

SUPPLEMENTARY FILE

Docking-Based Computational Analysis of Guava (*Psidium guajava*) Leaves Derived Bioactive Compounds as Coagulation Factor IXa Inhibitor

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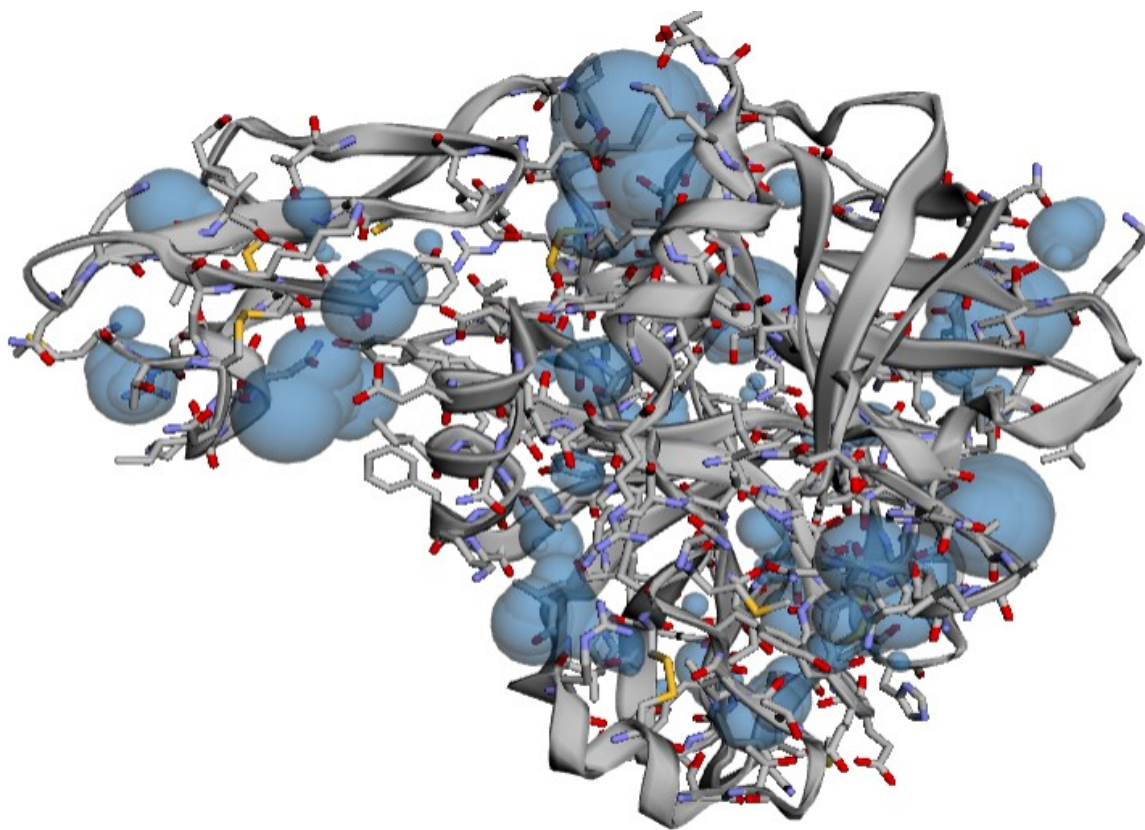


Figure S1. *Structural Representation of Identified CFIXa Active Sites*

Table S1. *Characteristic of All CFIXa Active Sites*

Active Site ID	Area (Å ²)	Volume (Å ³)	Active Site ID	Area (Å ²)	Volume (Å ³)	Active Site ID	Area (Å ²)	Volume (Å ³)
1	215.899	128.236	15	15.795	2.932	29	1.052	0.037
2	122.970	49.494	16	15.658	2.191	30	0.814	0.026
3	90.078	45.053	17	15.300	2.035	31	0.381	0.024
4	46.643	29.920	18	5.682	1.332	32	0.476	0.013
5	55.432	25.515	19	8.210	0.866	33	0.445	0.011
6	44.564	16.115	20	5.331	0.485	34	0.374	0.009
7	61.362	15.776	21	3.829	0.446	35	0.382	0.007
8	44.347	7.487	22	5.476	0.443	36	0.378	0.006
9	19.565	6.010	23	4.648	0.426	37	0.244	0.005
10	25.308	5.975	24	3.502	0.282	38	0.198	0.004
11	18.459	5.680	25	2.816	0.257	39	0.094	0.001
12	23.415	4.408	26	1.059	0.114	40	0.005	0.000
13	8.783	4.150	27	2.275	0.091	41	0.003	0.000
14	11.694	3.888	28	1.258	0.044	42	0.011	0.000

Table S2. Coagulation Factor IXa Chosen Active Sites Information

Active Site ID	Area (Å ²)	Volume (Å ³)	Amino Acid	Atom	Bond	Bond Length (Å)	Amino Acid	Atom	Bond	Bond Length (Å)
II	122.970	49.494	H:HIS57	NE2	CD2-NE2	1.36914	H:GLY216	N	N-CA	1.45804
			H:TYR99	CE1	CE1-CZ	1.37854		CA		
				OH	CZ-OH	1.37181		O		
			H:ASP189	CG	CG-OD1	1.25708	H:GLU217	N	N-CA	1.44962
				OD1				O	C-O	1.24217
				OD2	CG-OD2	1.27266		H:GLU219	O	C-O
			H:SER190	C	C-O	1.22405	H:CYS220	CA	CA-CB	1.52297
				O				SG	CB-SG	1.79269
				CB	CB-OG	1.41759	H:ALA221	N	N-CA	1.45148
				OG				CA		
			H:CYS191	N	N-CA	1.4671	H:MET221	N	N-CA	1.45196
				CA				C	C-O	1.24155
				C	C-O	1.22405	H:LYS224	O		
				O				CB		
			H:GLN192	N	N-CA	1.47061	H:TYR225	N	N-CA	1.4459
				CA				CA		
CG	CG-CD	1.50074		C	C-O	1.19949				
CD				O						

				NE2	CD-NE2	1.33314	H:GLY226	CA	CA-C	1.51956		
				H:SER195	OG	CB-OG	1.41511	H:ILE227	N	N-CA	1.45471	
				H:ILE213	CG2	CB-CG1	1.51821		O	C-O	1.24276	
				H:SER214	C	C-O	1.24355	H:TYR228	CE2	CE2-CZ	1.39065	
					O				CZ			
				H:TRP215	N	N-CA	1.48185					
					CA							
				C	C-O	1.22822						
				O								
				CB	CA-CB	1.55715						
				H:VAL32	CG1	CB-CG1	1.51508	H:THR76	N	N-CA	1.44632	
					CG2	CB-CG2	1.50644		CA			
				H:ASN34	OD1	CD-OD1	1.26025		OG1	CB-OG1	1.43909	
					ND2	CG-ND2	1.31645		CG2	CB-CG2	1.51245	
				H:ASP39	CB	CB-CG	1.52305	H:GLU80	CD	CG-CD	1.51131	
					CG				OE1	CD-OE1	1.25032	
					OD2	CG-OD2	1.23287		OE2	CD-OE2	1.24772	
				H:ALA40	CB	CA-CB	1.54078	H:LYS82	CE	CE-NZ	1.49946	
				H:VAL67	CG1	CB-CG1	1.51674		NZ			
					CG2	CB-CG2	1.52517					
III	90.078	45.053		H:GLU70	OE1	CD-OE1	1.2454					
					CA	CA-C	1.56548					
					O	C-O	1.22965					
					H:ILE73	CB	CB-CG2	1.51392				
						CG2						
						CD1	CG1-CD1	1.52142				
						CD	CG-CD	1.51932				
					H:GLU74	OE1	CD-OE1	1.25042				
						OE2	CD-OE2	1.25686				
						N	N-CA	1.46232				
		H:GLU75	C	C-O	1.2445							
			O									
V	55.432	25.515		H:ASP125	OD2	CG-OD2	1.25402	L:CYS88	C	C-O	1.21921	
				H:GLU127	CD	CD-CE	1.50033		O			

