

User: Vinci, David L.
Westchester Dept. of Labs & Research
Date: 02/25/2002 Time: 10:11:45

Sample: AE01230
Analysis: \$GCMS GCMS Scan For Unknowns
Units: ug
Started: 2/25/02 10:11 Ended: 2/25/02 10:11 Analyst: DLV
Page: 1

	Component Name	Viol	Result	Qualifier	MDL
1	Other unknown compounds		.		
2	Surr_2,4-DDD		90		5.0

Comments for Sample AE01230 - Analysis \$GCMS

Westchester Dept. of Labs & Research

User: Vinci, David L

Date: 02-25-2002 Time: 10:11:48

The GCMS scan, from 35 to 550 amu, does not reveal the presence of unusual compounds. The recovery for 1 of the 43 compounds in the LCS Standard was below its acceptance range.

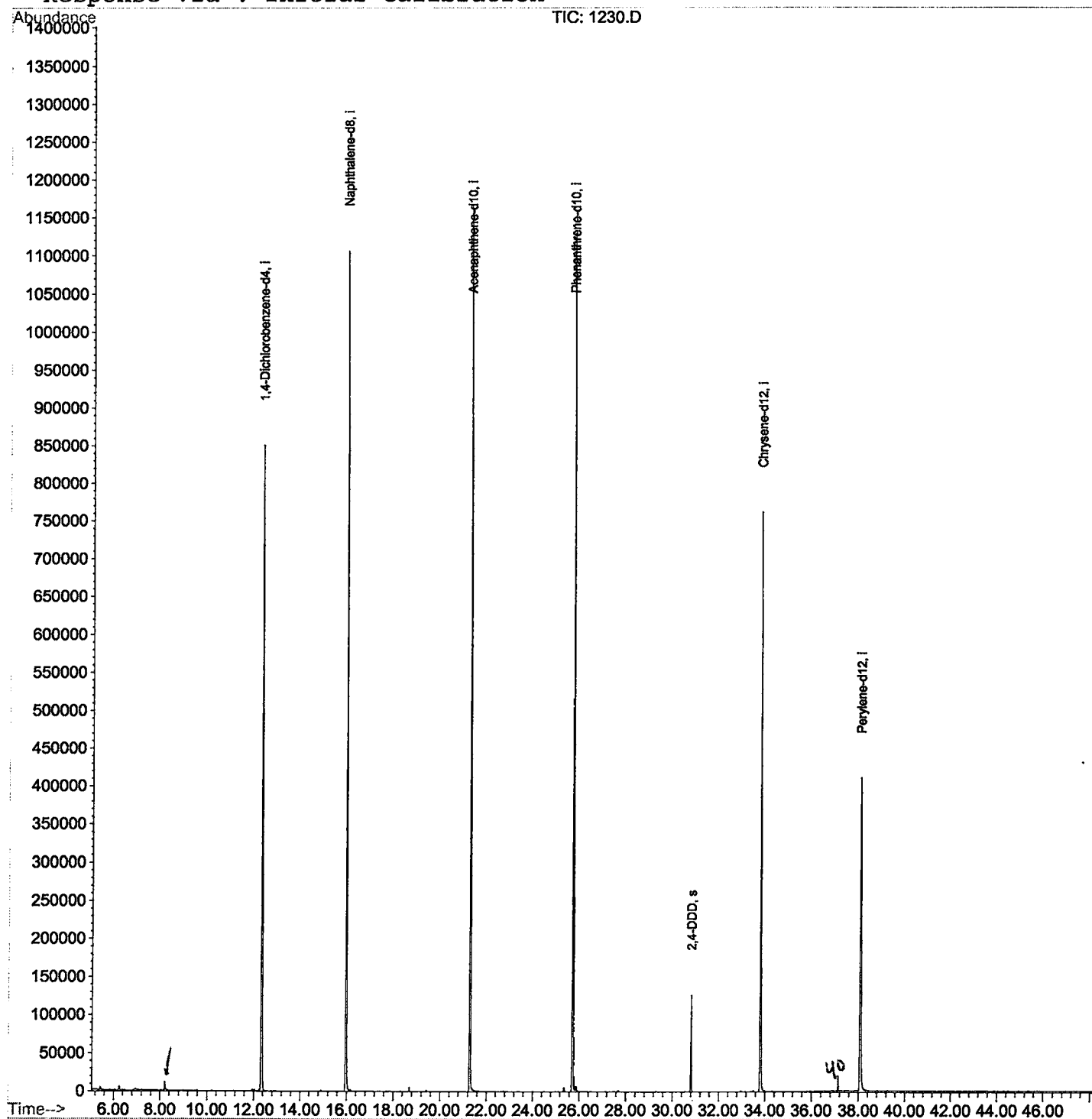
Quantitation Report

Data File : C:\HPCHEM\1\DATA\FEB1502\1230.D
Acq On : 17 Feb 2002 4:16 am
Sample : gc/ms scan 1/17
Misc :
MS Integration Params: GOOFY.P
Quant Time: Feb 20 15:33 2002

Vial: 37
Operator: 209 GALOS
Inst : GC/MS 209
Multiplr: 1.00

Quant Results File: GCMS0208.RES

Method : C:\HPCHEM\1\METHODS\GCMS0208.M (RTE Integrator)
Title : 209 625/TCLP calibration table
Last Update : Wed Feb 13 11:43:23 2002
Response via : Initial Calibration



LSC Area Percent Report

Data File : C:\HPCHEM\1\DATA\FEB1502\1230.D
 Acq On : 17 Feb 2002 4:16 am
 Sample : gc/ms scan 1/17
 Misc :
 MS Integration Params: LSCINT.P

Vial: 37
 Operator: 209 GALOS
 Inst : GC/MS 209
 Multiplr: 1.00

Method : C:\HPCHEM\1\METHODS\GCMS0208.M (RTE Integrator)
 Title : 209 625/TCLP calibration table
 Smoothing : OFF
 Sampling : 1
 Start Thrs: 0.2
 Stop Thrs : 0
 Filtering: 5
 Min Area: 1 % of largest Peak
 Max Peaks: 100
 Peak Location: TOP

If leading or trailing edge < 100 prefer < Baseline drop else tangent >
 Peak separation: 5

