

User: Vinci, David L
Westchester Dept. of Labs & Research
Date: 04/16/2002 Time: 15:56:58

Sample: AE07172
Analysis: \$GCMS GCMS Scan For Unknowns
Units: ug
Started: 4/16/02 15:56 Ended: 4/16/02 15:56 Analyst: DLV
Page: 1

	Component Name	Viol	Result	Qualifier	MDL
1	Other unknown compounds				
2	Surr_2,4-DDD		73		5.0

Comments for Sample AE07172 - Analysis \$GCMS

Westchester Dept. of Labs & Research

User: Vinci, David L

Date: 04-16-2002 Time: 15:57:28

The GCMS scan, from 35 to 550 amu, does not reveal the presence of unusual compounds. The recoveries for 2 of the 43 compounds in the LCS Standard were high.

Quantitation Report

(QT Reviewed)

Data File : C:\MSDCHEM\1\DATA\041002\7172.D

Acq On : 11 Apr 2002 9:43 am

Sample : SAMPLE 7172 3/27/02

Misc :

MS Integration Params: BENZO.P

Quant Time: Apr 15 18:34:07 2002

Vial: 25

Operator: Bohlk

Inst : Instrumen

Multiplr: 1.00

Quant Results File: 5080402BB.RES

Quant Method : C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)

Title : Quantitation For Method 508gcscan

Last Update : Mon Apr 15 15:56:12 2002

Response via : Initial Calibration

DataAcq Meth : 508SCAN

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev (Min)
1) 1,4-Dichlorobenzene-d4	9.41	152	511182	40.00	ug/L	0.00
10) Naphthalene-d8	12.88	136	2432523	40.00	ug/L	-0.01
17) Acenaphthene-d10	17.99	164	1383364	40.00	ug/L	-0.01
30) Phenanthrene-d10	22.34	188	1985825	40.00	ug/L	-0.01
39) Chrysene-d12	30.14	240	1450142	40.00	ug/L	-0.02
45) Perylene-d12	34.01	264	911607	40.00	ug/L	-0.02

System Monitoring Compounds

28) TCMX (SURR)	19.94	244	21354	7.29	ug/L	-0.01
Spiked Amount	10.000		Recovery	=	72.90%	
38) 2,4-DDD (SURR)	0.00	235	0	0.00	ug/L	
Spiked Amount	5.000		Recovery	=	0.00%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	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 - Dichlorobenze	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-Chloropr	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) methan	0.00	93	0	N.D.		
14) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.		
15) Naphthalene	0.00	128	0	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.	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	0.00	149	0	N.D.		
26) 4-Chlorophenyl-phenyl ethe	0.00	204	0	N.D.		
27) Fluorene	0.00	166	0	N.D.		
29) Azobenzene/ 1,2-Diphenylhyd	0.00	77	0	N.D.		
31) N-Nitrosodiphenylamine	0.00	169	0	N.D.		
32) 4-Bromophenyl-phenyl ether	0.00	248	0	N.D.		
33) Hexachlorobenzene	0.00	284	0	N.D.		
34) Phenanthrene	0.00	178	0	N.D.		
35) Anthracene	0.00	178	0	N.D.		
36) Di-n-butylphthalate	0.00	149	0	N.D.	d	
37) Fluoranthene	0.00	202	0	N.D.		
40) Pyrene	0.00	202	0	N.D.		
41) Butylbenzylphthalate	0.00	149	0	N.D.		
42) Benzo (a) anthracene	0.00	228	0	N.D.	d	
43) Bis (2-ethylhexyl) phthala	30.76	149	80942	3.10	ug/L	99
44) Chrysene	0.00	228	0	N.D.	d	

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

7172.D 5080402BB.M Mon Apr 15 18:35:39 2002

Quantitation Report

(QT Reviewed)

Data File : C:\MSDCHEM\1\DATA\041002\7172.D

Vial: 25

Acq On : 11 Apr 2002 9:43 am

Operator: Bohlk

Sample : SAMPLE 7172 3/27/02

Inst : Instrumen

Misc :

Multiplr: 1.00

MS Integration Params: BENZO.P

Quant Time: Apr 15 18:34:07 2002

Quant Results File: 5080402BB.RES

Quant Method : C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)

Title : Quantitation For Method 508gcscan

Last Update : Mon Apr 15 15:56:12 2002

Response via : Initial Calibration

DataAcq Meth : 508SCAN

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
46) Di-n-Octylphthalate	0.00	149	0	N.D.		
47) Benzo (b) fluoranthene	0.00	252	0	N.D.		
48) Benzo (k) fluoranthene	0.00	252	0	N.D.		
49) Benzo (a) pyrene	0.00	252	0	N.D.	d	
50) Indeno (1,2,3-cd) pyrene	0.00	276	0	N.D.		
51) Dibenz (a,h) anthracene	0.00	278	0	N.D.		
52) Benzo (g,h,i) perylene	0.00	276	0	N.D.		

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

7172.D 5080402BB.M Mon Apr 15 18:35:39 2002

Page 2

Data File : C:\MSDCHEM\1\DATA\041002\7172.D

Vial: 25

Acq On : 11 Apr 2002 9:43 am

Operator: Bohlk

Sample : SAMPLE 7172 3/27/02

Inst : Instrumen

Misc :