	OE1	CD-OE1	1.25442		CB	CA-CB	1.51823
	OE2	CD-OE2	1.25241		N	N-CA	1.46728
H:TYR128	CD2	CD2-CE2	1.39727	L:ASN89	CA		
	CE2				OD1	CG-OD1	1.2463
	CZ	CZ-OH	1.37223		L:ASN92	OD1	CG-OD1
OH	L:PHE98			CB	CA-CB	1.53733	
H:ILE129B	CD1	CG1-CD1	1.57649	L:CYS99	O	C-O	1.2262
				L:LYS100	CG	CG-CD	1.51948

Table S3. Extraction Set-Ups Percent Yield

Set-up	Solvent	Leaf Texture	Dry Sample Weight (g)	Extract Weight (g)	Percent Yield (%)
A	30% Ethanol	Powdered	50 g	15.83	31.66
B	95% Ethanol	Powdered	25 g	3.4	13.6
C	95% Ethanol	Shredded	200 g	78.541	39.27

Computation:

Extract Weight (g) = Flask w/ Crude Extract Weight (g) - Empty Flask Weight (g)

$$\text{Percent Yield (\%)} = \frac{\text{Extract Weight (g)}}{\text{Dry Sample Weight (g)}} \times 100$$

30% Ethanol | Powdered

ExtractWeight(g) = Flaskw/CrudeExtractWeight(g) – EmptyFlaskWeight(g)

$$= 403.61 \text{ g} - 387.78 \text{ g} = 15.83 \text{ g}$$

$$\text{Percent Yield (\%)} = \frac{\text{Extract Weight (g)}}{\text{Dry Sample Weight (g)}} \times 100$$

$$= \frac{15.83 \text{ g}}{50 \text{ g}} \times 100 = 31.66\%$$

95% Ethanol | Powdered

$$\begin{aligned} \text{ExtractWeight}(g) &= \text{Flaskw/CrudeExtractWeight}(g) - \text{EmptyFlaskWeight}(g) \\ &= 391.18 \text{ g} - 387.78 \text{ g} = 3.4 \text{ g} \end{aligned}$$

$$\begin{aligned} \text{Percent Yield (\%)} &= \frac{\text{Extract Weight (g)}}{\text{Dry Sample Weight (g)}} \times 100 \\ &= \frac{3.4 \text{ g}}{25 \text{ g}} \times 100 = 13.6\% \end{aligned}$$

95% Ethanol | Shredded

$$\begin{aligned} \text{ExtractWeight}(g) &= \text{Flaskw/CrudeExtractWeight}(g) - \text{EmptyFlaskWeight}(g) \\ &= 186.26 \text{ g} - 107.719 \text{ g} = 78.541 \text{ g} \end{aligned}$$

$$\begin{aligned} \text{Percent Yield (\%)} &= \frac{\text{Extract Weight (g)}}{\text{Dry Sample Weight (g)}} \times 100 \\ &= \frac{78.541 \text{ g}}{200 \text{ g}} \times 100 = 39.271\% \end{aligned}$$

Figure S2. GC-MS Mass Spectra and Chromatogram

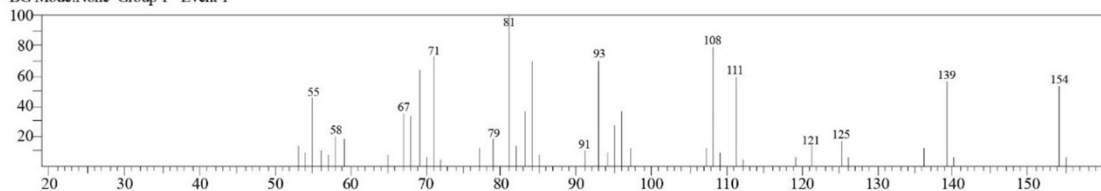
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:3.910(Scan#:92)

MassPeaks:41

RawMode:Single 3.910(92) BasePeak:81.10(10000)

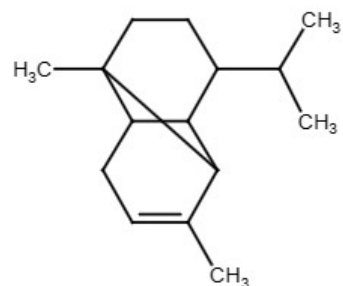
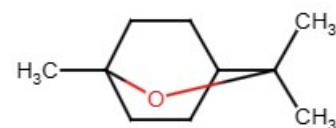
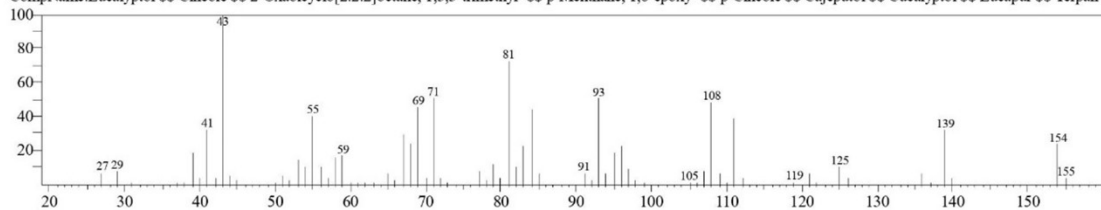
BG Mode:None Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:9937 Formula:C10H18O CAS:470-82-6 MolWeight:154

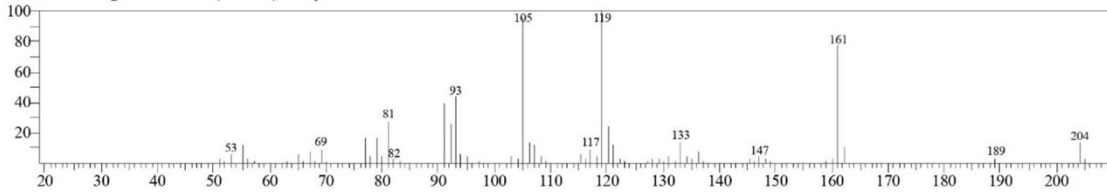
MassPeaks:72 BasePeak:43.00(10000)

CompName:Eucalyptol SS Cineole SS 2-Oxabicyclo[2.2.2]octane, 1,3,3-trimethyl- SS p-Menthane, 1,8-epoxy- SS p-Cineole SS Cajepulol SS Cucalyptol SS Eucapur SS Terpan S

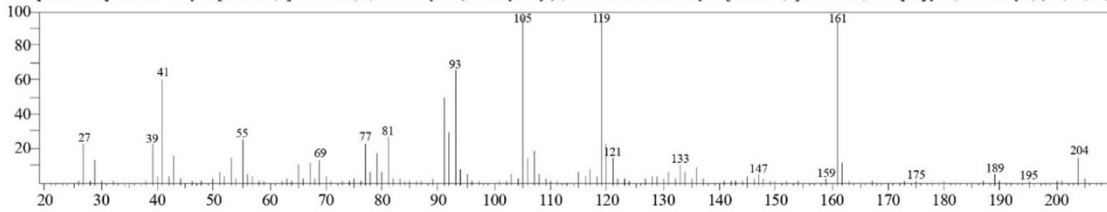


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:8.630(Scan#:564)
 MassPeaks:65
 RawMode:Single 8.630(564) BasePeak:119.15(10000)
 BG Mode:Averaged 8.520-8.740(553-575) Group 1 - Event 1

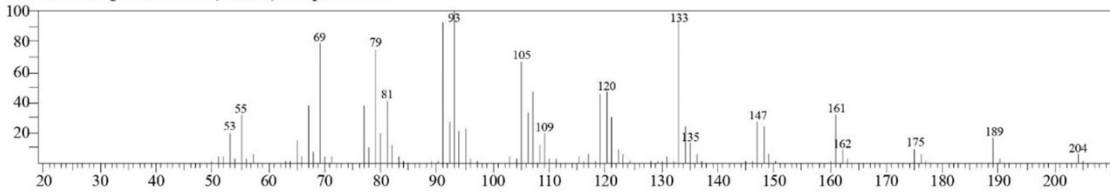


Spectrum2 #Library# NIST11s.lib Entry:18104 Formula:C15H24 CAS:3856-25-5 MolWeight:204
 MassPeaks:118 BasePeak:119.00(10000)
 CompName:Copaene SS Tricyclo[4.4.0.02,7]dec-3-ene, 1,3-dimethyl-8-(1-methylethyl)-, stereoisomer SS Tricyclo[4.4.0.02,7]dec-3-ene, 8-isopropyl-1,3-dimethyl-, (1R,2S,6S,7

