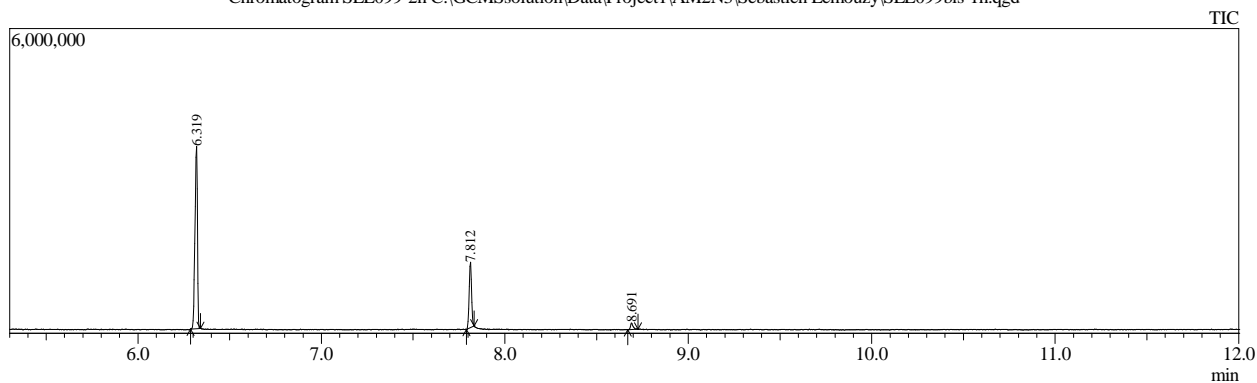


Sample Information

Sample Name : SLE099bis-1h
Vial # : 26
Injection Volume : 1.00
Data File : C:\GCMSsolution\Data\Project1\AM2N3\Sébastien Lemouzy\SLE099bis-1h.qgd
Method File : C:\GCMSsolution\Data\Project1\AM2N3\Sébastien Lemouzy\50-280 (split 10) début f
Tuning File : C:\GCMSsolution\System\Tune1\2021.01.04.qgt

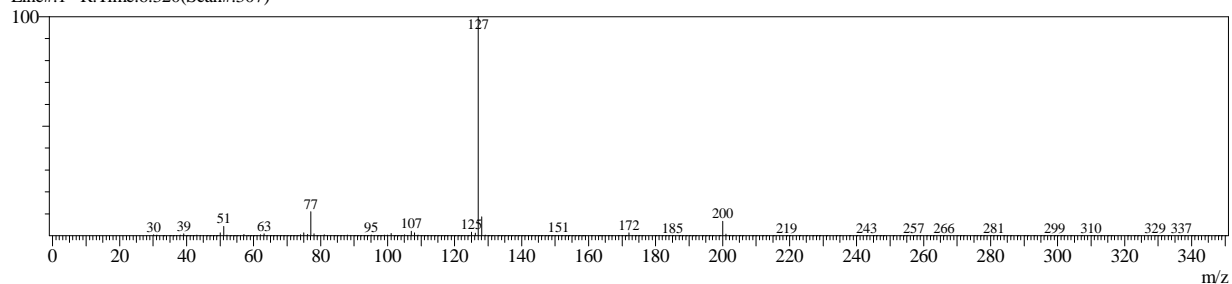
Chromatogram SLE099-2h C:\GCMSsolution\Data\Project1\AM2N3\Sébastien Lemouzy\SLE099bis-1h.qgd