Multiplr: 1.00

MS Integration Params: BENZO.P

Quant Time: Apr 15 18:35 2002

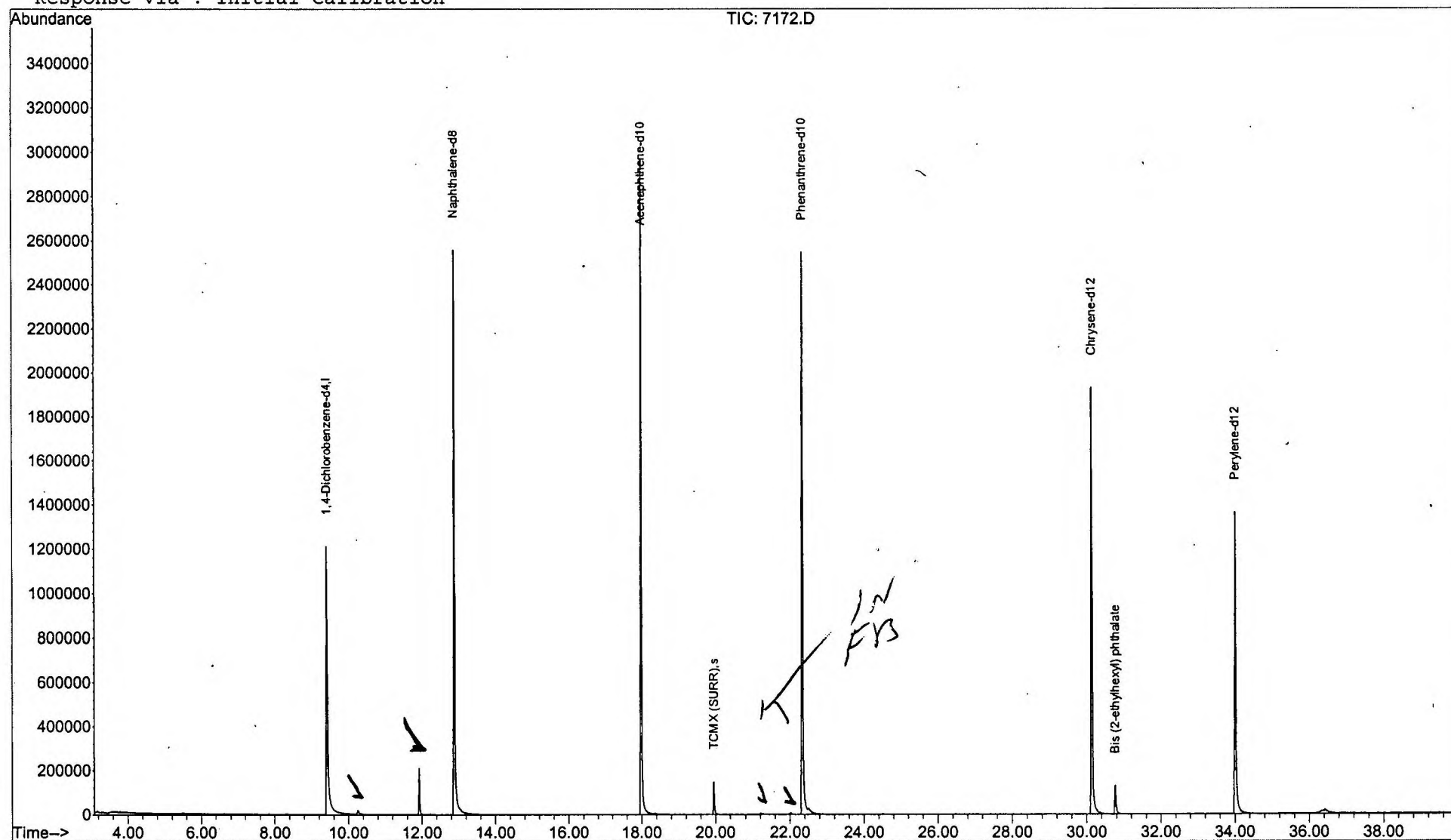
Quant Results File: 5080402BB.RES

Method : C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)

Title : Quantitation For Method 508gcscan

Last Update : Mon Apr 15 15:56:12 2002

Response via : Initial Calibration



LSC Area Percent Report

Data File : C:\MSDCHEM\1\DATA\041002\7172.D

Vial: 25

Acq On : 11 Apr 2002 9:43 am

Operator: Bohlk

Sample : SAMPLE 7172 3/27/02

Inst : Instrumen

Misc :

Multiplr: 1.00

MS Integration Params: LSCINT.P

Method : C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)