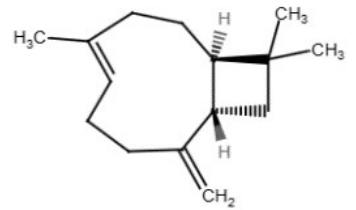
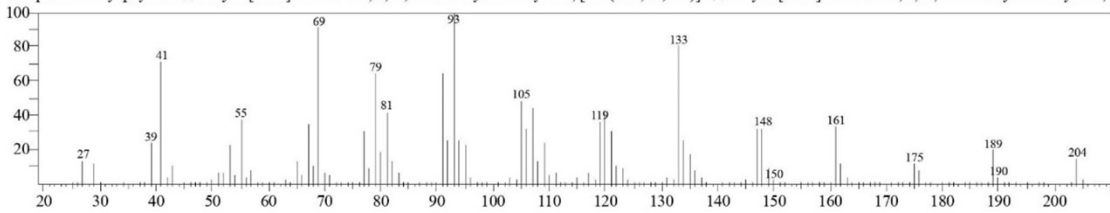


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:9.260(Scan#:627)
 MassPeaks:81
 RawMode:Single 9.260(627) BasePeak:93.10(10000)
 BG Mode:Averaged 9.180-9.360(619-637) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:18069 Formula:C15H24 CAS:87-44-5 MolWeight:204
 MassPeaks:176 BasePeak:93.00(10000)
 CompName:Caryophyllene SS Bicyclo[7.2.0]undec-4-ene, 4,11,11-trimethyl-8-methylene-, [1R-(1R*,4E,9S*)]- SS Bicyclo[7.2.0]undec-4-ene, 4,11,11-trimethyl-8-methylene-, (



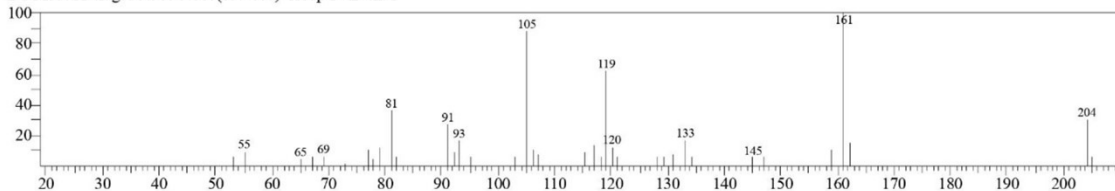
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:9.620(Scan#:663)

MassPeaks:37

RawMode:Single 9.620(663) BasePeak:161.15(10000)

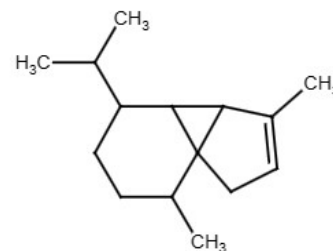
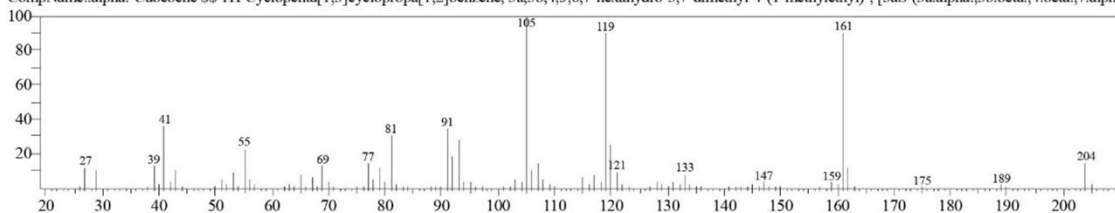
BG Mode:Averaged 9.560-9.660(657-667) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:18090 Formula:C15H24 CAS:17699-14-8 MolWeight:204

MassPeaks:99 BasePeak:105.00(10000)

CompName:.alpha.-Cubebene S5 1H-Cyclopenta[1,3]cyclopropa[1,2]benzene, 3a,3b,4,5,6,7-hexahydro-3,7-dimethyl-4-(1-methylethyl)-, [3aS-(3a.alpha.,3b.beta.,4.beta.,7.alpha.



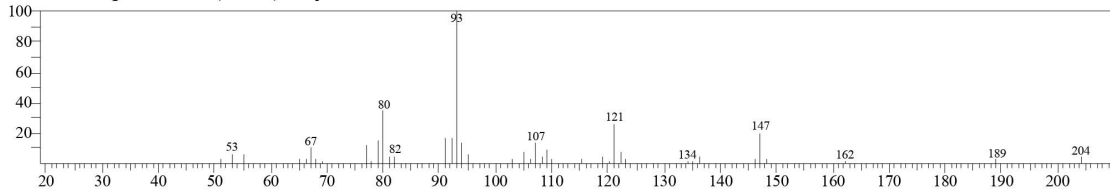
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:9.720(Scan#:673)

MassPeaks:42

RawMode:Single 9.720(673) BasePeak:93.10(10000)

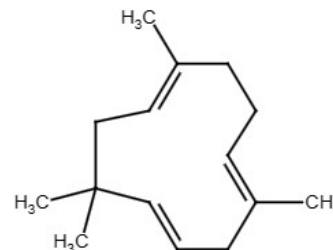
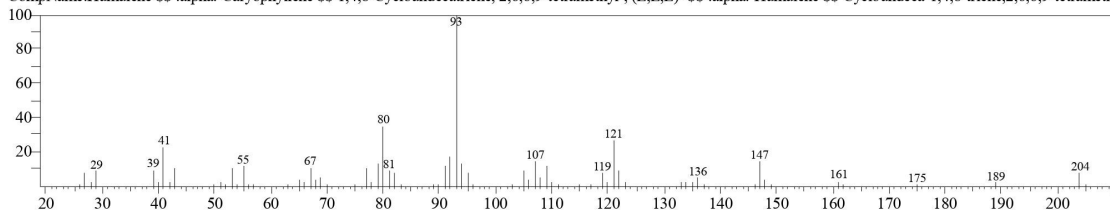
BG Mode:Averaged 9.680-9.800(669-681) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:18070 Formula:C15H24 CAS:6753-98-6 MolWeight:204

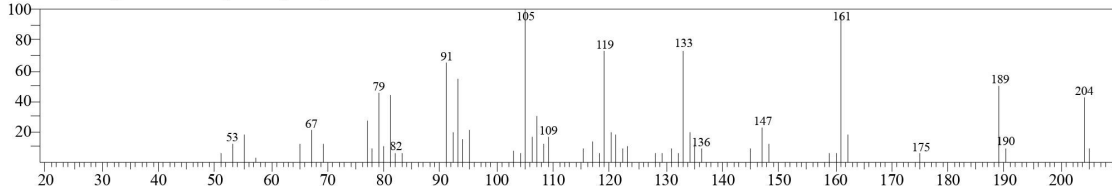
MassPeaks:70 BasePeak:93.00(10000)

CompName:Humulene S5 .alpha.-Caryophyllene S5 1,4,8-Cycloundecatriene, 2,6,6,9-tetramethyl-, (E,E,E)- S5 .alpha.-Humulene S5 Cycloundeca-1,4,8-triene,2,6,6,9-tetramethyl-

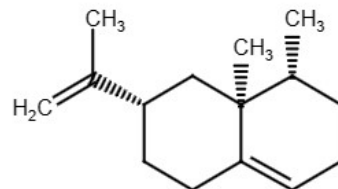
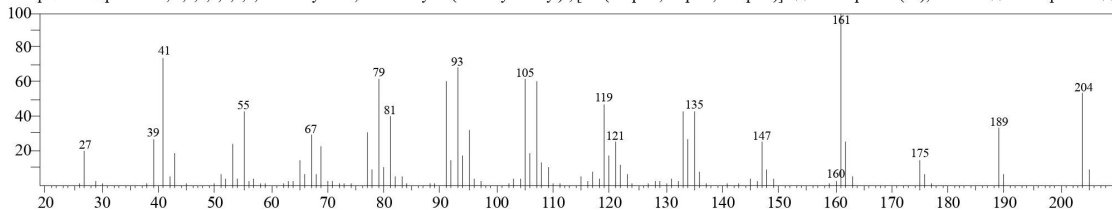