Signal : TIC

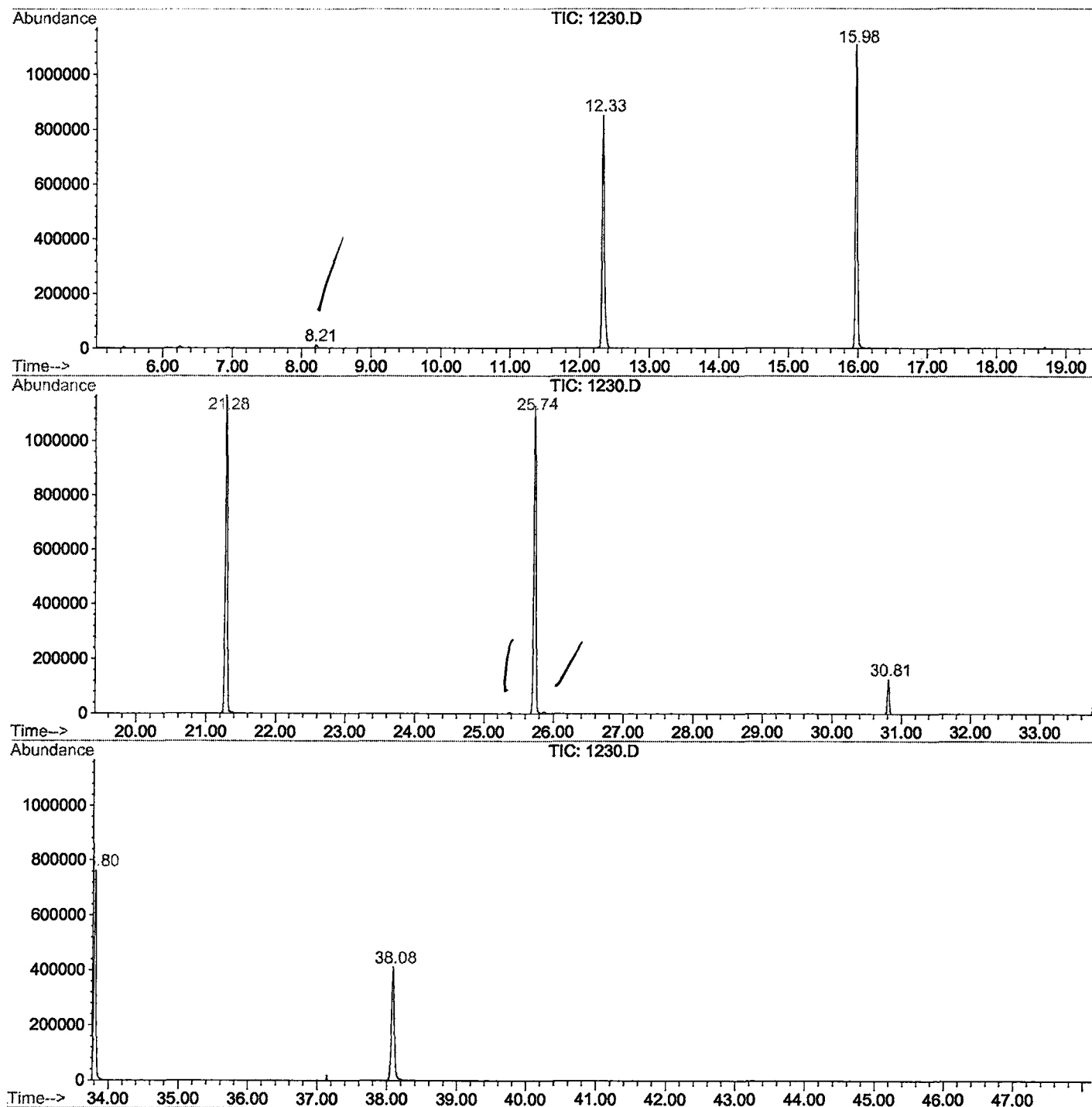
peak #	R.T. min	first scan	max scan	last scan	PK TY	peak height	peak area	peak % max.	% of total
1	8.210	340	345	351	rBV	11748	24952	1.03%	0.209%
2	12.332	788	794	807	rVB	849970	1855696	76.42%	15.525%
3	15.977	1184	1191	1207	rBV	1106193	2278469	93.83%	19.063%
4	21.283	1763	1769	1777	rBV	1167513	2428400	100.00%	20.317%
5	25.735	2247	2254	2265	rBV2	1128415	2310019	95.13%	19.327%
6	30.812	2802	2807	2813	rBV	126717	229246	9.44%	1.918%
7	33.805	3126	3133	3148	rBV	762121	1632310	67.22%	13.657%
8	38.083	3591	3599	3614	rBV2	410261	1193485	49.15%	9.985%

Sum of corrected areas: 11952577

1230.D GCMS0208.M Thu Feb 21 09:30:21 2002

LSC Report - Integrated Chromatogram

File : C:\HPCHEM\1\DATA\FEB1502\1230.D
 Operator : 209 GALOS
 Acquired : 17 Feb 2002 4:16 am using AcqMethod GCMS0208
 Instrument : GC/MS 209
 Sample Name: gc/ms scan 1/17
 Misc Info :
 Vial Number: 37
 Quant File :GCMS0208.RES (RTE Integrator)