Title : Quantitation For Method 508gcscan

Smoothing : OFF

Filtering: 5

Sampling : 1

Min Area: 1 % of largest Peak

Start Thrs: 0.2

Max Peaks: 100

Stop Thrs : 0

Peak Location: TOP

If leading or trailing edge < 100 prefer < Baseline drop else tangent >

Peak separation: 5

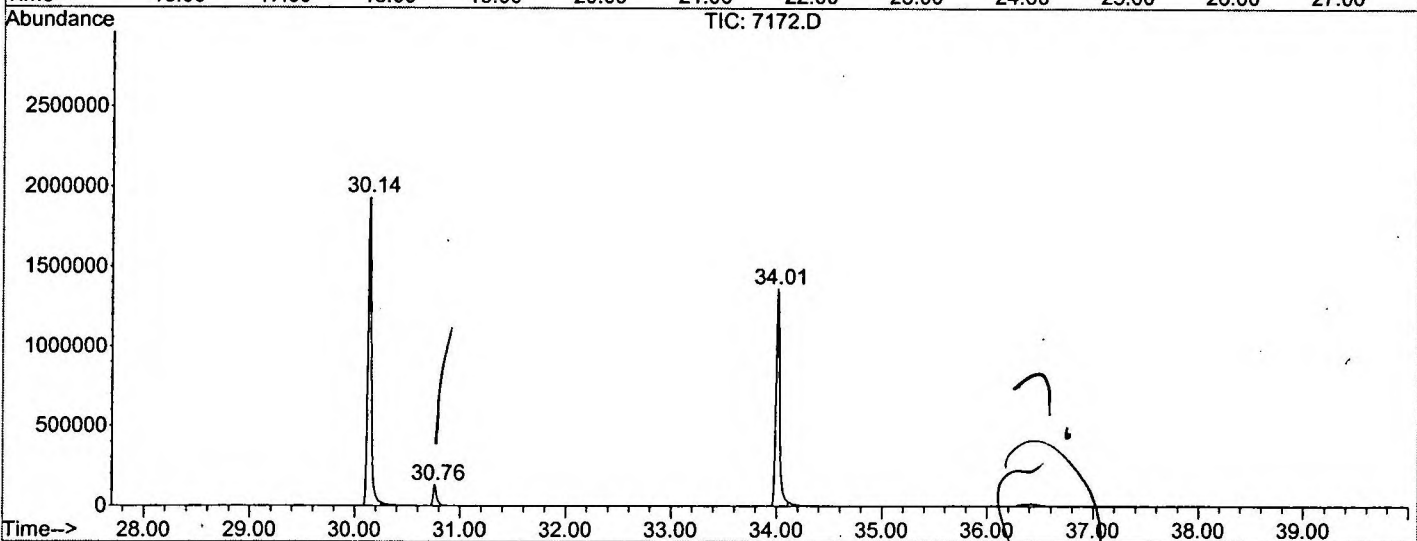
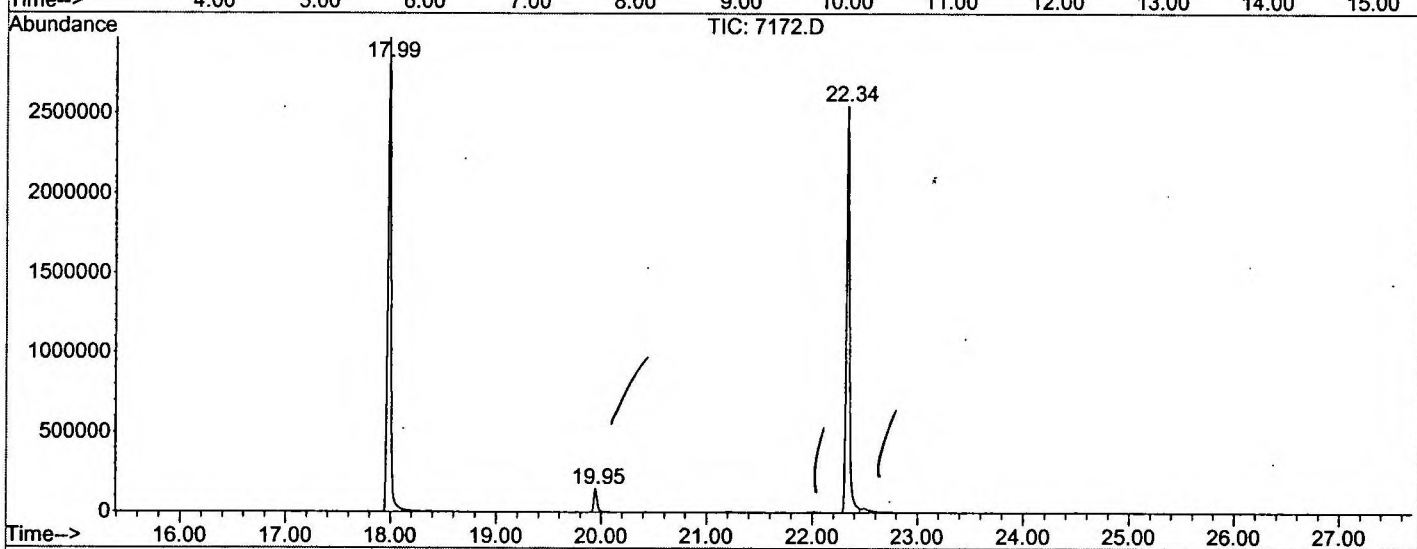
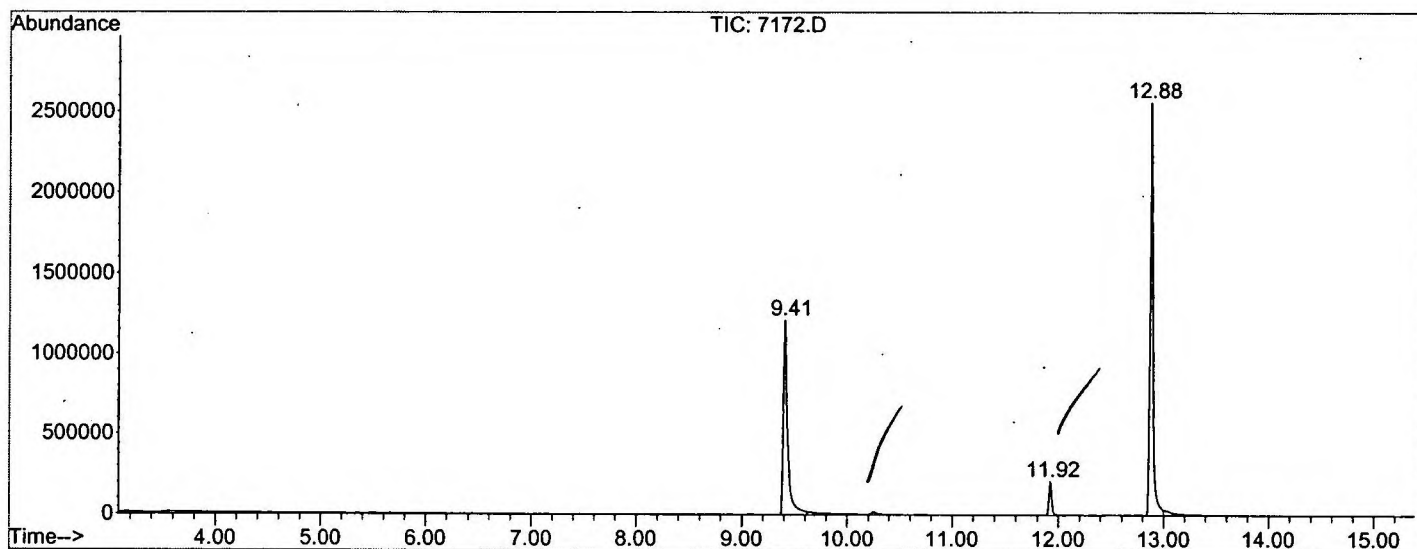
Signal : TIC

peak #	R.T. min	first scan	max scan	last scan	PK TY	peak height	corr. area	corr. % max.	% of total
1	9.407	1070	1077	1113	rBV	1210125	3333277	55.76%	11.413%
2	11.922	1498	1505	1514	rBV	209162	372961	6.24%	1.277%
3	12.879	1661	1668	1688	rBV	2558115	5326561	89.10%	18.238%
4	17.985	2528	2537	2559	rBV	2966822	5978329	100.00%	20.470%
5	19.948	2864	2871	2884	rBV3	147365	308231	5.16%	1.055%
6	22.339	3269	3278	3297	rBV2	2548575	5669692	94.84%	19.413%
7	30.142	4596	4606	4625	rBV2	1932435	4558480	76.25%	15.608%
8	30.765	4703	4712	4727	rBV	129489	309531	5.18%	1.060%
9	34.008	5255	5264	5281	rBV2	1363649	3348556	56.01%	11.465%

