-1242 (2)	3.74	3.83	458682	336058	625.360m	645.505
22) Aroclor-1242 (3)	4.20	4.21	975685	690977	672.443	665.042
23) Aroclor-1242 (4)	4.56	4.53	633568	359402	650.516	615.872
24) Aroclor-1242 (5)	4.81	5.17	473756	249080	614.079	665.124
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 (2)	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 (3)	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 (4)	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 (5)	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	1355222	775395	67.314m	59.810

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10424.D\ECD1A.CH Vial 139
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10424.D\ECD2B.CH
Acq On : 3 Aug 2005 10:50 Operator: JK
Sample : CAL 1242@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 11:07 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10423.D\ECD1A.CH Vial: 13
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10423.D\ECD2B.CH
 Acq On : 3 Aug 2005 10:35 Operator: JK
 Sample : CAL 1248@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 11:05 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

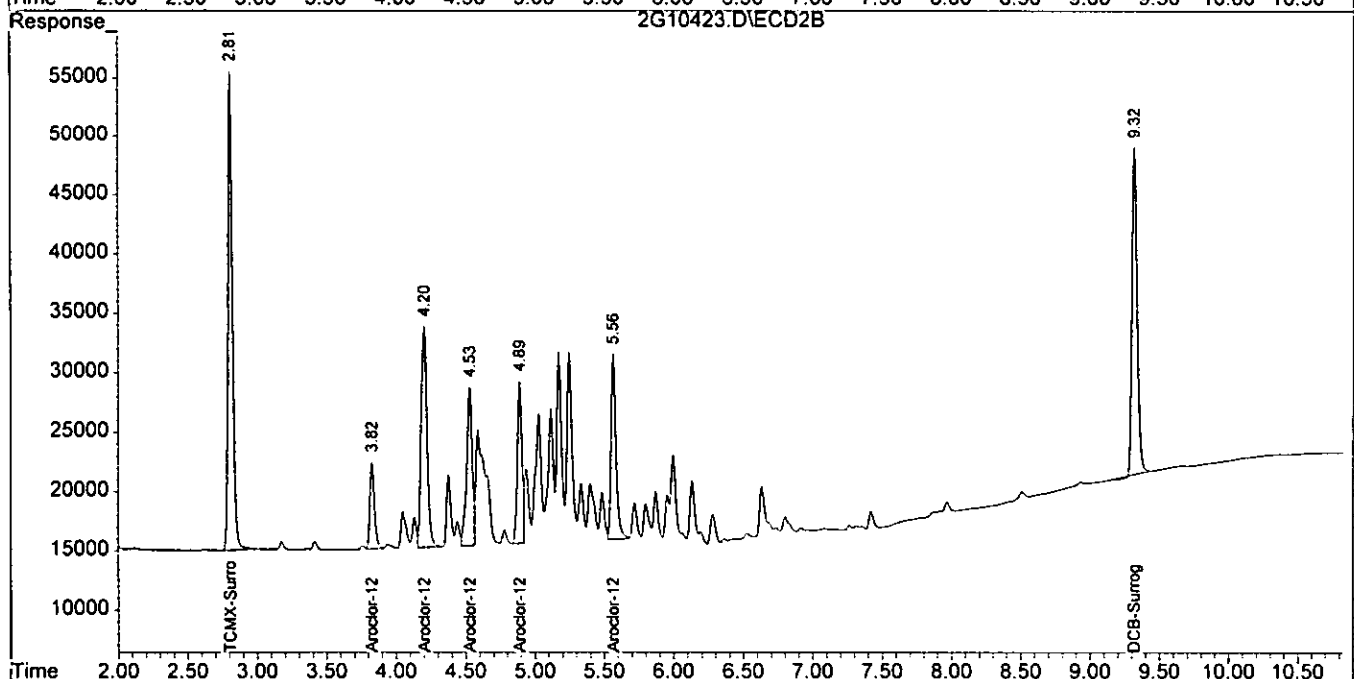
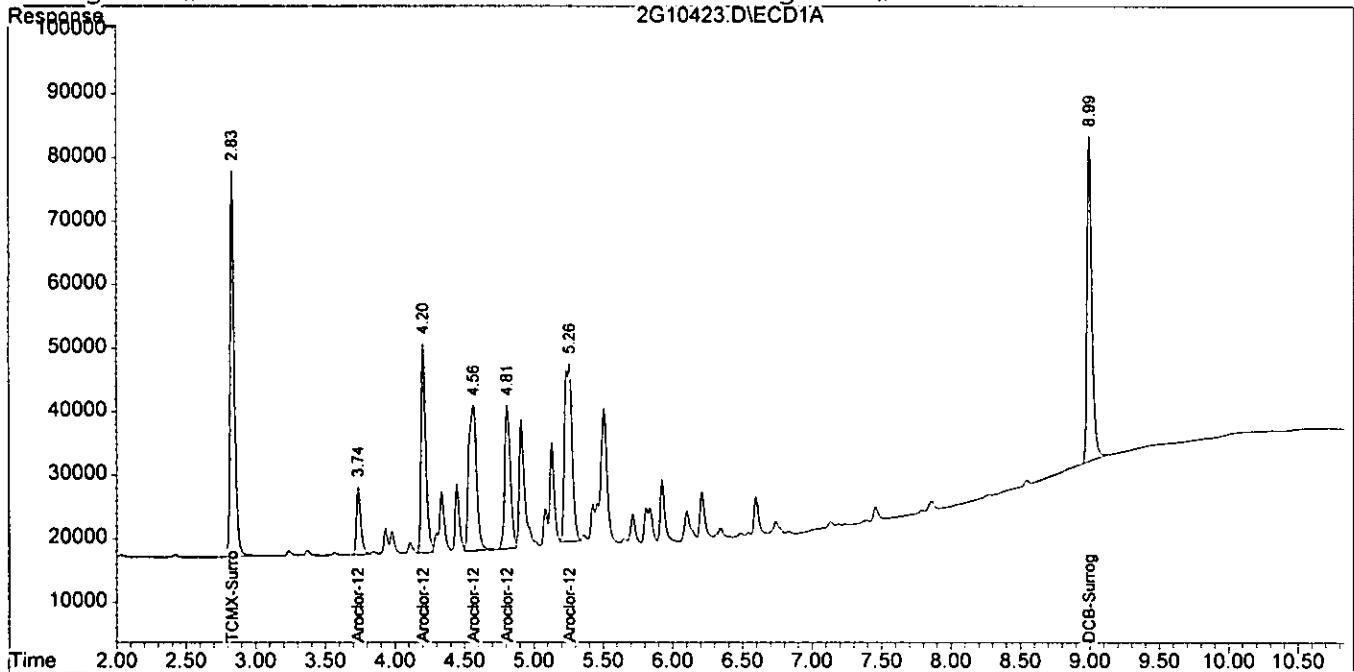
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.83	2.81	1257020	874297	69.557	65.214
2) Aroclor-1016	1 0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016	2 0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016	3 0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016	4 0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016	5 0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260	1 0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260	2 0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260	3 0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260	4 0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260	5 0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221	1 0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221	2 0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221	3 0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232	1 0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232	2 0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232	3 0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232	4 0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232	5 0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242	1 0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242	2 0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242	3 0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242	4 0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242	5 0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248	1 3.74	3.82	229337	166328	604.622	611.519
26) Aroclor-1248	2 4.20	4.20	765969	547895	624.800	621.781
27) Aroclor-1248	3 4.56	4.53	906202	369883	608.513	499.113
28) Aroclor-1248	4 4.81	4.89	639875	313320	511.400	613.906
29) Aroclor-1248	5 5.26	5.56	1044668	388338	617.110m	646.019
30) Aroclor-1254	1 0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254	2 0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254	3 0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254	4 0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254	5 0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.99	9.32	1217054	701709	60.451	54.126

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10423.D\ECD1A.CH Vial: 50
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10423.D\ECD2B.CH
Acq On : 3 Aug 2005 10:35 Operator: JK
Sample : CAL 1248@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 11:05 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info:



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10422.D\ECD1A.CH Vial: 117
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10422.D\ECD2B.CH
 Acq On : 3 Aug 2005 10:20 Operator: JK
 Sample : CAL 2154@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 10:41 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:06:41 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

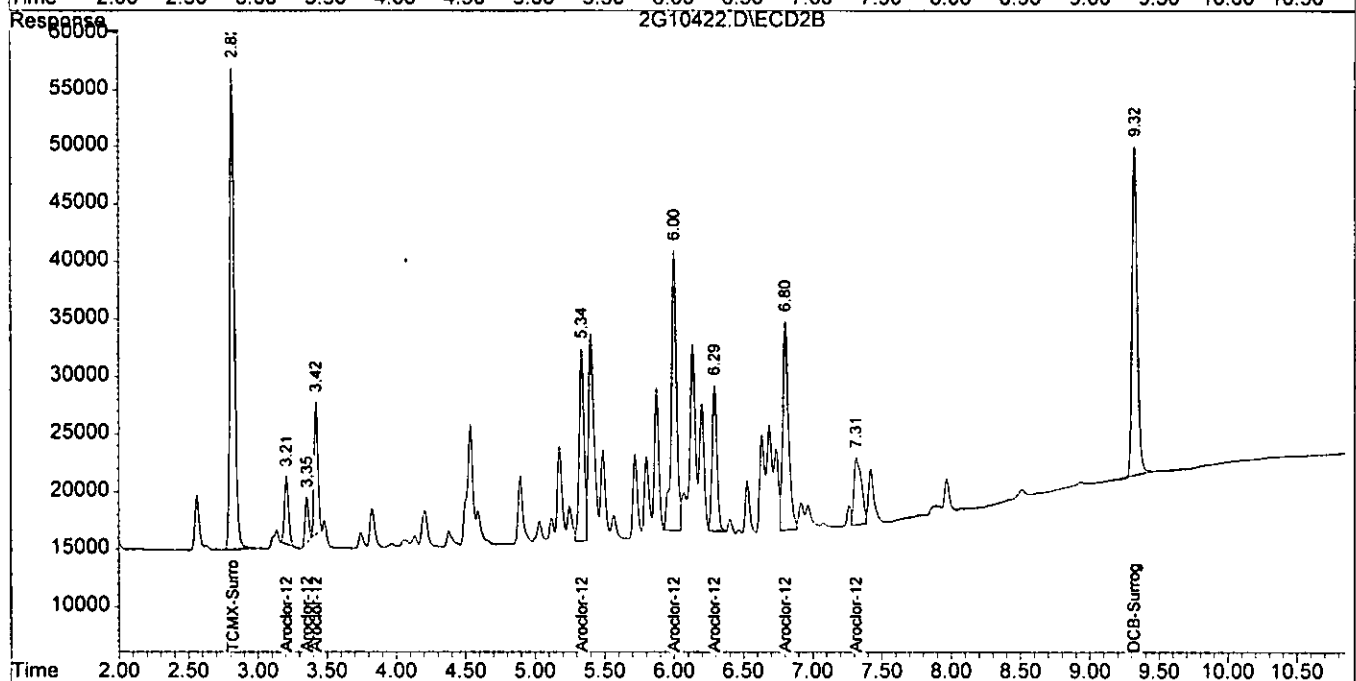
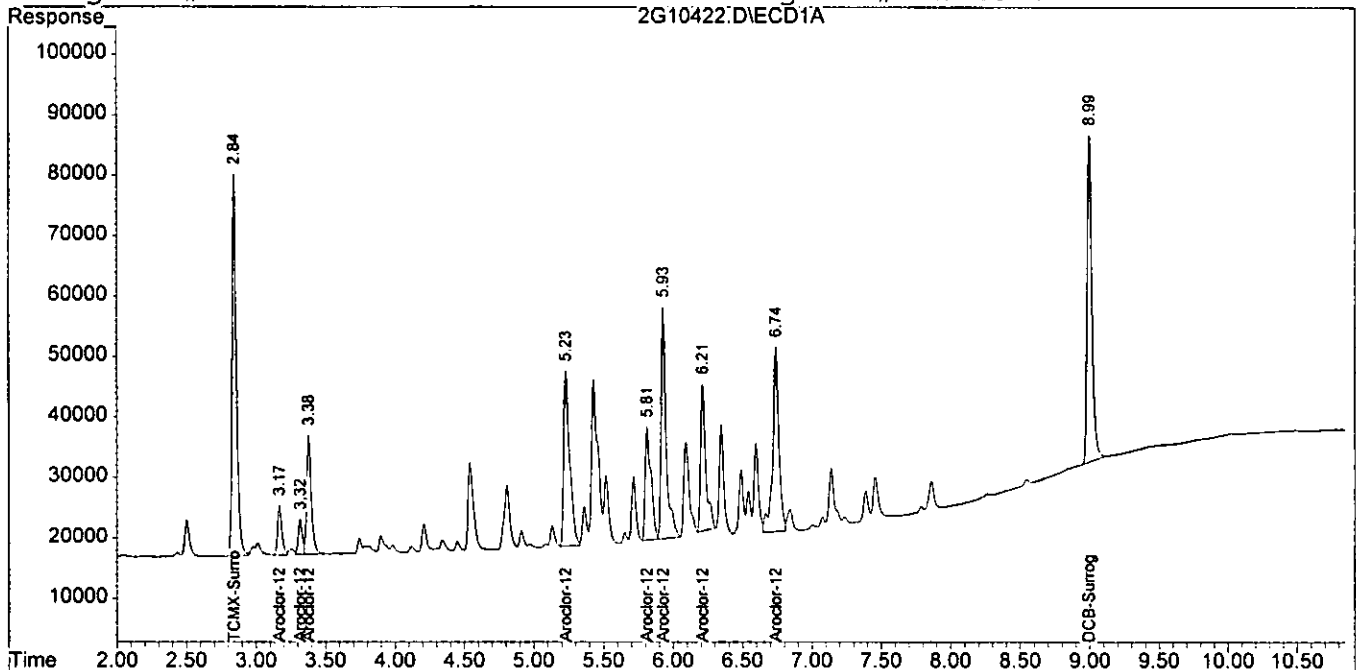
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	1349033	933307	74.648	69.616
2) Aroclor-1016 (1)	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 (2)	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 (3)	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 (4)	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 (5)	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 (1)	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 (2)	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 (3)	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 (4)	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 (5)	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 (1)	3.17	3.21	174516	120549	647.362	686.717
13) Aroclor-1221 (2)	3.32	3.35	116570	69671	663.060	728.528
14) Aroclor-1221 (3)	3.38	3.42	444728	216972	656.103	681.956
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 (2)	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 (3)	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 (4)	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 (5)	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	5.23	5.34	866409	398342	653.495	654.635
31) Aroclor-1254 (2)	5.81	6.00	548467	633765	639.625	656.357
32) Aroclor-1254 (3)	5.93	6.29	952615	307204	631.222	615.415
33) Aroclor-1254 (4)	6.21	6.80	632981	522429	621.569	598.055
34) Aroclor-1254 (5)	6.74	7.32	936339	222126	644.184	603.584
35) DCB-Surrogate	9.00	9.32	1286543	732068	63.903	56.468

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10422.D\ECD1A.CH Vial: 17
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10422.D\ECD2B.CH
Acq On : 3 Aug 2005 10:20 Operator: JK
Sample : CAL 2154@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 10:41 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:06:41 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form 6

Initial Calibration

Instrument: GC_2

Level #:	Data File:	Cal Identifier:	Analysis Date/Time	Level #:	Data File:	Cal Identifier:	Analysis Date/Time	Calibration Level Concentrations																
								Lvl1	Lvl2	Lvl3	Lvl4	Lvl5	Lvl6	Lvl7	Lvl8									
1	2G10503	CAL 1660@50PPB	08/05/05 02:48	2	2G10504	CAL 1660@200PPB	08/05/05 03:02	50.00	200.0	500.0	1000.	2000.	4000.											
3	2G10505	CAL 1660@500PPB	08/05/05 03:17	4	2G10506	CAL 1660@1000PPB	08/05/05 03:31	50.00	200.0	500.0	1000.	2000.	4000.											
5	2G10507	CAL 1660@2000PPB	08/05/05 03:46	6	2G10508	CAL 1660@4000PPB	08/05/05 04:00	50.00	200.0	500.0	1000.	2000.	4000.											
7	2G10512	CAL 1232@500PPB	08/05/05 04:58	8	2G10511	CAL 1242@500PPB	08/05/05 04:43	50.00	200.0	500.0	1000.	2000.	4000.											
9	2G10510	CAL 1248@500PPB	08/05/05 04:29	10	2G10509	CAL 2154@500PPB	08/05/05 04:15	50.00	200.0	500.0	1000.	2000.	4000.											
Compound	Col	Mr	Fit	RF1	RF2	RF3	RF4	RF5	RF6	RF7	RF8	AvgRf	RT	Corr1	Corr2	%Rsd	Lvl1	Lvl2	Lvl3	Lvl4	Lvl5	Lvl6	Lvl7	Lvl8
Aroclor-1016	2	2	Avg	0.0756	0.0727	0.0669	0.0612	0.0526	0.0489	---	---	0.0627	3.80	0.994	0.999	18	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1016	2	3	Avg	0.1554	0.1510	0.1372	0.1268	0.1099	0.1009	---	---	0.130	4.18	0.996	0.999	17	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1016	2	4	Qua	0.1097	0.0888	0.0749	0.0655	0.0541	0.0481	---	---	0.0735	4.50	0.993	0.998	31	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1016	2	5	Avg	0.0485	0.0501	0.0455	0.0416	0.0353	0.0316	---	---	0.0421	4.86	0.993	0.999	18	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1260	2	1	Avg	0.0879	0.0884	0.0832	0.0768	0.0661	0.0622	---	---	0.0775	6.17	0.997	0.999	14	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1260	2	2	Avg	0.0986	0.0973	0.0923	0.0850	0.0736	0.0735	---	---	0.0867	6.26	0.998	0.999	13	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1260	2	3	Avg	0.1549	0.1768	0.1799	0.1790	0.1649	0.1601	---	---	0.169	7.39	0.999	1.00	6.3	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1260	2	4	Avg	0.0826	0.0898	0.0924	0.0867	0.0776	0.0732	---	---	0.0838	7.94	0.998	0.999	8.8	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1260	2	5	Avg	0.0489	0.0533	0.0504	0.0621	0.0583	0.0518	---	---	0.0558	8.48	0.995	1.00	9.4	50.00	200.0	500.0	1000.	2000.	4000.		
Aroclor-1221	2	1	Avg	---	---	---	---	---	---	---	---	0.0196	3.18	-1	-1	Lvl=10	500.0							
Aroclor-1221	2	2	Avg	---	---	---	---	---	---	---	---	0.0146	3.33	-1	-1	Lvl=10	500.0							
Aroclor-1221	2	3	Avg	---	---	---	---	---	---	---	---	0.0461	3.39	-1	-1	Lvl=10	500.0							
Aroclor-1232	2	1	Avg	---	---	---	---	---	---	---	---	0.0317	3.39	-1	-1	Lvl=7	500.0							
Aroclor-1232	2	2	Avg	---	---	---	---	---	---	---	---	0.0374	3.81	-1	-1	Lvl=7	500.0							
Aroclor-1232	2	3	Avg	---	---	---	---	---	---	---	---	0.0716	4.18	-1	-1	Lvl=7	500.0							
Aroclor-1232	2	4	Avg	---	---	---	---	---	---	---	---	0.0470	4.50	-1	-1	Lvl=7	500.0							
Aroclor-1232	2	5	Avg	---	---	---	---	---	---	---	---	0.0295	4.86	-1	-1	Lvl=7	500.0							
Aroclor-1242	2	1	Avg	---	---	---	---	---	---	---	---	0.0286	3.40	-1	-1	Lvl=8	500.0							
Aroclor-1242	2	2	Avg	---	---	---	---	---	---	---	---	0.0584	3.81	-1	-1	Lvl=8	500.0							
Aroclor-1242	2	3	Avg	---	---	---	---	---	---	---	---	0.121	4.18	-1	-1	Lvl=8	500.0							
Aroclor-1242	2	4	Avg	---	---	---	---	---	---	---	---	0.0664	4.50	-1	-1	Lvl=8	500.0							
Aroclor-1242	2	5	Avg	---	---	---	---	---	---	---	---	0.0536	5.22	-1	-1	Lvl=8	500.0							
Aroclor-1248	2	1	Avg	---	---	---	---	---	---	---	---	0.0288	3.81	-1	-1	Lvl=9	500.0							
Aroclor-1248	2	2	Avg	---	---	---	---	---	---	---	---	0.0947	4.18	-1	-1	Lvl=9	500.0							
Aroclor-1248	2	3	Avg	---	---	---	---	---	---	---	---	0.0671	4.51	-1	-1	Lvl=9	500.0							
Aroclor-1248	2	4	Avg	---	---	---	---	---	---	---	---	0.0529	4.87	-1	-1	Lvl=9	500.0							
Aroclor-1248	2	5	Avg	---	---	---	---	---	---	---	---	0.0679	5.54	-1	-1	Lvl=9	500.0							
Aroclor-1254	2	1	Avg	---	---	---	---	---	---	---	---	0.0697	5.31	-1	-1	Lvl=10	500.0							
Aroclor-1254	2	2	Avg	---	---	---	---	---	---	---	---	0.112	5.97	-1	-1	Lvl=10	500.0							
Aroclor-1254	2	3	Avg	---	---	---	---	---	---	---	---	0.0565	6.26	-1	-1	Lvl=10	500.0							
Aroclor-1254	2	4	Avg	---	---	---	---	---	---	---	---	0.100	6.77	-1	-1	Lvl=10	500.0							
Aroclor-1254	2	5	Avg	---	---	---	---	---	---	---	---	0.0424	7.29	-1	-1	Lvl=10	500.0							
DCB-Surrogate	2	0	Avg	1.6701	1.6611	1.5902	1.4994	1.3495	1.2895	---	---	1.51	9.28	0.998	0.999	11	5.00	20.00	50.00	100.0	200.0	400.0		

Avg Rsd Col 1: 10.7 Avg Rsd Col 2: 13.9

Flags
c - failed the initial calibration criteria(if applicable)

Note:
Col = Column Number
Mir = MultiPeak Analyte 0=single peak analyte, >0=multi peak analyte (i.e. pcb/chlordane etc.)
Fit = Indicates whether Avg RF, Linear, or Quadratic Curve was used for compound.
Corr 1 = Correlation Coefficient for linear Eq.
Corr 2 = Correlation Coefficient for quad Eq.
^Lvl: These compounds use a single pt calibration as specified by the method. The file used to update this calibration point is listed in the header under level #.

All Response Factors = Response Factors / 10000
Initial Calibration Criteria: either %RSD <=20 or Corr >= .995
Columns: Signal #1 db-1701 : Signal #2 db-608

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10503.D\ECD1A.CH Vial: 113
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10503.D\ECD2B.CH
 Acq On : 5 Aug 2005 2:48 Operator: JK
 Sample : CAL 1660@50PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

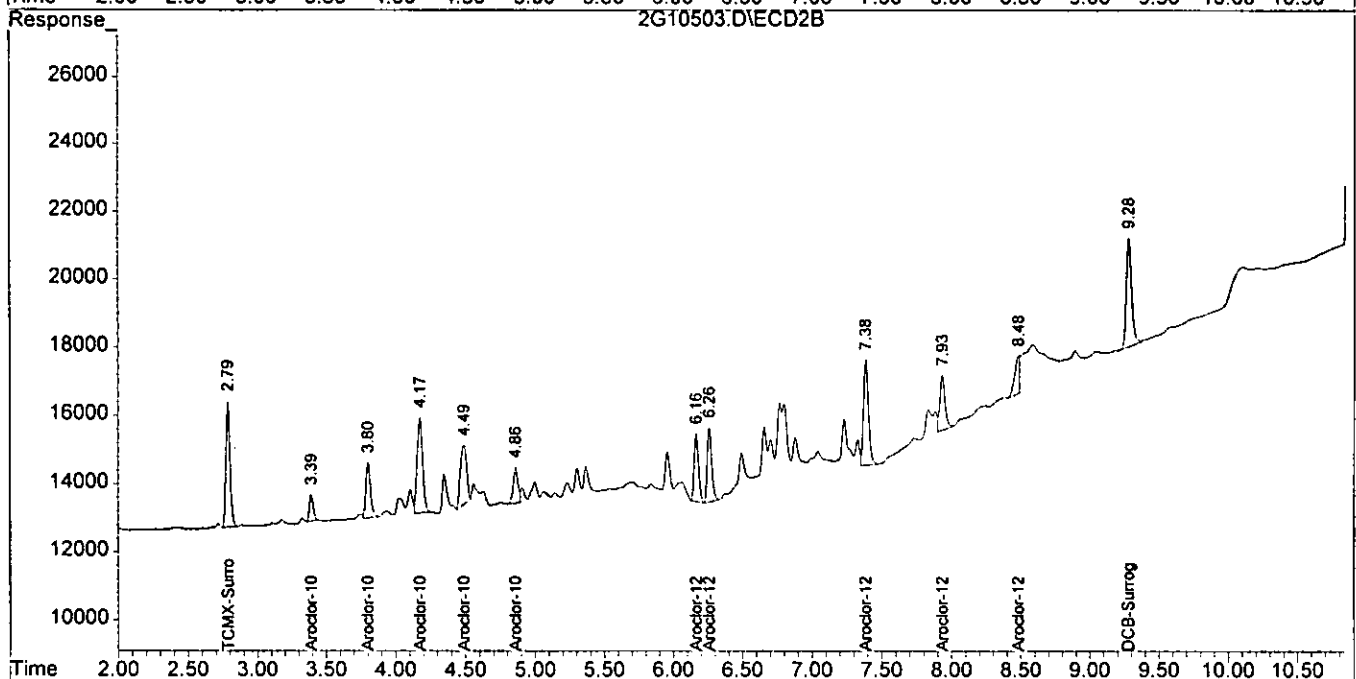
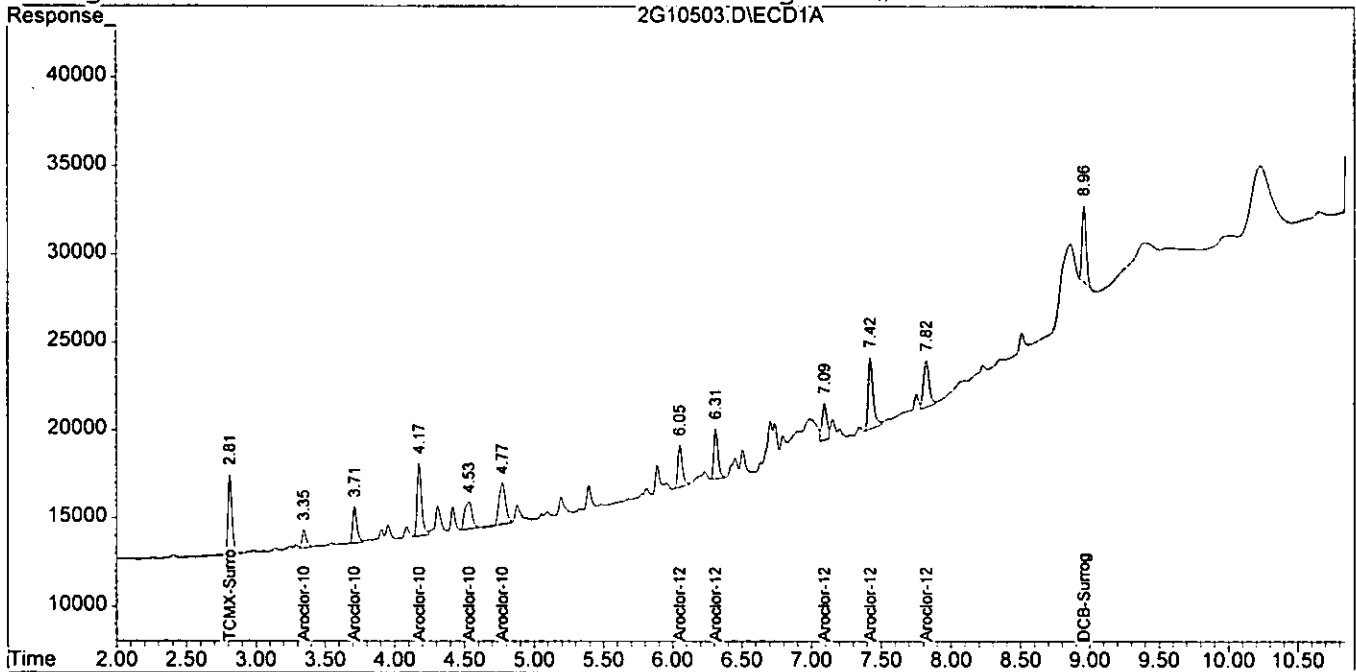
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	88466	75385	4.895	5.623
2) Aroclor-1016	3.35	3.39	21539	15232	53.325	59.289
3) Aroclor-1016	3.71	3.80	42548	37813	54.799	66.430m
4) Aroclor-1016	4.18	4.17	92812	77737	58.609	65.785m
5) Aroclor-1016	4.53	4.49	56400	54868	54.057	78.172m#
6) Aroclor-1016	4.77	4.86	76224	24263	82.529	62.450m
7) Aroclor-1260	6.05	6.16	54373	43985	59.216	64.646
8) Aroclor-1260	6.31	6.26	61214	49300	53.994	64.855
9) Aroclor-1260	7.09	7.38	49747	77459	65.311m	53.851m
10) Aroclor-1260	7.42	7.93	97932	41326	49.401	58.940m
11) Aroclor-1260	7.82	8.48	76258	24467	55.202	52.246m
12) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	89242	83509	4.433	6.441m#