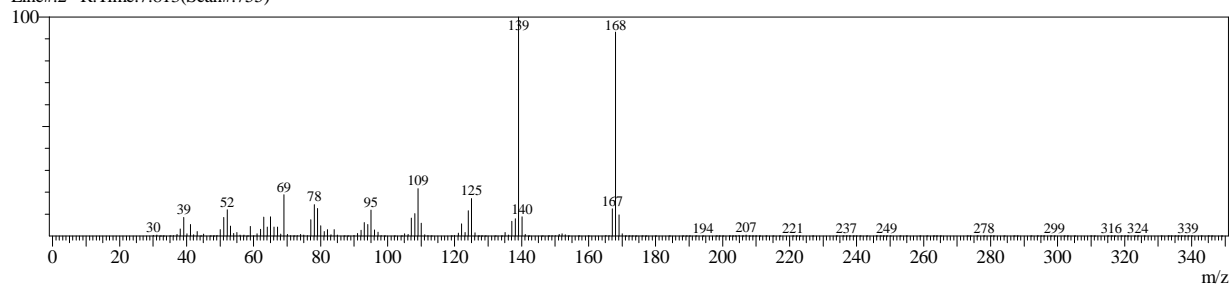
Peak Report							Name
Peak#	R.Time	I.Time	F.Time	Area	Area%	Height	
1	6.319	6.287	6.340	3335104	72.49	3620532	Ethyl 2,2-difluoro phenylacetate
2	7.812	7.790	7.833	1133611	24.64	1298900	Benzene, 1,3,5-trimethoxy-
3	8.691	8.670	8.727	132310	2.88	127274	Hexyl 2,2-difluoro phenylacetate
				4601025	100.00	5046706	

Spectrum

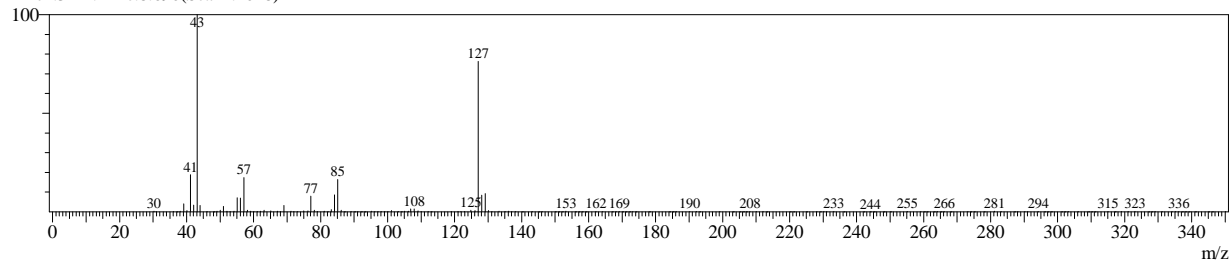
Line#1 R.Time:6.320(Scan#:307)



Line#2 R.Time:7.813(Scan#:755)



Line#3 R.Time:8.690(Scan#:1018)



Method

[Comment]

===== Analytical Line 1 =====

[AOC-20i+s]

of Rinses with Presolvent :6
 # of Rinses with Solvent(post) :6
 # of Rinses with Sample :2
 Plunger Speed(Suction) :High
 Viscosity Comp. Time :0.2 sec
 Plunger Speed(Injection) :High
 Syringe Insertion Speed :High
 Injection Mode :Normal
 Pumping Times :5
 Inj. Port Dwell Time :0.0 sec
 Terminal Air Gap :No
 Plunger Washing Speed :High
 Washing Volume :8uL
 Syringe Suction Position :0.0 mm
 Syringe Injection Position :0.0 mm
 Solvent Selection :All A,B,C

[GC-2010]

Column Oven Temp.	:50.0 °C	
Injection Temp.	:250.00 °C	
Injection Mode	:Split	
Flow Control Mode	:Linear Velocity	
Pressure	:108.3 kPa	
Total Flow	:11.1 mL/min	
Column Flow	:0.74 mL/min	
Linear Velocity	:38.2 cm/sec	
Purge Flow	:3.0 mL/min	
Split Ratio	:10.0	
High Pressure Injection	:OFF	
Carrier Gas Saver	:ON	
Carrier Gas Saver Split Ratio	:10.0	
Carrier Gas Saver Time	:1.00 min	
Splitter Hold	:OFF	
Oven Temp. Program		
Rate	Temperature(°C)	Hold Time(min)
-	50.0	2.00
22.00	280.0	2.00

< Ready Check Heat Unit >

Column Oven : Yes
 SPL1 : Yes
 MS : Yes

< Ready Check Detector(FTD) >

< Ready Check Baseline Drift >

< Ready Check Injection Flow >

SPL1 Carrier : Yes
 SPL1 Purge : Yes

< Ready Check APC Flow >

< Ready Check Detector APC Flow >

External Wait :No

Equilibrium Time :1.0 min

[GC Program]