Sum of corrected areas: 29205618

LSC Report - Integrated Chromatogram

File : C:\MSDCHEM\1\DATA\041002\7172.D
 Operator : Bohlk
 Acquired : 11 Apr 2002 9:43 am using AcqMethod 508SCAN
 Instrument : Instrumen
 Sample Name: SAMPLE 7172 3/27/02
 Misc Info :
 Vial Number: 25
 Quant File :5080402BB.RES (RTE Integrator)



Library Search Compound Report

Data File : C:\MSDCHEM\1\DATA\041002\7172.D

Acq On : 11 Apr 2002 9:43 am

Sample : SAMPLE 7172 3/27/02

Misc :

MS Integration Params: LSCINT.P

Vial: 25

Operator: Bohlk

Inst : Instrumen

Multiplr: 1.00

Quant Method : C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)

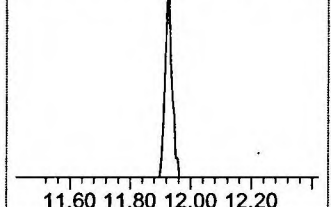
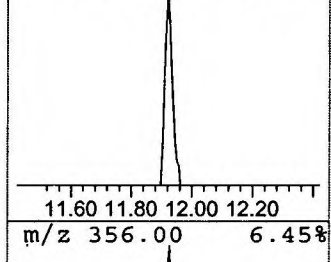
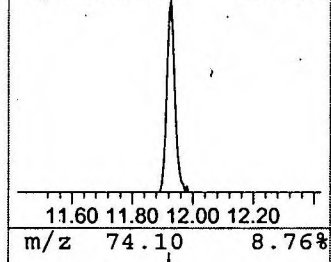
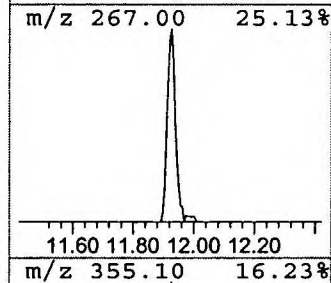
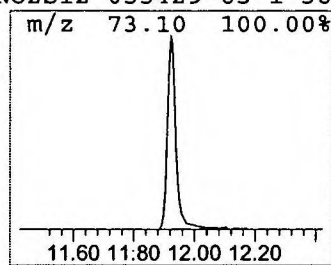
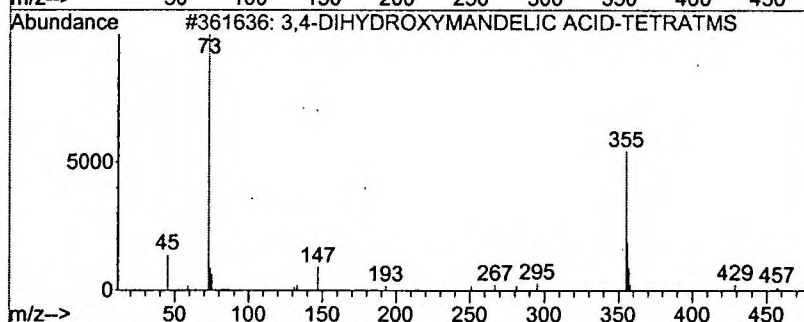
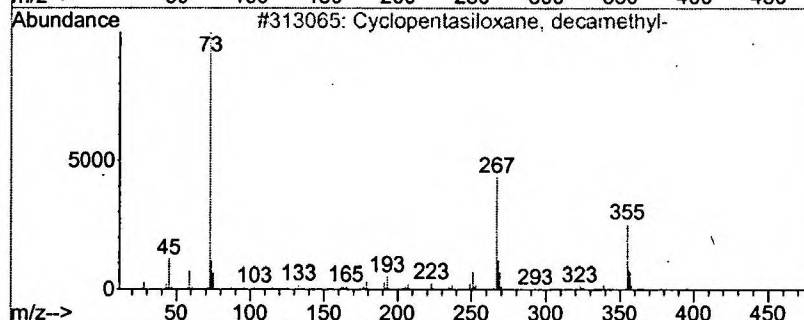
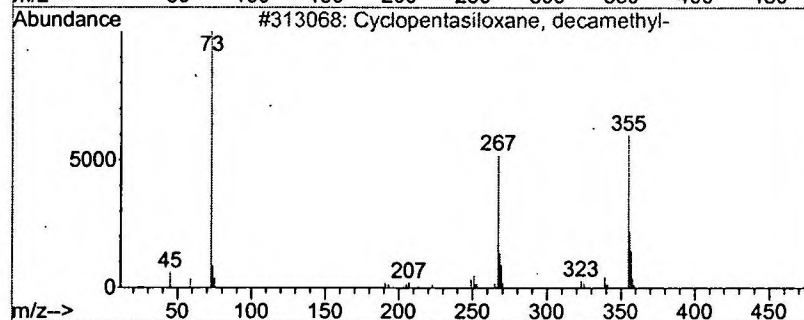
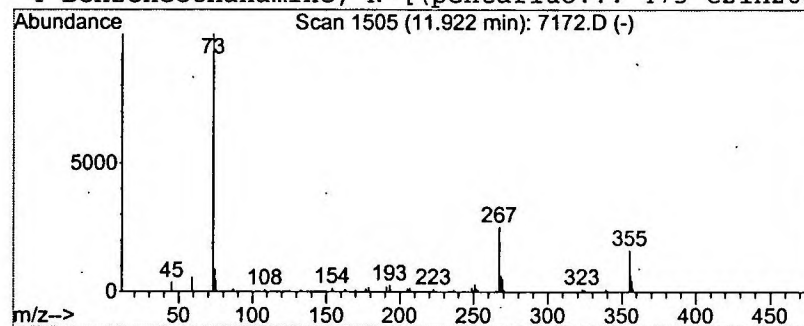
Title : Quantitation For Method 508gcscan

Library : C:\DATABASE\WILEY7N.L

Peak Number 1 Cyclopentasiloxane, decamet... Concentration Rank 1

R.T.	EstConc	Area	Relative to ISTD	R.T.
11.92	2.80 ug/L	372961	Naphthalene-d8	12.88

Hit#	of	Tentative ID	MW	MolForm	CAS#	Qual.
1		Cyclopentasiloxane, decamethyl-	370	C10H30O5Si5	000541-02-6	90
2		Cyclopentasiloxane, decamethyl-	370	C10H30O5Si5	000541-02-6	59
3		3,4-DIHYDROXYMANDELIC ACID-TETRATMS	472	C20H40O5Si4	000000-00-0	38
4		Benzeneethanamine, N-[(pentafluoroethyl)amino]-	475	C21H26F5NO2Si2	055429-85-1	38