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10503.D\ECD1A.CH Vial: 1
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10503.D\ECD2B.CH
Acq On : 5 Aug 2005 2:48 Operator: JK
Sample : CAL 1660@50PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 7:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10504.D\ECD1A.CH Vial: 32
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10504.D\ECD2B.CH
 Acq On : 5 Aug 2005 3:02 Operator: JK
 Sample : CAL 1660@200PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:29 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

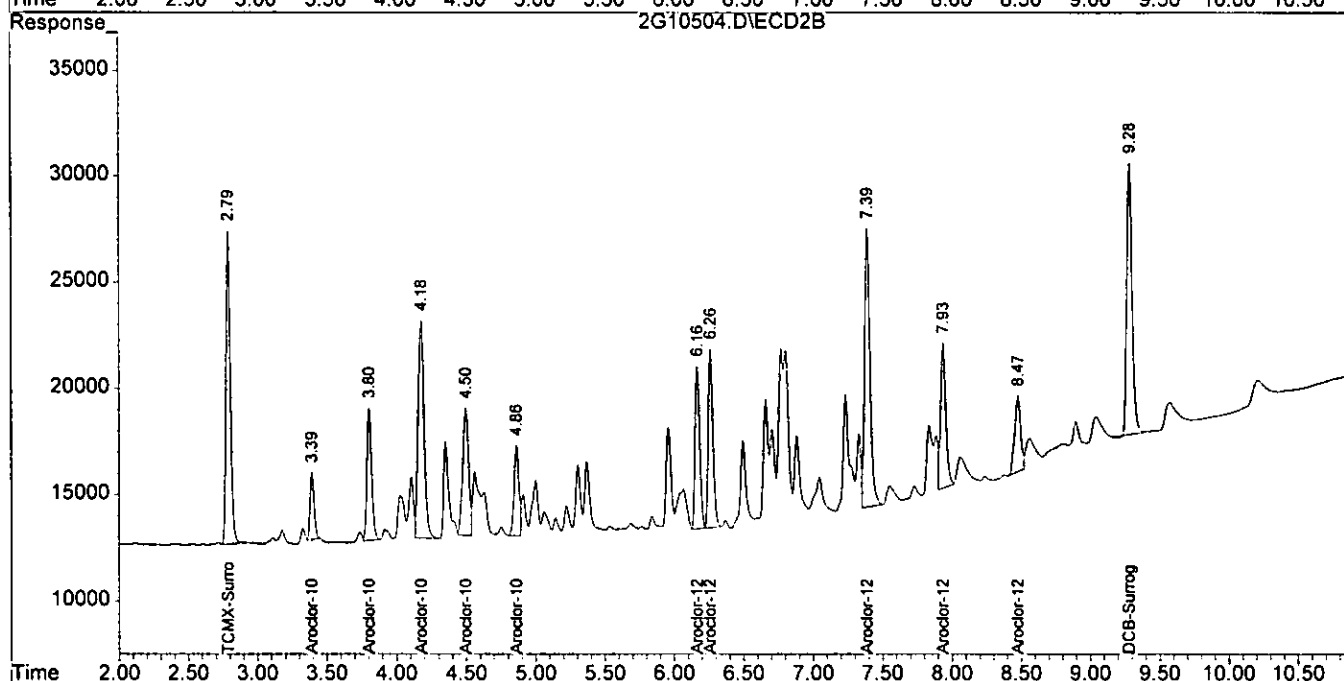
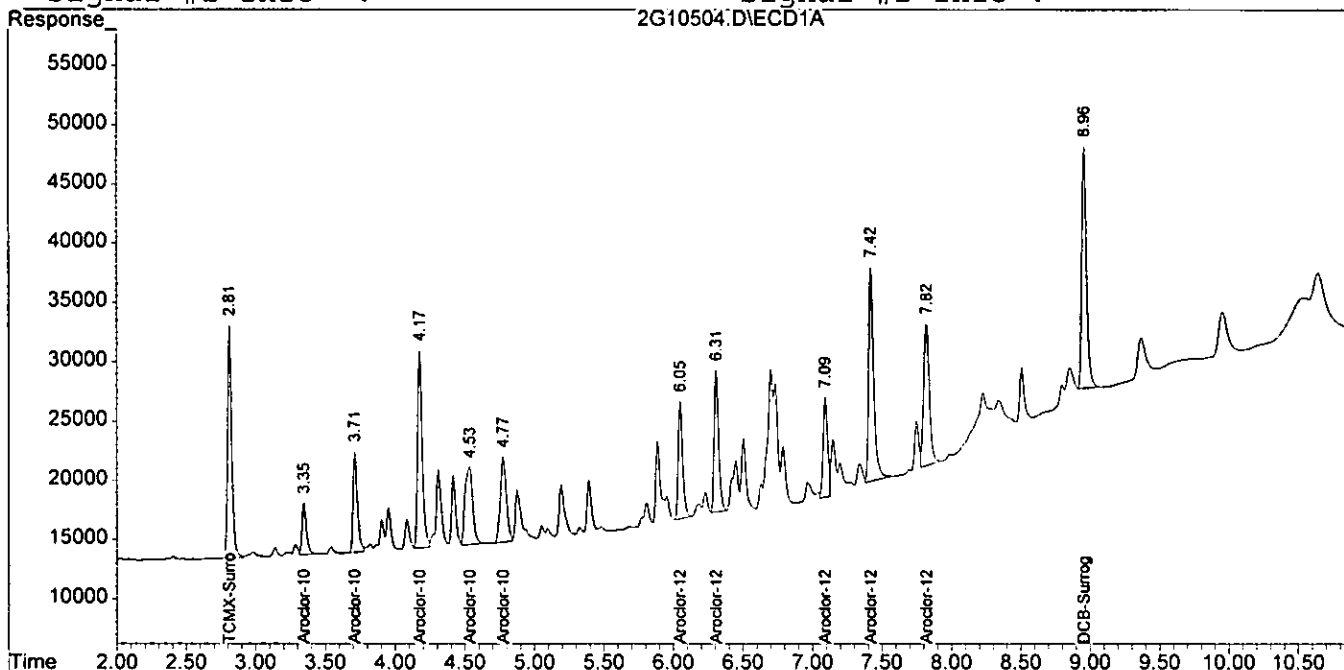
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	400628	316064	22.169	23.575
2) Aroclor-1016 {1}	3.35	3.39	95231	63077	235.769	245.521m
3) Aroclor-1016 {2}	3.71	3.80	185634	145451	239.081	255.527m
4) Aroclor-1016 {3}	4.17	4.18	386473	302110	244.049	255.659m
5) Aroclor-1016 {4}	4.53	4.50	253932	177599	243.385	253.029m
6) Aroclor-1016 {5}	4.77	4.86	219239	100330	237.375	258.241m
7) Aroclor-1260 {1}	6.05	6.16	221856	176861	241.618	259.937
8) Aroclor-1260 {2}	6.31	6.26	268892	194671	237.175	256.096
9) Aroclor-1260 {3}	7.09	7.39	193759	353615	254.378	245.838
10) Aroclor-1260 {4}	7.42	7.93	454220	179747	229.126	256.363m
11) Aroclor-1260 {5}	7.82	8.47	344436	106662	249.335	227.766
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	478104	332233	23.747	25.627

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10504.D\ECD1A.CH Vial 152
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10504.D\ECD2B.CH
Acq On : 5 Aug 2005 3:02 Operator: JK
Sample : CAL 1660@200PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 7:29 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10505.D\ECD1A.CH Vial 133
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10505.D\ECD2B.CH
 Acq On : 5 Aug 2005 3:17 Operator: JK
 Sample : CAL 1660@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

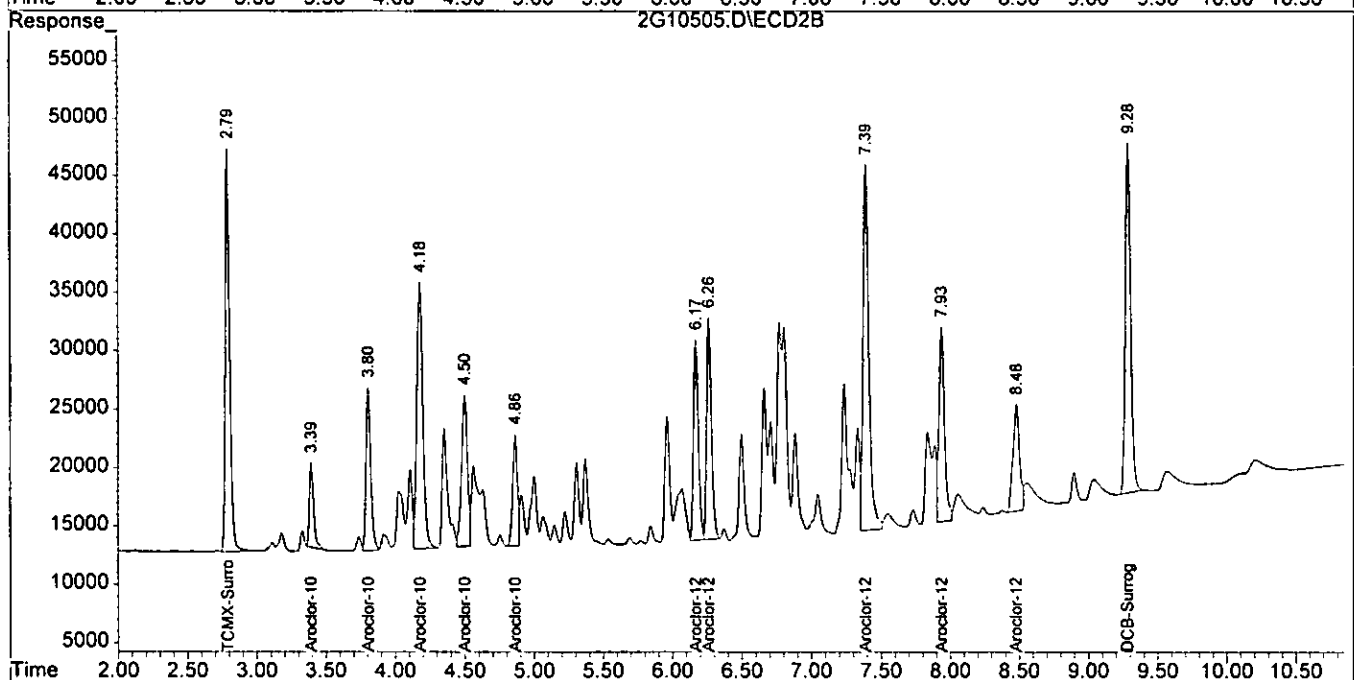
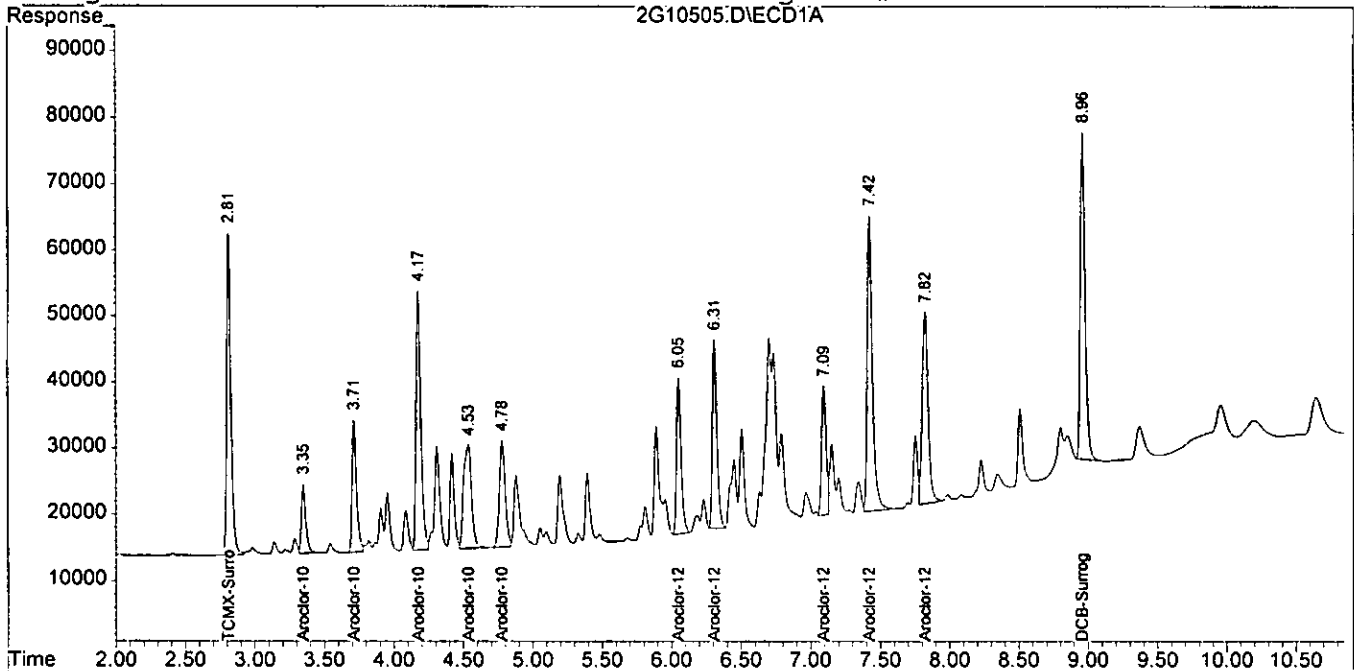
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	1023547	760239	56.637	56.707
2) Aroclor-1016 (1)	3.35	3.39	235649	156420	583.411	608.852
3) Aroclor-1016 (2)	3.71	3.80	442530	334842	569.940	588.249
4) Aroclor-1016 (3)	4.17	4.18	928646	686220	586.420	580.709
5) Aroclor-1016 (4)	4.53	4.50	619130	374590	593.414	533.686
6) Aroclor-1016 (5)	4.78	4.86	473041	227870	512.172	586.518m
7) Aroclor-1260 (1)	6.05	6.17	534771	416463	582.405	612.085
8) Aroclor-1260 (2)	6.31	6.26	645067	461648	568.979	607.311
9) Aroclor-1260 (3)	7.09	7.39	434727	899498	570.735	625.344
10) Aroclor-1260 (4)	7.42	7.94	1158203	462227	584.242	659.247
11) Aroclor-1260 (5)	7.82	8.48	865349	301977	626.421	644.844
12) Aroclor-1221 (1)	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 (2)	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 (3)	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 (2)	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 (3)	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 (4)	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 (5)	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 (2)	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 (3)	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 (4)	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 (5)	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	1176610	795135	58.442	61.333m

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10505.D\ECD1A.CH Vial 33
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10505.D\ECD2B.CH
Acq On : 5 Aug 2005 3:17 Operator: JK
Sample : CAL 1660@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 7:44 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



444

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10506.D\ECD1A.CH Vial
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10506.D\ECD2B.CH
 Acq On : 5 Aug 2005 3:31 Operator: JK
 Sample : CAL 1660@1000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 6:43 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

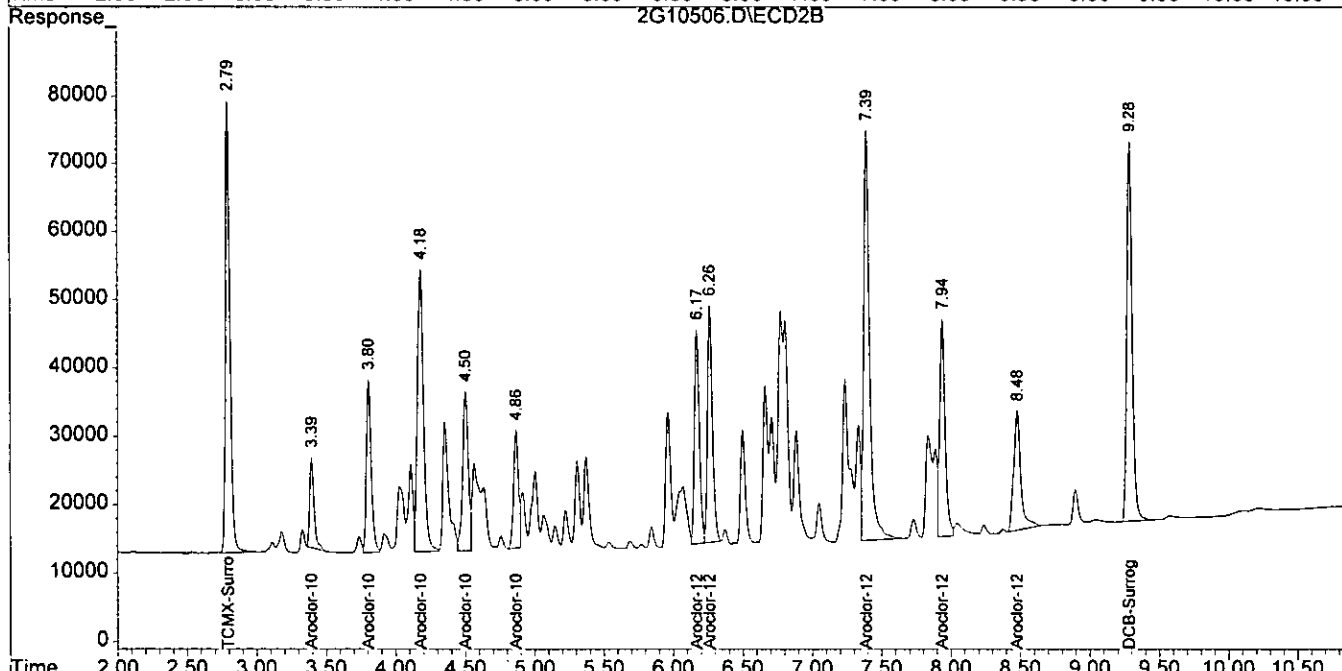
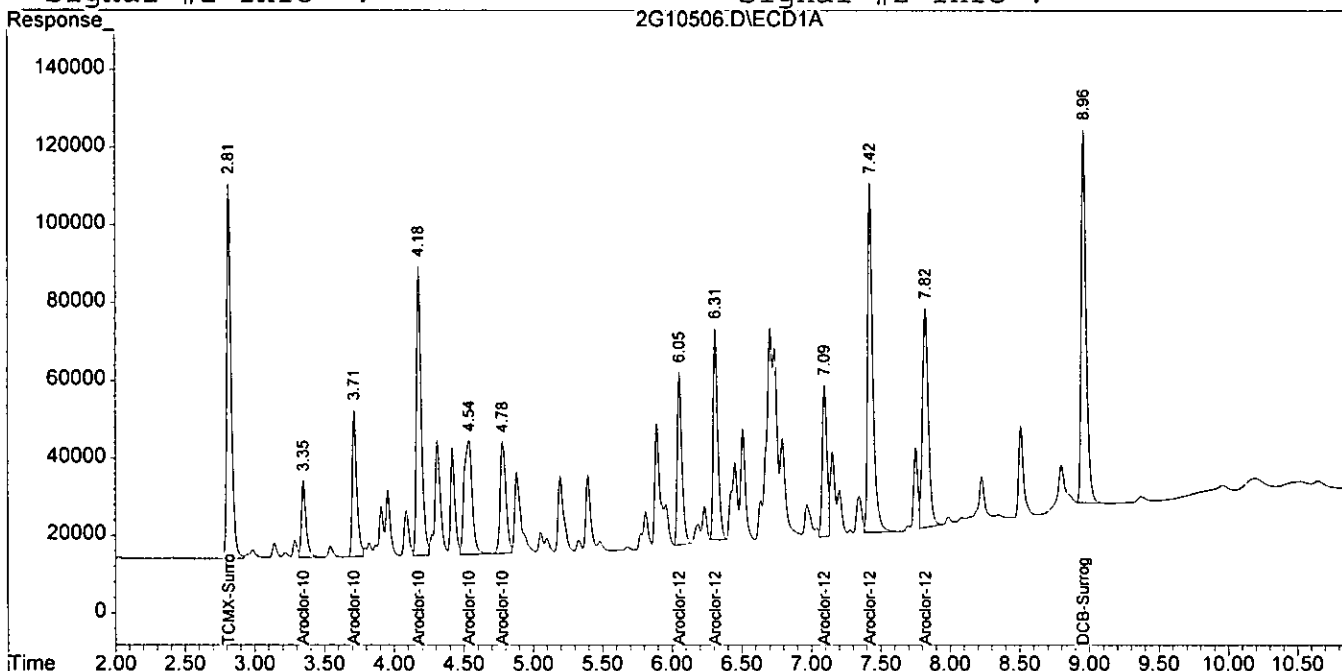
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	2068098	1461943	114.437	109.047
2) Aroclor-1016	3.35	3.39	459024	286799	1136.434	1116.345
3) Aroclor-1016	3.71	3.80	844073	612504	1087.093	1076.045
4) Aroclor-1016	4.18	4.18	1779339	1268838	1123.614	1073.745
5) Aroclor-1016	4.54	4.50	1182689	655086	1133.566	933.314
6) Aroclor-1016	4.78	4.86	834006	416639	902.996	1072.394m
7) Aroclor-1260	6.05	6.17	1020008	768784	1110.864	1129.900
8) Aroclor-1260	6.31	6.26	1236936	850794	1091.035	1119.245
9) Aroclor-1260	7.09	7.39	901419	1790164	1183.435	1244.548
10) Aroclor-1260	7.42	7.94	2336939	866983	1178.840	1236.525
11) Aroclor-1260	7.82	8.48	1664672	621168	1205.046	1326.446
12) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	2348809	1499448	116.666	115.660

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10506.D\ECD1A.CH Vial:
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10506.D\ECD2B.CH
Acq On : 5 Aug 2005 3:31 Operator: JK
Sample : CAL 1660@1000PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 6:43 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10507.D\ECD1A.CH Vial: 113
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10507.D\ECD2B.CH
 Acq On : 5 Aug 2005 3:46 Operator: JK
 Sample : CAL 1660@2000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

OP/11/0

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

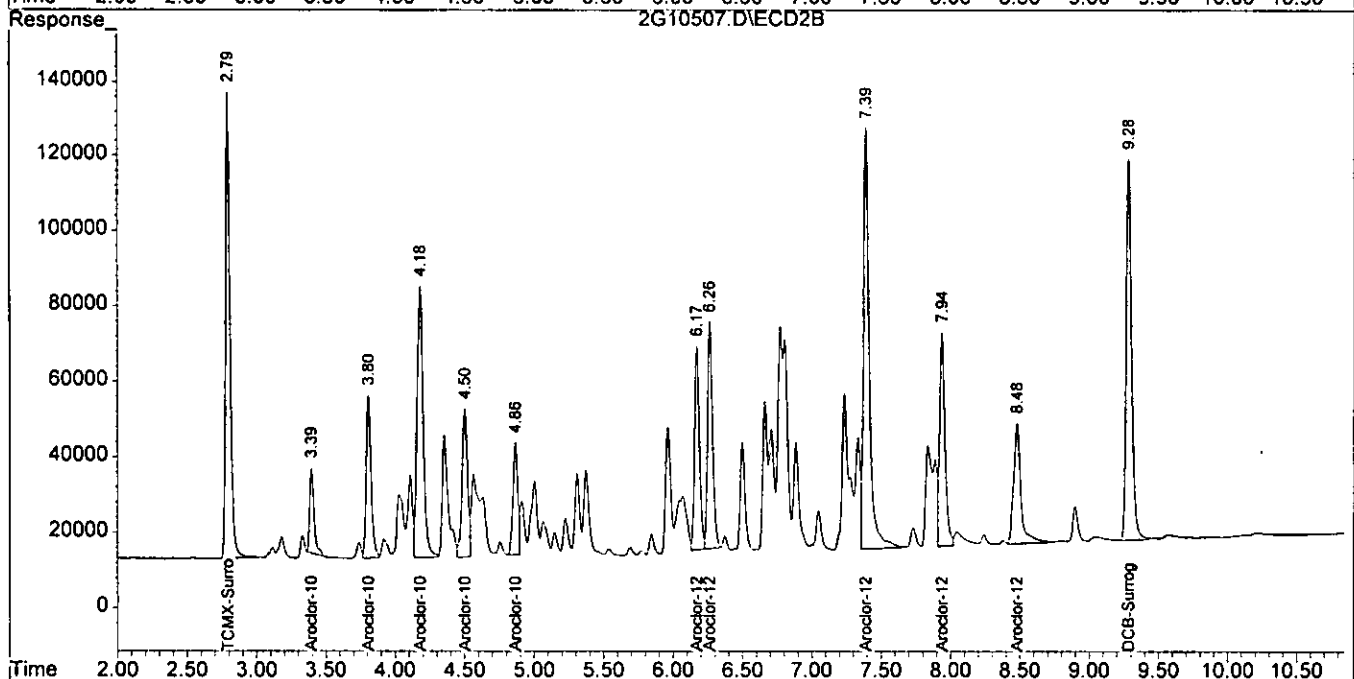
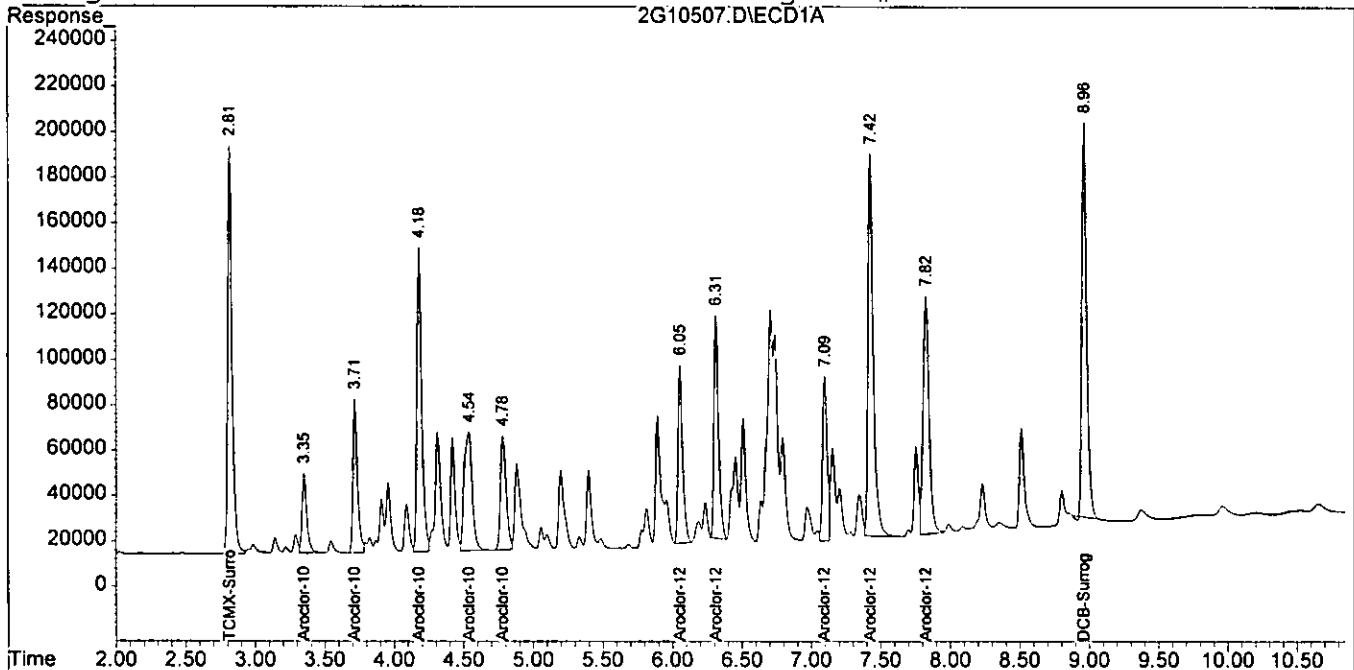
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	3911873	2714639	.216.461	202.486
2) Aroclor-1016 (1)	3.35	3.39	828392	491354	2050.904	1912.560
3) Aroclor-1016 (2)	3.71	3.81	1514692	1052607	1950.793	1849.214
4) Aroclor-1016 (3)	4.18	4.18	3199834	2198953	2020.625	1860.848
5) Aroclor-1016 (4)	4.54	4.50	2114810	1082399	2026.971	1542.116
6) Aroclor-1016 (5)	4.78	4.86	1430841	707369	1549.203	1820.709m
7) Aroclor-1260 (1)	6.05	6.17	1814321	1322679	1975.930	1943.973
8) Aroclor-1260 (2)	6.31	6.26	2235502	1472104	1971.816	1936.596
9) Aroclor-1260 (3)	7.09	7.39	1679772	3299506	2205.303	2293.863
10) Aroclor-1260 (4)	7.42	7.94	4397510	1553699	2218.270	2215.947
11) Aroclor-1260 (5)	7.82	8.48	3149933	1167070	2280.218	2492.167
12) Aroclor-1221 (1)	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 (2)	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 (3)	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 (2)	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 (3)	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 (4)	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 (5)	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 (2)	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 (3)	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 (4)	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 (5)	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	4172369	2699181	207.242	208.201m

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10507.D\ECD1A.CH Vial 335
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10507.D\ECD2B.CH
Acq On : 5 Aug 2005 3:46 Operator: JK
Sample : CAL 1660@2000PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 7:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10508.D\ECD1A.CH Vial 1136
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10508.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:00 Operator: JK
 Sample : CAL 1660@4000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:02 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