[GCMS-QP2010 SE]

IonSourceTemp :200.00 °C
 Interface Temp. :280.00 °C
 Solvent Cut Time :1.50 min
 Detector Gain Mode :Relative
 Detector Gain :0.89 kV +0.00 kV
 Threshold :0

[MS Table]

--Group 1 - Event 1--

Start Time :5.30min
 End Time :14.45min
 ACQ Mode :Scan
 Event Time :0.20sec
 Scan Speed :5000
 Start m/z :30.00
 End m/z :800.00

Sample Inlet Unit :GC

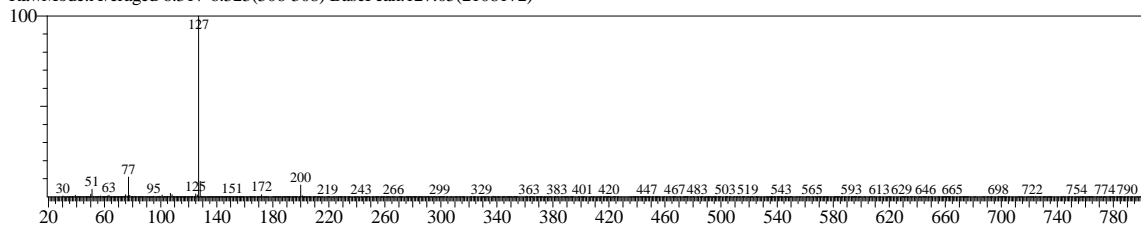
[MS Program]

Use MS Program :OFF

Library

Line#:1 R.Time:6.320(Scan#:307) MassPeaks:474

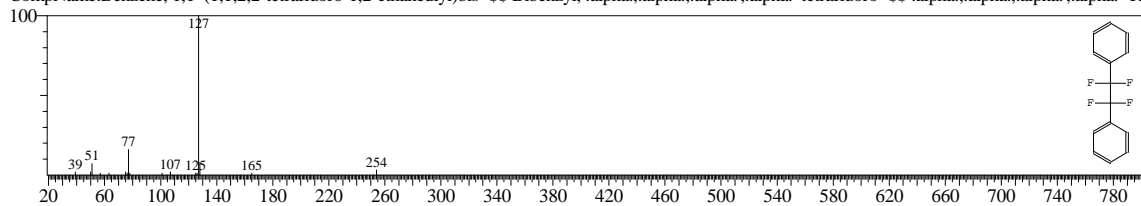
RawMode:Averaged 6.317-6.323(306-308) BasePeak:127.05(2108172)



Hit#1 Entry:78799 Library:NIST08.LIB

SI:91 Formula:C14H10F4 CAS:425-32-1 MolWeight:254 RetIndex:1289

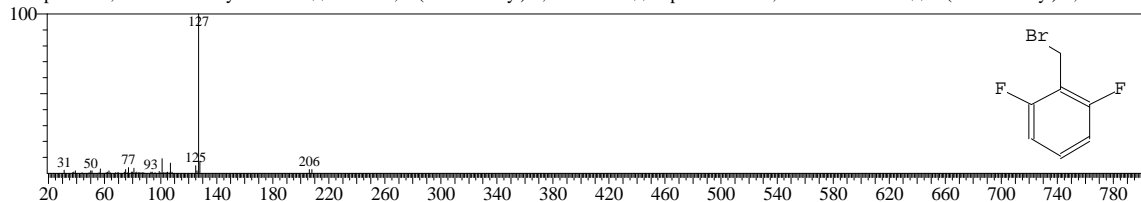
CompName:Benzen, 1,1'-(1,1,2,2-tetrafluoro-1,2-ethanediyl)bis- \$Bibenzyl, .alpha.,.alpha.,.alpha.',.alpha.'-tetrafluoro- \$\$.alpha.,.alpha.,.alpha.',.alpha.'-Te