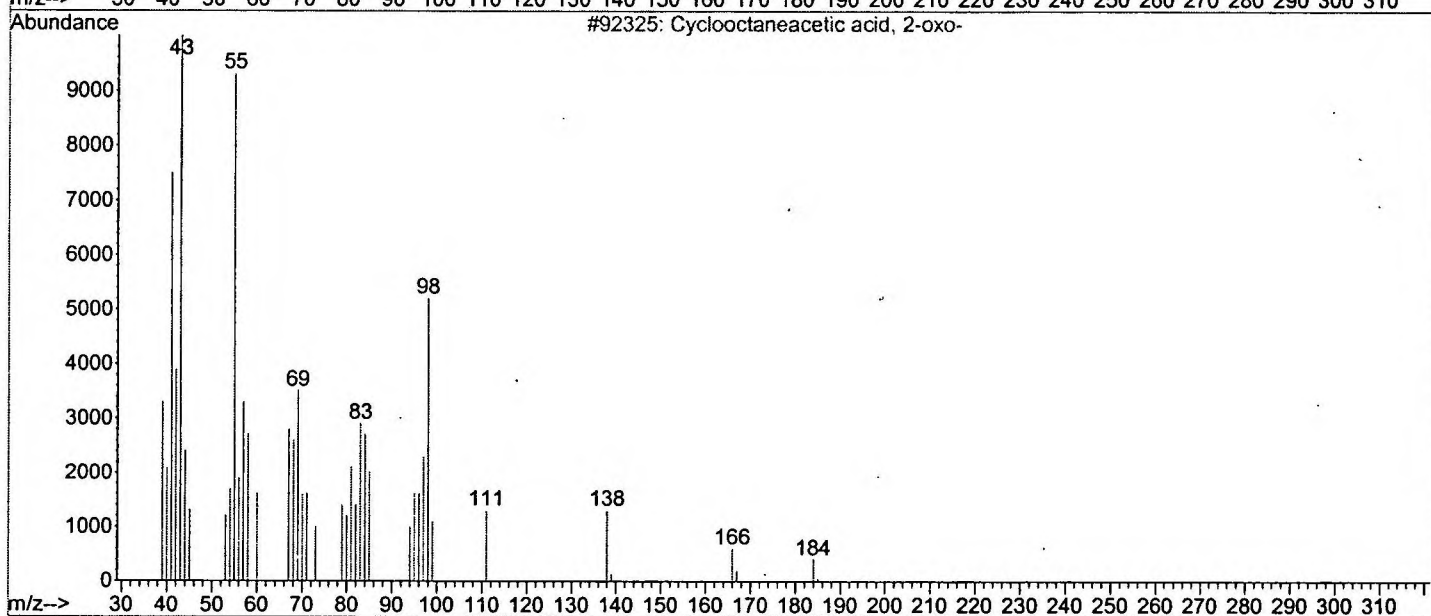
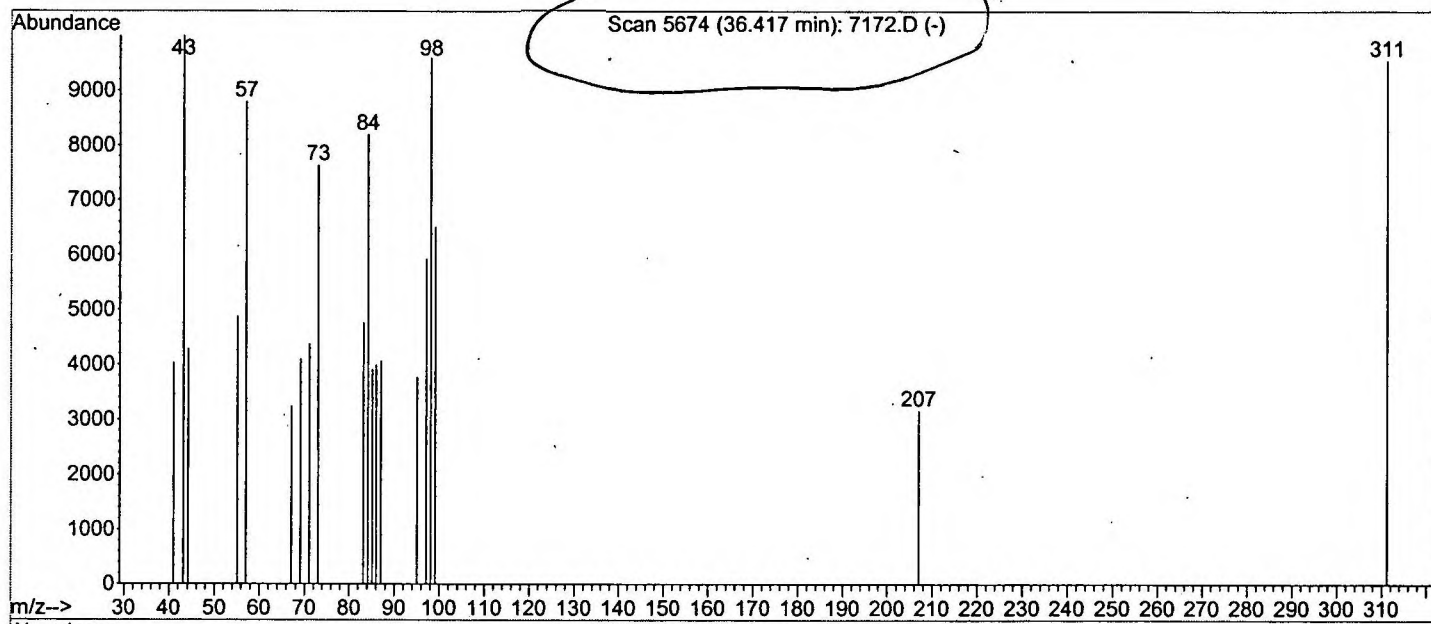