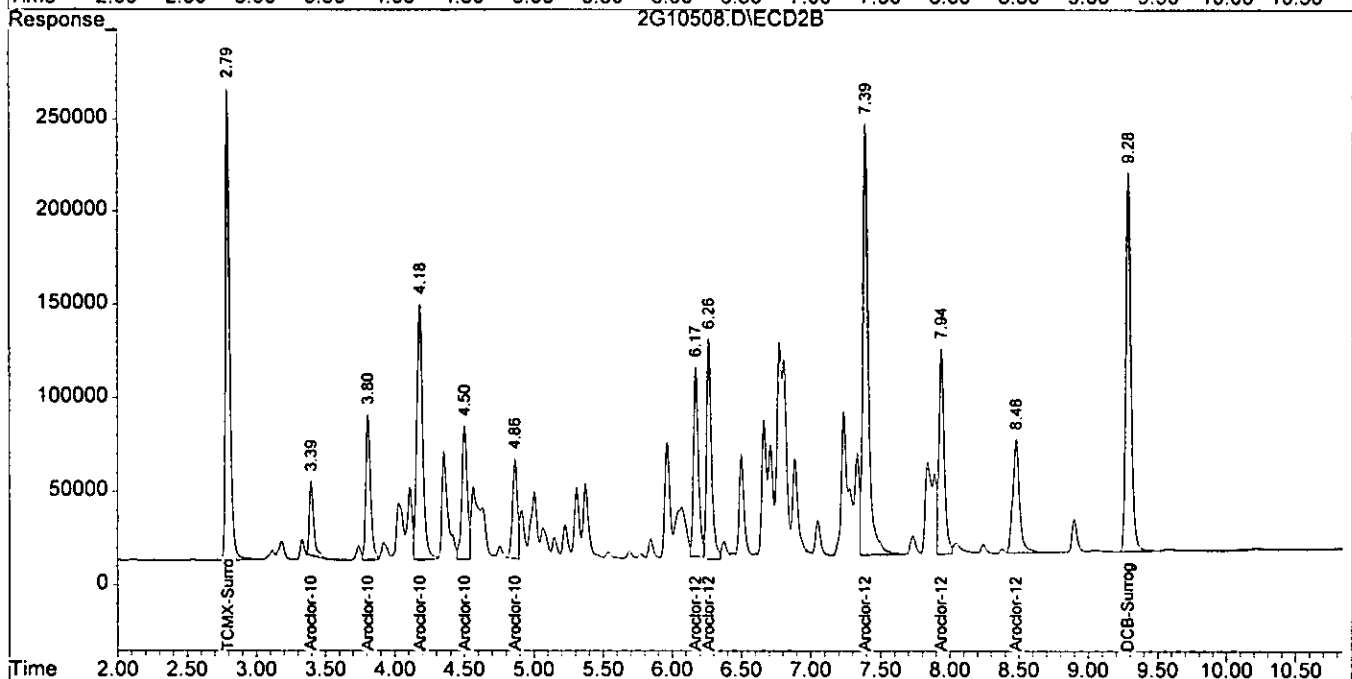
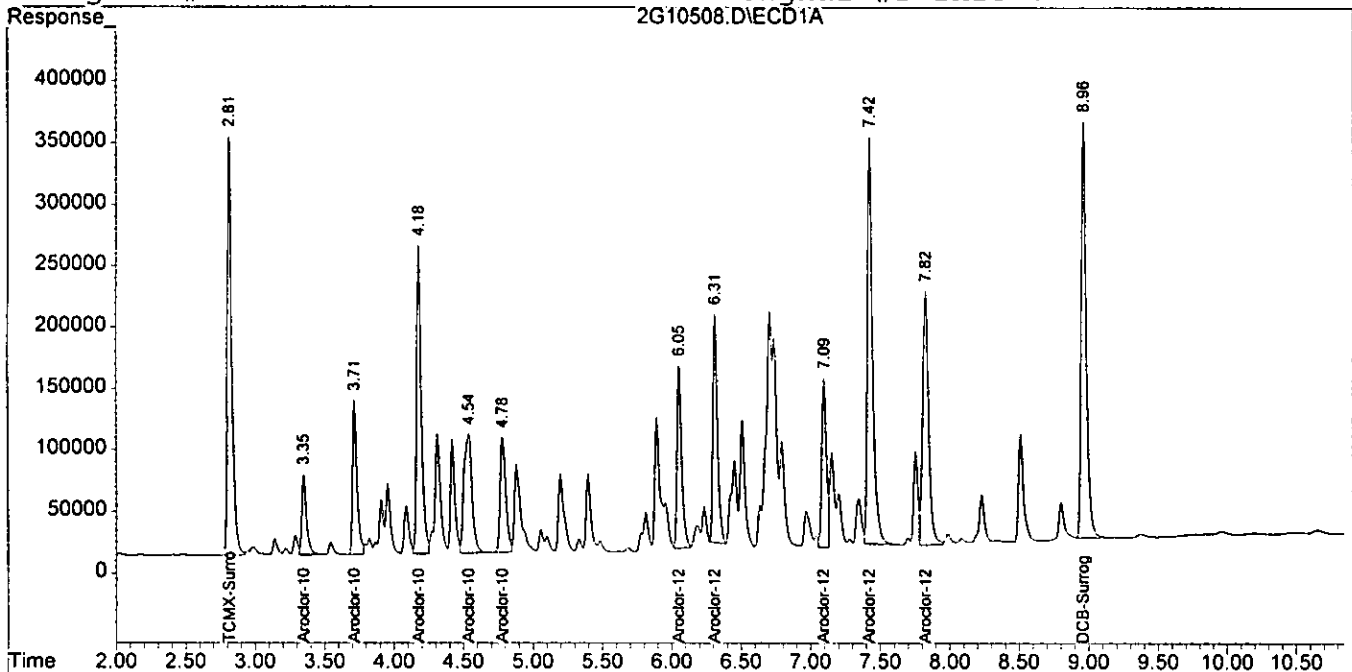
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.81	2.79	7503126	5326951	415.181	397.340
2) Aroclor-1016 (1)	3.35	3.39	1517580	868690	3757.168	3381.310
3) Aroclor-1016 (2)	3.71	3.81	2784603	1876068	3586.328	3295.866
4) Aroclor-1016 (3)	4.18	4.18	5958515	4036906	3762.672	3416.203
5) Aroclor-1016 (4)	4.54	4.50	3883086	1923953	3721.801	2741.096 #
6) Aroclor-1016 (5)	4.78	4.86	2647417	1264584	2866.417	3254.931m
7) Aroclor-1260 (1)	6.05	6.17	3411714	2490447	3715.609m	3660.268m
8) Aroclor-1260 (2)	6.31	6.26	4153018	2940352	3663.154	3868.118m
9) Aroclor-1260 (3)	7.09	7.39	3166218	6406382	4156.797	4453.807
10) Aroclor-1260 (4)	7.42	7.94	8571854	2930093	4323.967	4179.016
11) Aroclor-1260 (5)	7.82	8.48	6268932	2074593	4538.043	4430.096
12) Aroclor-1221 (1)	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 (2)	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 (3)	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 (2)	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 (3)	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 (4)	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 (5)	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 (2)	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 (3)	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 (4)	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 (5)	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	8351630	5158203	414.826	397.878

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10508.D\ECD1A.CH Vial: 111
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10508.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:00 Operator: JK
 Sample : CAL 1660@4000PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 7:02 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Multiple Level Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10512.D\ECD1A.CH Vial: 110
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10512.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:58 Operator: JK
 Sample : CAL 1232@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 6:56 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/0

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

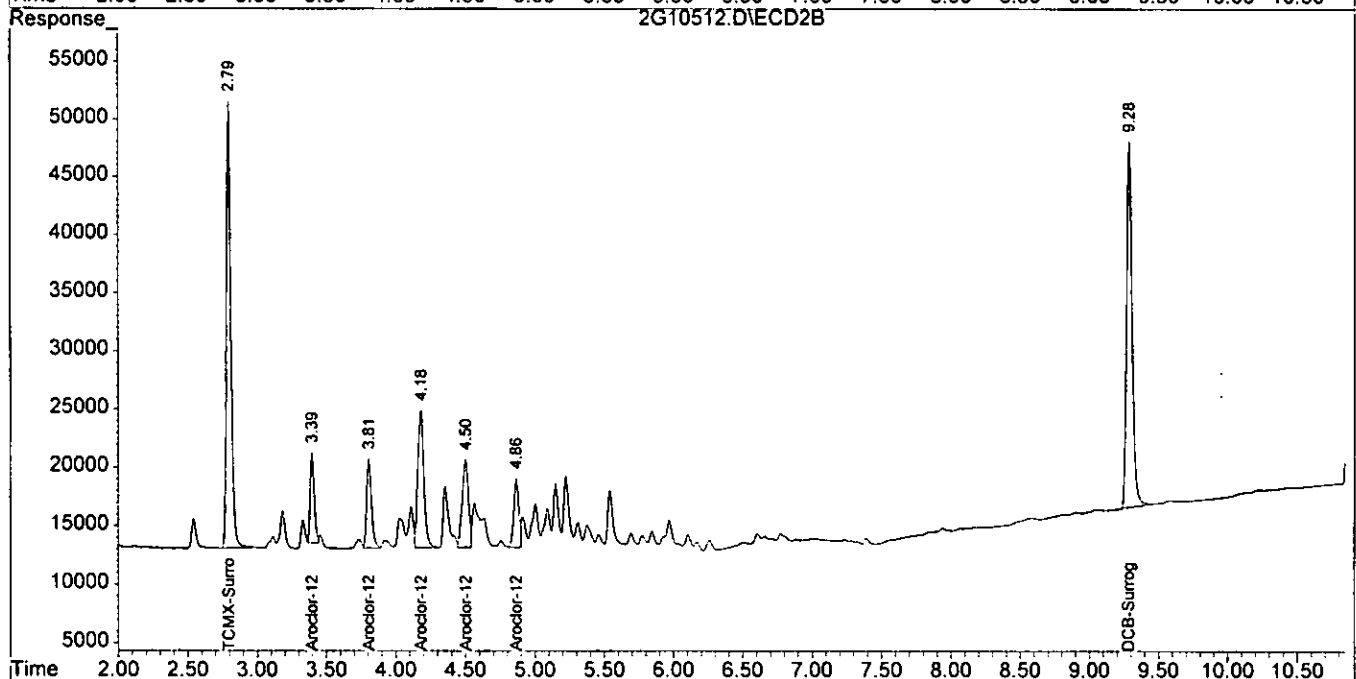
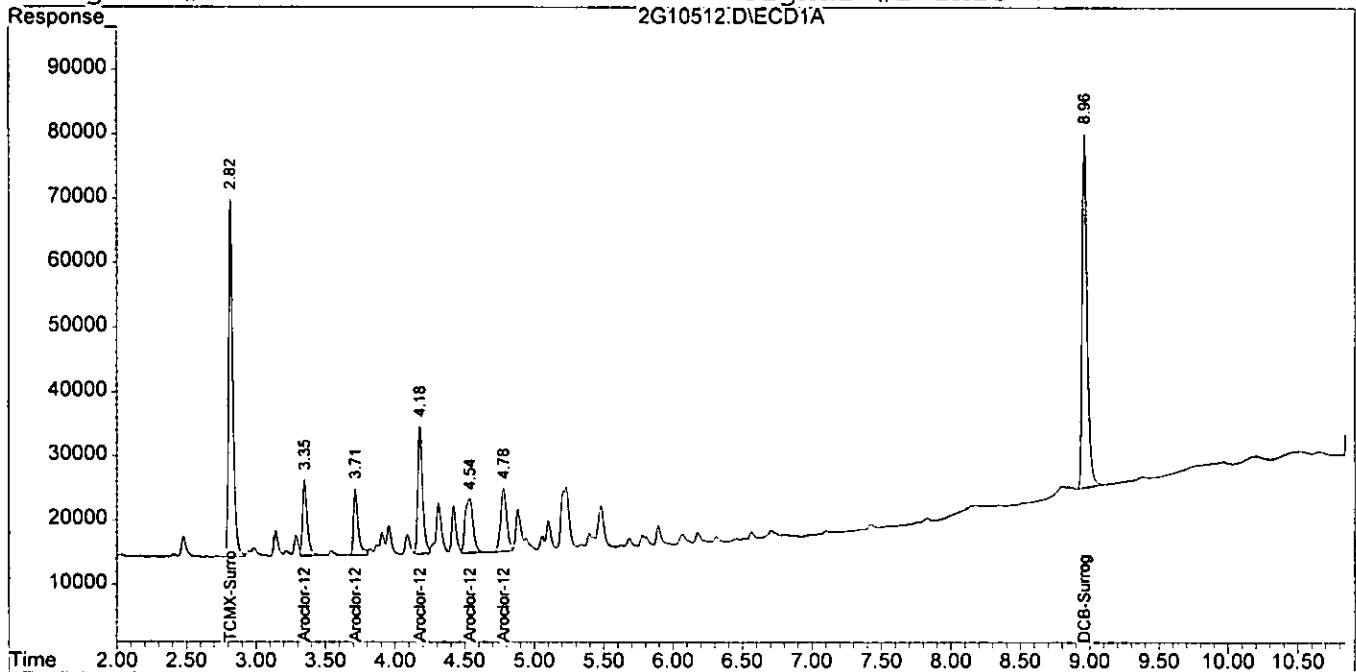
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.79	1188612	852532	65.771	63.591
2) Aroclor-1016 1	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 2	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 3	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 4	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 5	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 1	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 2	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 3	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 4	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 5	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 1	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 2	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 3	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 1	3.35	3.39	273407	158520	552.897	655.790m
16) Aroclor-1232 2	3.71	3.81	242868	187082	561.539	577.282
17) Aroclor-1232 3	4.18	4.18	475031	358078	581.903	595.144
18) Aroclor-1232 4	4.54	4.50	331372	235186	566.210	574.826
19) Aroclor-1232 5	4.78	4.86	301271	147438	557.471	593.934m
20) Aroclor-1242 1	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 2	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 3	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 4	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 5	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 1	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 2	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 3	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 4	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 5	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 1	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 2	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 3	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 4	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 5	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	1352374	860464	67.173	66.372

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10512.D\ECD1A.CH Vial 110
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10512.D\ECD2B.CH
Acq On : 5 Aug 2005 4:58 Operator: JK
Sample : CAL 1232@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 6:56 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10511.D\ECD1A.CH Vial 1149
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10511.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:43 Operator: JK
 Sample : CAL 1242@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 6:55 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

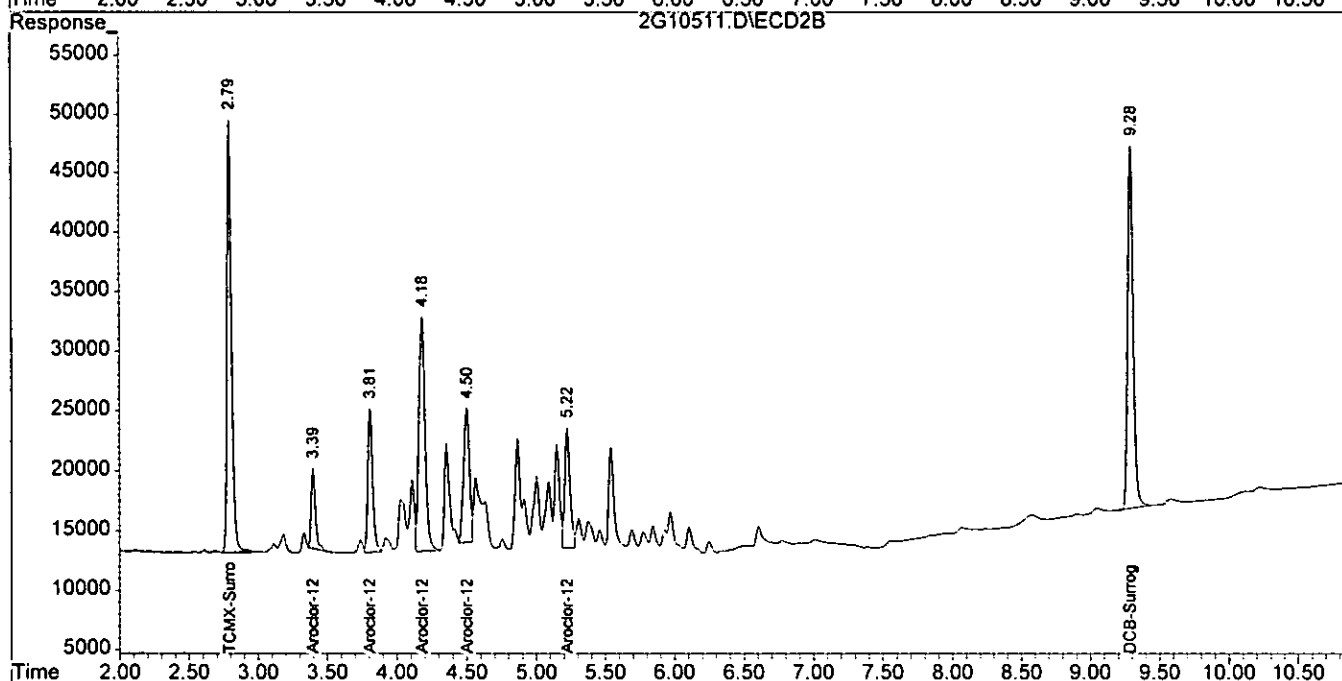
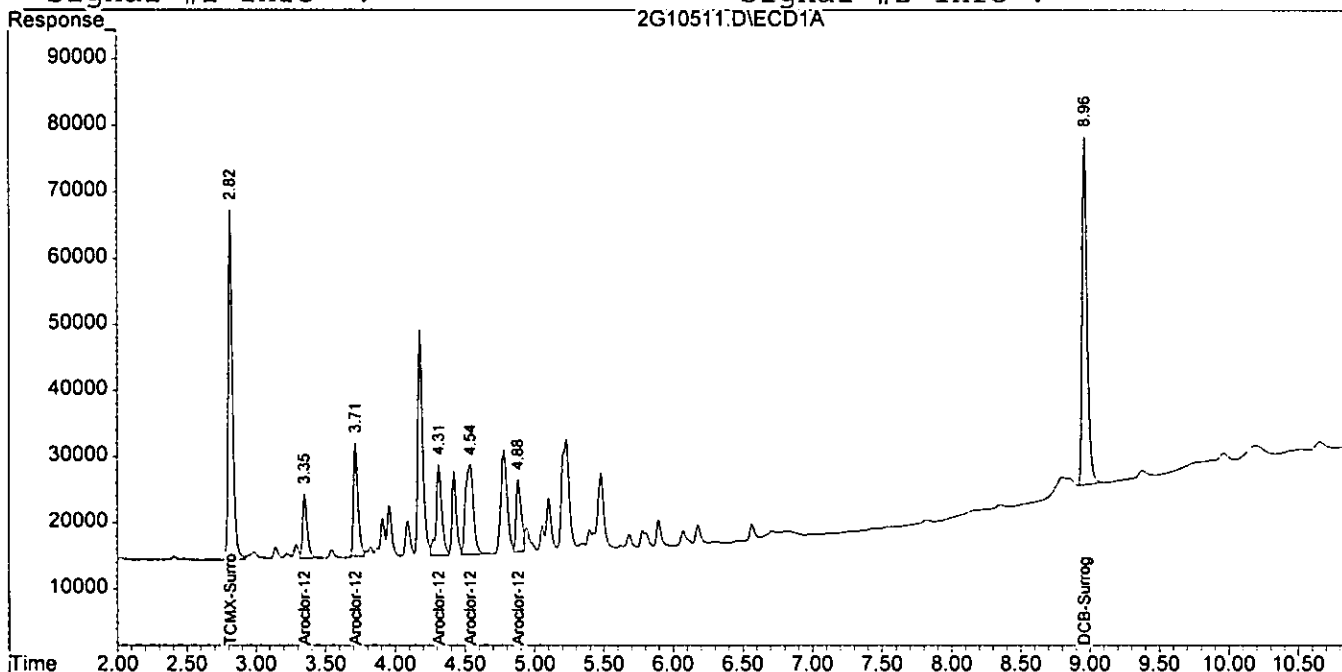
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.79	1134180	818918	62.759	61.083
2) Aroclor-1016 (1)	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 (2)	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 (3)	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 (4)	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 (5)	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 (1)	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 (2)	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 (3)	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 (4)	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 (5)	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 (1)	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 (2)	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 (3)	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 (1)	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 (2)	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 (3)	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 (4)	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 (5)	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 (1)	3.35	3.40	224274	143157	530.686	579.799
21) Aroclor-1242 (2)	3.71	3.81	377487	292006	514.661m	560.890
22) Aroclor-1242 (3)	4.31	4.18	386631	603217	266.466	580.575 #
23) Aroclor-1242 (4)	4.54	4.50	540268	331788	554.719	568.553m
24) Aroclor-1242 (5)	4.88	5.22	276767	267919	358.743	715.430 #
25) Aroclor-1248 (1)	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 (2)	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 (3)	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 (4)	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 (5)	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 (1)	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 (2)	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 (3)	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 (4)	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 (5)	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	1283612	823159	63.757	63.494

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10511.D\ECD1A.CH Vial:
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10511.D\ECD2B.CH
Acq On : 5 Aug 2005 4:43 Operator: JK
Sample : CAL 1242@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 6:55 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info:



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10510.D\ECD1A.CH Vial:
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10510.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:29 Operator: JK
 Sample : CAL 1248@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 6:54 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

08/11/01

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

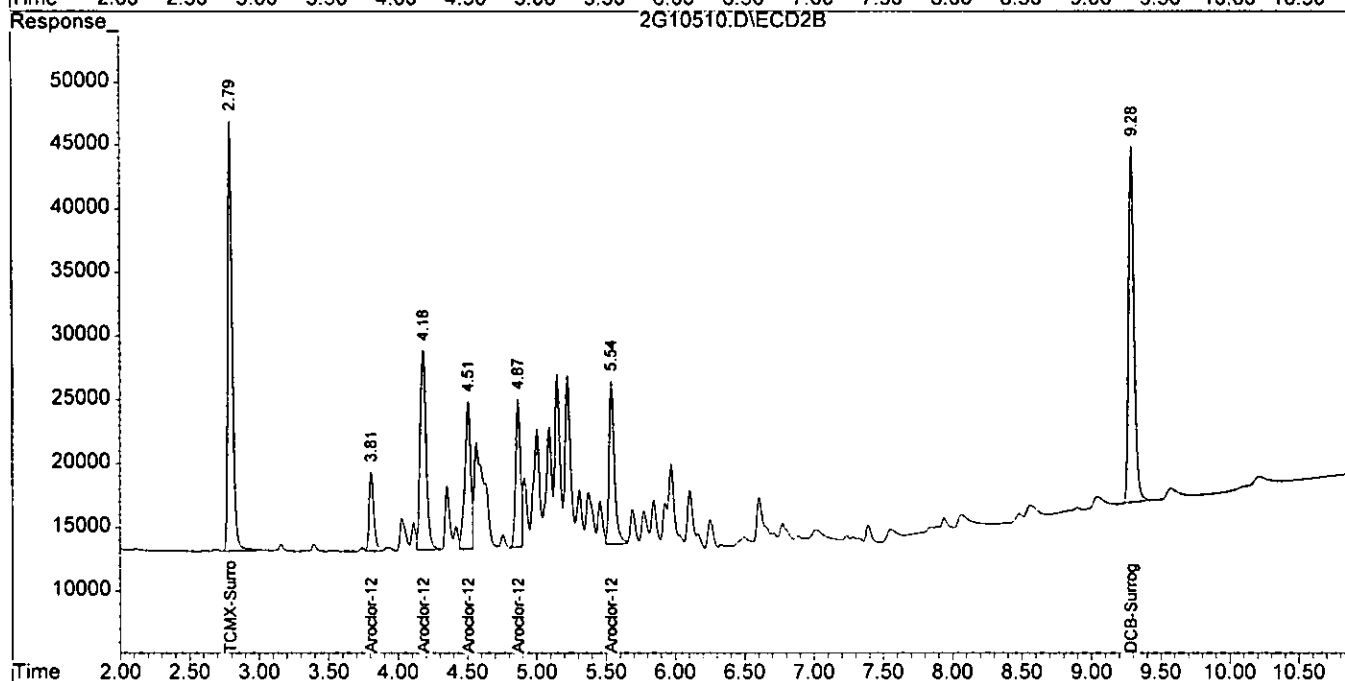
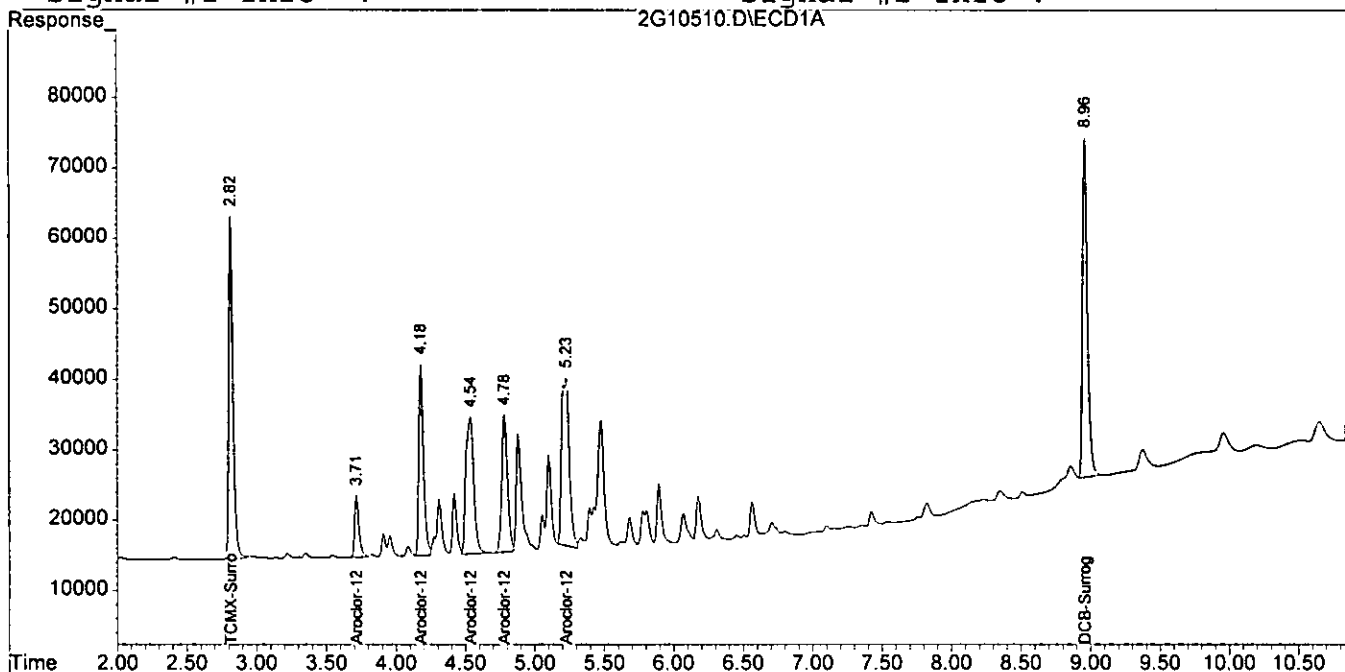
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.79	1018596	752396	56.363	56.122
2) Aroclor-1016 {1}	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 {2}	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 {3}	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 {4}	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 {5}	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 {1}	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 {2}	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 {3}	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 {4}	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 {5}	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 {1}	0.00	0.00	0	0	N.D. d	N.D. d
13) Aroclor-1221 {2}	0.00	0.00	0	0	N.D. d	N.D. d
14) Aroclor-1221 {3}	0.00	0.00	0	0	N.D. d	N.D. d
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	3.71	3.81	195115	143813	514.400	528.738
26) Aroclor-1248 {2}	4.18	4.18	645166	473566	526.261	537.429
27) Aroclor-1248 {3}	4.54	4.51	774276	335728	519.925	453.024
28) Aroclor-1248 {4}	4.78	4.87	568856	264234	454.640	517.730m
29) Aroclor-1248 {5}	5.23	5.54	916025	339244	541.117	564.349
30) Aroclor-1254 {1}	0.00	0.00	0	0	N.D. d	N.D. d
31) Aroclor-1254 {2}	0.00	0.00	0	0	N.D. d	N.D. d
32) Aroclor-1254 {3}	0.00	0.00	0	0	N.D. d	N.D. d
33) Aroclor-1254 {4}	0.00	0.00	0	0	N.D. d	N.D. d
34) Aroclor-1254 {5}	0.00	0.00	0	0	N.D. d	N.D. d
35) DCB-Surrogate	8.96	9.28	1163484	754638	57.790	58.209

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10510.D\ECD1A.CH Vial 118
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10510.D\ECD2B.CH
Acq On : 5 Aug 2005 4:29 Operator: JK
Sample : CAL 1248@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 6:54 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10509.D\ECD1A.CH Vial:
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10509.D\ECD2B.CH
 Acq On : 5 Aug 2005 4:15 Operator: JK
 Sample : CAL 2154@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 6:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Jul 22 08:05:17 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