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:9.920(Scan#:693)
 MassPeaks:54
 RawMode:Single 9.920(693) BasePeak:105.10(10000)
 BG Mode:Averaged 9.880-10.000(689-701) Group 1 - Event 1

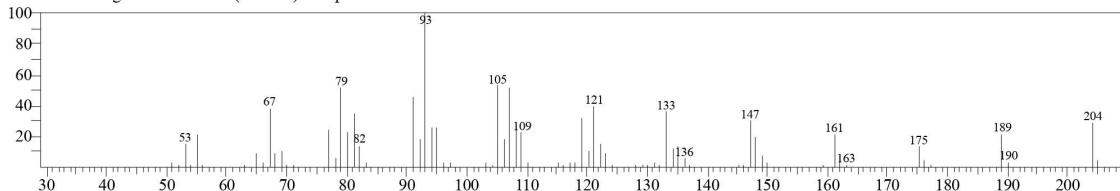


Spectrum2 #Library# NIST11s.lib Entry:18118 Formula:C15H24 CAS:10219-75-7 MolWeight:204
 MassPeaks:100 BasePeak:161.00(10000)
 CompName:Naphthalene, 1,2,3,5,6,7,8,8a-octahydro-1,8a-dimethyl-7-(1-methylethenyl)-, [1S-(1.alpha.,7.alpha.,8a.alpha.)]- \$\$ Eremophila-1(10),11-diene \$\$ Eremophilene \$\$:

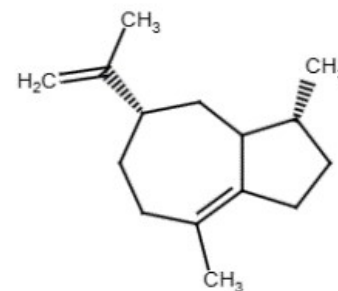
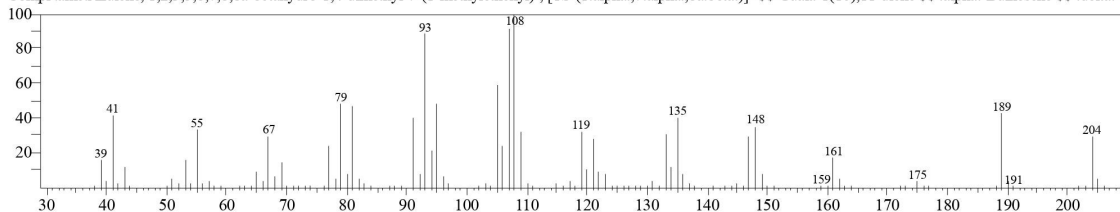


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.170(Scan#:718)
 MassPeaks:75
 RawMode:Single 10.170(718) BasePeak:93.10(10000)
 BG Mode:Averaged 10.120-10.210(713-722) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:18097 Formula:C15H24 CAS:3691-11-0 MolWeight:204
 MassPeaks:117 BasePeak:108.00(10000)
 CompName:Azulene, 1,2,3,5,6,7,8,8a-octahydro-1,4-dimethyl-7-(1-methylethenyl)-, [1S-(1.alpha.,7.alpha.,8a.beta.)]- \$\$ Guaia-1(10),11-diene \$\$.alpha.-Bulnesene \$\$.delta.-G



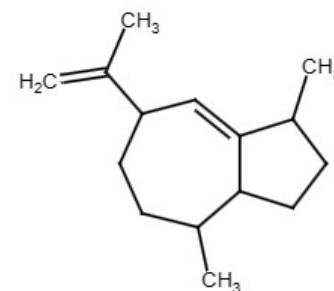
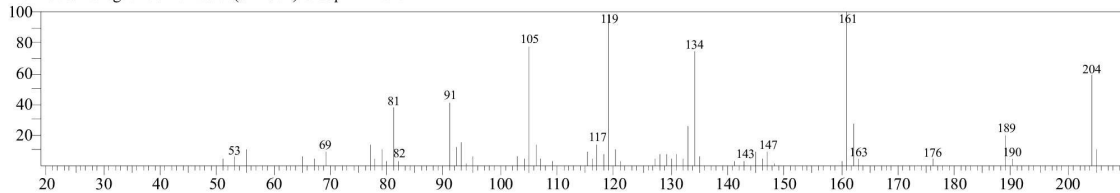
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.470(Scan#:748)

MassPeaks:54

RawMode:Single 10.470(748) BasePeak:161.15(10000)

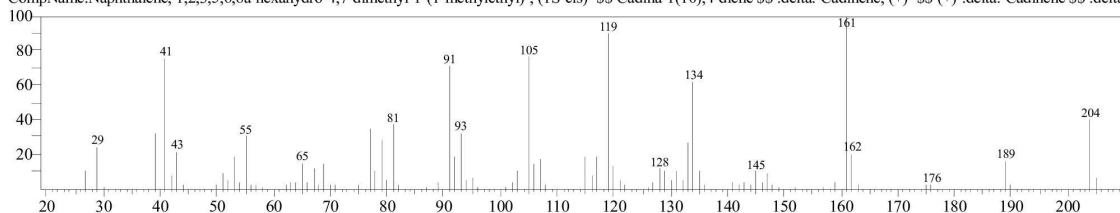
BG Mode:Averaged 10.410-10.500(742-751) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:18135 Formula:C15H24 CAS:483-76-1 MolWeight:204

MassPeaks:88 BasePeak:161.00(10000)

CompName:Naphthalene, 1,2,3,5,6,8a-hexahydro-4,7-dimethyl-1-(1-methylethyl)-, (1S-cis)- SS Cadina-1(10),4-diene SS .delta.-Cadinene, (+)- SS (+)-.delta.-Cadinene SS .delta.-



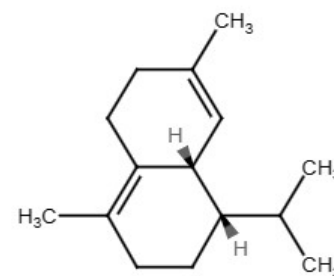
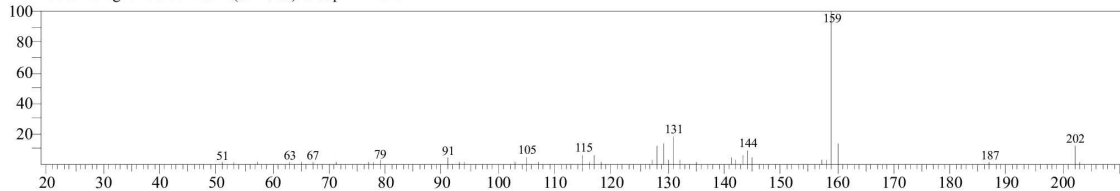
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.540(Scan#:755)

MassPeaks:48

RawMode:Single 10.540(755) BasePeak:159.15(10000)

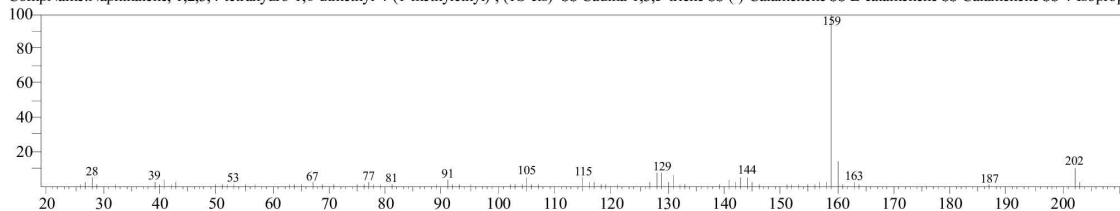
BG Mode:Averaged 10.500-10.580(751-759) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:17784 Formula:C15H22 CAS:483-77-2 MolWeight:202

MassPeaks:74 BasePeak:159.00(10000)

CompName:Naphthalene, 1,2,3,4-tetrahydro-1,6-dimethyl-4-(1-methylethyl)-, (1S-cis)- SS Cadina-1,3,5-triene SS (-)-Calamenene SS L-calamenene SS Calamenene SS 4-Isoprop