Hit#2 Entry:46037 Library:NIST08.LIB

SI:86 Formula:C7H5BrF2 CAS:85118-00-9 MolWeight:206 RetIndex:1040

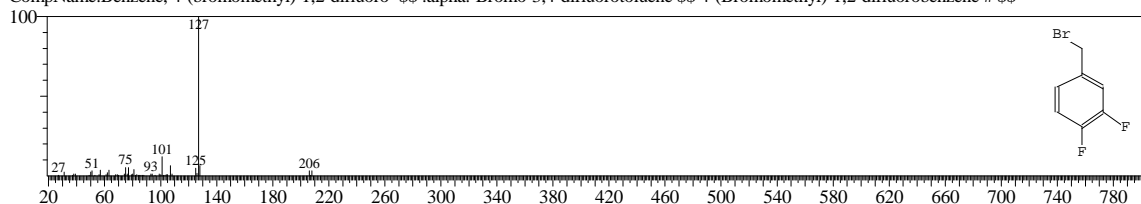
CompName:2,6-Difluorobenzyl bromide \$Benzene, 2-(bromomethyl)-1,3-difluoro- \$\$.alpha.-Bromo-2,6-difluorotoluene \$2-(Bromomethyl)-1,3-difluor



Hit#3 Entry:46038 Library:NIST08.LIB

SI:84 Formula:C7H5BrF2 CAS:85118-01-0 MolWeight:206 RetIndex:1040

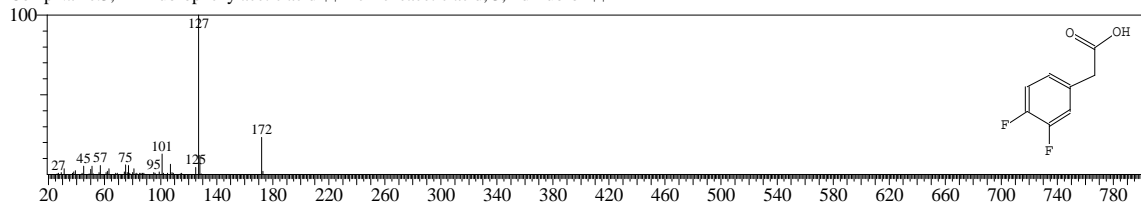
CompName:Benzen, 4-(bromomethyl)-1,2-difluoro- \$\$.alpha.-Bromo-3,4-difluorotoluene \$4-(Bromomethyl)-1,2-difluorobenzene # \$



Hit#4 Entry:26302 Library:NIST08.LIB

SI:84 Formula:C8H6F2O2 CAS:658-93-5 MolWeight:172 RetIndex:1199

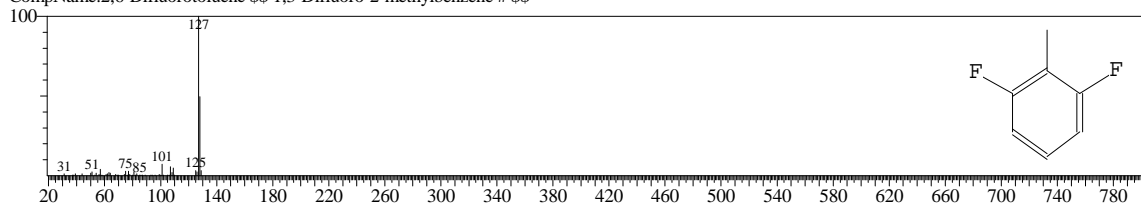
CompName:3,4-Difluorophenylacetic acid \$Benzeneacetic acid, 3,4-difluoro- \$