28/11/05

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

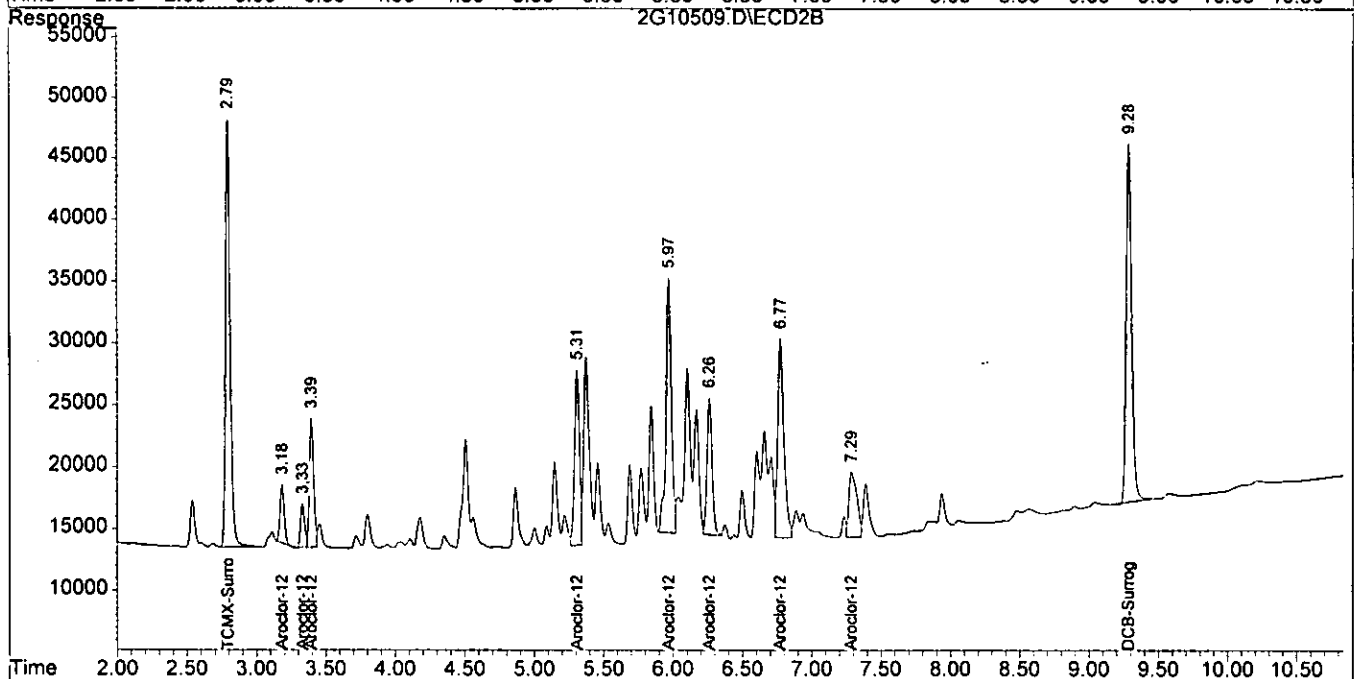
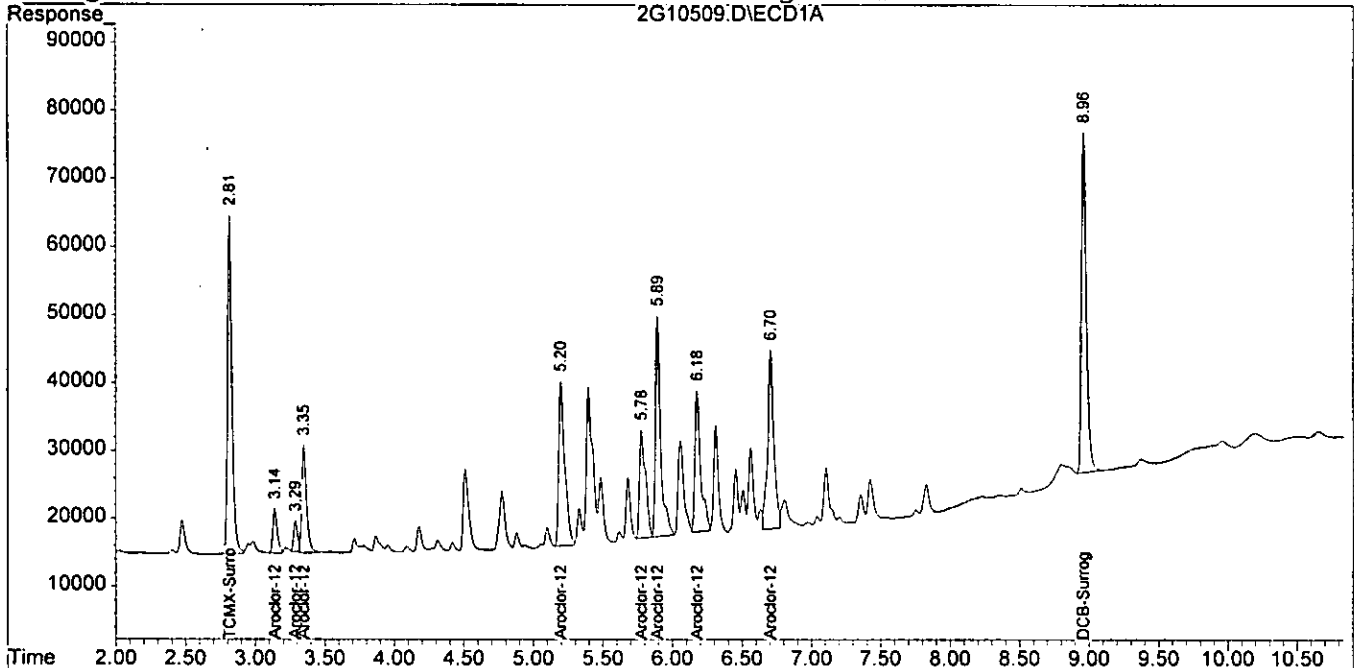
Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.79	1093345	796785	60.500	59.433
2) Aroclor-1016 {1}	0.00	0.00	0	0	N.D. d	N.D. d
3) Aroclor-1016 {2}	0.00	0.00	0	0	N.D. d	N.D. d
4) Aroclor-1016 {3}	0.00	0.00	0	0	N.D. d	N.D. d
5) Aroclor-1016 {4}	0.00	0.00	0	0	N.D. d	N.D. d
6) Aroclor-1016 {5}	0.00	0.00	0	0	N.D. d	N.D. d
7) Aroclor-1260 {1}	0.00	0.00	0	0	N.D. d	N.D. d
8) Aroclor-1260 {2}	0.00	0.00	0	0	N.D. d	N.D. d
9) Aroclor-1260 {3}	0.00	0.00	0	0	N.D. d	N.D. d
10) Aroclor-1260 {4}	0.00	0.00	0	0	N.D. d	N.D. d
11) Aroclor-1260 {5}	0.00	0.00	0	0	N.D. d	N.D. d
12) Aroclor-1221 {1}	3.14	3.18	142349	98182	528.040m	559.302
13) Aroclor-1221 {2}	3.29	3.33	90182	73005	512.964m	763.393m#
14) Aroclor-1221 {3}	3.35	3.39	369875	230567	545.673	724.684m#
15) Aroclor-1232 {1}	0.00	0.00	0	0	N.D. d	N.D. d
16) Aroclor-1232 {2}	0.00	0.00	0	0	N.D. d	N.D. d
17) Aroclor-1232 {3}	0.00	0.00	0	0	N.D. d	N.D. d
18) Aroclor-1232 {4}	0.00	0.00	0	0	N.D. d	N.D. d
19) Aroclor-1232 {5}	0.00	0.00	0	0	N.D. d	N.D. d
20) Aroclor-1242 {1}	0.00	0.00	0	0	N.D. d	N.D. d
21) Aroclor-1242 {2}	0.00	0.00	0	0	N.D. d	N.D. d
22) Aroclor-1242 {3}	0.00	0.00	0	0	N.D. d	N.D. d
23) Aroclor-1242 {4}	0.00	0.00	0	0	N.D. d	N.D. d
24) Aroclor-1242 {5}	0.00	0.00	0	0	N.D. d	N.D. d
25) Aroclor-1248 {1}	0.00	0.00	0	0	N.D. d	N.D. d
26) Aroclor-1248 {2}	0.00	0.00	0	0	N.D. d	N.D. d
27) Aroclor-1248 {3}	0.00	0.00	0	0	N.D. d	N.D. d
28) Aroclor-1248 {4}	0.00	0.00	0	0	N.D. d	N.D. d
29) Aroclor-1248 {5}	0.00	0.00	0	0	N.D. d	N.D. d
30) Aroclor-1254 {1}	5.20	5.31	731638	348480	551.843	572.692
31) Aroclor-1254 {2}	5.78	5.97	479310	558827	558.974	578.747
32) Aroclor-1254 {3}	5.89	6.26	837784	282672	555.133	566.270
33) Aroclor-1254 {4}	6.18	6.77	603667	501776	592.784m	574.412
34) Aroclor-1254 {5}	6.70	7.29	824594	211975	567.306	576.001
35) DCB-Surrogate	8.96	9.28	1222387	782957	60.716	60.393

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10509.D\ECD1A.CH Vial: 1001
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10509.D\ECD2B.CH
Acq On : 5 Aug 2005 4:15 Operator: JK
Sample : CAL 2154@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 6:45 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Jul 22 08:05:17 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form7
Continuing Calibration

1149

Data File:
Method:
Calibration Name:
Calibration Date/Time

2G10446.D 8082 CAL 1660@1000PP 08/03/05 16:41	2G10533.D 8082 CAL1660@1000PP 08/05/05 10:59	2G10580.D 8082 CAL 1660@500PP 08/08/05 08:12	2G10600.D 8082 CAL 1660@1000PP 08/08/05 13:25	2G10622.D 8082 CAL 1660@2000PP 08/08/05 18:42
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Compound	Limit	Col	Mr	Conc			Conc			Conc			Conc			Conc		
				Conc	Exp	%Diff	Conc	Exp	%Diff	Conc	Exp	%Diff	Conc	Exp	%Diff	Conc	Exp	%Diff
TCMX-Surrogate	15	1	0	106.5	100	6.5	112.2	100	12.2	50.51	50	1.0	109.3	100	9.3	231.5	200	15.7*
Aroclor-1016	15	1	1	1098	1000	9.8	1115	1000	11.5	534.1	500	6.8	1067	1000	6.7	2196	2000	9.8
Aroclor-1016	15	1	2	1096	1000	9.6	1088	1000	8.8	523.1	500	4.6	1010	1000	1.0	2197	2000	9.8
Aroclor-1016	15	1	3	1106	1000	10.6	1085	1000	8.5	510.5	500	2.1	984.3	1000	1.6	2122	2000	6.1
Aroclor-1016	15	1	4	1102	1000	10.2	1116	1000	11.6	533.8	500	6.8	996.7	1000	0.3	2170	2000	8.5
Aroclor-1016	15	1	5	1163	1000	16.3*	1091	1000	9.1	491.3	500	1.7	1020	1000	2.0	2355	2000	17.8*
Aroclor-1260	15	1	1	1119	1000	11.9	1097	1000	9.7	523.4	500	4.7	947.3	1000	5.3	1958	2000	2.1
Aroclor-1260	15	1	2	1119	1000	11.9	1104	1000	10.4	522.6	500	4.5	946.8	1000	5.3	1962	2000	1.9
Aroclor-1260	15	1	3	1067	1000	6.7	1108	1000	10.8	522.3	500	4.5	935	1000	6.5	1928	2000	3.6
Aroclor-1260	15	1	4	1098	1000	9.8	1119	1000	11.9	516	500	3.2	952.1	1000	4.8	2059	2000	3.0
Aroclor-1260	15	1	5	1152	1000	15.2	1122	1000	12.2	498	500	0.4	1081	1000	8.1	2001	2000	0.1
DCB-Surrogate	15	1	0	108.1	100	8.1	115.8	100	15.8*	51.66	50	3.3	95.47	100	4.5	213.8	200	6.9
Average Difference	15	1	0			10.5			11.0			3.6			4.6			7.1
TCMX-Surrogate	15	2	0	104	100	4.0	100.8	100	0.8	47.59	50	4.8	96.54	100	3.5	208.5	200	4.2
Aroclor-1016	15	2	1	1025	1000	2.5	1076	1000	7.6	489.2	500	2.2	944.1	1000	5.6	2003	2000	0.2
Aroclor-1016	15	2	2	1126	1000	12.6	999.1	1000	0.1	502.7	500	0.5	930.4	1000	7.0	1906	2000	4.7
Aroclor-1016	15	2	3	1117	1000	11.7	989.9	1000	1.0	488.7	500	2.3	911.5	1000	8.9	1910	2000	4.5
Aroclor-1016	15	2	4	1156	1000	15.6*	1053	1000	5.3	489.4	500	2.1	1013	1000	1.3	2292	2000	14.6
Aroclor-1016	15	2	5	1021	1000	2.1	1050	1000	5.0	507.7	500	1.5	945	1000	5.5	1990	2000	0.5
Aroclor-1260	15	2	1	1060	1000	6.0	997	1000	0.3	475.4	500	4.9	862.3	1000	13.8	1832	2000	8.4
Aroclor-1260	15	2	2	1049	1000	4.9	985.7	1000	1.4	467.1	500	6.6	845.1	1000	15.5	1796	2000	10.2
Aroclor-1260	15	2	3	1178	1000	17.8*	1025	1000	2.5	491.6	500	1.7	833.3	1000	16.7*	1935	2000	3.2
Aroclor-1260	15	2	4	1197	1000	19.7*	1025	1000	2.5	489.4	500	2.1	808.5	1000	19.2*	1828	2000	8.6
Aroclor-1260	15	2	5	1176	1000	17.6*	1091	1000	9.1	505.4	500	1.1	1020	1000	2.0	1925	2000	3.8
DCB-Surrogate	15	2	0	116.0	100	16.0*	96.97	100	3.0	48.2	50	3.6	79.37	100	20.6*	168.2	200	15.9*
Average Difference	15	2	0			10.9			3.2			2.8			9.9			6.6

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10446.D\ECD1A.CH Vial: 1
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10446.D\ECD2B.CH
 Acq On : 3 Aug 2005 16:41 Operator: JK
 Sample : CAL 1660@1000PPB Inst : gc_2
 Misc : S,PCB:0.5 Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 4 7:28 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 11:13:53 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	2739028	1843450	106.456	103.969
2) Aroclor-1016 {1}	3.37	3.42	593641	325997	1097.532	1025.318
3) Aroclor-1016 {2}	3.74	3.83	1090721	749070	1095.504	1125.626
4) Aroclor-1016 {3}	4.21	4.21	2323251	1537697	1106.256	1116.533
5) Aroclor-1016 {4}	4.57	4.53	1522467	816509	1101.587	1155.733
6) Aroclor-1016 {5}	4.81	4.89	1121446	514434	1162.577	1021.418
7) Aroclor-1260 {1}	6.08	6.20	1298848	894339	1118.532	1060.481
8) Aroclor-1260 {2}	6.34	6.29	1566344	991035	1119.475	1048.545
9) Aroclor-1260 {3}	7.13	7.42	1095067	2042149	1067.362	1177.659
10) Aroclor-1260 {4}	7.45	7.97	2785656	989156	1098.399	1197.037
11) Aroclor-1260 {5}	7.86	8.51	2081970	633230	1151.948m	1175.856m
35) DCB-Surrogate	8.99	9.32	2704636	1640013	108.091	116.022

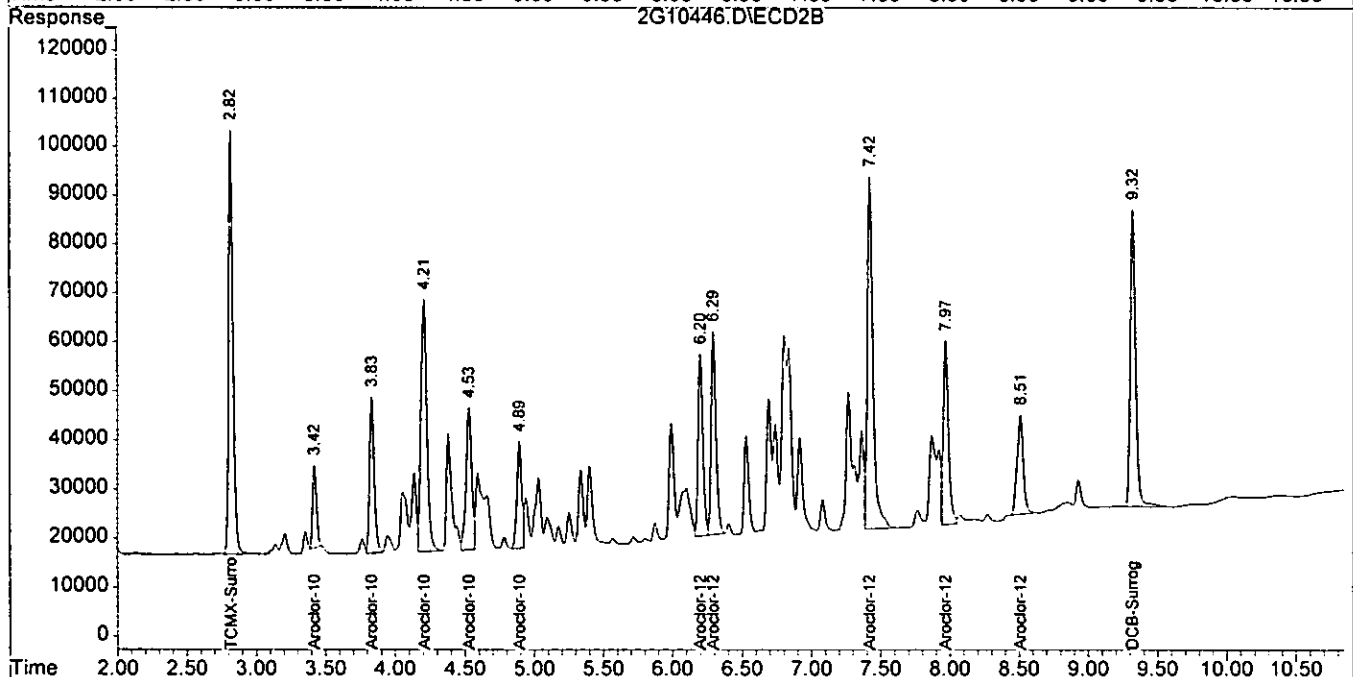
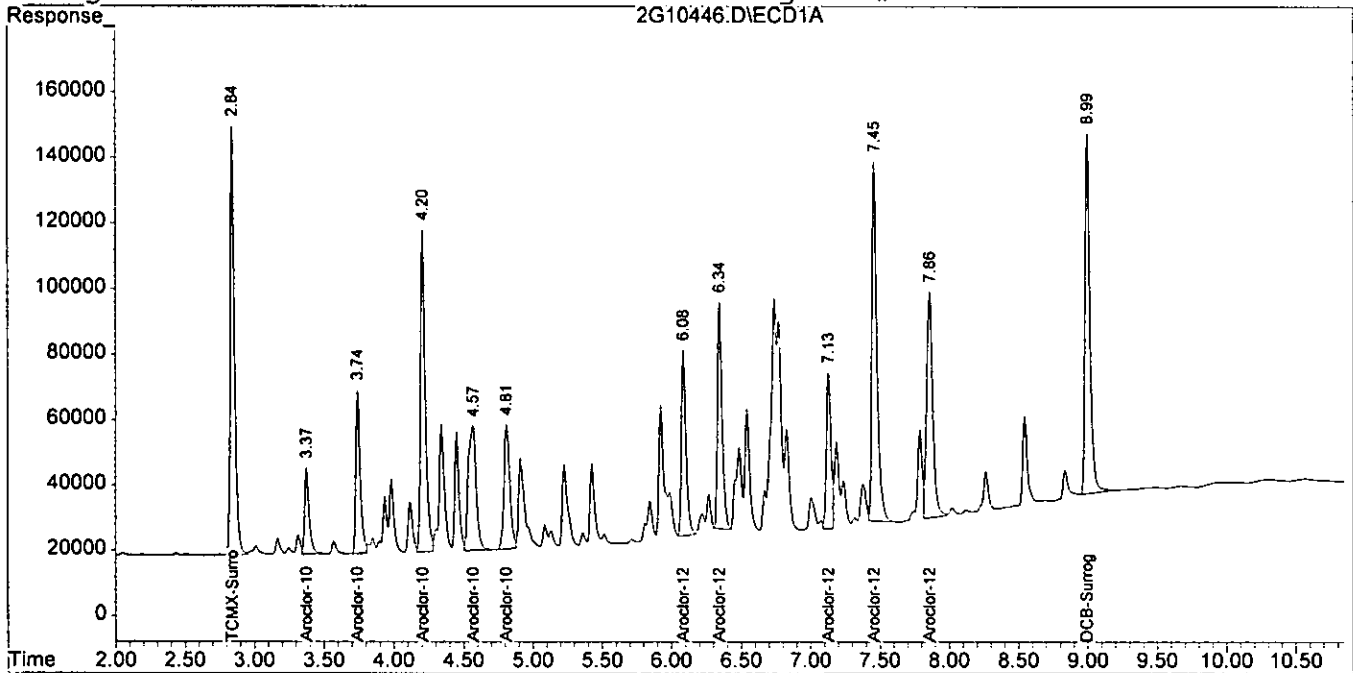
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10446.D\ECD1A.CH Vial: 881
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10446.D\ECD2B.CH
Acq On : 3 Aug 2005 16:41 Operator: JK
Sample : CAL 1660@1000PPB Inst : gc_2
Misc : S,PCB:0.5 Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 4 7:28 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 11:13:53 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10533.D\ECD1A.CH Vial: 31
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10533.D\ECD2B.CH
 Acq On : 5 Aug 2005 10:59 Operator: JK
 Sample : CAL1660@1000PPB Inst : gc_2
 Misc : S,PCB:0.5 Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 11:26 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	2191890	1471432	112.219	100.789
2) Aroclor-1016 {1}	3.35	3.40	488983	301656	1115.194	1075.733
3) Aroclor-1016 {2}	3.72	3.81	899332	626256	1087.514	999.072
4) Aroclor-1016 {3}	4.18	4.19	1900759	1289330	1084.621	989.853
5) Aroclor-1016 {4}	4.54	4.51	1273682	649201	1116.159	1053.460
6) Aroclor-1016 {5}	4.78	4.87	861709	442786	1090.812	1050.453
7) Aroclor-1260 {1}	6.06	6.17	1105145	772583	1096.668	996.954
8) Aroclor-1260 {2}	6.31	6.27	1334060	855044	1103.773	985.721
9) Aroclor-1260 {3}	7.10	7.39	991316	1735055	1108.432	1024.864
10) Aroclor-1260 {4}	7.43	7.94	2466416	858656	1118.995	1025.045
11) Aroclor-1260 {5}	7.83	8.48	1829640	609293	1121.917	1091.288
35) DCB-Surrogate	8.96	9.29	2492922	1464294	115.791	96.971

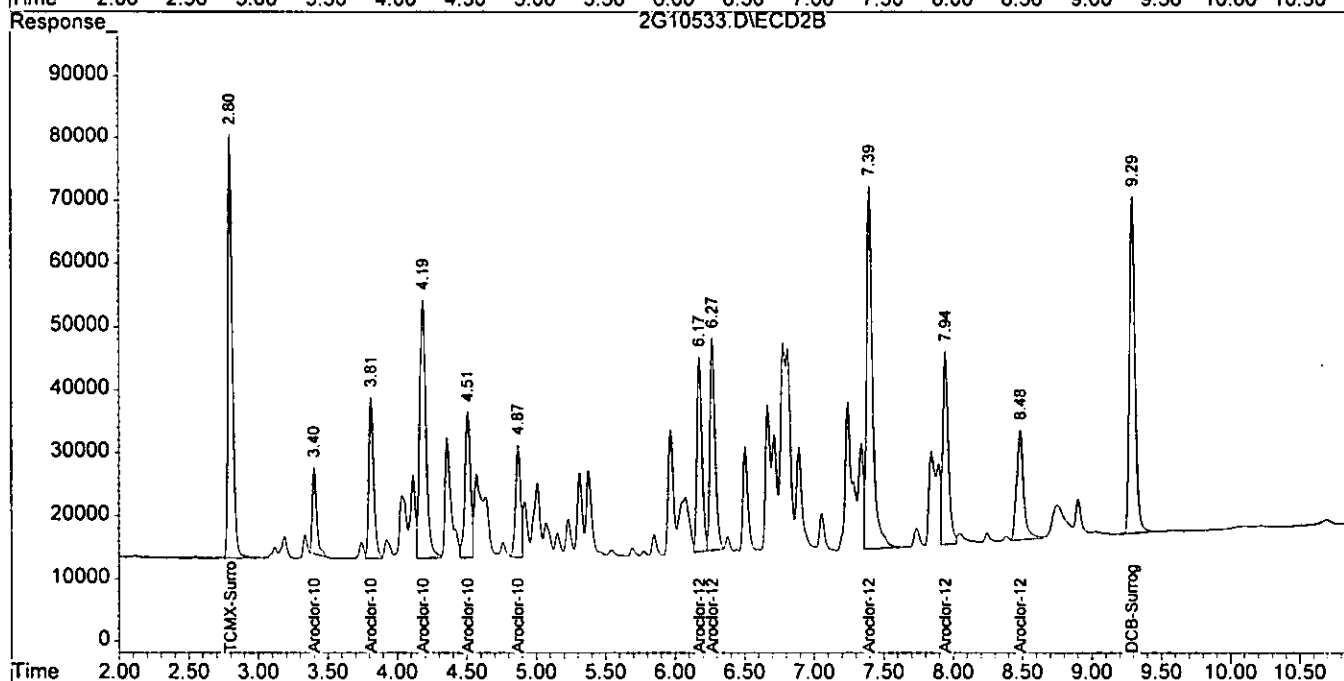
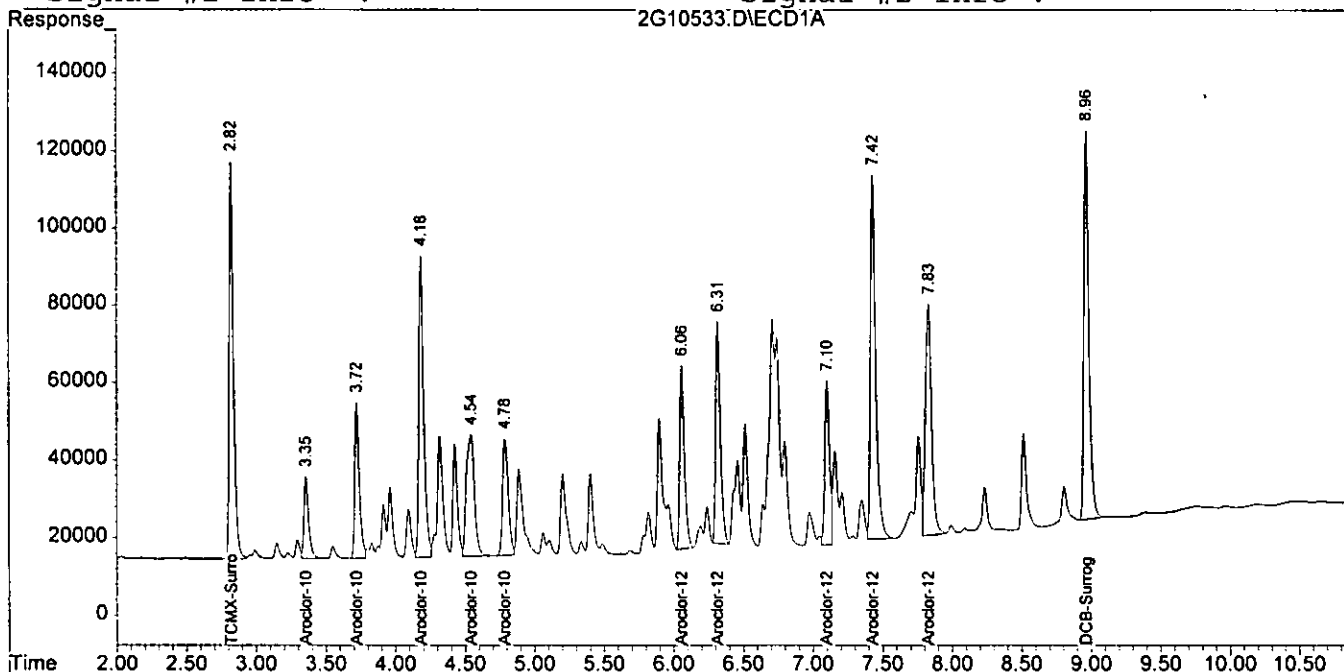
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10533.D\ECD1A.CH Vial: 11
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10533.D\ECD2B.CH
Acq On : 5 Aug 2005 10:59 Operator: JK
Sample : CAL1660@1000PPB Inst : gc_2
Misc : S,PCB:0.5 Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 11:26 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10580.D\ECD1A.CH Vial: 11
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10580.D\ECD2B.CH
 Acq On : 8 Aug 2005 8:12 Operator: JK
 Sample : CAL 1660@500PPB Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 8:20 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.82	2.79	986508	694718	50.507	47.586
2) Aroclor-1016 {1}	3.35	3.39	234180	137188	534.079	489.226
3) Aroclor-1016 {2}	3.71	3.80	432585	315082	523.101	502.653
4) Aroclor-1016 {3}	4.18	4.18	894698	636577	510.537	488.718
5) Aroclor-1016 {4}	4.54	4.50	609135	341761	533.800	489.377
6) Aroclor-1016 {5}	4.78	4.86	435755	214001	491.322	507.691
7) Aroclor-1260 {1}	6.05	6.17	527458	368422	523.412	475.417
8) Aroclor-1260 {2}	6.31	6.26	631644	405175	522.609	467.098
9) Aroclor-1260 {3}	7.09	7.39	467090	832177	522.273	491.551
10) Aroclor-1260 {4}	7.42	7.94	1137336	409966	516.001	489.409
11) Aroclor-1260 {5}	7.82	8.48	812101	282163	497.972	505.374
35) DCB-Surrogate	8.96	9.28	1166954	727865	51.655	48.202