1230.D GCMS0208.M

Thu Feb 21 09:30:26 2002

Library Search Compound Report

Data File : C:\HPCHEM\1\DATA\FEB1502\1230.D
 Acq On : 17 Feb 2002 4:16 am
 Sample : gc/ms scan 1/17
 Misc :
 MS Integration Params: LSCINT.P

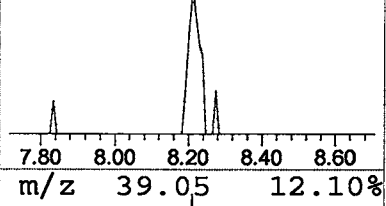
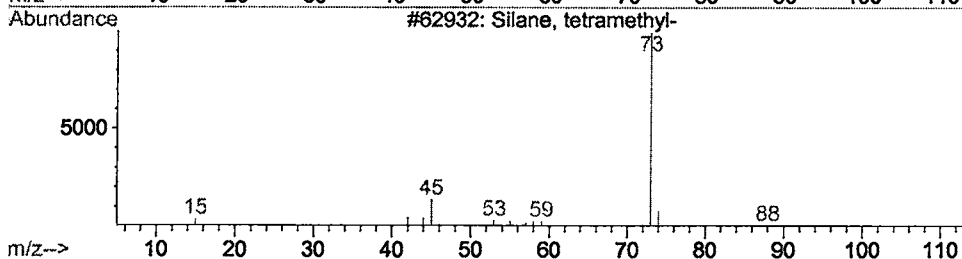
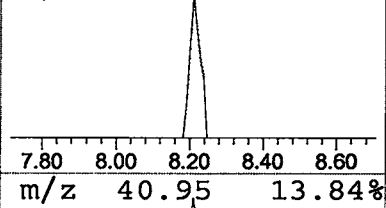
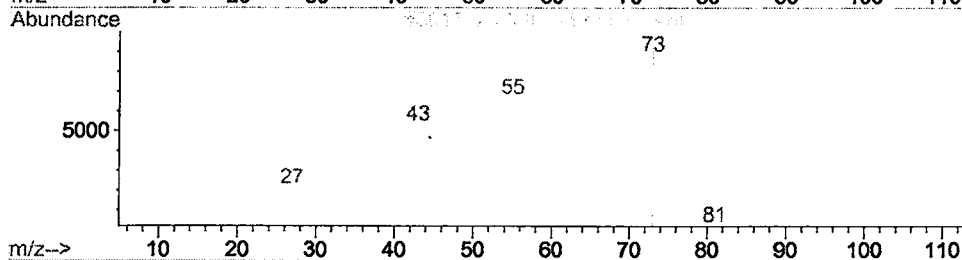
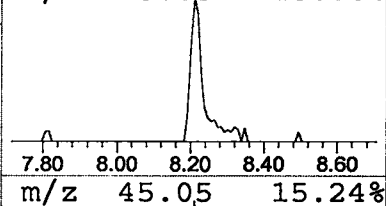
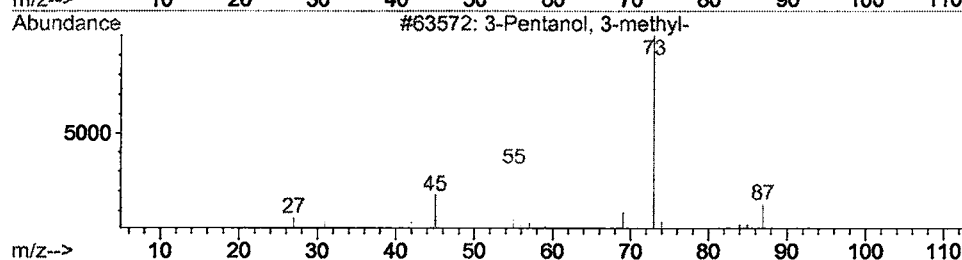
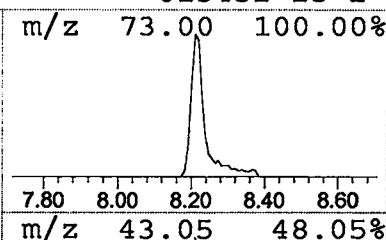
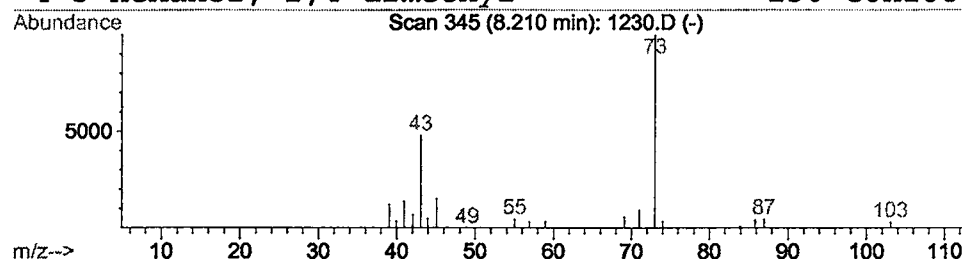
Vial: 37
 Operator: 209 GALOS
 Inst : GC/MS 209
 Multiplr: 1.00