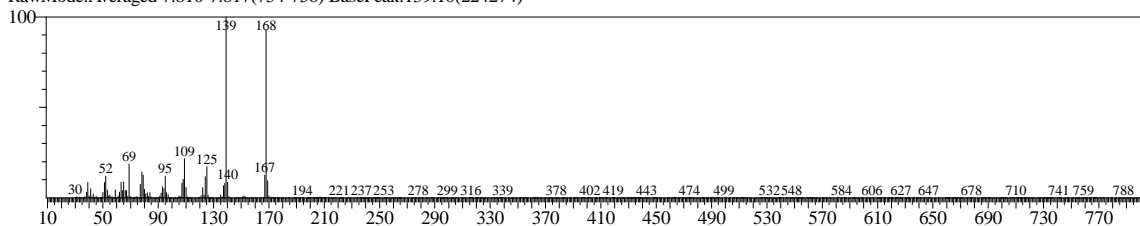
Hit#5 Entry:7081 Library:NIST08.LIB

SI:83 Formula:C7H6F2 CAS:443-84-5 MolWeight:128 RetIndex:743

CompName:2,6-Difluorotoluene \$1,3-Difluoro-2-methylbenzene # \$



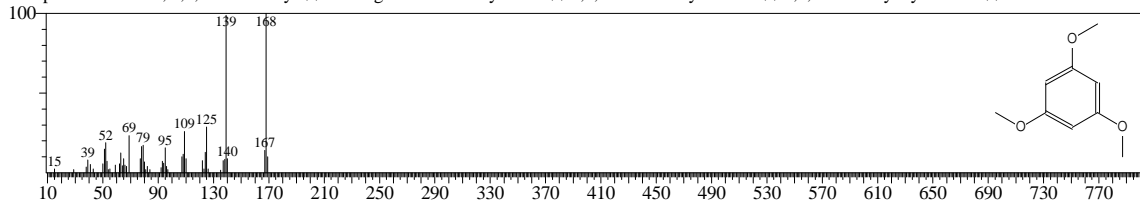
Line#:2 R.Time:7.813(Scan#:755) MassPeaks:452
RawMode:Averaged 7.810-7.817(754-756) BasePeak:139.10(224274)



Hit#:1 Entry:24091 Library:NIST08.LIB

SI:96 Formula:C₉H₁₂O₃ CAS:621-23-8 MolWeight:168 RetIndex:1248

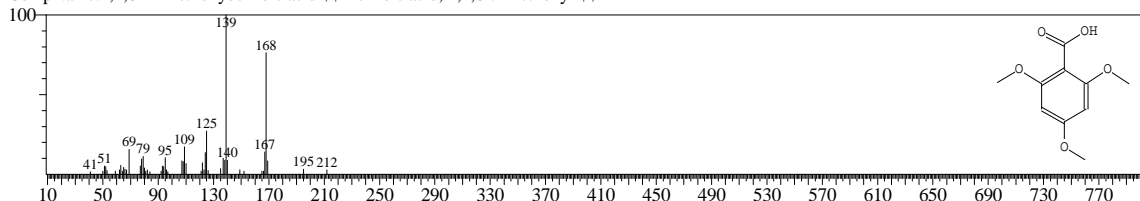
CompName:Benzene, 1,3,5-trimethoxy- \$\$ Phloroglucinol trimethyl ether \$\$ 1,3,5-Trimethoxybenzene \$\$ 1,3,5-Trimethoxybenzene \$\$



Hit#:2 Entry:50345 Library:NIST08.LIB

SI:92 Formula:C₁₀H₁₂O₅ CAS:570-02-5 MolWeight:212 RetIndex:1717

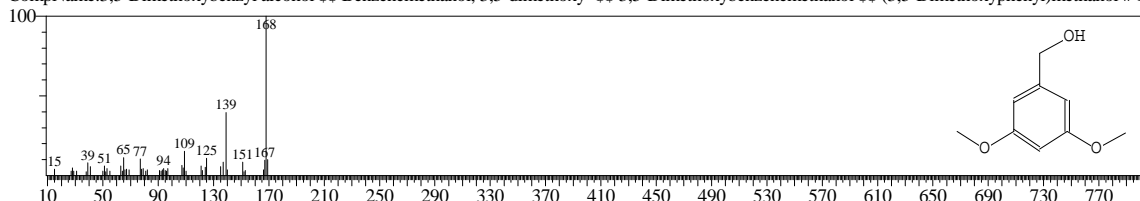
CompName:2,4,6-Trimethoxybenzoic acid \$\$ Benzoic acid, 2,4,6-trimethoxy- \$\$