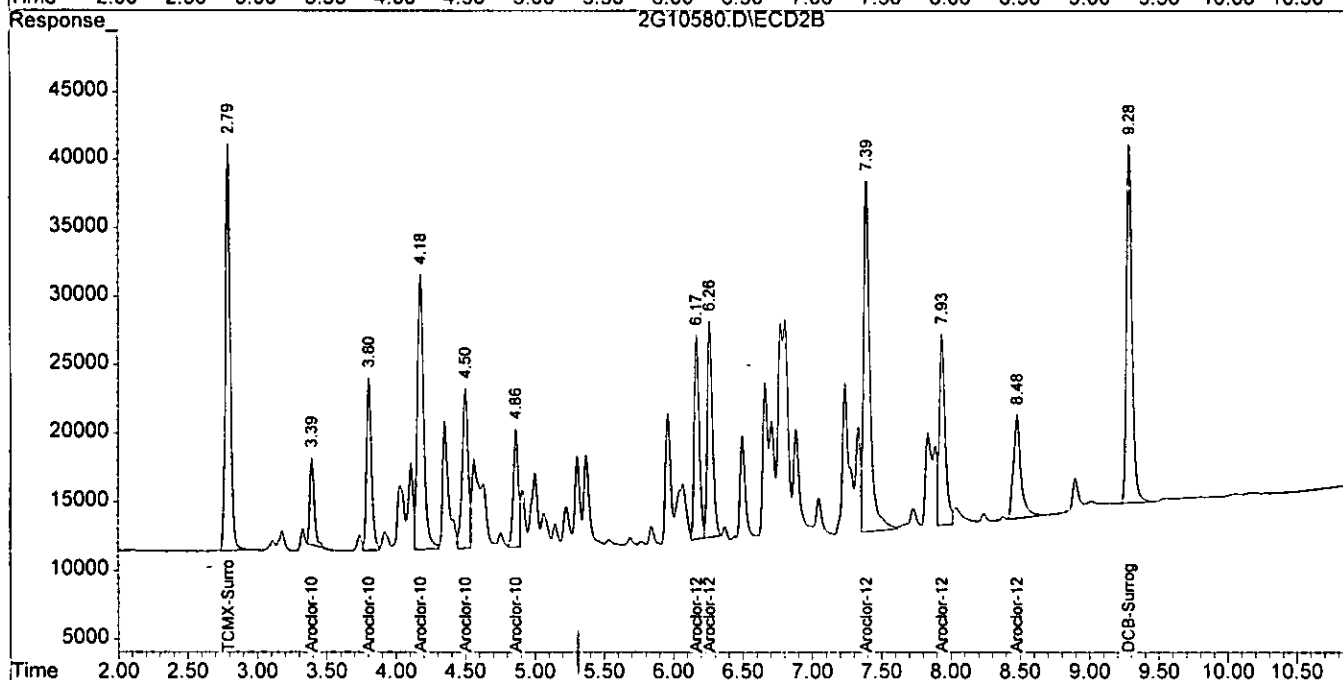
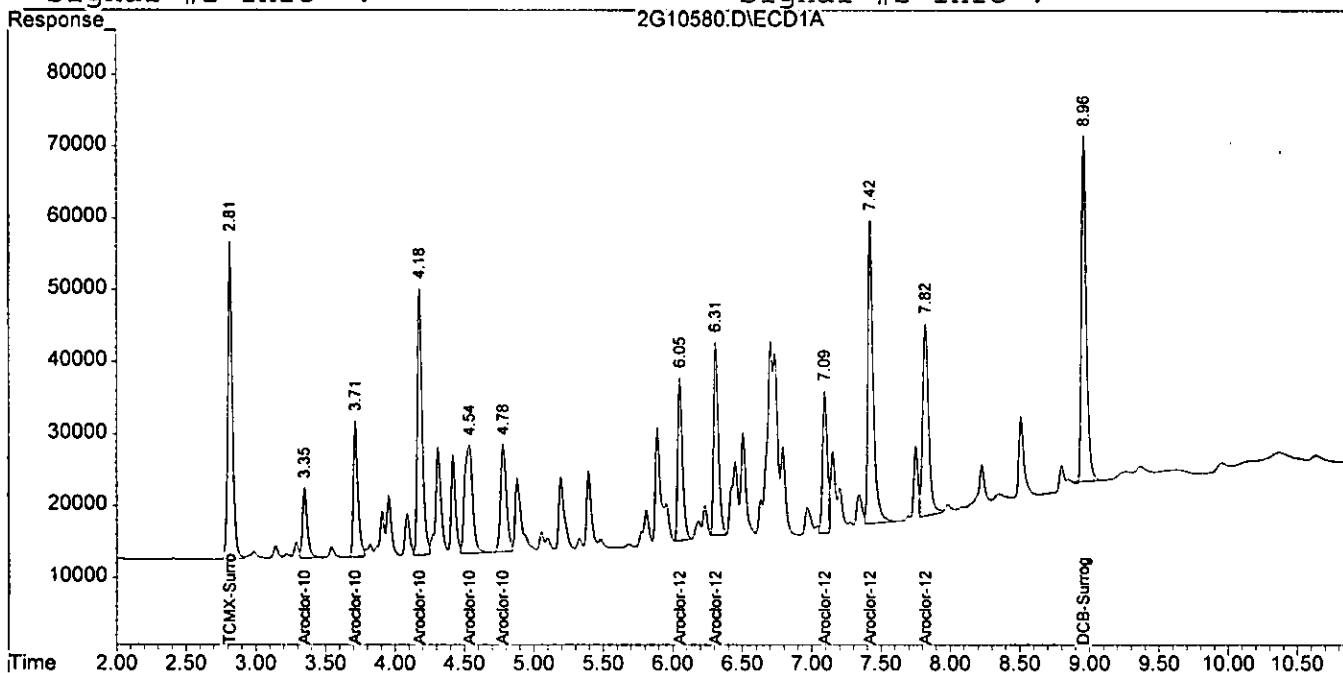
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10580.D\ECD1A.CH Vial: 7511
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10580.D\ECD2B.CH
Acq On : 8 Aug 2005 8:12 Operator: JK
Sample : CAL 1660@500PPB Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 8:20 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10600.D\ECD1A.CH Vial: 122
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10600.D\ECD2B.CH
 Acq On : 8 Aug 2005 13:25 Operator: JK
 Sample : CAL 1660@1000PPB Inst : gc_2
 Misc : S,PCB:0.5 Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 13:33 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	2135531	1409416	109.333	96.541
2) Aroclor-1016 {1}	3.36	3.40	467808	264741	1066.900	944.093
3) Aroclor-1016 {2}	3.72	3.81	835120	583176	1009.865	930.347
4) Aroclor-1016 {3}	4.18	4.19	1724915	1187242	984.280	911.478
5) Aroclor-1016 {4}	4.54	4.51	1137377	627979	996.712	1013.261
6) Aroclor-1016 {5}	4.79	4.87	812404	398333	1019.931	944.995
7) Aroclor-1260 {1}	6.06	6.18	954663	668252	947.340	862.323
8) Aroclor-1260 {2}	6.31	6.27	1144290	733068	946.762	845.103
9) Aroclor-1260 {3}	7.10	7.40	836173	1410734	934.960	833.294
10) Aroclor-1260 {4}	7.43	7.94	2098634	677240	952.134	808.475
11) Aroclor-1260 {5}	7.83	8.48	1763345	569231	1081.265	1019.535
35) DCB-Surrogate	8.96	9.29	2072699	1198590	95.465	79.375m

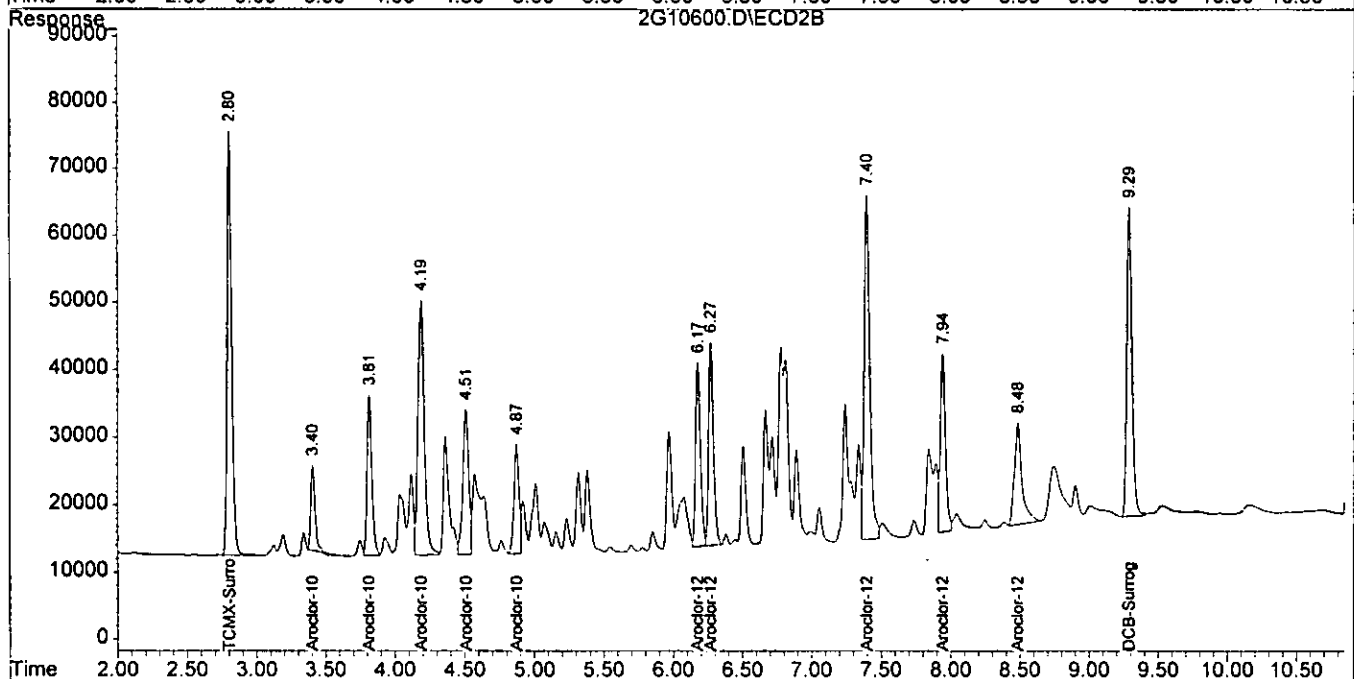
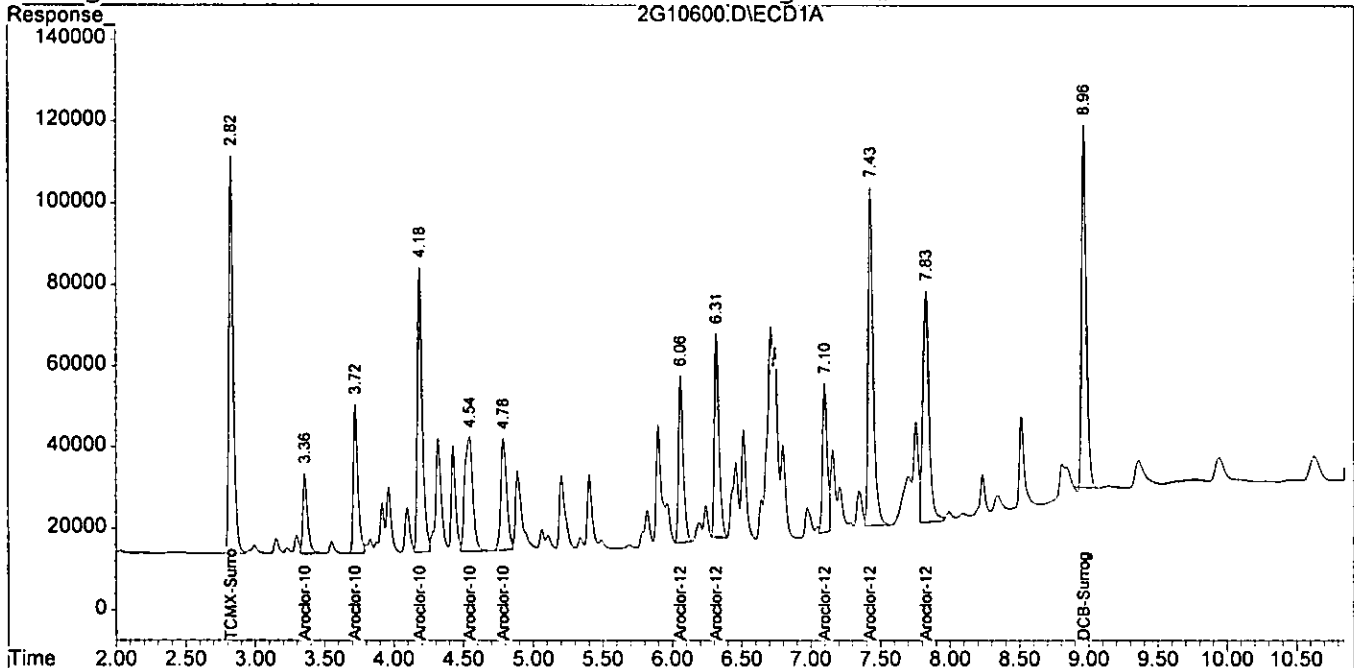
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10600.D\ECD1A.CH Vial: 22
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10600.D\ECD2B.CH
Acq On : 8 Aug 2005 13:25 Operator: JK
Sample : CAL 1660@1000PPB Inst : gc_2
Misc : S,PCB:0.5 Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 13:33 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10622.D\ECD1A.CH Vial: 21
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10622.D\ECD2B.CH
 Acq On : 8 Aug 2005 18:42 Operator: JK
 Sample : CAL 1660@2000PPB Inst : gc_2
 Misc : S,PCB:0.25 Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 5:03 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.82	2.80	4521259	3043659	231.476	208.482
2) Aroclor-1016 {1}	3.35	3.40	962762	561762	2195.714	2003.299
3) Aroclor-1016 {2}	3.72	3.81	1816712	1194654	2196.852	1905.843
4) Aroclor-1016 {3}	4.18	4.19	3719556	2488301	2122.473	1910.335
5) Aroclor-1016 {4}	4.54	4.51	2476483	1251530	2170.203	2292.372
6) Aroclor-1016 {5}	4.78	4.87	1693525	838998	2355.156	1990.415
7) Aroclor-1260 {1}	6.06	6.17	1972692	1419820	1957.561	1832.158
8) Aroclor-1260 {2}	6.31	6.27	2370794	1557956	1961.545	1796.059
9) Aroclor-1260 {3}	7.10	7.39	1724502	3276282	1928.238	1935.237
10) Aroclor-1260 {4}	7.42	7.94	4538544	1531126	2059.104	1827.825
11) Aroclor-1260 {5}	7.83	8.48	3263547	1074701	2001.173	1924.868
35) DCB-Surrogate	8.96	9.29	4518135	2539216	213.748	168.156

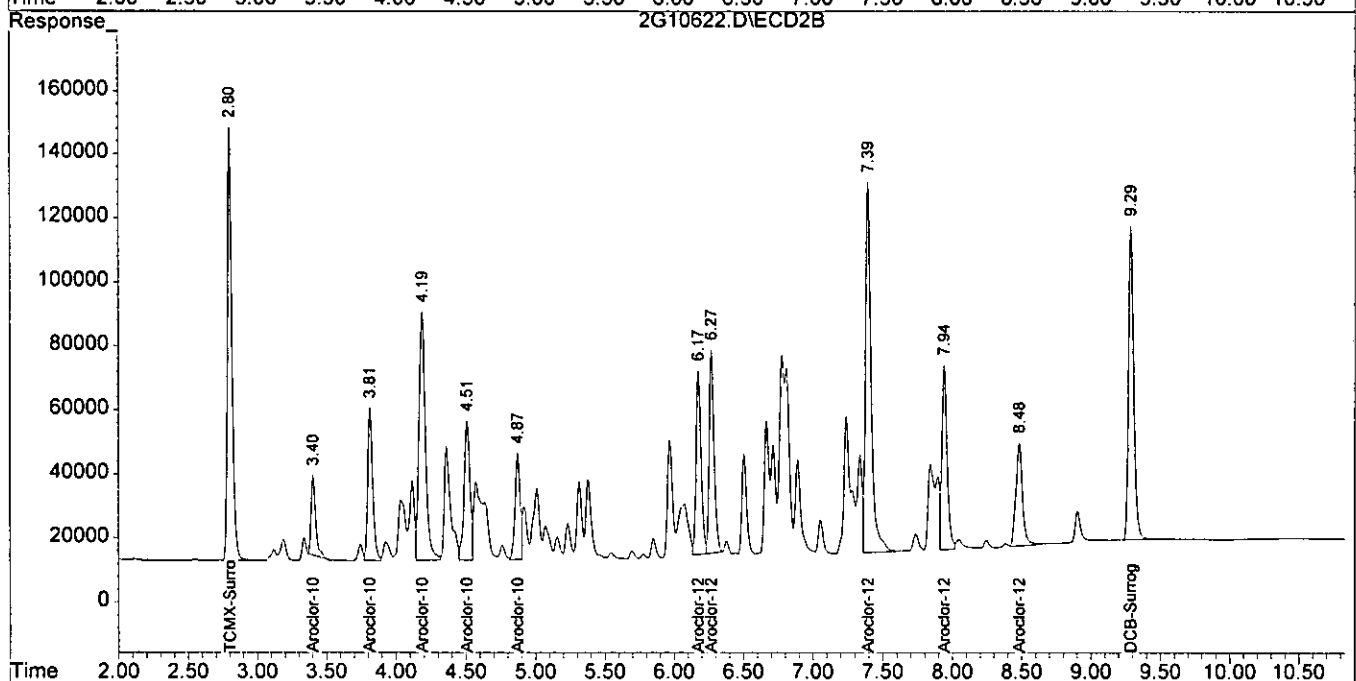
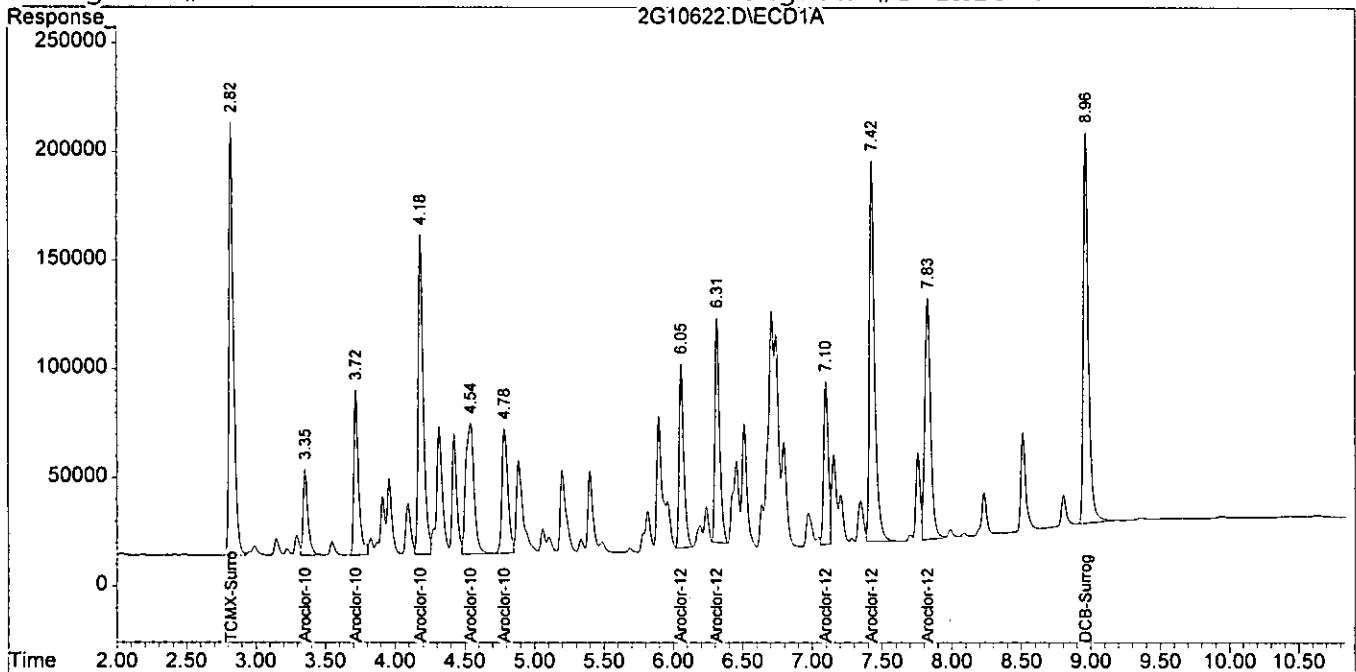
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10622.D\ECD1A.CH Vial: 921
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10622.D\ECD2B.CH
 Acq On : 8 Aug 2005 18:42 Operator: JK
 Sample : CAL 1660@2000PPB Inst : gc_2
 Misc : S,PCB:0.25 Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 9 5:03 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Multiple Level Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :



GC PCB Data
Raw QC Data

Form1

ORGANICS PCB REPORT

Sample Number: WMB2305
Client Id:
Data File: 2G10427.D
Analysis Date: 08/03/05 11:33
Date Rec/Extracted: NA-08/02/05

Matrix: Aqueous
Initial Vol: 1000ml
Final Vol: 5ml
Dilution: 1
Solids: 0

Units: ug/L

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.25	U	12672-29-6	Aroclor-1248	0.25	U
11104-28-2	Aroclor-1221	0.25	U	11097-69-1	Aroclor-1254	0.25	U
11141-16-5	Aroclor-1232	0.25	U	11096-82-5	Aroclor-1260	0.25	U
53469-21-9	Aroclor-1242	0.25	U				

Worksheet #: 18089

Total Target Concentration 0

*U - Indicates the compound was analyzed but not detected.
B - Indicates the analyte was found in the blank as well as in the sample.
E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10427.D\ECD1A.CH Vial: 1111
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10427.D\ECD2B.CH
 Acq On : 3 Aug 2005 11:33 Operator: JK
 Sample : WMB2305 Inst : gc_2
 Misc : A,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 3 11:49 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GCADATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 11:18:05 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.84	2.82	2643664	1681443	102.749	94.832
35) DCB-Surrogate	8.99	9.32	2846193	1601800	113.748	113.319

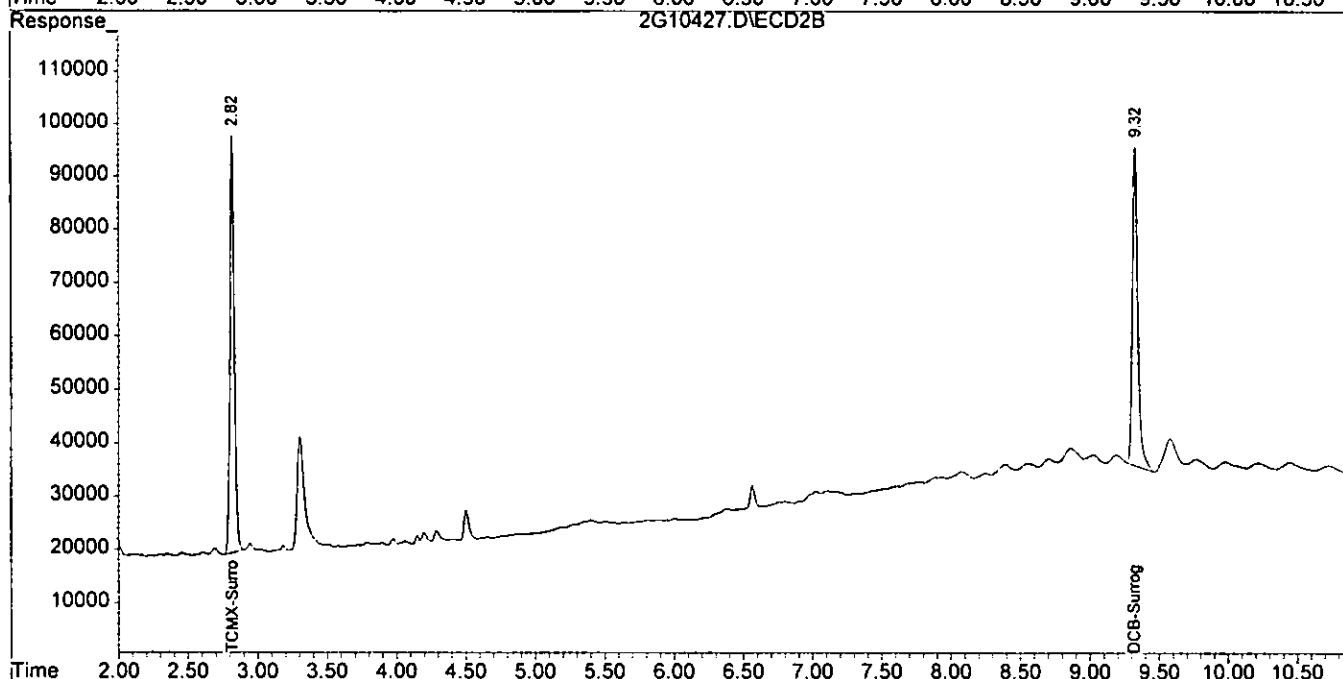
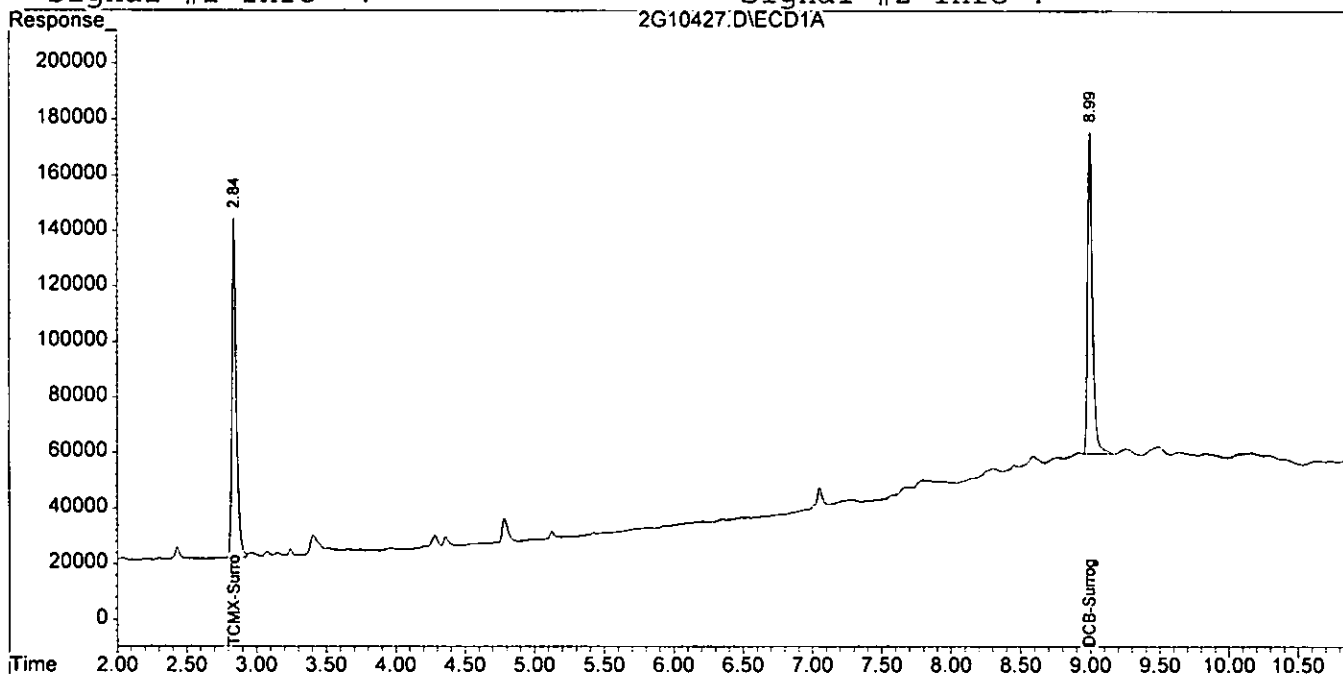
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10427.D\ECD1A.CH Vial: 111
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10427.D\ECD2B.CH
Acq On : 3 Aug 2005 11:33 Operator: JK
Sample : WMB2305 Inst : gc_2
Misc : A,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 3 11:49 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 11:18:05 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form1

ORGANICS PCB REPORT

Sample Number: SMB729B
 Client Id:
 Data File: 2G10591.D
 Analysis Date: 08/08/05 11:06
 Date Rec/Extracted: NA-08/07/05

Matrix: Soil
 Initial Vol: 20g
 Final Vol: 10ml
 Dilution: 1
 Solids: 100

Units: mg/Kg

Cas #	Compound	RL	Conc	Cas #	Compound	RL	Conc
12674-11-2	Aroclor-1016	0.025	U	12672-29-6	Aroclor-1248	0.025	U
11104-28-2	Aroclor-1221	0.025	U	11097-69-1	Aroclor-1254	0.025	U
11141-16-5	Aroclor-1232	0.025	U	11096-82-5	Aroclor-1260	0.025	U
53469-21-9	Aroclor-1242	0.025	U				

Worksheet #: 18089

Total Target Concentration 0

*U - Indicates the compound was analyzed but not detected.
 B - Indicates the analyte was found in the blank as well as in the sample.
 E - Indicates the analyte concentration exceeds the calibration range of the instrument.*