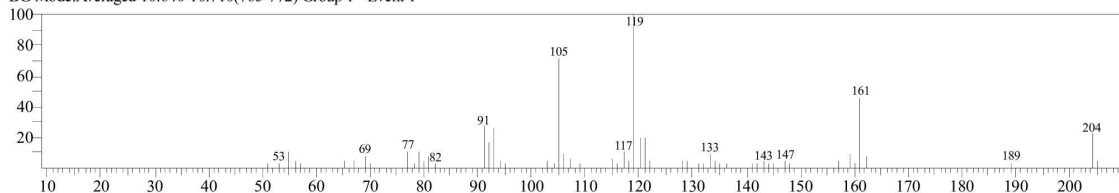
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.680(Scan#:769)

MassPeaks:57

RawMode:Single 10.680(769) BasePeak:119.10(10000)

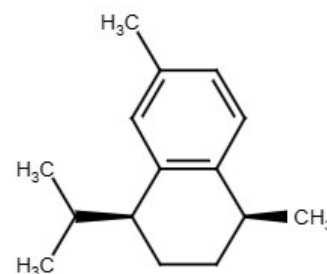
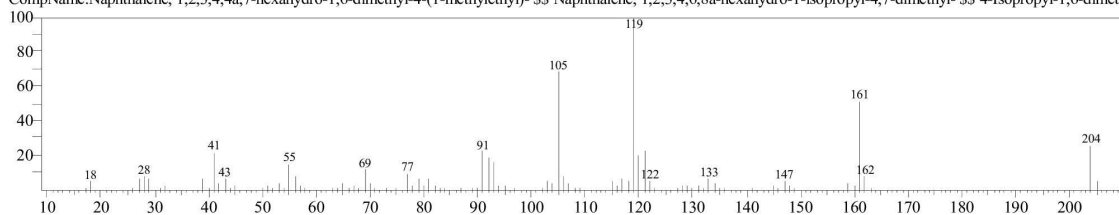
BG Mode:Averaged 10.640-10.710(765-772) Group 1 - Event 1



Spectrum2 #Library# NIST1s.lib Entry:18105 Formula:C15H24 CAS:16728-99-7 MolWeight:204

MassPeaks:92 BasePeak:119.00(10000)

CompName:Naphthalene, 1,2,3,4,4a,7-hexahydro-1,6-dimethyl-4-(1-methylethyl)- SS Naphthalene, 1,2,3,4,6,8a-hexahydro-1-isopropyl-4,7-dimethyl- SS 4-Isopropyl-1,6-dimethyl-



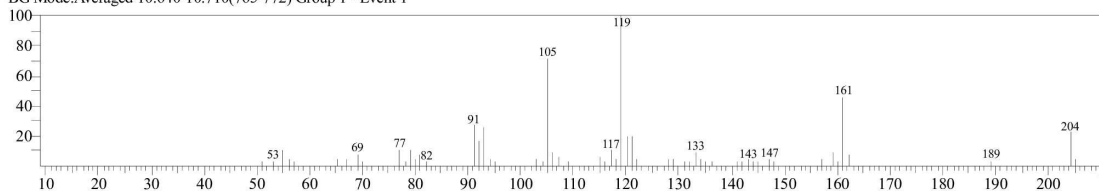
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.680(Scan#:769)

MassPeaks:57

RawMode:Single 10.680(769) BasePeak:119.10(10000)

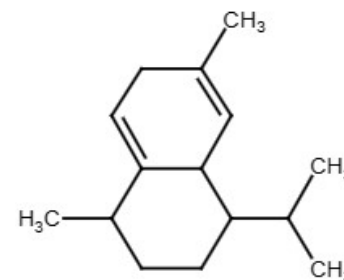
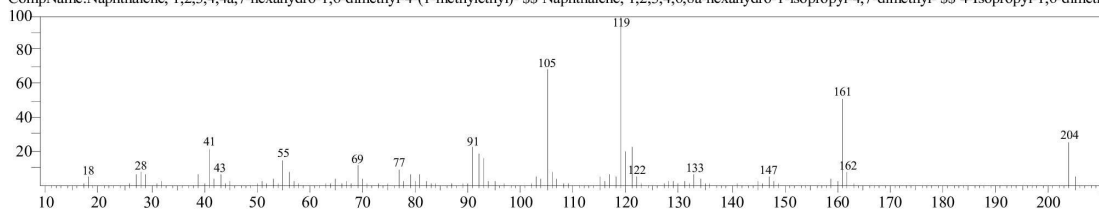
BG Mode:Averaged 10.640-10.710(765-772) Group 1 - Event 1



Spectrum2 #Library# NIST1s.lib Entry:18105 Formula:C15H24 CAS:16728-99-7 MolWeight:204

MassPeaks:92 BasePeak:119.00(10000)

CompName:Naphthalene, 1,2,3,4,4a,7-hexahydro-1,6-dimethyl-4-(1-methylethyl)- SS Naphthalene, 1,2,3,4,6,8a-hexahydro-1-isopropyl-4,7-dimethyl- SS 4-Isopropyl-1,6-dimethyl-



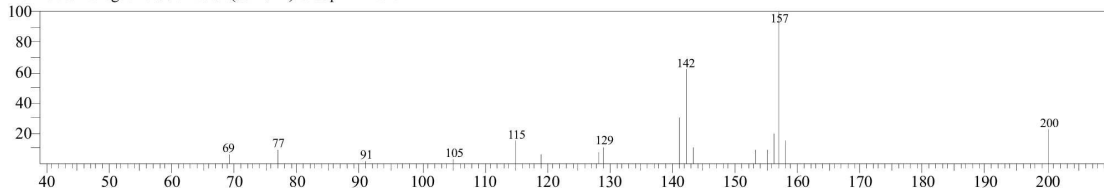
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:10.800(Scan#:781)

MassPeaks:17

RawMode:Single 10.800(781) BasePeak:157.10(10000)

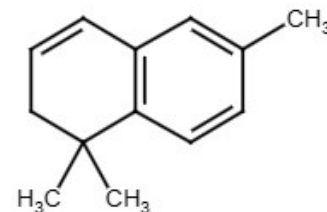
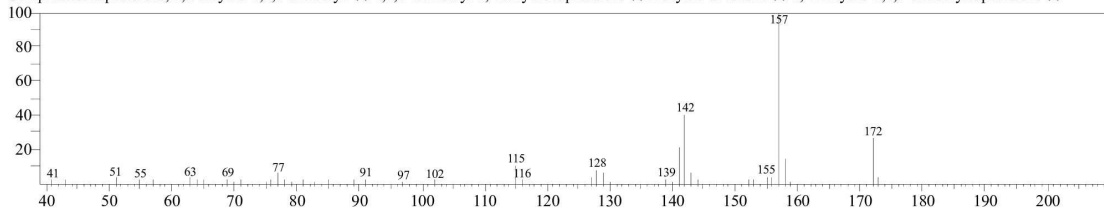
BG Mode:Averaged 10.770-10.830(778-784) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:13177 Formula:C13H16 CAS:30364-38-6 MolWeight:172

MassPeaks:44 BasePeak:157.00(10000)

CompName:Naphthalene, 1,2-dihydro-1,1,6-trimethyl- SS 1,1,6-Trimethyl-1,2-dihydronaphthalene SS Dehydro-ar-ionene SS 1,2-Dihydro-1,1,6-trimethylnaphthalene SS



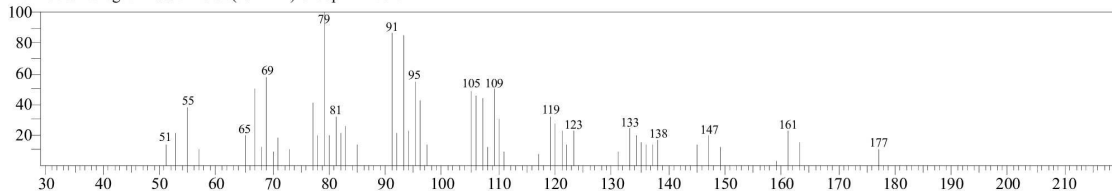
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:11.390(Scan#:840)

MassPeaks:53

RawMode:Single 11.390(840) BasePeak:79.05(10000)