Hit#:3 Entry:24090 Library:NIST08.LIB

SI:84 Formula:C₉H₁₂O₃ CAS:705-76-0 MolWeight:168 RetIndex:1415

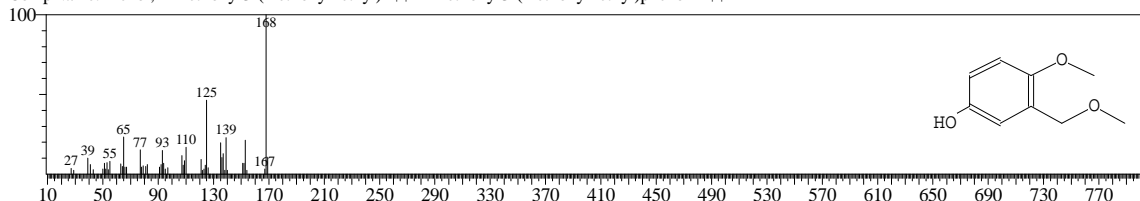
CompName:3,5-Dimethoxybenzyl alcohol \$\$ Benzenemethanol, 3,5-dimethoxy- \$\$ 3,5-Dimethoxybenzenemethanol \$\$ (3,5-Dimethoxyphenyl)methanol # \$



Hit#:4 Entry:24087 Library:NIST08.LIB

SI:76 Formula:C₉H₁₂O₃ CAS:59907-65-2 MolWeight:168 RetIndex:1379

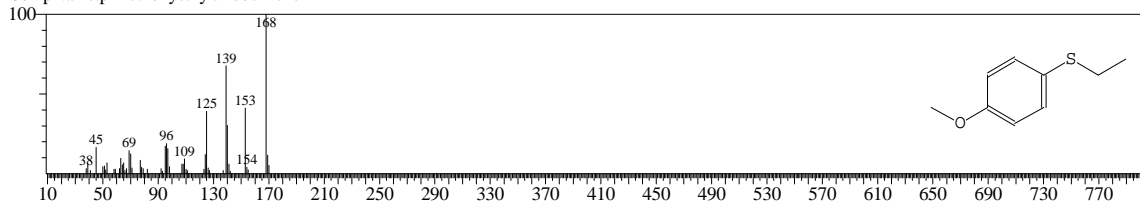
CompName:Phenol, 4-methoxy-3-(methoxymethyl)- \$\$ 4-Methoxy-3-(methoxymethyl)phenol # \$\$