*R - Retention Time Out
 J - Indicates an estimated value when a compound is detected at less than the specified detection limit.*

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10591.D\ECD1A.CH Vial: 512
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10591.D\ECD2B.CH
 Acq On : 8 Aug 2005 11:06 Operator: JK
 Sample : SMB729B Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 11:39 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1912707	1229762	97.925	84.235
35) DCB-Surrogate	8.96	9.28	2400691	1351488	111.330	89.500

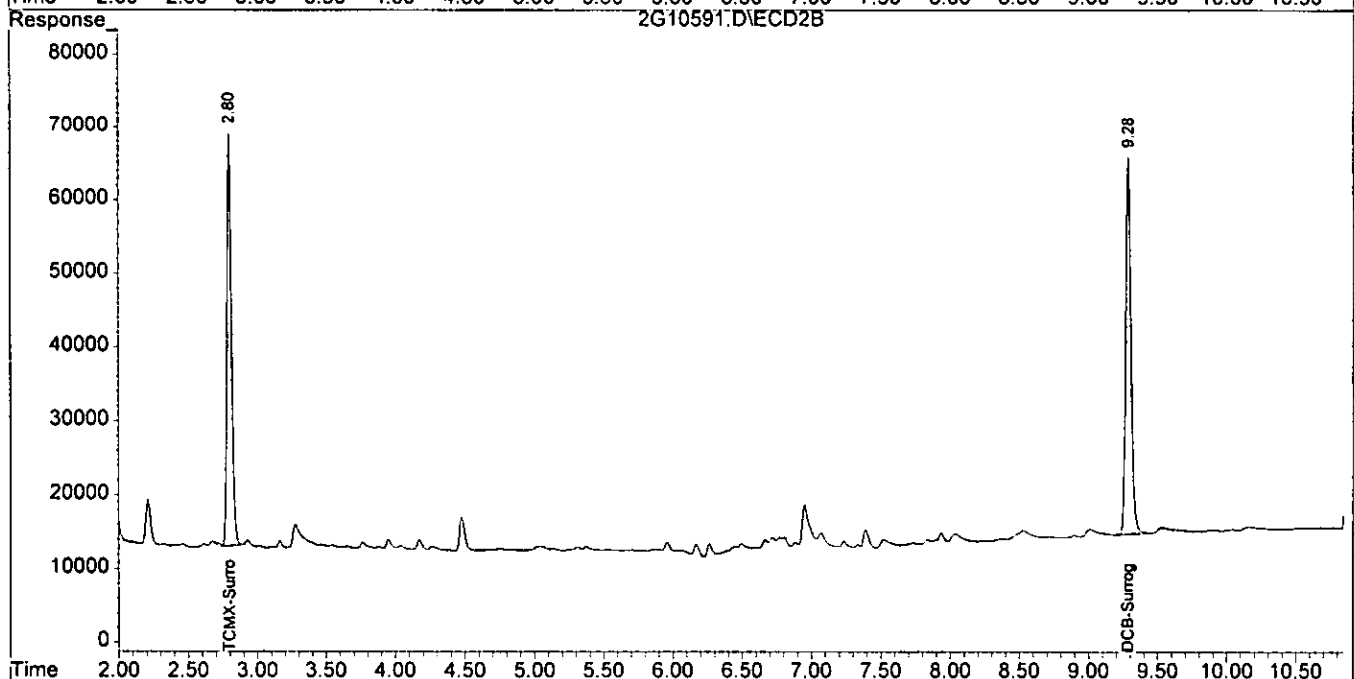
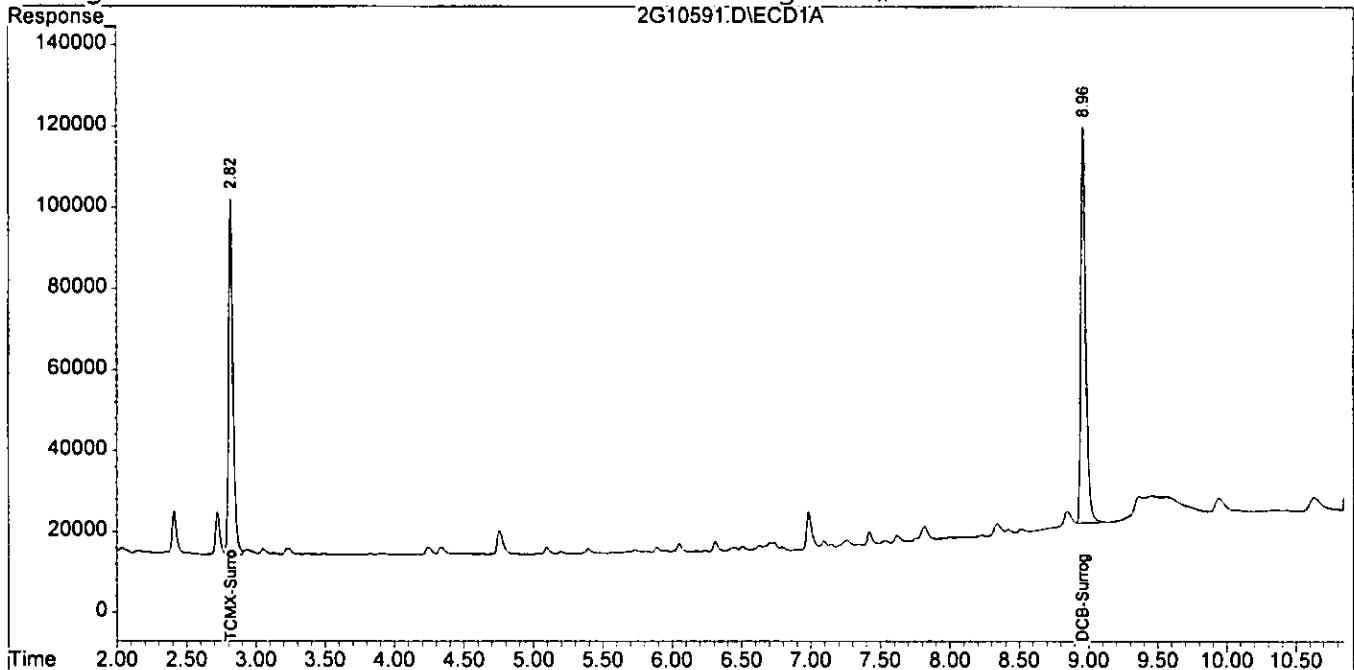
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10591.D\ECD1A.CH Vial: 12
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10591.D\ECD2B.CH
Acq On : 8 Aug 2005 11:06 Operator: JK
Sample : SMB729B Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 11:39 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Form3
MBS Data
Method: 8082

1155

Data File:====>		2G10592.D			2G10433.D								
Data/Batch/Sample ID:====>		SMB729B(MS)			WMB2305(MS)								
Date/Time:====>		08/08/05 11:20			08/03/05 13:14								
Compound	Limit(s)				Conc %			Conc %			Conc %		
	Soil	Aq	Col	Mr	Conc	Exp	Rec	Conc	Exp	Rec	Conc	Exp	Rec
Aroclor-1016	29-131	29-131	1	0	1039	1000	104	982.1	1000	98			
Aroclor-1260	29-131	29-131	1	0	1096	1000	110	957.8	1000	96			

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10592.D\ECD1A.CH Vial:
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10592.D\ECD2B.CH
 Acq On : 8 Aug 2005 11:20 Operator: JK
 Sample : SMB729B(MS) Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 8 11:40 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1916853	1233764	98.138	84.509
2) Aroclor-1016 {1}	3.35	3.40	445455	248489	1015.923	886.137
3) Aroclor-1016 {2}	3.72	3.81	823532	561969	995.853	896.515
4) Aroclor-1016 {3}	4.18	4.18	1784382	1174668	1018.214	901.824
5) Aroclor-1016 {4}	4.54	4.50	1199999	618439	1051.589	995.254
6) Aroclor-1016 {5}	4.78	4.87	876534	403590	1112.206	957.467
7) Aroclor-1260 {1}	6.05	6.17	1096337	733164	1087.928	946.086
8) Aroclor-1260 {2}	6.31	6.26	1315979	798882	1088.814	920.976
9) Aroclor-1260 {3}	7.09	7.39	958890	1673479	1072.175	988.492
10) Aroclor-1260 {4}	7.42	7.94	2467281	825799	1119.387	985.820
11) Aroclor-1260 {5}	7.82	8.48	1812968	595753	1111.694	1067.038
35) DCB-Surrogate	8.96	9.28	2440836	1356393	113.271	89.825

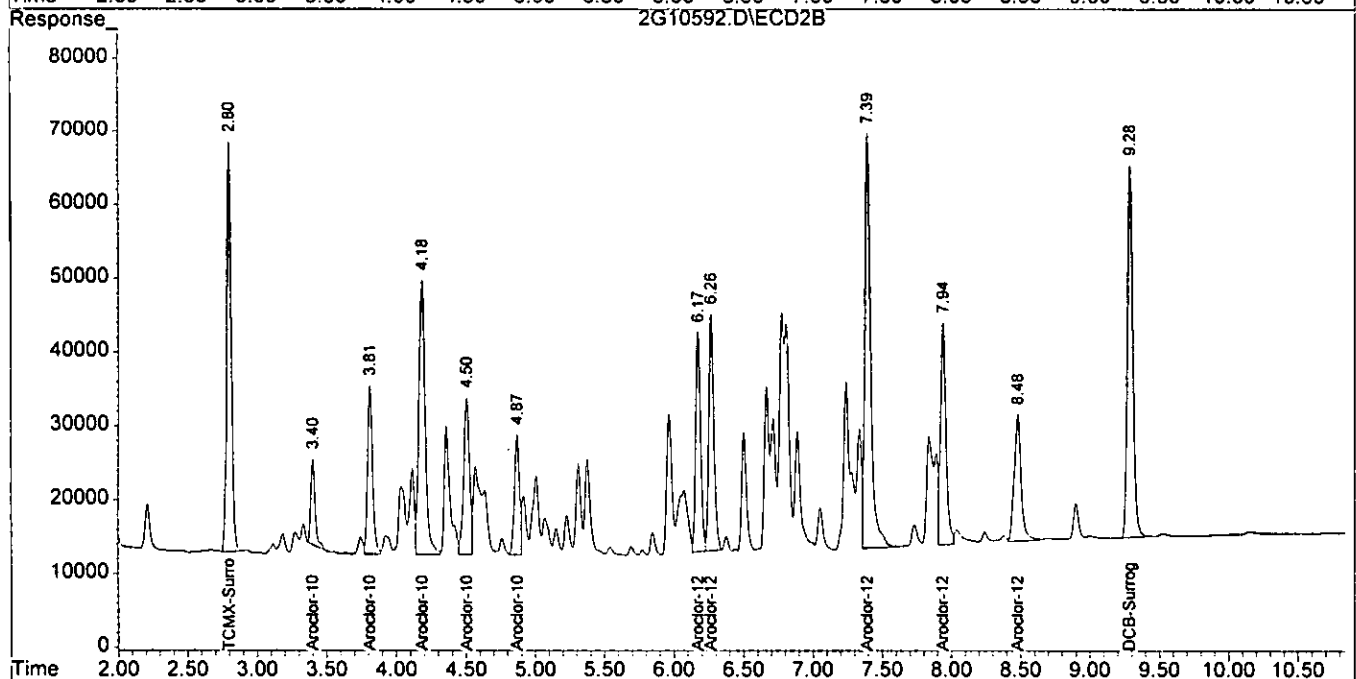
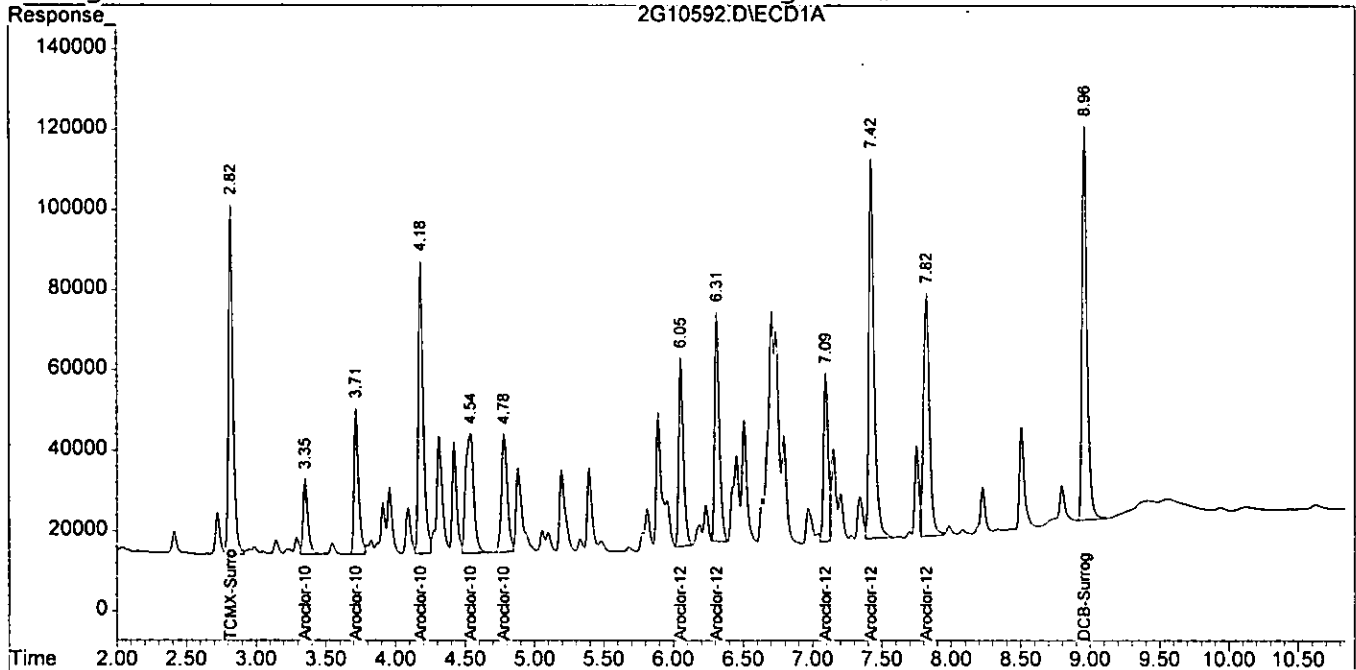
08/11/01

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10592.D\ECD1A.CH Vial:
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-08-05\2G10592.D\ECD2B.CH
Acq On : 8 Aug 2005 11:20 Operator: JK
Sample : SMB729B(MS) Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 8 11:40 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Data File : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10433.D\ECD1A.CH Vial: 1130
 Acq On : 3 Aug 2005 13:14 Operator: JK
 Sample : WMB2305 (MS) Inst : gc_2
 Misc : A,PCB Multiplr: 1.00
 IntFile : AUTOINT1.E

Data File : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10433.D\ECD2B.CH Vial: 30
 Acq On : 3 Aug 2005 13:14 Operator: JK
 Sample : WMB2305 Inst : gc_2
 Misc : A,PCB Multiplr: 1.00
 IntFile : AUTOINT2.E

Quant Time: Aug 3 13:35 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Wed Aug 03 11:18:05 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.84	2.82	2514332	1651288	97.723	93.131
2) Aroclor-1016 {1}	3.37	3.42	671196	398485	1254.906	1285.331m
3) Aroclor-1016 {2}	3.74	3.83	872804	610736	858.567	894.587
4) Aroclor-1016 {3}	4.20	4.21	1976892	1241781	927.545	879.181
5) Aroclor-1016 {4}	4.57	4.53	1272740	677762	906.582	925.538
6) Aroclor-1016 {5}	4.81	4.89	952439	433308	962.724	860.340
7) Aroclor-1260 {1}	6.08	6.20	1101281	768043	934.781	910.722
8) Aroclor-1260 {2}	6.34	6.29	1354597	865151	956.546	915.356
9) Aroclor-1260 {3}	7.13	7.42	981654	1952765	949.551	1126.114
10) Aroclor-1260 {4}	7.45	7.97	2517867	797353	992.809	964.924
11) Aroclor-1260 {5}	7.86	8.51	1726477	543036	955.255	1008.374m
35) DCB-Surrogate	8.99	9.32	2587969	1364670	103.428	96.543

02/11/05

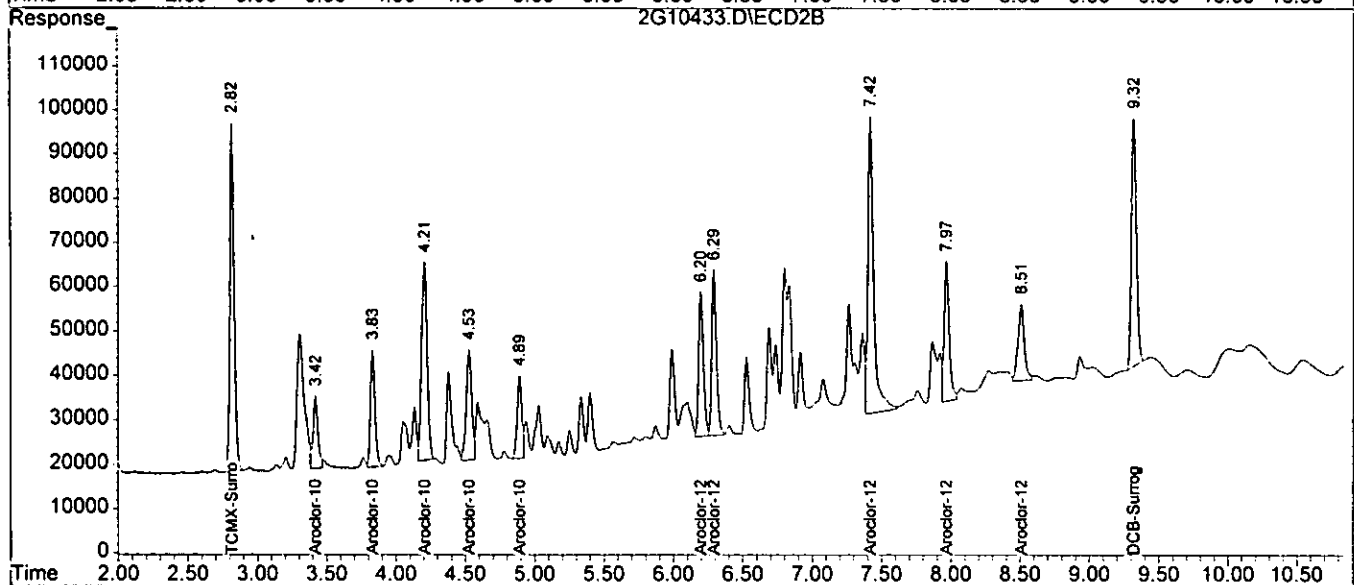
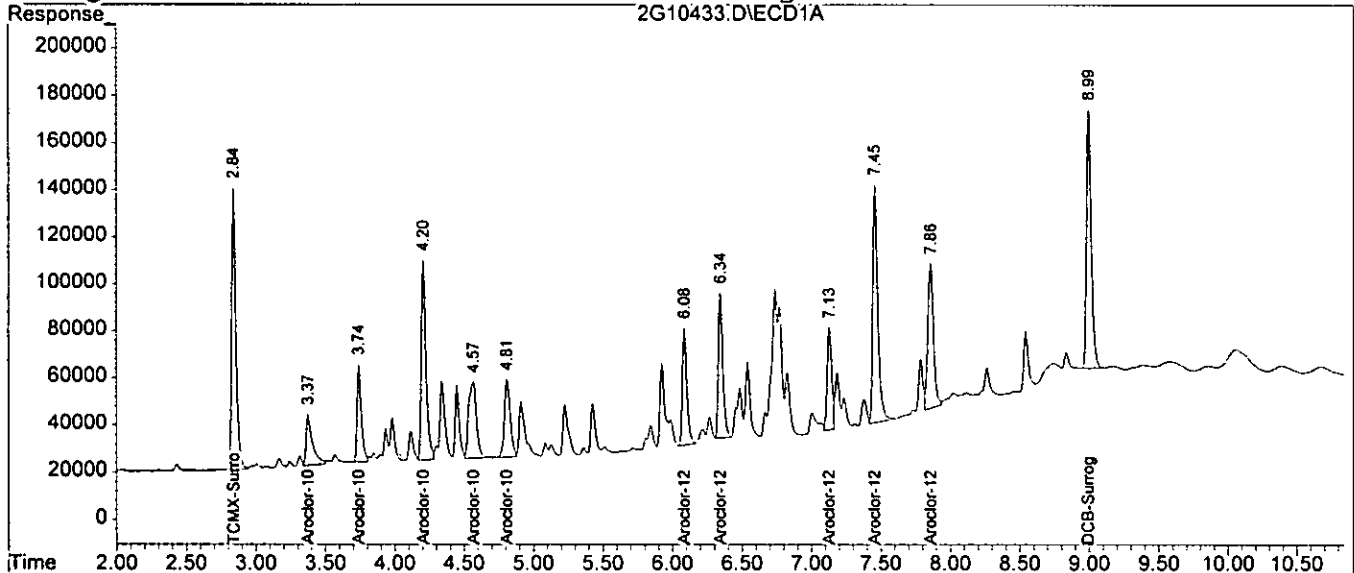
Quantitation Report

Data File : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10433.D\ECD1A.CH Vial: 30
Acq On : 3 Aug 2005 13:14 Operator: JK
Sample : WMB2305 (MS) Inst : gc_2
Misc : A, PCB Multiplr: 1.00
IntFile : AUTOINT1.E

Data File : G:\Gcdata\2005\Gc_2\Data\08-03-05\2G10433.D\ECD2B.CH Vial: 30
Acq On : 3 Aug 2005 13:14 Operator: JK
Sample : WMB2305 Inst : gc_2
Misc : A, PCB Multiplr: 1.00
IntFile : AUTOINT2.E
Quant Time: Aug 3 13:35 2005 Quant Results File: 2G_C0803.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0803.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Wed Aug 03 11:18:05 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10514.D\ECD1A.CH Vial: 112
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10514.D\ECD2B.CH
 Acq On : 5 Aug 2005 6:25 Operator: JK
 Sample : SMB727B(MS) Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 8:20 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2

Target Compounds						
1) TCMX-Surrogate	2.81	2.79	1469870	1054964	75.253	72.262
2) Aroclor-1016 {1}	3.35	3.39	385104	234399	878.284	835.889
3) Aroclor-1016 {2}	3.71	3.80	717397	536631	867.509	856.093
4) Aroclor-1016 {3}	4.18	4.18	1579748	1156528	901.444	887.897
5) Aroclor-1016 {4}	4.53	4.50	1069947	619128	937.621	996.553
6) Aroclor-1016 {5}	4.78	4.86	792697	401314	991.714	952.066
7) Aroclor-1260 {1}	6.05	6.17	976740	748553	969.248	965.945
8) Aroclor-1260 {2}	6.31	6.26	1186735	815592	981.880	940.239
9) Aroclor-1260 {3}	7.09	7.39	883809	1644705	988.225	971.496
10) Aroclor-1260 {4}	7.42	7.93	2216322	858649	1005.529	1025.036
11) Aroclor-1260 {5}	7.82	8.48	1620862	665603	993.896	1192.144
35) DCB-Surrogate	8.96	9.28	2156267	1439376	99.507	95.321

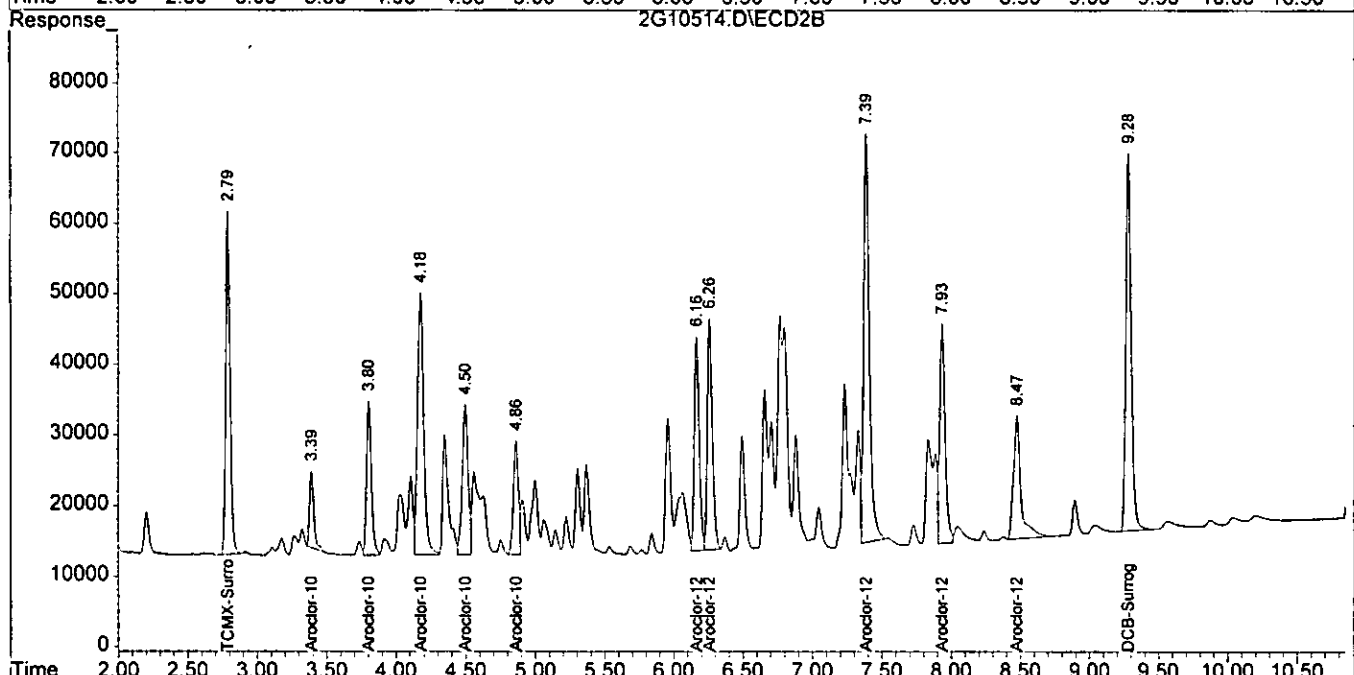
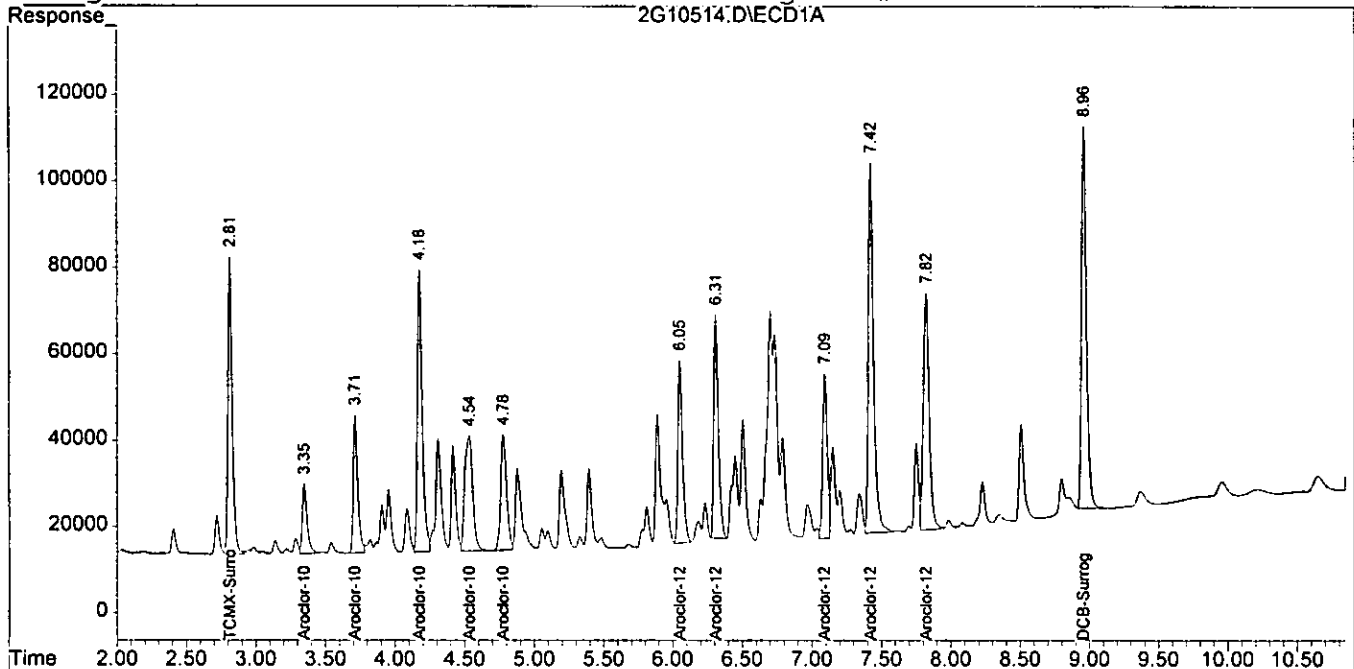
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10514.D\ECD1A.CH Vial: 2
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10514.D\ECD2B.CH Vial: 2
Acq On : 5 Aug 2005 6:25 Operator: JK
Sample : SMB727B(MS) Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 8:20 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