Tentatively Identified Compound (LSC) summary

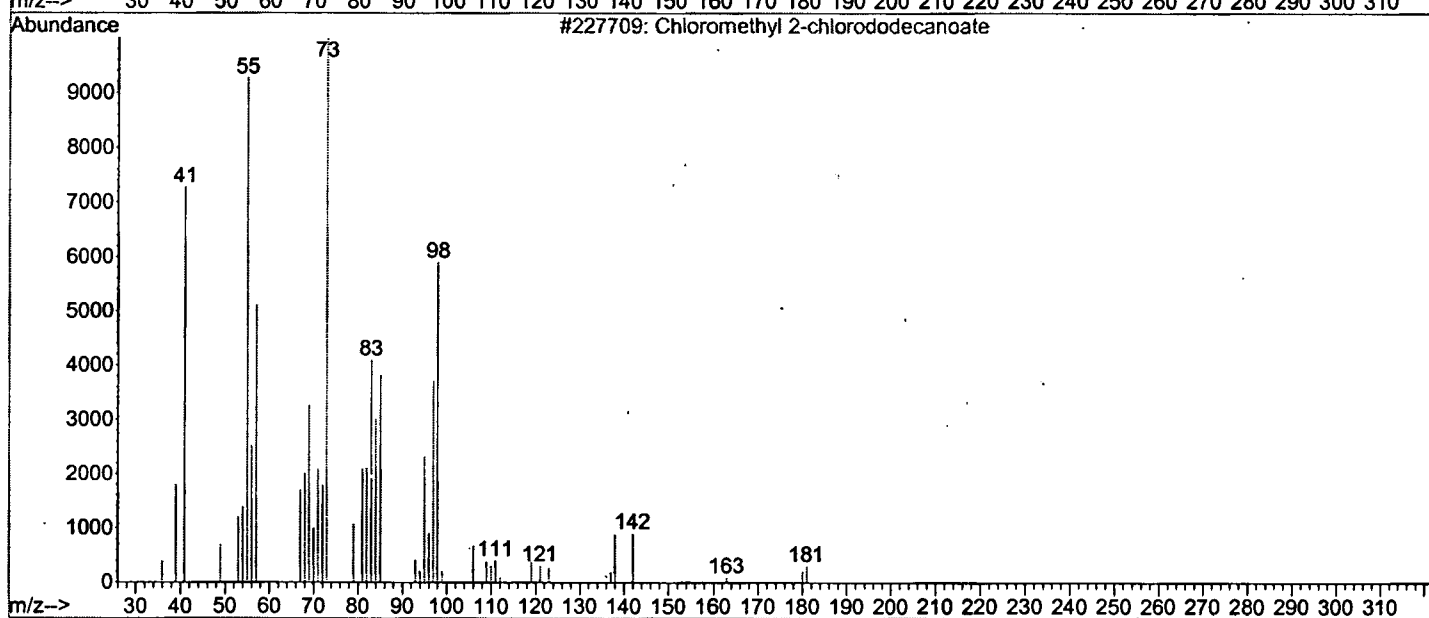
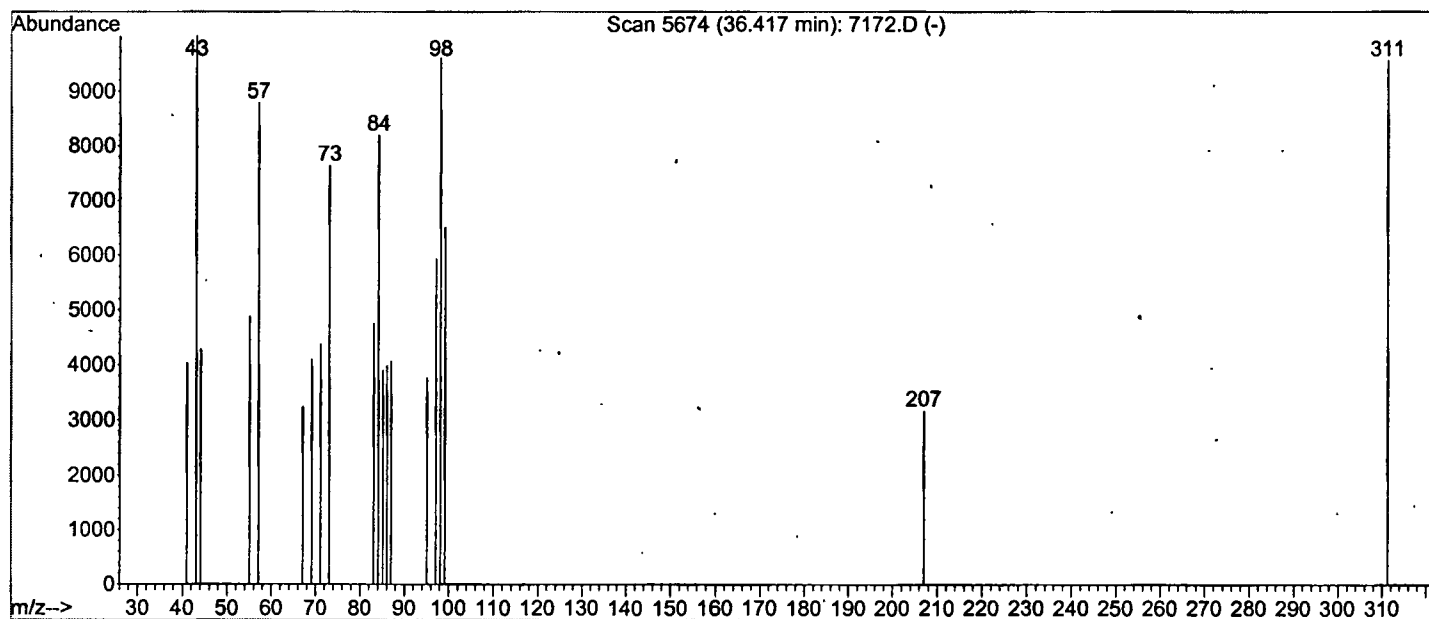
Operator ID: Bohlk Date Acquired: 11 Apr 2002 9:43 am
 Data File: C:\MSDCHEM\1\DATA\041002\7172.D
 Name: SAMPLE 7172 3/27/02
 Misc:
 Method: C:\MSDCHEM\1\5973N\5080402BB.M (RTE Integrator)
 Title: Quantitation For Method 508gcscan
 Library Searched: C:\DATABASE\WILEY7N.L

TIC Top Hit name	RT	EstConc	Units	Response	---Internal Standard---			
					#	RT	Resp	Conc
Cyclopentasiloxan...	11.92	2.8 ug/L		372961	2	12.88	5326560	40.0