Quant Method : C:\HPCHEM\1\METHODS\GCMS0208.M (RTE Integrator)
 Title : 209 625/TCLP calibration table
 Library : C:\DATABASE\NBS75K.L

 Peak Number 1 3-Pentanol, 3-methyl- Concentration Rank 1

R.T.	EstConc	Area	Relative to ISTD	R.T.
8.21	0.27 ug/l	24952	1,4-Dichlorobenzene-d4	12.33

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			3-Pentanol, 3-methyl-	102	C6H14O	000077-74-7	25
2			2-Methyl-5-hexen-3-ol	114	C7H14O	032815-70-6	23
3			Silane, tetramethyl-	88	C4H12Si	000075-76-3	23
4			3-Hexanol, 2,4-dimethyl-	130	C8H18O	013432-25-2	9



Tentatively Identified Compound (LSC) summary

Operator ID: 209 GALOS Date Acquired: 17 Feb 2002 4:16 am
 Data File: C:\HPCHEM\1\DATA\FEB1502\1230.D
 Name: gc/ms scan 1/17
 Misc:
 Method: C:\HPCHEM\1\METHODS\GCMS0208.M (RTE Integrator)
 Title: 209 625/TCLP calibration table
 Library Searched: C:\DATABASE\NBS75K.L

TIC Top Hit name	RT	EstConc	Units	Area	IntStd	ISRT	ISArea	ISConc
3-Pentanol, 3-methyl	8.21	0.3	ug/l	24952	ISTD01	12.33	1855700	20.0
1230.D GCMS0208.M	Thu Feb 21 09:30:35 2002							

Quantitation Report

(QT Reviewed)

Data File : C:\HPCHEM\1\DATA\FEB1502\1230.D
 Acq On : 17 Feb 2002 4:16 am
 Sample : gc/ms scan 1/17
 Misc :
 MS Integration Params: GOOFY.P
 Quant Time: Feb 20 15:33 2002

Vial: 37
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Quant Results File: GCMS0208.RES

Quant Method : C:\HPCHEM\1\METHODS\GCMS0208.M (RTE Integrator)
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 Last Update : Wed Feb 13 11:43:23 2002
 Response via : Initial Calibration
 DataAcq Meth : GCMS0208

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	12.33	152	286293	20.00	ug/l	-0.02
10) Naphthalene-d8	15.98	136	1091956	20.00	ug/l	-0.02
17) Acenaphthene-d10	21.28	164	575952	20.00	ug/l	-0.03
29) Phenanthrene-d10	25.74	188	825182	20.00	ug/l	-0.03
38) Chrysene-d12	33.80	240	531150	20.00	ug/l	-0.04
44) Perylene-d12	38.08	264	347215	20.00	ug/l	-0.04