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Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10525.D\ECD1A.CH Vial:
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10525.D\ECD2B.CH
 Acq On : 5 Aug 2005 9:04 Operator: JK
 Sample : AC18778-020 (MS) Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 9:10 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GCDATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1983864	1391349	101.568	95.303
2) Aroclor-1016 {1}	3.35	3.40	483369	264524	1102.390	943.319
3) Aroclor-1016 {2}	3.72	3.81	848952	610658	1026.592	974.189
4) Aroclor-1016 {3}	4.18	4.18	1834085	1264634	1046.576	970.893
5) Aroclor-1016 {4}	4.54	4.51	1207256	609110	1057.948	977.685
6) Aroclor-1016 {5}	4.78	4.87	794615	436713	994.458	1036.047
7) Aroclor-1260 {1}	6.05	6.17	1093500	785353	1085.113	1013.431
8) Aroclor-1260 {2}	6.31	6.26	1326582	979138	1097.586	1128.781
9) Aroclor-1260 {3}	7.10	7.39	966849	1750251	1081.075	1033.840
10) Aroclor-1260 {4}	7.42	7.94	2487261	863412	1128.452	1030.723
11) Aroclor-1260 {5}	7.82	8.48	1856944	597086	1138.659	1069.425
35) DCB-Surrogate	8.96	9.29	2448695	1425849	113.652	94.425

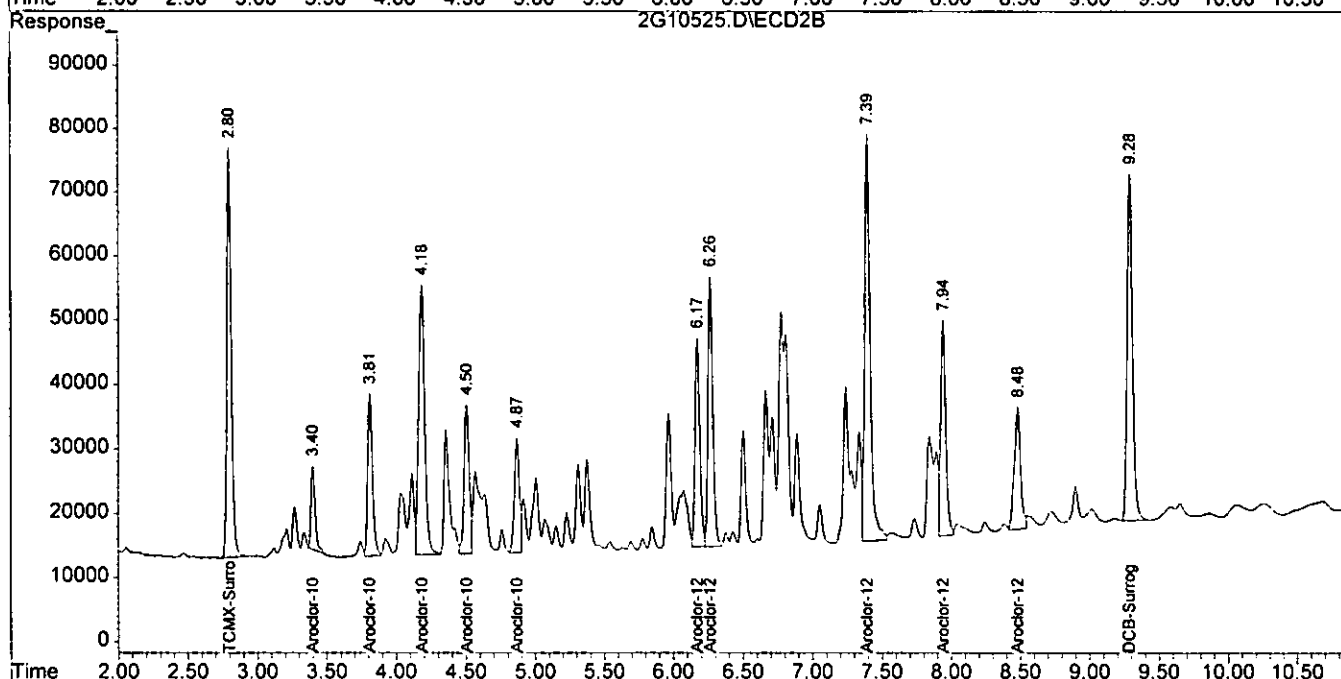
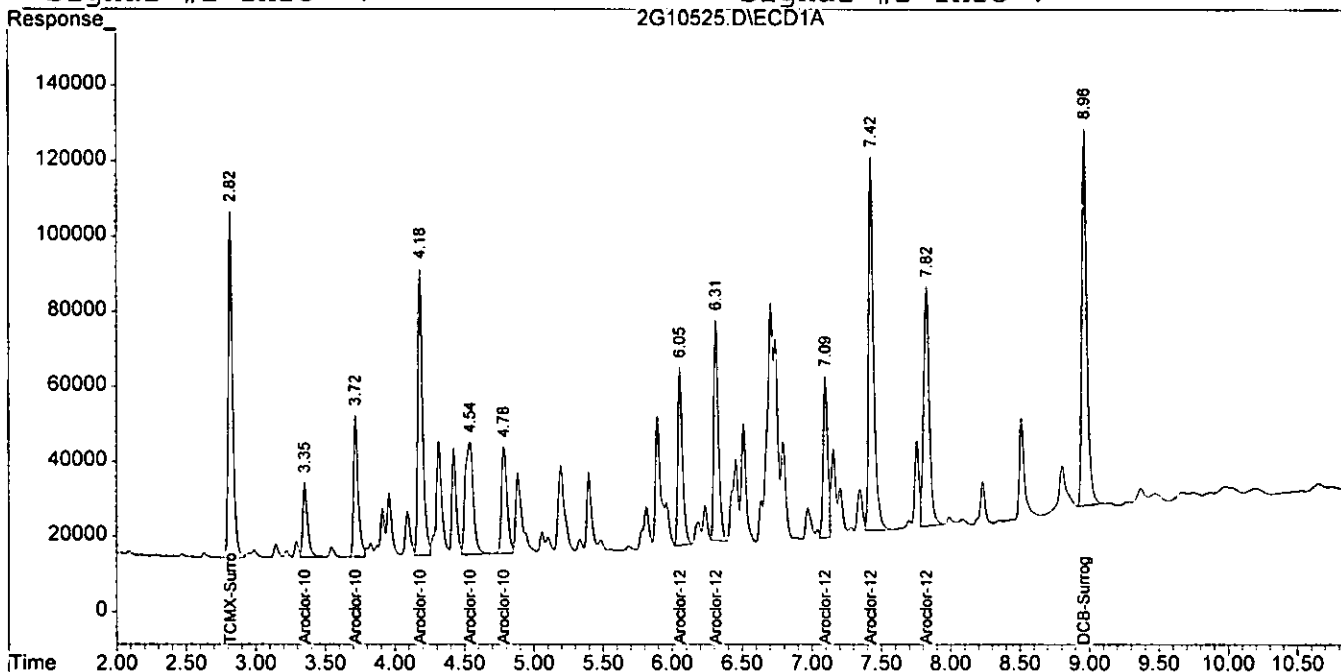
08/11/05

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10525.D\ECD1A.CH Vial: 23
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10525.D\ECD2B.CH
Acq On : 5 Aug 2005 9:04 Operator: JK
Sample : AC18778-020 (MS) Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 9:10 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10526.D\ECD1A.CH Vial: 1424
 Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10526.D\ECD2B.CH
 Acq On : 5 Aug 2005 9:18 Operator: JK
 Sample : AC18778-020 (MSD) Inst : gc_2
 Misc : S,PCB Multiplr: 1.00
 IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
 Quant Time: Aug 5 9:27 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
 Title : @GC_2,ug,608,8082
 Last Update : Fri Aug 05 07:46:38 2005
 Response via : Initial Calibration
 DataAcq Meth : 2G_8081.M

Volume Inj. :
 Signal #1 Phase : Signal #2 Phase:
 Signal #1 Info : Signal #2 Info :

Compound	RT#1	RT#2	Resp#1	Resp#2	pg#1	pg#2
Target Compounds						
1) TCMX-Surrogate	2.82	2.80	1060991	760305	54.320	52.079
2) Aroclor-1016 {1}	3.35	3.40	279655	138820	637.793	495.046
3) Aroclor-1016 {2}	3.72	3.81	466926	343393	564.628	547.818
4) Aroclor-1016 {3}	4.18	4.18	989761	709620	564.783	544.794
5) Aroclor-1016 {4}	4.54	4.51	631606	333612	553.492	474.927
6) Aroclor-1016 {5}	4.78	4.87	425635	243588	477.416	577.882
7) Aroclor-1260 {1}	6.06	6.17	598479	446659	593.889	576.375
8) Aroclor-1260 {2}	6.31	6.26	726746	756012	601.294	871.554 #
9) Aroclor-1260 {3}	7.10	7.39	519320	998068	580.674	589.540
10) Aroclor-1260 {4}	7.42	7.94	1304473	489413	591.830	584.250
11) Aroclor-1260 {5}	7.83	8.48	988796	340338	606.319	609.571
35) DCB-Surrogate	8.96	9.29	1264018	768070	56.350	50.864

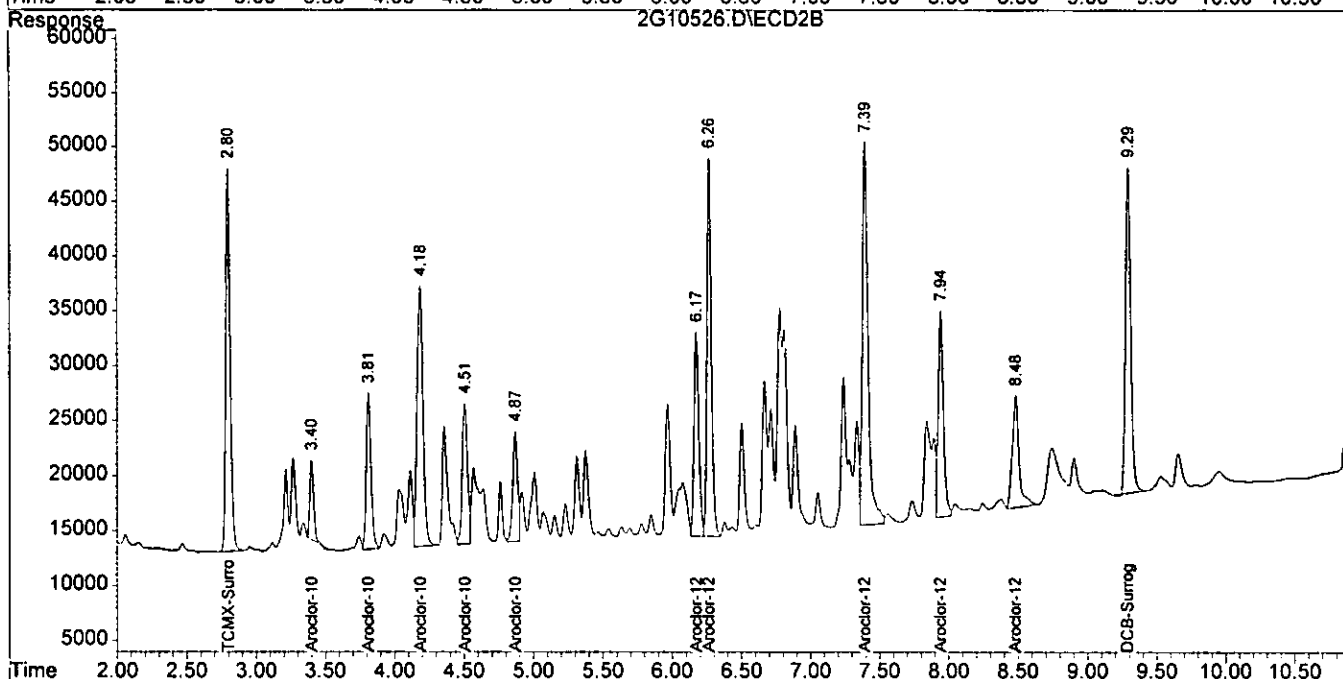
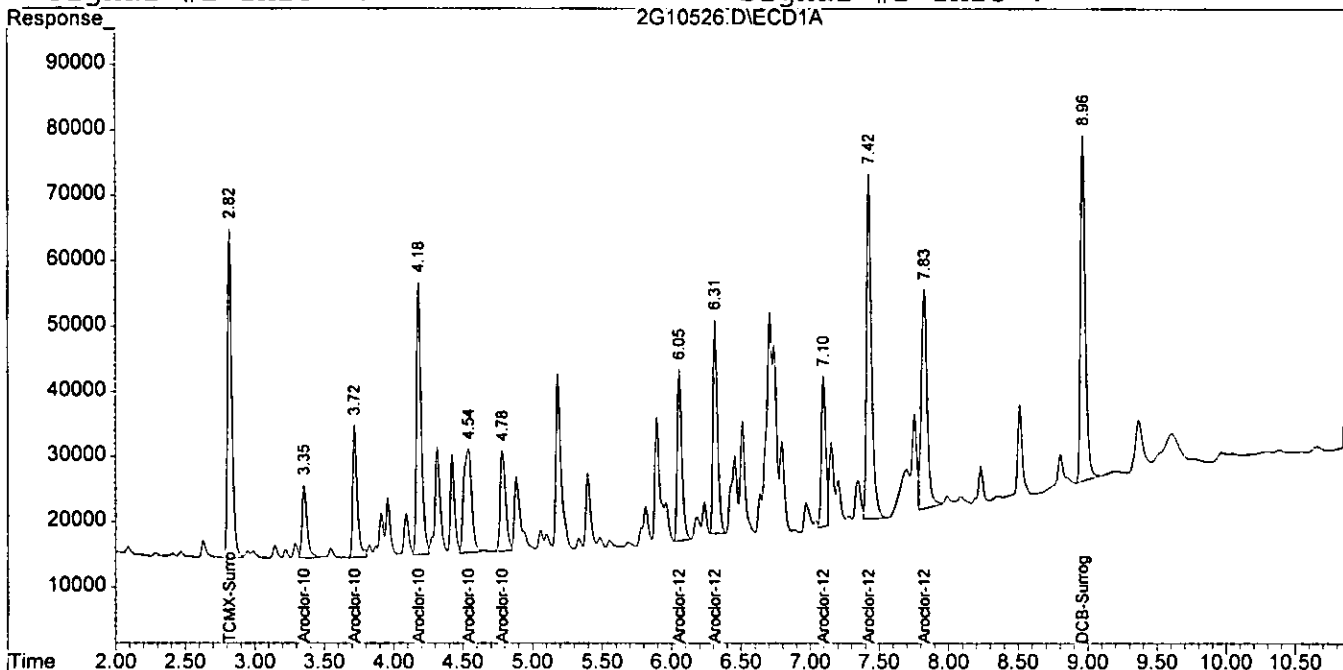
03/11/01

Quantitation Report

Signal #1 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10526.D\ECD1A.CH Vial: 1014
Signal #2 : G:\Gcdata\2005\Gc_2\Data\08-05-05\2G10526.D\ECD2B.CH
Acq On : 5 Aug 2005 9:18 Operator: JK
Sample : AC18778-020 (MSD) Inst : gc_2
Misc : S,PCB Multiplr: 1.00
IntFile Signal #1: AUTOINT1.E IntFile Signal #2: AUTOINT2.E
Quant Time: Aug 5 9:27 2005 Quant Results File: 2G_C0805.RES

Quant Method : G:\GC\DATA\2005\GC_2\METHODS\2G_C0805.M (Chemstation Integr
Title : @GC_2,ug,608,8082
Last Update : Fri Aug 05 07:46:38 2005
Response via : Multiple Level Calibration
DataAcq Meth : 2G_8081.M

Volume Inj. :
Signal #1 Phase : Signal #2 Phase:
Signal #1 Info : Signal #2 Info :



**GC PCB Data
Extraction/Logbook Data**

RUN LOG

Instrument: GC_2 Year: 2005

Analyst: JK

8000

Main data table with columns: Data File, Sample Number, Flags, Comments, Test Group, Matrix, Surr Dil, Sam Dil, Method(s), Analysis Date, IniCal, Cal 600, Beg Cal, End Cal, BlkFile. Rows include sample IDs like 2G10408, 2G10410, etc., and various chemical analysis details.

Summary table with columns: Abc, Area, B5m, B8m, Bnf, C16, C8i, Cme, Cn, D1o,D2o, Dnc, Do, Eba, Emp, En. Rows list various flags and their corresponding descriptions, such as 'Area Not Checked' and 'Extraction Performed Past Hold'.

RUN LOG

Instrument: GC_2

Year: 2005

Analyst: JK

8000

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	Cal 600	8000 Beg Cal	End Cal	BlkFile
2G10502	CAL 1660@50PPB	I26	IsC16C26C18C28		Soil	1	1	608 8082	08/05 02:34	2G10062				
2G10503	CAL 1660@50PPB	I26			Soil	1	1	608 8082	08/05 02:48	2G10503				
2G10504	CAL 1660@200PPB	I26			Soil	1	1	608 8082	08/05 03:02	2G10503				
2G10505	CAL 1660@500PPB	I26			Soil	1	1	608 8082	08/05 03:17	2G10503				
2G10506	CAL 1660@1000PPB	I26			Soil	1	1	608 8082	08/05 03:31	2G10503				
2G10507	CAL 1660@2000PPB	I26			Soil	1	1	608 8082	08/05 03:46	2G10503				
2G10508	CAL 1660@4000PPB	I26			Soil	1	1	608 8082	08/05 04:00	2G10503				
2G10509	CAL 2154@500PPB	I26			Soil	1	1	608 8082	08/05 04:15	2G10503				
2G10510	CAL 1248@500PPB	I26			Soil	1	1	608 8082	08/05 04:29	2G10503				
2G10511	CAL 1242@500PPB	I26			Soil	1	1	608 8082	08/05 04:43	2G10503				
2G10512	CAL 1232@500PPB	I26			Soil	1	1	608 8082	08/05 04:58	2G10503				
2G10513	SMB727B				Soil	1	1	8082	08/05 06:11	2G10503		2G10503	2G10533	
2G10514	SMB727B(MS)		SMB727B		Soil	1	1	8082	08/05 06:25	2G10503		2G10503	2G10533	
2G10515	AC18737-033			PCB-8082	Soil	1	1	8082	08/05 06:40	2G10503		2G10503	2G10533	
2G10516	AC18919-001			PCB-8082	Soil	1	1	8082	08/05 06:54	2G10503		2G10503	2G10533	
2G10517	AC18919-002			PCB-8082	Soil	1	1	8082	08/05 07:09	2G10503		2G10503	2G10533	
2G10518	AC18919-003			PCB-8082	Soil	1	1	8082	08/05 07:23	2G10503		2G10503	2G10533	
2G10519	SMB2405				Soil	1	1	8082	08/05 07:37	2G10503		2G10503	2G10533	
2G10520	SMB2405(MS)		SMB2405		Soil	1	1	8082	08/05 07:52	2G10503		2G10503	2G10533	
2G10521	AC18876-002(MS)	M18	SMB2405		Soil	1	1	8082	08/05 08:06	2G10503		2G10503	2G10533	
2G10522	AC18876-002(MSD)	M18	SMB2405		Soil	1	1	8082	08/05 08:21	2G10503		2G10503	2G10533	
2G10523	AC18876-002		SMB2405		Soil	1	1	8082	08/05 08:35	2G10503		2G10503	2G10533	
2G10524	AC18876-001			PCB-8082	Soil	1	1	8082	08/05 08:50	2G10503		2G10503	2G10533	
2G10525	AC18778-020(MS)			PCB-8082	Soil	1	1	8082	08/05 09:04	2G10503		2G10503	2G10533	
2G10526	AC18778-020(MSD)	R18R28	SMB727B		Soil	1	1	8082	08/05 09:18	2G10503		2G10503	2G10533	
2G10527	AC18778-020		SMB727B		Soil	1	1	8082	08/05 09:33	2G10503		2G10503	2G10533	
2G10528	AC18737-033(10X)			PCB-8082	Soil	10	10	8082	08/05 09:47	2G10503		2G10503	2G10533	
2G10529	AC18778-010			PCB-8082	Soil	1	1	8082	08/05 10:02	2G10503		2G10503	2G10533	
2G10530	AC18778-011			PCB-8082	Soil	1	1	8082	08/05 10:16	2G10503		2G10503	2G10533	
2G10531	AC18778-012			PCB-8082	Soil	1	1	8082	08/05 10:30	2G10503		2G10503	2G10533	
2G10532	AC18778-013			PCB-8082	Soil	1	1	8082	08/05 10:45	2G10503		2G10503	2G10533	
2G10533	CAL 1660@1000PPB	I26			Soil	0.5	1	608 8082	08/05 10:59	2G10503				
2G10534	AC18778-014			PCB-8082	Soil	1	1	8082	08/05 11:17	2G10503		2G10533	2G10547	
2G10535	AC18778-003(R)			PCB-8082	Soil	1	1	8082	08/05 11:32	2G10503		2G10533	2G10547	
2G10536	TEST0805				Soil	1	1	8082	08/05 11:46	2G10503		2G10533	2G10547	
2G10537	AC18778-024			PCB-8082	Soil	1	1	8082	08/05 12:01	2G10503		2G10533	2G10547	
2G10538	AC18778-016			PCB-8082	Soil	1	1	8082	08/05 12:15	2G10503		2G10533	2G10547	
2G10539	18786-009				Soil	1	1	8082	08/05 12:30	2G10503		2G10533	2G10547	
2G10540	AC18778-018			PCB-8082	Soil	1	1	8082	08/05 13:17	2G10503		2G10533	2G10547	
2G10541	AC18778-019			PCB-8082	Soil	1	1	8082	08/05 13:46	2G10503		2G10533	2G10547	
2G10542	AC18919-001			PCB-8082	Soil	1	1	8082	08/05 14:00	2G10503		2G10533	2G10547	
2G10543	AC18919-002			PCB-8082	Soil	1	1	8082	08/05 14:14	2G10503		2G10533	2G10547	
2G10544	AC18919-003			PCB-8082	Soil	1	1	8082	08/05 14:29	2G10503		2G10533	2G10547	
2G10545	AC18778-021			PCB-8082	Soil	1	1	8082	08/05 14:43	2G10503		2G10533	2G10547	
2G10546	AC18778-023			PCB-8082	Soil	1	1	8082	08/05 14:58	2G10503		2G10533	2G10547	
2G10547	AC18778-022			PCB-8082	Soil	1	1	8082	08/05 15:12	2G10503		2G10533	2G10547	
2G10548	AC18778-015			PCB-8082	Soil	1	1	8082	08/05 15:29	2G10503		2G10533	2G10547	
2G10549	AC18778-017			PCB-8082	Soil	1	1	8082	08/05 15:43	2G10503		2G10533	2G10547	
2G10550	CAL 1660@2000PPB	I26			Soil	0.25	1	608 8082	08/05 15:58	2G10503				
2G10551	2000PPB				Soil	0.25	1	8082	08/05 16:12	2G10503		2G10547	2G10569	
2G10552	2000PPB				Soil	0.25	1	8082	08/05 16:26	2G10503		2G10547	2G10569	
2G10553	AC18778-014(R)			PCB-8082	Soil	1	1	8082	08/05 16:41	2G10503		2G10547	2G10569	
2G10554	AC18778-024(R)			PCB-8082	Soil	1	1	8082	08/05 16:55	2G10503		2G10547	2G10569	
2G10555	test0805				Soil	1	1	8082	08/05 17:10	2G10503		2G10547	2G10569	
2G10556	AC18778-003(R)			PCB-8082	Soil	1	1	8082	08/05 17:24	2G10503		2G10547	2G10569	
2G10557	AC18778-010			PCB-8082	Soil	1	1	8082	08/05 17:39	2G10503		2G10547	2G10569	
2G10558	AC18778-011			PCB-8082	Soil	1	1	8082	08/05 17:53	2G10503		2G10547	2G10569	
2G10559	AC18778-012			PCB-8082	Soil	1	1	8082	08/05 18:07	2G10503		2G10547	2G10569	
2G10560	AC18778-013			PCB-8082	Soil	1	1	8082	08/05 18:22	2G10503		2G10547	2G10569	
2G10561	AC18778-014			PCB-8082	Soil	1	1	8082	08/05 18:36	2G10503		2G10547	2G10569	
2G10562	AC18778-015			PCB-8082	Soil	1	1	8082	08/05 18:51	2G10503		2G10547	2G10569	
2G10563	AC18778-016			PCB-8082	Soil	1	1	8082	08/05 19:05	2G10503		2G10547	2G10569	
2G10564	AC18778-017			PCB-8082	Soil	1	1	8082	08/05 19:19	2G10503		2G10547	2G10569	
2G10565	AC18778-018			PCB-8082	Soil	1	1	8082	08/05 19:34	2G10503		2G10547	2G10569	
2G10566	AC18778-019			PCB-8082	Soil	1	1	8082	08/05 19:48	2G10503		2G10547	2G10569	
2G10567	AC18778-020			PCB-8082	Soil	1	1	8082	08/05 20:03	2G10503		2G10547	2G10569	

Acc	Area Not Checked	Fn	Filtration Performed Post Hold	Ca	Warning Possible Carry Over
Am	Area Out	Fm	Solvent Extraction Date Missing/Not checked	R1 R2R	Ret Out on Method (roll) and/or not? 8000 series
RAm	Blank 8000 series missing	Fin	Tdn/Solvent Extraction Date Missing/Not checked	R1 R2A	Retention Time Out Or %Dfif Out
BAm	Blank 8000 series missing	Elo	Tdo Extraction Performed Outside of Hold	Rn	Can't Calculate Dfif
Br	Blank Not Found/Assigned	Ev	Eval Time Exceeded	S6	8000 series surrogate out
Brf	Blank Not Found/Assigned	Hb	Analysis Before Collection Date	S8	8000 series surrogate out
CAI	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	Sa6 Sb6	Acid and or BN Surrogate Out (800 series)
CAI	Calibration Column 1 Out (8000 Series)	H18 I28	Initial cal 8000 series failed: Column 1 and/or 2	Sa8 Sb8	Acid and or BN Surrogate Out (8000 series)
CAI	Calibration Column 2 Out (8000 Series)	Is	Initial Cal Not Checked	Sr	Surrogate Diluted Out
CAI	Calibration Column 2 Out (8000 Series)	Iv	Prob with extra cov for init calibration check r/s	Sm	Surrogate Not Checked
CAI	8000 series sample/blank did not have baseline cal	Iw	Initial cal warning. Ini cal file > method	T15	Outside of 500 series: Time Time
CAI	8000 series sample/blank did not have baseline cal	Iy	Initial Cal File Not Indexed Properly for a sample	T1R	Outside of 800 series: Time Time/Cal Time
CAI	Calibration Not checked for sample/blank/eval	M18 M28	Snike Out Col 1 and or Col 2 8000 series	T1A	Outside of 8000 series: Time Time/Cal Time
CAI	Calibration Not checked for sample/blank/eval	M18a M18b	Snike Out Col 1 8000 series Acid and or BN	Tm	Too Many Samples/for beginning Calibration
CAI	Calibration Not checked for sample/blank/eval	M18a M28	Snike Out Col 1 and or Col 2 8000 series	Tmw	If for 800 ser Too many samples begin Calibration
CAI	Calibration Not checked for sample/blank/eval	M18a M18b	Snike Out Col 1 8000 series Acid and or BN	Tn	Time Not Checked
CAI	Calibration Not checked for sample/blank/eval	M18a M28	Snike Out Col 1 8000 series Acid and or BN	Tn	Time File Failed
CAI	Calibration Not checked for sample/blank/eval	Ms	Snike Not Checked for this method	Wa	Warning Instrument not in Tdl or field
CAI	Calibration Not checked for sample/blank/eval	Oc	Warning Compounds/Check Calibration		

RUN LOG

Instrument: GC_2 Year: 2005

Analyst: JK

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	Cal 600	8000 Beg Cal	End Cal	BlkFile
2G10565	AC18778-021			PCB-8082	Soil	1	1	8082	08/05 20:17	2G10503		2G10547	2G10569	
10566	AC18778-022			PCB-8082	Soil	1	1	8082	08/05 20:32	2G10503		2G10547	2G10569	
10567	AC18778-023			PCB-8082	Soil	1	1	8082	08/05 20:46	2G10503		2G10547	2G10569	
2G10568	AC18778-024	Tm		PCB-8082	Soil	1	1	8082	08/05 21:01	2G10503		2G10547	2G10569	
2G10569	CAL 1660@1000PPB I26				Soil	0.5	1	608 8082	08/05 21:15	2G10503				
2G10570	1000PPB	Cme			Soil	0.5	1	8082	08/05 21:29	2G10503			2G10569	
2G10571	2000PPB	Cme			Soil	0.25	1	8082	08/05 21:44	2G10503			2G10569	
2G10572	2000PPB	Cme			Soil	0.25	1	8082	08/05 21:58	2G10503			2G10569	