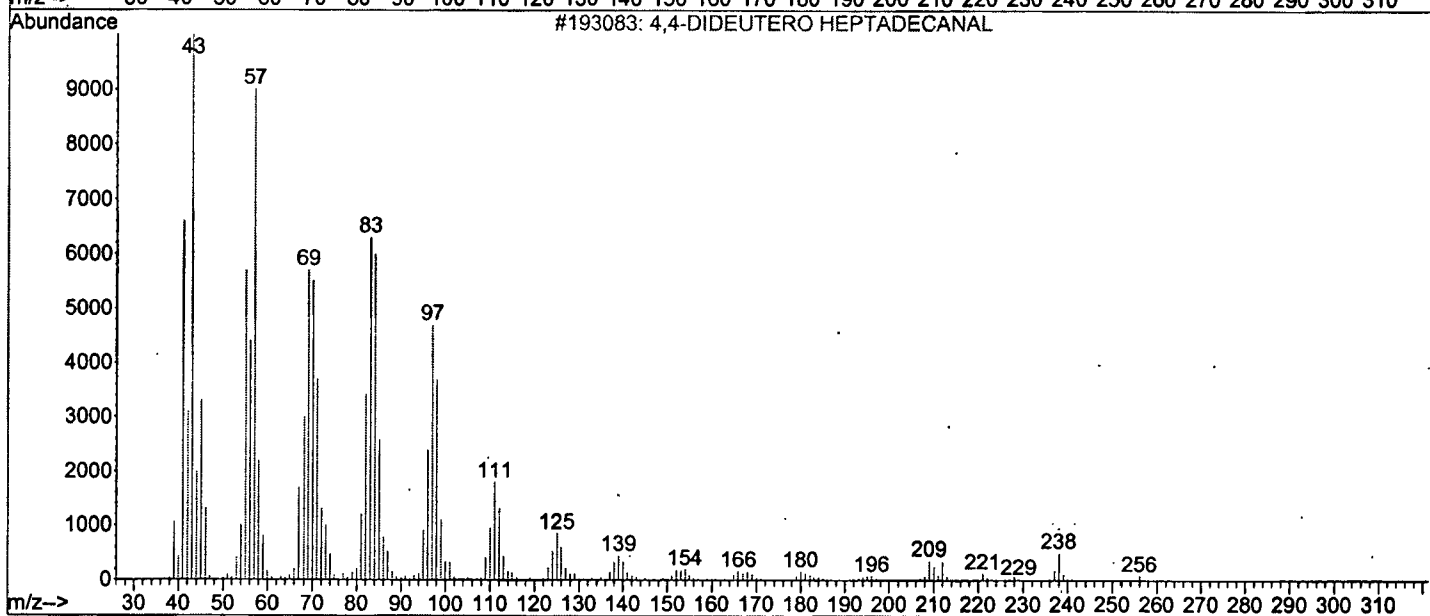
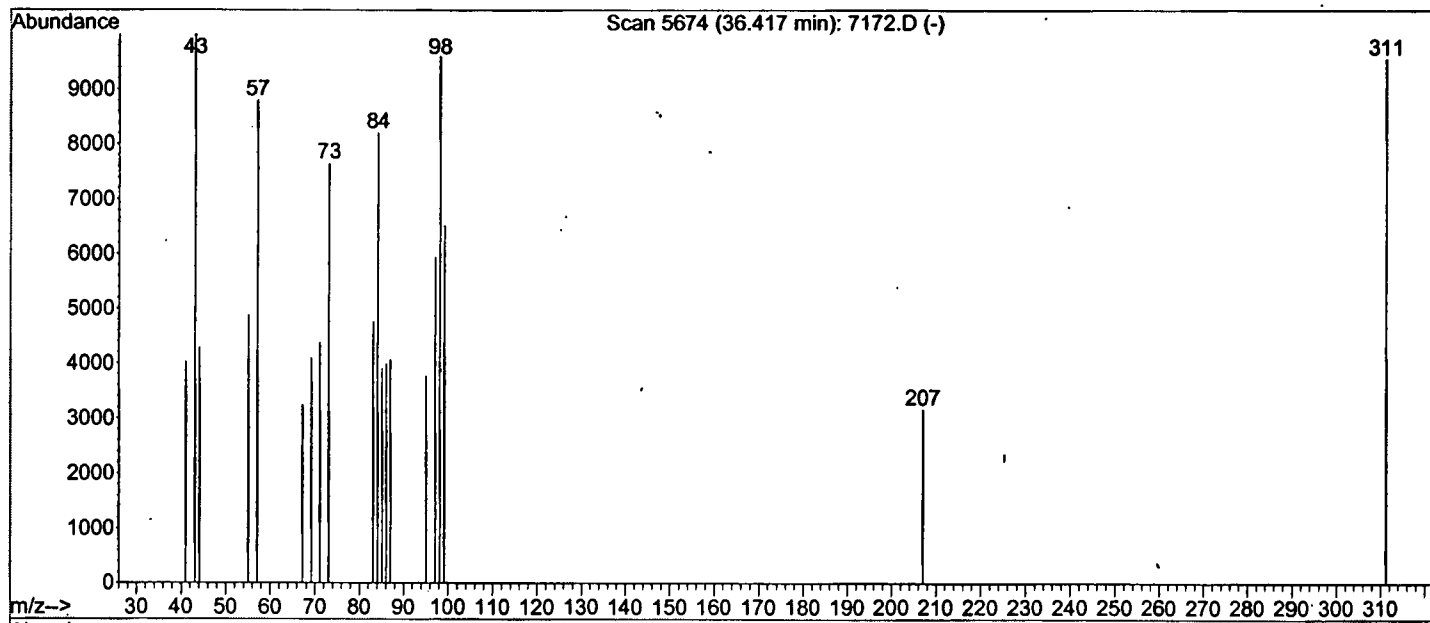
Library Searched : C:\Database\wiley7n.1
 Quality : 38
 ID : Cyclooctaneacetic acid, 2-oxo



Library Searched : C:\Database\wiley7n.1
 Quality : 27
 ID : Chloromethyl 2-chlorododecanoate