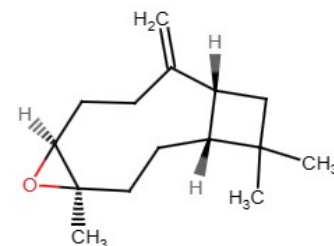
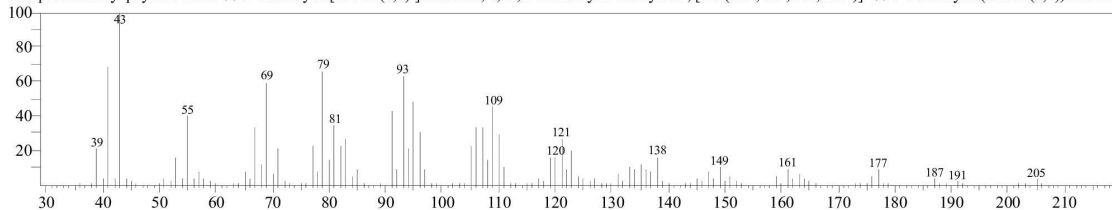
BG Mode:Averaged 11.330-11.430(834-844) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:20008 Formula:C15H24O CAS:1139-30-6 MolWeight:220

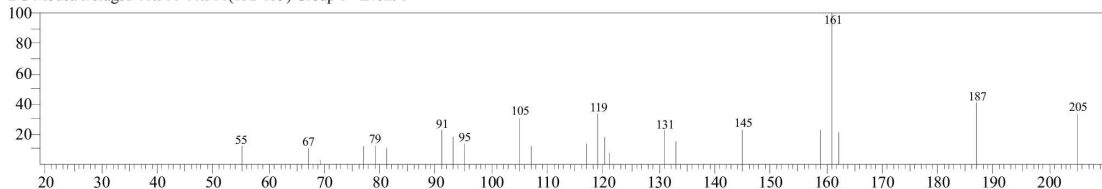
MassPeaks:130 BasePeak:43.00(10000)

CompName:Caryophyllene oxide SS 5-Oxatricyclo[8.2.0.0(4,6)]dodecane, 4,12,12-trimethyl-9-methylene-, [1R-(1R*,4R*,6R*,10S*)]- SS 5-Oxatricyclo(8.2.0.0(4,6))dodecane.

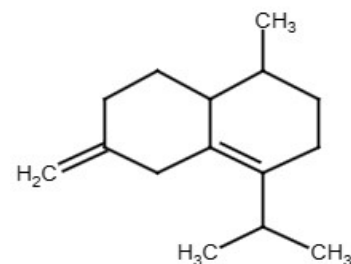
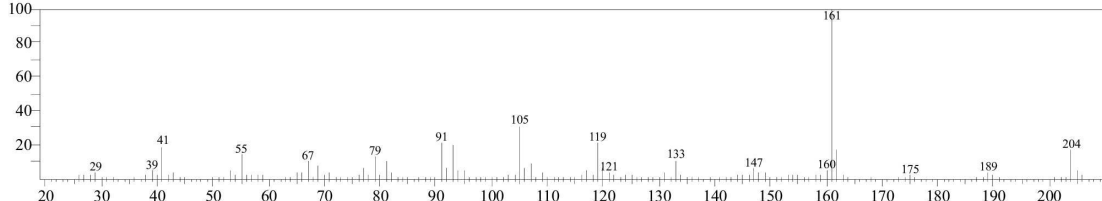


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:11.540(Scan#:855)
 MassPeaks:23
 RawMode:Single 11.540(855) BasePeak:161.15(10000)
 BG Mode:Averaged 11.510-11.580(852-859) Group 1 - Event 1

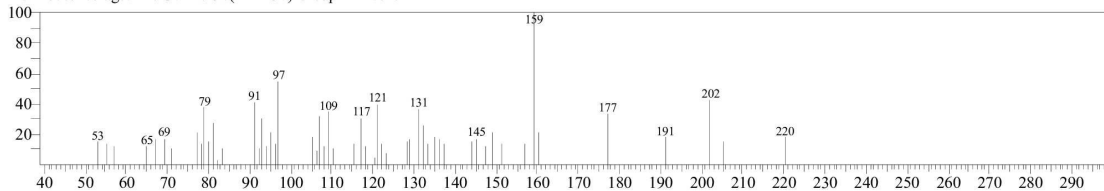


Spectrum2 #Library# NIST11s.lib Entry:18129 Formula:C15H24 CAS:150320-52-8 MolWeight:204
 MassPeaks:141 BasePeak:161.00(10000)
 CompName:Bicyclo[4.4.0]dec-1-ene, 2-isopropyl-5-methyl-9-methylene- SS 8-Isopropyl-5-methyl-2-methylene-1,2,3,4,4a,5,6,7-octahydronaphthalene # SS 2-Isopropyl-5-methyl-

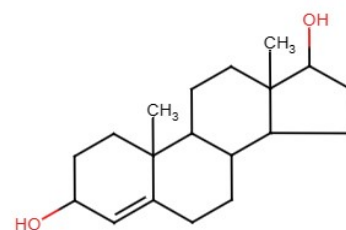
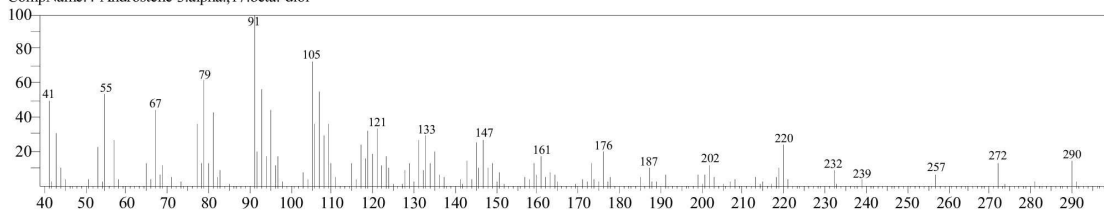


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:11.860(Scan#:887)
 MassPeaks:55
 RawMode:Single 11.860(887) BasePeak:159.15(10000)
 BG Mode:Averaged 11.830-11.890(884-890) Group 1 - Event 1

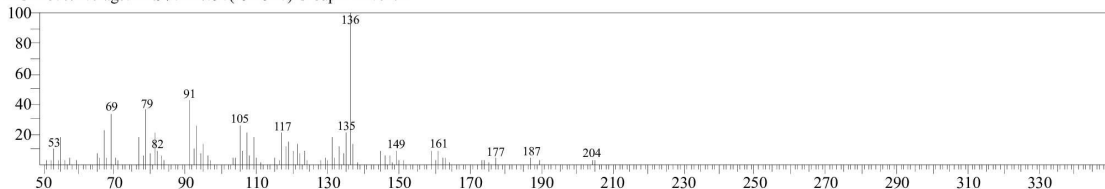


Spectrum2 #Library# SWGDRUG MS LIBRARY VERSION 3.10.lib Entry:2918 Formula:C19H30O2 CAS:1852-61-5 MolWeight:290
 MassPeaks:125 BasePeak:91.00(10000)
 CompName:4-Androstene-3.alpha.,17.beta.-diol

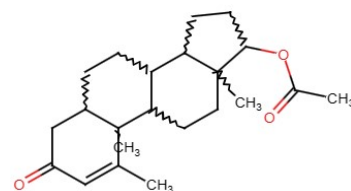
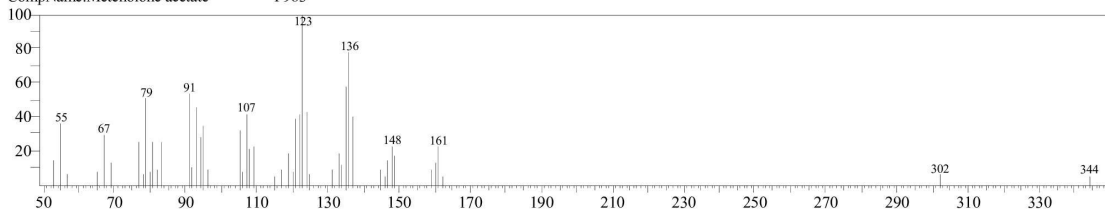


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:12.040(Scan#:905)
 MassPeaks:81
 RawMode:Single 12.040(905) BasePeak:136.15(10000)
 BG Mode:Averaged 11.970-12.090(898-910) Group 1 - Event 1