Acc	Area Not Checked	Fa	Extraction Performed Past Hold	Ca	Warning Possible Carry Over
Ac	Area Out	Fem	Solvent Extraction Date Missing/Not check'd	R1A R2A	Rnd Out on Mkdir (col1) and/or col2) 8000 series
RAm	Blank 8000 series missing	Ffn	Trin/Solvent Extraction Date Missing/Not check'd	R1A R2A	Rnd Out on Mkdir (col1) and/or col2) 8000 series
BBm	Blank 8000 series missing	Eto	Tota Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rtn	Can't Calculate Drift
C1A	Calibration Column 1 Out (8000 Series)	Hb	Analysis Before Collection Date	S8	800 series surrogate out
	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
	Calibration Column 2 Out (800 Series)	I16 I26	Initial cal 8000 series failed Column 1 and or 2	Sa8 Sb8	Acid and or BN Surrogate Out (800 series)
	Calibration Column 2 Out (8000 Series)	I18 I28	Initial cal 8000 series failed Column 1 and or 2	Sa8 Sb8	Acid and or BN Surrogate Out (8000 series)
	8000 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Failed Out
LCr	8000 series sample/blank did not have passing cal	Iv	Prtn with nskol rev for init calibration check rfc	Snc	Surrogate Not Checked
CAI	Findng Cal missing for sample (8000 series)	Iw	Initial cal warning: ini cal file <> method	T15	Outside of 800 series Tune time
Cme	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a sample	T1A	Outside of 800 series Tune time/Cal Time
Cn	Drift Out Column 1 or Column 2 Calc or Init Calc	M1A M2A	Snake Out Col 1 and/or Col 2 800 series	T1B	Outside of 8000 series Tune time/Cal Time
D1A D2A	Drift Not Checked	M1Aa M1Ab	Snake Out Col 1 800 series Acid and/or BN	Tm	Too Many Samples for beginning Calibration
Dm	Drift Out	M1Aa M1Ab	Snake Out Col 1 and/or Col 2 8000 series	Tmw	If for 800 ser Too many samples begin Calibration
Dn	An Extraction Before Collection Date	M1Aa M1Ab	Snake Out Col 1 8000 series Acid and/or BN	Tn	Tune Not Checked
Eba	Problem Checking Parameters/matchcheck/reports	Mnc	Snake Not Checked for this method	Tn	Tune File Failed
Ebn	Eval Time Not Checked	Op	Warning Compound(s) Over Calibration	Wa	Warning Instrument Id not in Tol or field

RUN LOG

Instrument: GC_2 Year: 2005

Analyst: JK

1182

Data File	Sample Number	Flags	Comments	Test Group	Matrix	Surr Dil	Sam Dil	Method(s)	Analysis Date	IniCal	Cal 600	8000 Beg Cal	End Cal	BkFile
2G10573.	CAL HERB@100PPB				Aqueou	1	1	8151	08/08 05:55	2G10086				
2G10574.	100PPB				Aqueou	1	1	8151	08/08 06:24	2G10086		2G10573	2G10579	
	'0575. WMB2311				Aqueou	1	1	8151	08/08 06:42	2G10086		2G10573	2G10579	
	_10576. WMB2311(MS)		WMB2311		Aqueou	1	1	8151	08/08 07:01	2G10086		2G10573	2G10579	
2G10577.	AC18907-005(T)			HETCLP-815	Aqueou	1	1	8151	08/08 07:19	2G10086		2G10573	2G10579	
2G10578.	AC18737-026(R)	EO		HE-8151	Aqueou	1	1	8151	08/08 07:37	2G10086		2G10573	2G10579	
2G10579.	CAL HERB@200PPB C28				Aqueou	0.5	1	8151	08/08 07:55	2G10086				
2G10580.	CAL 1660@500PPB				Soil	1	1	608 8082	08/08 08:12	2G10503				
2G10581.	AC18920-001			PCB-8082	Soil	1	1	8082	08/08 08:27	2G10503		2G10580	2G10600	
2G10582.	AC18907-005			PCB-8082	Soil	1	1	8082	08/08 08:41	2G10503		2G10580	2G10600	
2G10583.	WMB2310				Aqueou	1	1	608 8082	08/08 08:56	2G10503	2G10580	2G10580	2G10600	
2G10584.	WMB2310(MS)		WMB2310		Aqueou	1	1	608 8082	08/08 09:10	2G10503	2G10580	2G10580	2G10600	
2G10585.	AC18873-014			PCB-8082	Aqueou	1	1	8082	08/08 09:25	2G10503		2G10580	2G10600	
2G10586.	AC18886-009			PCB-8082	Aqueou	1	1	8082	08/08 09:39	2G10503		2G10580	2G10600	
2G10587.	AC18888-001			PCB-8082	Aqueou	1	1	8082	08/08 09:53	2G10503		2G10580	2G10600	
2G10588.	AC18916-025			PCB-8082	Aqueou	1	1	8082	08/08 10:08	2G10503		2G10580	2G10600	
2G10588.	test				Aqueou	1	1	608 8082	08/08 10:22	2G10503	2G10580	2G10580	2G10600	
2G10589.	SMB728B				Soil	1	1	8082	08/08 10:37	2G10503		2G10580	2G10600	
2G10590.	SMB728B(MS)		SMB728B		Soil	1	1	8082	08/08 10:51	2G10503		2G10580	2G10600	
2G10591.	SMB729B				Soil	1	1	8082	08/08 11:06	2G10503		2G10580	2G10600	
2G10592.	SMB729B(MS)		SMB729B		Soil	1	1	8082	08/08 11:20	2G10503		2G10580	2G10600	
2G10593.	AC18820-005		SMB729B	PCB-8082	Soil	1	1	8082	08/08 11:34	2G10503		2G10580	2G10600	
2G10594.	AC18820-005(MS)	M18	SMB729B	PCB-8082	Soil	1	1	8082	08/08 11:49	2G10503		2G10580	2G10600	
2G10595.	AC18820-005(MSD)	M18	SMB729B	PCB-8082	Soil	1	1	8082	08/08 12:03	2G10503		2G10580	2G10600	
2G10596.	AC18939-001			PCB-8082	Soil	1	1	8082	08/08 12:18	2G10503		2G10580	2G10600	
2G10597.	AC18774-029			PCB-8082	Soil	1	1	8082	08/08 12:32	2G10503		2G10580	2G10600	
2G10598.	AC18807-001			PCB-8082	Soil	1	1	8082	08/08 12:47	2G10503		2G10580	2G10600	
2G10599.	AC18807-004			PCB-8082	Soil	1	1	8082	08/08 13:10	2G10503		2G10580	2G10600	
2G10600.	CAL 1660@1000PPB				Soil	0.5	1	608 8082	08/08 13:25	2G10503				
2G10601.	SMB730B(MS)		SMB730B		Soil	1	1	8082	08/08 13:39	2G10503		2G10600	2G10622	
2G10602.	SMB730B				Soil	1	1	8082	08/08 13:54	2G10503		2G10600	2G10622	
2G10603.	AC18820-001			PCB-8082	Soil	1	1	8082	08/08 14:08	2G10503		2G10600	2G10622	
2G10604.	AC18820-002			PCB-8082	Soil	1	1	8082	08/08 14:22	2G10503		2G10600	2G10622	
2G10605.	AC18820-003			PCB-8082	Soil	1	1	8082	08/08 14:37	2G10503		2G10600	2G10622	
2G10606.	AC18820-004			PCB-8082	Soil	1	1	8082	08/08 14:51	2G10503		2G10600	2G10622	
2G10607.	AC18807-023			PCB-8082	Soil	1	1	8082	08/08 15:06	2G10503		2G10600	2G10622	
2G10608.	AC18807-014			PCB-8082	Soil	1	1	8082	08/08 15:20	2G10503		2G10600	2G10622	
2G10609.	AC18807-017			PCB-8082	Soil	1	1	8082	08/08 15:34	2G10503		2G10600	2G10622	
2G10610.	AC18807-020			PCB-8082	Soil	1	1	8082	08/08 15:49	2G10503		2G10600	2G10622	
2G10611.	AC18807-008			PCB-8082	Soil	1	1	8082	08/08 16:03	2G10503		2G10600	2G10622	
2G10612.	AC18848-006			PCB-8082	Soil	1	1	8082	08/08 16:18	2G10503		2G10600	2G10622	
2G10613.	AC18848-007			PCB-8082	Soil	1	1	8082	08/08 16:32	2G10503		2G10600	2G10622	
2G10614.	AC18848-008			PCB-8082	Soil	1	1	8082	08/08 16:47	2G10503		2G10600	2G10622	
2G10615.	AC18845-002			PCB-8082	Soil	1	1	8082	08/08 17:01	2G10503		2G10600	2G10622	
2G10616.	AC18845-004			PCB-8082	Soil	1	1	8082	08/08 17:16	2G10503		2G10600	2G10622	
2G10617.	AC18845-007			PCB-8082	Soil	1	1	8082	08/08 17:30	2G10503		2G10600	2G10622	
2G10618.	AC18845-010			PCB-8082	Soil	1	1	8082	08/08 17:44	2G10503		2G10600	2G10622	
2G10619.	AC18845-012			PCB-8082	Soil	1	1	8082	08/08 17:59	2G10503		2G10600	2G10622	
2G10620.	500PPB				Soil	1	1	8082	08/08 18:13	2G10503		2G10600	2G10622	
2G10621.	500PPB	Tm			Soil	1	1	8082	08/08 18:28	2G10503		2G10600	2G10622	
2G10622.	CAL 1660@2000PPB				Soil	0.25	1	608 8082	08/08 18:42	2G10503				
2G10623.	2000PPB	Cme			Soil	0.25	1	8082	08/08 18:56	2G10503		2G10622		

Anc	Area Not Checked	EO	Extraction Performed Past Hold	Co	Warning Possible Carry Over
Ad	Area Out	Estm	Solvent Extraction Date Missing/Not check'd	R16,R26	Rpd Out on MsMsd (col1 and or col2) 8000 series
Bdm	Blank 8000 series missing	Ein	Totp/Solvent Extraction Date Missing/Not check'd	R16,R26	Rpd Out on MsMsd (col1 and or col2) 8000 series
Bfm	Blank 8000 series missing	Eio	Totp Extraction Performed Outside of Hold	Ro	Retention Time Out Or %Diff Out
Bnf	Blank Not Found/Assigned	Ev	Eval Time Exceeded	Rin	Can't Calculate Dm
C16	Calibration Column 1 Out (800 Series)	Hb	Analysis Before Collection Date	S6	800 series surrogate out
18	Calibration Column 1 Out (8000 Series)	Ho	Sample Analyzed outside of hold time	S8	8000 series surrogate out
1	Calibration Column 2 Out (800 Series)	(16,28	Initial cal 8000 series failed Column 1 and or 2	Sa6,Sb6	Acid and or BN Surrogate Out (800 series)
1	Calibration Column 2 Out (8000 Series)	(18,28	Initial cal 8000 series failed Column 1 and or 2	Sa8,Sb8	Acid and or BN Surrogate Out (8000 series)
1	800 series sample/blank did not have passing cal	Is	Initial Cal Not Checked	Sd	Surrogate Diluted Out
1	8000 series sample/blank did not have passing cal	Iv	Prob with calprc.csv for int calibration check rts	Snc	Surrogate Not Checked
Cme	Ending Cal missing for sample (8000 series)	Iw	Initial Cal warning.. ini cal file <- method..	T15	Outside of 500 series Tune time
Cn	Calibration Not Checked for sample/blank/eval	Ix	Initial Cal Files Not Updated Properly for a samp	T6	Outside of 8000 series Tune time/Cal Time
D1o,D2o	Drift Out Column 1 or Column 2 Cals or Init Cals	M16,M26	Spike Out Col 1 and or Col 2 8000 series	T8	Too Many Samples/ for beginning Calibration
Dec	Drift Not Checked	M16a,M16b	Spike Out Col 1 and or Col 2 8000 series	Tm	If for 600 ser Too many samples begin Calibration
De	Drift Out	M18,M28	Spike Out Col 1 and or Col 2 8000 series	Tmw	Tune Not Checked
Eba	An Extraction Before Collection Date	M18a,M18b	Spike Out Col 1 8000 series Acid and or BN	Tn	Tune File Failed
Emp	Problem Checking Prep/rundates modcheck/preprund	Mnc	Spike Not Checked for this ms/msd	To	Warning... Instrument Id not in TdLoc field
En	Eval Time Not Checked	Oc	Warning Compound(s) Over Calibration	Wie	

Veritech Internally Prepared Standard Log

1483

Veritech Lot Number: V-210

Prepared By: Yarka		Department: Organics		
Description: PEST/PCB SURR		BatchNumber:		
Prep Date: 9/20/04		Concentration: 200 ppm		
Expiration Date: 9/30/05		Final Volume: 100 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
480	TCMX	20 mg	neat	200 ppm
481	DCB	20 mg	neat	200 ppm
485	Acetone Neat	100 ml		

Veritech Lot Number: V-2874

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1232 INTERM		BatchNumber: B-331		
Prep Date: 5/3/05		Concentration: 100 ppm		
Expiration Date: 9/30/05		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
813	Aroclor 1232	100 ul	1000 ppm	100 ppm
V-210	PEST/PCB SURR	50 ul	200 ppm	10 ppm
478	HEXANE	850 ul	NEAT	

Veritech Lot Number: V-2875

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1242 inter		BatchNumber:		
Prep Date: 5/3/05		Concentration: 100 ppm		
Expiration Date: 9/30/05		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
814	Aroclor 1242	100 ul	1000 ppm	100 ppm
V-210	PEST/PCB SURR	50 ul	200 ppm	10 ppm
478	HEXANE	850 ul	NEAT	

Veritech Lot Number: V-2876

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1248 inter		BatchNumber:		
Prep Date: 5/3/05		Concentration: 100 ppm		
Expiration Date: 9/30/05		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
815	Aroclor 1248	100 ul	1000 ppm	100 ppm
478	HEXANE	850 ul	NEAT	
V-210	PEST/PCB SURR	50 ul	200 ppm	10 ppm

Veritech Lot Number: V-2877

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 2154 inter		BatchNumber:		
Prep Date: 5/3/05		Concentration: 100 ppm		
Expiration Date: 9/30/05		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
816	Aroclor 1254	100 ul	1000 ppm	100 ppm
833	Aroclor 1221	100 ul	1000 ppm	100 ppm
478	HEXANE	750 ul	NEAT	
V-210	PEST/PCB SURR	50 ul	200 ppm	10 ppm

Veritech Internally Prepared Standard Log

Veritech Lot Number: V-2878

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1232 ws		BatchNumber:		
Prep Date: 5/3/05		Concentration: 500 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
478	HEXANE	9900 ul	NEAT	
V-2874	1232 INTERM	100 ul	100 ppm	500 ppb

Veritech Lot Number: V-2879

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1242 ws		BatchNumber:		
Prep Date: 5/3/05		Concentration: 500 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
478	HEXANE	9900 ul	NEAT	
V-2875	1242 inter	100 ul	100 ppm	500 ppb

Veritech Lot Number: V-2880

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 1248 ws		BatchNumber:		
Prep Date: 5/3/05		Concentration: 500 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
478	HEXANE	9950 ul	NEAT	
V-2876	1248 inter	50 ul	100 ppm	500 ppb

Veritech Lot Number: V-2882

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: 2154 ws		BatchNumber:		
Prep Date: 5/3/05		Concentration: 500 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
478	HEXANE	9950 ul	NEAT	
V-2877	2154 inter	50 ul	100 ppm	500 ppb

Veritech Lot Number: V-3166

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: TCMX/DCB SGT		BatchNumber:		
Prep Date: 5/12/05		Concentration: 10 ppm		
Expiration Date: 9/30/05		Final Volume: 200 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
950	Acetone	190 ml	Neat ml	
V-210	PEST/PCB SURR	10 ml	200 ppm	

Veritech Internally Prepared Standard Log

Veritech Lot Number: V-4707

Prepared By: Quimby, Richard		Department: Organics		
Description: PCB Spike		BatchNumber:		
Prep Date: 7/8/05		Concentration: 100 ppm		
Expiration Date: 1/7/06		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
950	Acetone	8 ml	Neat	
1074	AROCLOR 1016	1 ml	1000 ppm	100 ppm
1075	AROCLOR 1260	1 ml	1000 ppm	100 ppm

Veritech Lot Number: V-4986

Prepared By: Desai, Kinjal		Department: Organics		
Description: 1660-INTERMEDIATE		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 100PPM		
Expiration Date: 9/30/05		Final Volume: 1 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
817	Aroclor 1260	100 ul	1000 ppm	100 ppm
V-210	PEST/PCB SURR	50 ul	200PPM	100 ppm
802	n-Hexane	750 ul		neat
855	Aroclor 1016	100 ul	1000 ppm	100 ppm

Veritech Lot Number: V-4987

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@4000PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 4000 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	400 ul	100PPM	4000 ppb
802	n-Hexane	9600 ul		neat

Veritech Lot Number: V-4988

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@2000PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 2000 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	200 ul	100PPM	2000 ppb
802	n-Hexane	9800 ul		neat

Veritech Lot Number: V-4989

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@1000PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 1000 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	100 ul	100PPM	1000 ppb
802	n-Hexane	9900 ul		neat

Veritech Internally Prepared Standard Log

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Veritech Lot Number: V-4990

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@500PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 500 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	50 ul	100PPM	500 ppb
802	n-Hexane	9950 ul		neat

Veritech Lot Number: V-4991

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@200PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 200 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	20 ul	100PPM	200 ppb
802	n-Hexane	9980 ul		neat

Veritech Lot Number: V-4992

Prepared By: Desai, Kinjal		Department: Organics		
Description: CAL 1660@50PPB		BatchNumber: B-527		
Prep Date: 7/20/05		Concentration: 50 ppb		
Expiration Date: 9/30/05		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
V-4986	1660-INTERMEDIATE	5 ul	100PPM	50 ppb
802	n-Hexane	9995 ul		neat

Veritech Standard Receipt Log

1187

Veritech Control/Receipt Number: 478

Description
HEXANE

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
FISHER	H300-4	023660	07/14/03	01/04/07	Yarka	1	4L	NEAT	

Veritech Control/Receipt Number: 480

Description
TCMX

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2298	LB07127	10/24/02	09/30/05	Yarka	1	1g	neat	

Veritech Control/Receipt Number: 481

Description
DCB

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2537	LB07636	10/24/02	10/31/05	Yarka	1	0.1g	neat	

Veritech Control/Receipt Number: 485

Description
Acetone Neat

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher	a40-4	038587	04/14/04	01/19/10	richq	1	4L	neat	

Veritech Control/Receipt Number: 802

Description
n-Hexane

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Pharmco	35900HPLC	3002069	05/20/04	10/13/10	Yarka	1	4L	neat	

Veritech Control/Receipt Number: 813

Description
Aroclor 1232

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-4805	1b21183	10/15/04	03/31/07	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 814

Description
Aroclor 1242

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-4806	1b18566	10/15/04	03/31/07	jean	1	1ml	1000	ppm

Veritech Standard Receipt Log

5199

Veritech Control/Receipt Number: 815

Description

Aroclor 1248

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-4807	lb14850	10/15/04	09/30/06	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 816

Description

Aroclor 1254

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-4808	lb19887	10/15/04	04/30/07	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 817

Description

Aroclor 1260

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-4809	lb20748	10/15/04	06/30/07	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 833

Description

Aroclor 1221

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-8098	lb19357	10/20/04	03/31/07	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 855

Description

Aroclor 1016

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	4-8097	lb20874	11/11/04	06/30/07	jean	1	1ml	1000	ppm

Veritech Control/Receipt Number: 950

Description

Acetone

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher Scientific	A40-4	043780	12/13/04	11/17/10	Akmal	1	4L	Neat	

Veritech Control/Receipt Number: 1074

Description

AROCLOR 1016

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-216S-H-10X-PAK	B3100245	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Veritech Standard Receipt Log

Veritech Control/Receipt Number: 1075

Description
AROCLOR 1260

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-260S-H-10X-PAK	B3060001	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Veritech Standard Receipt Log

1158

Veritech Control/Receipt Number: 480

Description
TCMX

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2298	LB07127	10/24/02	09/30/05	Yarka	1	1g	neat	

Veritech Control/Receipt Number: 481

Description
DCB

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2537	LB07636	10/24/02	10/31/05	Yarka	1	0.1g	neat	

Veritech Control/Receipt Number: 485

Description
Acetone Neat

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher	a40-4	038587	04/14/04	01/19/10	richq	1	4L	neat	

Veritech Control/Receipt Number: 950

Description
Acetone

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher Scientific	A40-4	043780	12/13/04	11/17/10	Akmal	1	4L	Neat	

Veritech Control/Receipt Number: 1074

Description
AROCLOR 1016

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-216S-H-10X-PAK	B3100245	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Veritech Control/Receipt Number: 1075

Description
AROCLOR 1260

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-260S-H-10X-PAK	B3060001	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Veritech Internally Prepared Standard Log

Veritech Lot Number: V-210

Prepared By: Yarka		Department: Organics		
Description: PEST/PCB SURR		BatchNumber:		
Prep Date: 9/20/04		Concentration: 200 ppm		
Expiration Date: 9/30/05		Final Volume: 100 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
480	TCMX	20 mg	neat	200 ppm
481	DCB	20 mg	neat	200 ppm
485	Acetone Neat	100 ml		

Veritech Lot Number: V-3166

Prepared By: Korytova, Jaroslava		Department: Organics		
Description: TCMX/DCB SGT		BatchNumber:		
Prep Date: 5/12/05		Concentration: 10 ppm		
Expiration Date: 9/30/05		Final Volume: 200 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
950	Acetone	190 ml	Neat ml	
V-210	PEST/PCB SURR	10 ml	200 ppm	

Veritech Lot Number: V-4707

Prepared By: Quimby, Richard		Department: Organics		
Description: PCB Spike		BatchNumber:		
Prep Date: 7/8/05		Concentration: 100 ppm		
Expiration Date: 1/7/06		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
950	Acetone	8 ml	Neat	
1074	AROCLOR 1016	1 ml	1000 ppm	100 ppm
1075	AROCLOR 1260	1 ml	1000 ppm	100 ppm

Veritech Lot Number: V-5154

Prepared By: Quimby, Richard		Department: Organics		
Description: PEST/PCB SURR		BatchNumber:		
Prep Date: 7/26/05		Concentration: 10 ppm		
Expiration Date: 9/30/05		Final Volume: 200 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
950	Acetone	190 ml	Neat	
V-210	PEST/PCB SURR	10 ml	200 ppm	10 ppm

Veritech Lot Number: V-5452

Prepared By: Quimby, Richard		Department: Organics		
Description: PCB SPK		BatchNumber:		
Prep Date: 8/1/05		Concentration: 100 ppm		
Expiration Date: 1/31/06		Final Volume: 10 ml		
Veritech Lot# /Rec#	Lot Description	Amount Used	Conc of Std	Final Conc
1074	AROCLOR 1016	1 ml	1000 ppm	100 ppm
1075	AROCLOR 1260	1 ml	1000 ppm	100 ppm
950	Acetone	8 ml	Neat	

Veritech Standard Receipt Log

1153

Veritech Control/Receipt Number: 480									
Description									
TCMX									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2298	LB07127	10/24/02	09/30/05	Yarka	1	1g	neat	

Veritech Control/Receipt Number: 481									
Description									
DCB									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
supelco	44-2537	LB07636	10/24/02	10/31/05	Yarka	1	0.1g	neat	

Veritech Control/Receipt Number: 485									
Description									
Acetone Neat									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher	a40-4	038587	04/14/04	01/19/10	richq	1	4L	neat	

Veritech Control/Receipt Number: 950									
Description									
Acetone									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
Fisher Scientific	A40-4	043780	12/13/04	11/17/10	Akmal	1	4L	Neat	

Veritech Control/Receipt Number: 1074									
Description									
AROCLOR 1016									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-216S-H-10X-PAK	B3100245	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Veritech Control/Receipt Number: 1075									
Description									
AROCLOR 1260									

Manufacturer	Catalog Num:	Lot Num:	Date Rec:	Exp Date:	Rec By:	Num of Cont	Volume /Cont	Conc:	Units:
ACCUSTANDAR	C-260S-H-10X-PAK	B3060001	03/29/05	03/28/08	Revolus, Jean	5	1ml	1000	PPM

Method Blank No. SMB- 7278
Blank Spike (SMBS): 7268, 7278 PEST
Blank Spike (SMBS): 7268, 7278 PCB

Date: 8/4/05
Matrix Spike: 18855-002, 18778-011
Matrix Spike: 18778-008, 18778-020

Analysis: Pest / PCB / Herb / Other

Sample Number	No. in batch				Initial Volume	Final Volume	Extracted By/Position/ Comments
	Pest	PCB	Herb	Other			
MB 7278	x	x			20g	10.0ml	/ 1, 1 / ASE I Rack 15
MB S 7278	x	x					/ 2, 3 /
18778-010	13	12					/ 1 / ASE II
18778-012	14	13					/ 2 /
18778-013	15	14					/ 3 /
18778-014	16	15					/ 4 /
18778-015	17	16					/ 5 /
18778-016	18	17					/ 6 /
18778-017	19	18					/ 7 /
18778-019	20	19					/ 8 /
18778-D11 MS	x						/ 4 / ASE I
18778-D11 MSD	x						/ 5 /
18778-D11	1	20					/ 6 /
18778-D20MS		x					/ 7 /
18778-D20MSD		x					/ 8 /
18778-020	2	1					/ 9 / ASE II
18778-021	3	2					/ 10 /
18778-022	4	3					/ 11 /
18778-023	5	4					/ 12 / ASE I
18778-024	6	5					/ 10 / ASE I
18919-001		6					/ 11 /
18919-002		7					/ 12 /
18919-003		8					/ 13 /
18778-003	R	R					/ 14 /
18778-009	R	R					/ 15 /
18778-018	7	9					/ /
							/ /
							/ /
							/ /
							/ /

Cleanup: Acid TBA Copper Florisil Other

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	100	V-4707	Pest <u>PCB</u> / Herb / Other
100	10	V-4044	PEST

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	
100	10	V-5754	<u>Pest / PCB</u> / Herb / Other

Reagent Lots: MeCl2 _____ Acetone 050776 Hexane 044526 Na2SO4 _____ Ether _____
MTBE _____ Other _____

Relinquished By: LTKN
Received By: _____

Date: 8/4/05
Date: 08/05/05

Method Blank No. SMB- 7298
Blank Spike (SMBS): 7278, 7298 PEST
Blank Spike (SMBS): 7278, 7298 PCB

Date: 8/7/05
Matrix Spike: 18778-011, 18230-011
Matrix Spike: 18778-020, 18220-005

Analysis (Pest / PCB) Herb / Other

Sample Number	No. in batch				Initial Volume	Final Volume	Extracted By/Position/ Comments
	Pest	PCB	Herb	Other			
MB 7298	X	X			20g	10 ml	/ 1,1 / ASE I
MB5 7298	X	X					/ 2,3 /
18807-001	9	14					/ 1 / ASE II
18807-004	10	15					/ 2 /
18807-008	11	16					/ 3 /
18807-014	12	17					/ 4 /
18807-017	13	18					/ 5 /
18807-020	14	19					/ 6 /
18807-023	15	20					/ 7 /
18820-005 MS		X					/ 4 / ASE I
18820-005 MS		X					/ 5 /
18820-005	16	1					/ 6 /
18820-001	17	2					/ 8 / ASE R
18820-002	18	3					/ 9 /
18820-003	19	4					/ 10 /
18820-004	20	5					/ 11 /
18830-001	16						/ 10 / ASE I
18830-002	17						/ 11 /
18830-009	18						/ 12 /
18830-010	19						/ 13 /
18830-012	20						/ 14 /
18830-011 MS	X						/ 7 /
18830-011 MS	X						/ 8 /
18830-011	1						/ 9 /
18830-019	2						/ 15 /
18830-020	3						/ 16 /

Cleanup: Acid TBA Copper Florisil Other

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	Pest (PCB) Herb / Other
100	100	V-5452	Pest (PCB) Herb / Other
100	10	V-4044	PEST

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	Pest (PCB) Herb / Other
100	10	V-5154	Pest / PCB Herb / Other

Reagent Lots: MeCL2 _____ Acetone 050776 Hexane 044526 Na2SO4 _____ Ether _____
MTBE _____ Other _____

Relinquished By: GKN
Received By: Kozell

Date: 8/7/05
Date: 8/8/05

Method Blank No. WMB- 2305
 Blank Spike (WMBS): 2287, 2305
 Blank Spike (WMBS): 2288 PCB

Date: 8/2/05
 Matrix Spike: 18499-008, 18808-001
 Matrix Spike: 18550-001

Analysis: Pest / PCB / Herb / Other(list):

Sample Number	No. in batch				Initial Vol	Final Vol	Comments	TCLP QC	Extraction Fluid
	Pest	PCB	Herb	Other					
Mb 2305	X	X			1000ml	5ml		18499-008	V5084
MBS 2305	X	X			↓				
18766-001	19				100ml			15	4
18766-002	20				↓			16	5
18807-007									
MS 18808-001	X				100ml				
M21 18808-001	X				↓				
18807-007	1	18			1000ml			18808-001	EF1 V5263
18808-001	2				100ml			1	1
18819-002	3				↓			2	2
18820-012		19			930ml			(RACK 16)	
18863-001	X	20			1000ml				
EF1 V5263	X				100ml				
18819-004	4				100ml	5ml	MSL Rack #1	3	3
18819-006	5							4	4
18819-008	6							5	5
18819-010	7							6	6
18819-012	8							7	7
18819-014	9							8	8
18819-016	10							9	9
18819-018	11							10	10
	as								
	12								

Cleanup: Acid ___ TBA ___ Copper ___ Florisil ___ Other ___

Spike Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	Pest / PCB / Herb / Other
50	10	V4044	Pest / PCB / Herb / Other
↓	100	V4707	Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other

Surrogate Standard

Vol (ul's)	Conc. (ppm/ppb)	Lot No.	Pest / PCB / Herb / Other
50	10	V3166	Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other
			Pest / PCB / Herb / Other

Reagent Lots: MeCL₂ 07907 Acetone _____ Hexane 04526 Na₂SO₄ 052002 Ether _____
 MTBE _____ Other _____

Relinquished By: alex. H. / MSL
 Received By: Kozel

Date: 8/2/05
 Date: 8/3/05