System Monitoring Compounds

37) 2,4-DDD	30.81	235	37398	4.51	ug/mL	0.31
Spiked Amount	5.000		Recovery	=	90.20%	

Target Compounds

				Qvalue
2) N-nitroso-dimethyl amine	0.00	74	0	N.D.
3) bis(2-Chloroethyl)ether	0.00	93	0	N.D.
4) 1,3-Dichlorobenzene	0.00	146	0	N.D.
5) 1,4-Dichlorobenzene	0.00	146	0	N.D.
6) 1,2-Dichlorobenzene	0.00	146	0	N.D.
7) 2,2'-oxybis(1-Chloropropan	0.00	45	0	N.D.
8) N-Nitroso-di-n-propylamine	0.00	70	0	N.D.
9) Hexachloroethane	0.00	117	0	N.D.
11) Nitrobenzene	0.00	77	0	N.D.
12) Isophorone	0.00	82	0	N.D.
13) bis(2-Chloroethoxy)methane	0.00	93	0	N.D.
14) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.
15) Naphthalene	16.02	128	585	N.D.
16) Hexachlorobutadiene	0.00	225	0	N.D.
18) Hexachlorocyclopentadiene	0.00	237	0	N.D.
19) 2-Chloronaphthalene	0.00	162	0	N.D.
20) Dimethylphthalate	0.00	163	0	N.D.
21) 2,6-Dinitrotoluene	0.00	165	0	N.D.
22) Acenaphthylene	0.00	152	0	N.D.
23) Acenaphthene	0.00	153	0	N.D.
24) 2,4-Dinitrotoluene	0.00	165	0	N.D.
25) Diethylphthalate	22.76	149	105	N.D.
26) 4-Chlorophenyl-phenylether	0.00	204	0	N.D.
27) Fluorene	0.00	166	0	N.D.
28) Azobenzen/ 1,2-Diphenylhyd	0.00	77	0	N.D.

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

1230.D GCMS0208.M Wed Feb 20 15:34:17 2002

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Quantitation Report

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Operator: 209 GALOS

Sample : gc/ms scan 1/17

Inst : GC/MS 209

Misc :

Multiplr: 1.00

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Title : 209 625/TCLP calibration table

Last Update : Wed Feb 13 11:43:23 2002

Response via : Initial Calibration

DataAcq Meth : GCMS0208

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
30) N-Nitrosodiphenylamine	0.00	168	0	N.D.		
31) 4-Bromophenyl-phenylether	0.00	248	0	N.D.		
32) Hexachlorobenzene	0.00	284	0	N.D.		
33) Phenanthrene	25.74	178	122	N.D.		
34) Anthracene	0.00	178	0	N.D.		
35) Di-n-butylphthalate	27.70	149	2344	N.D.		
36) Fluoranthene	0.00	202	0	N.D.		
39) Pyrene	0.00	202	0	N.D.		
40) Butylbenzylphthalate	0.00	149	0	N.D.		
41) Benzo(a)anthracene	33.80	228	1376	N.D.		
42) Bis(2-ethylhexyl)phthalate	34.01	149	635	N.D.		
43) Chrysene	33.80	228	1376	N.D.		
45) Di-n-Octylphthalate	0.00	149	0	N.D.		
46) Benzo(b)fluoranthene	0.00	252	0	N.D.		
47) Benzo(k)fluoranthene	0.00	252	0	N.D.		
48) Benzo(a)pyrene	38.08	252	1278	N.D.		
49) Indeno(1,2,3-cd)pyrene	0.00	276	0	N.D.		
50) Dibenzo(a,h)anthracene	0.00	278	0	N.D.		
51) Benzo(g,h,i)perylene	0.00	276	0	N.D.		

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

1230.D GCMS0208.M Wed Feb 20 15:34:21 2002

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