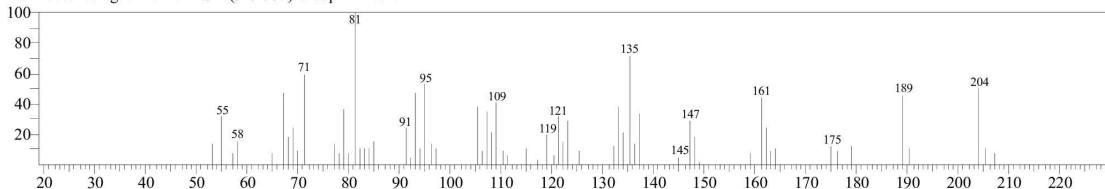


Spectrum2 #Library# MPW2011.lib Entry:5616 Formula:C22H32O3 CAS:0-00-0 MolWeight:344
 MassPeaks:50 BasePeak:123.00(10000)
 CompName:Metenolone acetate P983

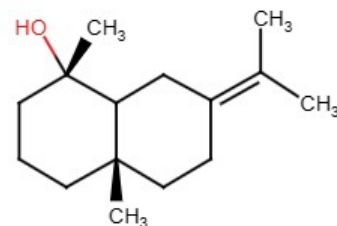
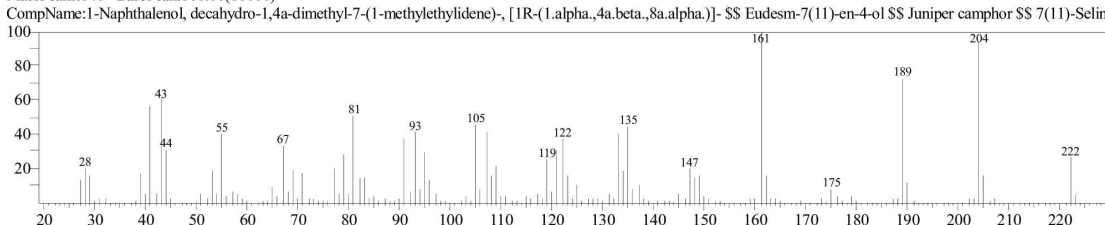


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:12.320(Scan#:933)
 MassPeaks:64
 RawMode:Single 12.320(933) BasePeak:81.10(10000)
 BG Mode:Averaged 12.280-12.340(929-935) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:20299 Formula:C15H26O CAS:473-04-1 MolWeight:222
 MassPeaks:140 BasePeak:161.00(10000)
 CompName:1-Naphthalenol, decahydro-1,4a-dimethyl-7-(1-methylethylidene)-, [1R-(1.alpha.,4a.beta.,8a.alpha.)]-SS Eudesm-7(11)-en-4-ol SS Juniper camphor SS 7(11)-Seline



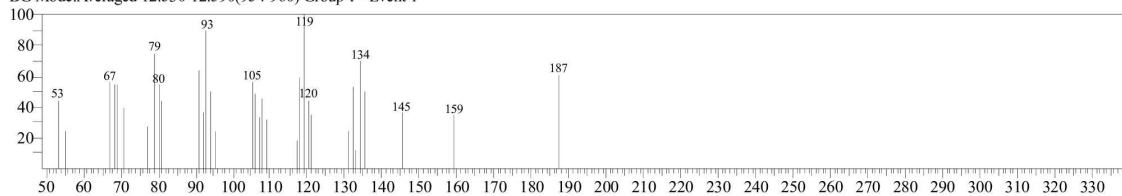
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:12.560(Scan#:957)

MassPeaks:33

RawMode:Single 12.560(957) BasePeak:119.10(10000)

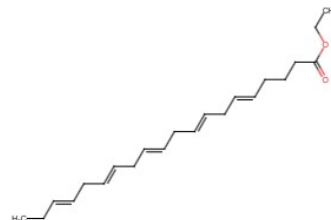
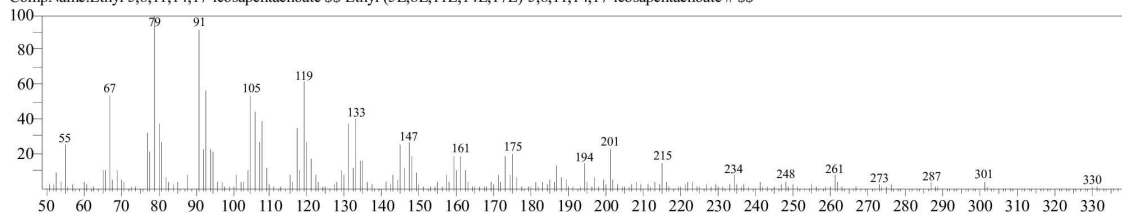
BG Mode:Averaged 12.530-12.590(954-960) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:27690 Formula:C22H34O2 CAS:84494-70-2 MolWeight:330

MassPeaks:192 BasePeak:79.00(10000)

CompName:Ethyl 5,8,11,14,17-icosapentaenoate \$\$ Ethyl (5E,8E,11E,14E,17E)-5,8,11,14,17-icosapentaenoate # \$\$



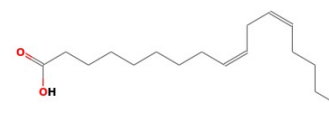
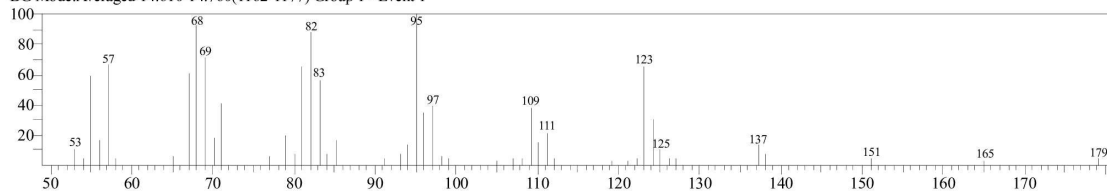
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:14.680(Scan#:1169)

MassPeaks:48

RawMode:Single 14.680(1169) BasePeak:95.10(10000)

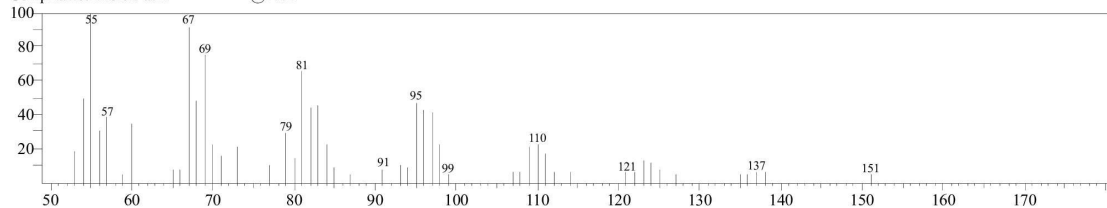
BG Mode:Averaged 14.610-14.760(1162-1177) Group 1 - Event 1



Spectrum2 #Library# MPW2011.lib Entry:3284 Formula:C18H32O2 CAS:60-33-3 MolWeight:280

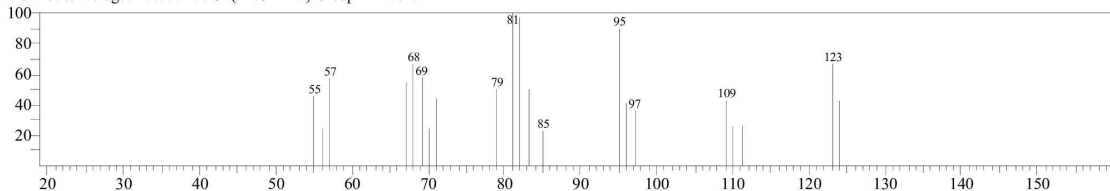
MassPeaks:50 BasePeak:55.00(10000)

CompName:Linoleic acid @P651

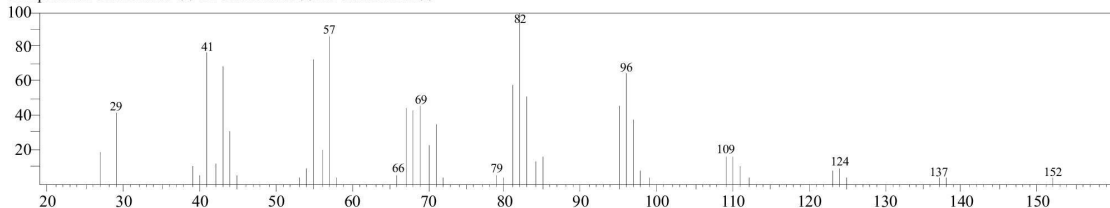


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:15.110(Scan#:1212)
 MassPeaks:21
 RawMode:Single 15.110(1212) BasePeak:81.10(10000)
 BG Mode:Averaged 15.060-15.190(1207-1220) Group 1 - Event 1