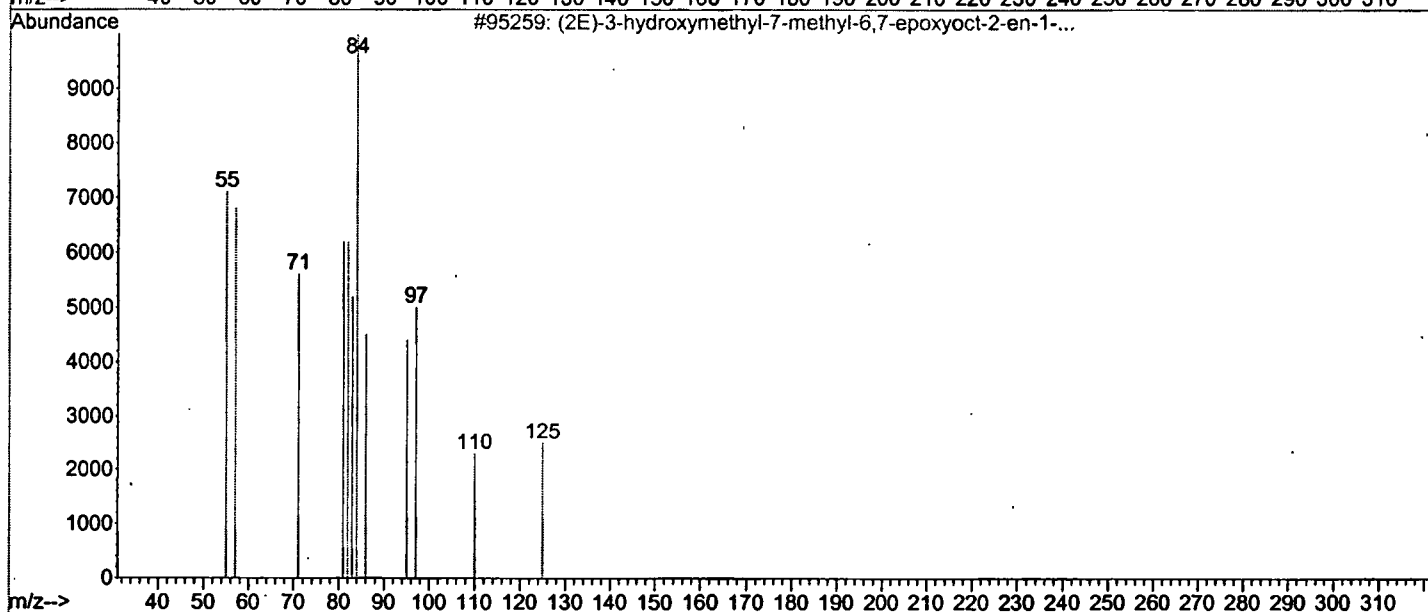
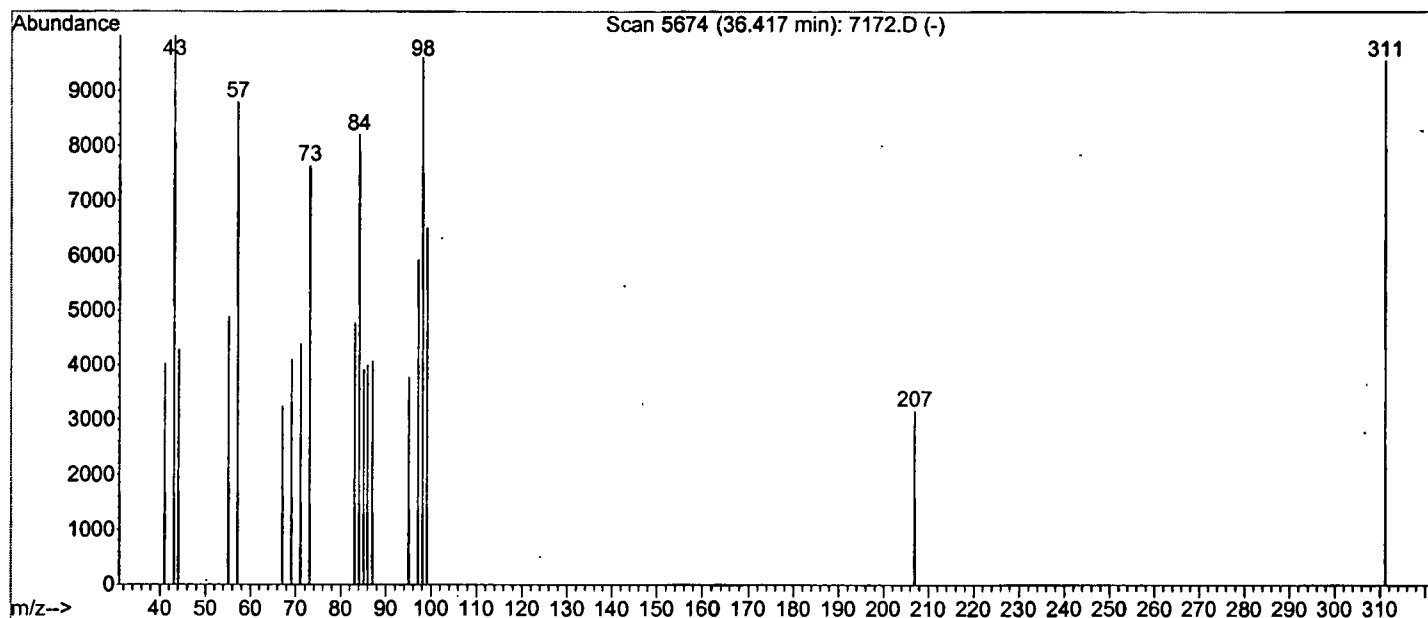
Library Searched : C:\Database\wiley7n.1
 Quality : 16
 ID : 4,4-DIDEUTERO HEPTADECANAL



Library Searched : C:\Database\wiley7n.1

Quality : 14

ID : (2E)-3-hydroxymethyl-7-methyl-6,7-epoxyoct-2-en-1-ol \$\$ 2-Butene-1,4-diol, 2-[2-(3,3-dimethyloxiranyl)ethyl]-, (E)- (CAS)



Library Searched : C:\Database\wiley7n.1

Quality : 14

ID : Piperidine, 4-methyl- (CAS) \$\$ 4-Pipecoline \$\$ 4-Methylpiperidine \$\$.
gamma.-Pipecoline \$\$.gamma.-Pipecoline-4-methylpiperidine