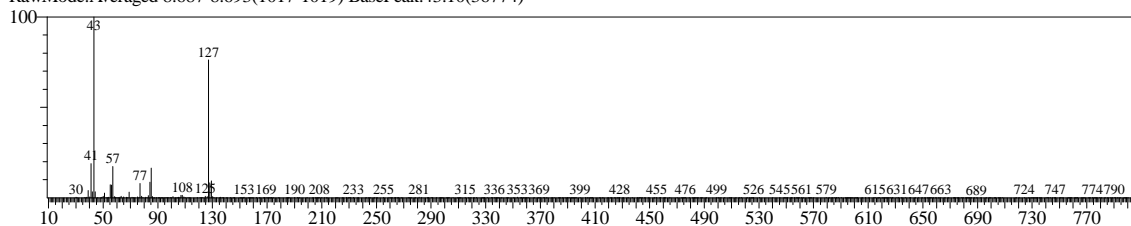
Hit#:5 Entry:24025 Library:NIST08.LIB

SI:76 Formula:C₉H₁₂O₂S CAS:0-00-0 MolWeight:168 RetIndex:1333

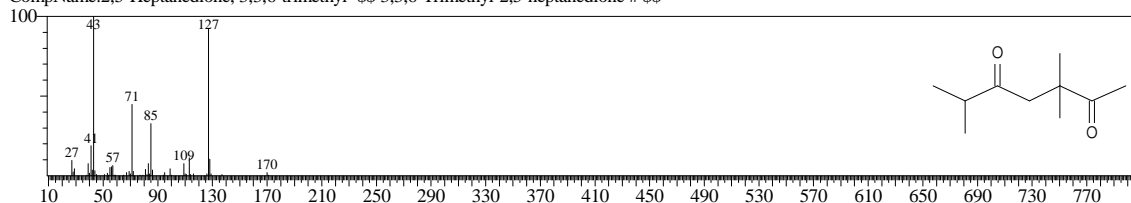
CompName:p-methoxyethylthiobenzene



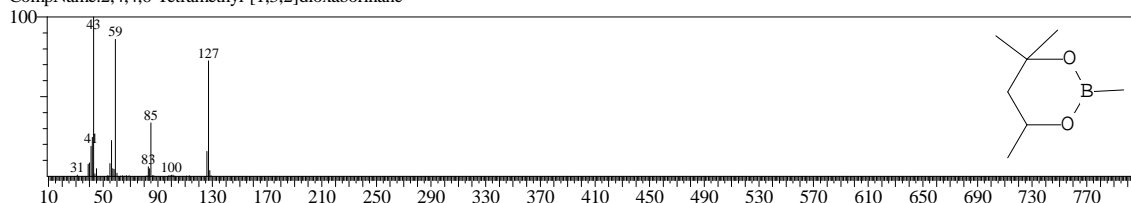
Line#3 R.Time:8.690(Scan#:1018) MassPeaks:422
RawMode:Averaged 8.687-8.693(1017-1019) BasePeak:43.10(36774)



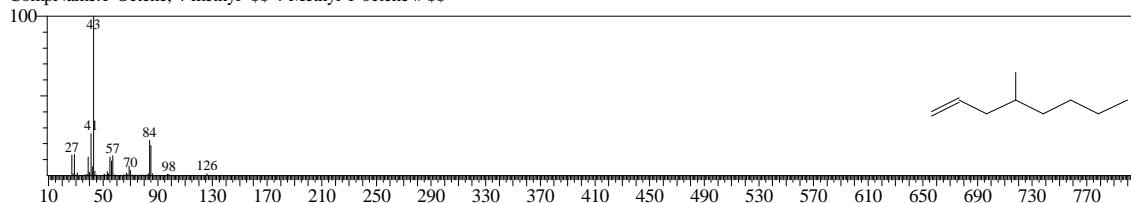
Hit#1 Entry:25451 Library:NIST08.LIB
SI:84 Formula:C10H18O2 CAS:51513-40-7 MolWeight:170 RetIndex:1139
CompName:2,5-Heptanedione, 3,3,6-trimethyl- \$\$ 3,3,6-Trimethyl-2,5-heptanedione # \$\$



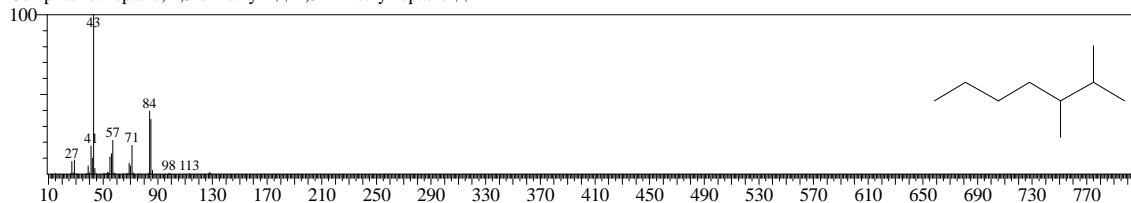
Hit#2 Entry:11931 Library:NIST08.LIB
SI:81 Formula:C7H15BO2 CAS:211613-23-9 MolWeight:142 RetIndex:0
CompName:2,4,4,6-Tetramethyl-[1,3,2]dioxaborinane



Hit#3 Entry:6629 Library:NIST08.LIB
SI:81 Formula:C9H18 CAS:13151-12-7 MolWeight:126 RetIndex:842
CompName:1-Octene, 4-methyl- \$\$ 4-Methyl-1-octene # \$\$



Hit#4 Entry:7400 Library:NIST08.LIB
SI:80 Formula:C9H20 CAS:3074-71-3 MolWeight:128 RetIndex:788
CompName:Heptane, 2,3-dimethyl- \$\$ 2,3-Dimethylheptane \$\$



Hit#5 Entry:78913 Library:NIST08.LIB
SI:80 Formula:C14H22O4 CAS:0-00-0 MolWeight:254 RetIndex:1806
CompName:5,7-Dodecadiene-4,9-dione, 6,7-dihydroxy-2,11-dimethyl-