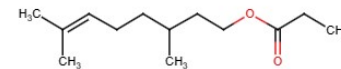
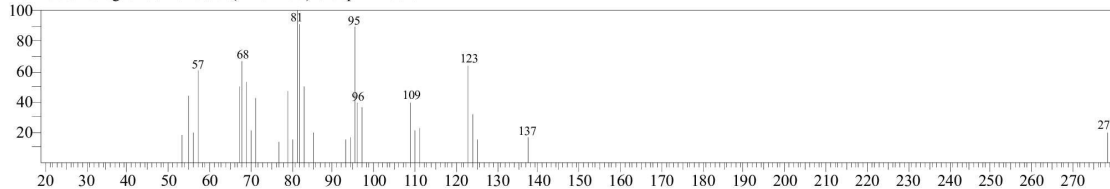


Spectrum2 #Library# NIST11s.lib Entry:20737 Formula:C15H30O CAS:2765-11-9 MolWeight:226
 MassPeaks:44 BasePeak:82.00(10000)
 CompName:1-Pentadecanal

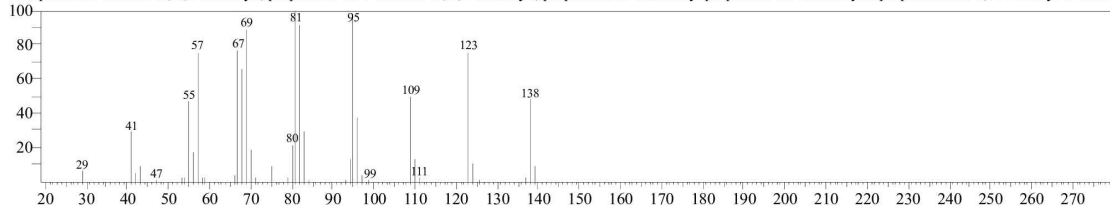


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:15.490(Scan#:1250)
 MassPeaks:29
 RawMode:Single 15.490(1250) BasePeak:81.10(10000)
 BG Mode:Averaged 15.440-15.550(1245-1256) Group 1 - Event 1

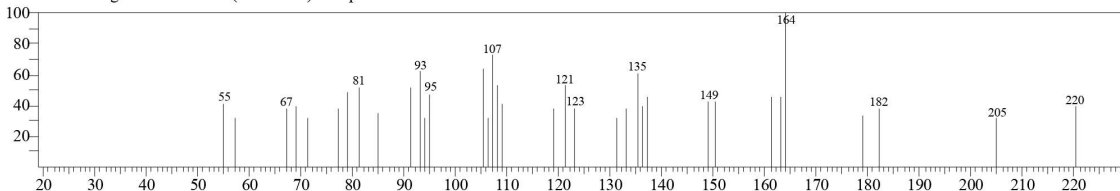


Spectrum2 #Library# NIST11s.lib Entry:19101 Formula:C13H24O2 CAS:141-14-0 MolWeight:212
 MassPeaks:40 BasePeak:81.00(10000)
 CompName:6-Octen-1-ol, 3,7-dimethyl-, propionate

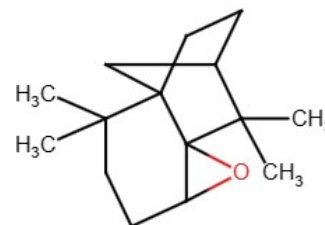
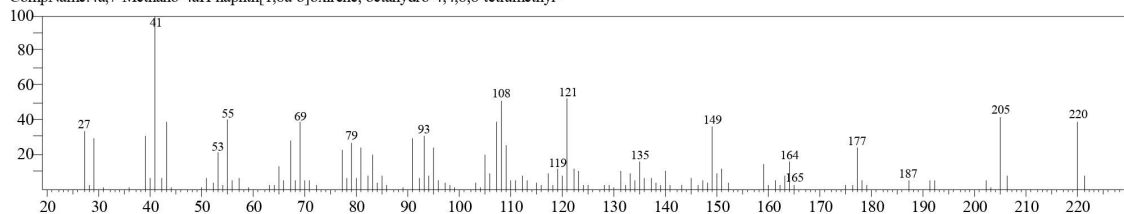


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:16.000(Scan#:1301)
MassPeaks:35
RawMode:Single 16.000(1301) BasePeak:164.10(10000)
BG Mode:Averaged 15.910-16.050(1292-1306) Group 1 - Event 1

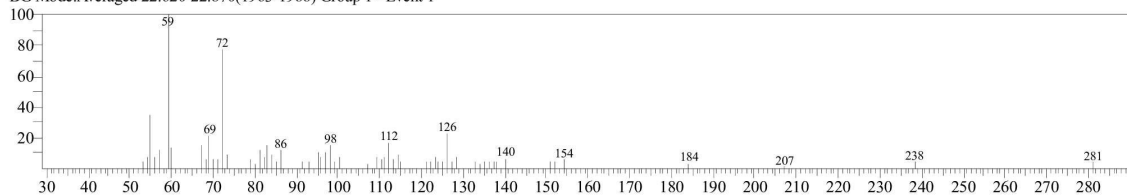


Spectrum2 #Library# NIST11s.lib Entry:20007 Formula:C15H24O CAS:67999-56-8 MolWeight:220
MassPeaks:116 BasePeak:41.00(10000)
CompName:4a,7-Methano-4aH-naphth[1,8a-b]oxirene, octahydro-4,4,8,8-tetramethyl-

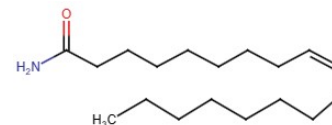
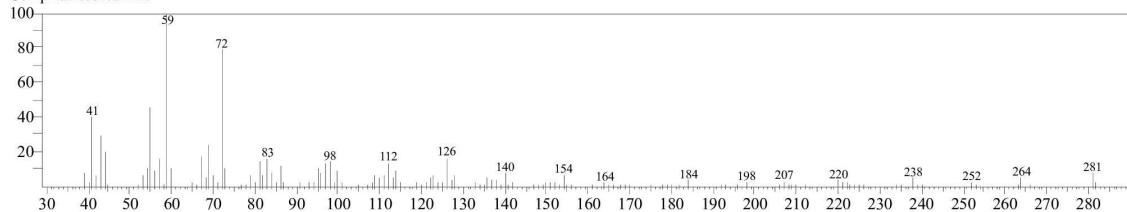


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:22.730(Scan#:1974)
MassPeaks:60
RawMode:Single 22.730(1974) BasePeak:59.05(10000)
BG Mode:Averaged 22.620-22.870(1963-1988) Group 1 - Event 1

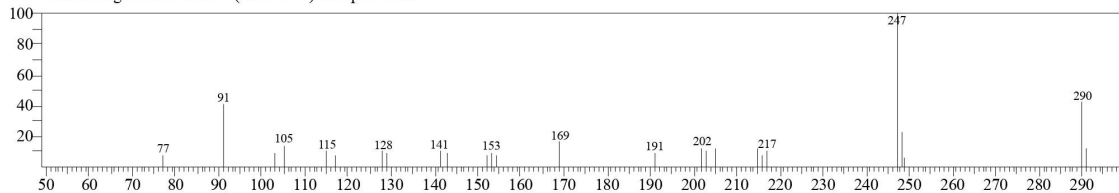


Spectrum2 #Library# SWGDRUG MS LIBRARY VERSION 3.10.lib Entry:424 Formula:C18H35NO CAS:301-02-0 MolWeight:281
MassPeaks:128 BasePeak:59.00(10000)
CompName:Oleamide

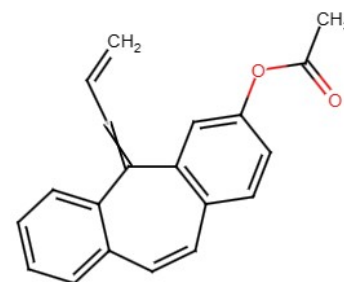
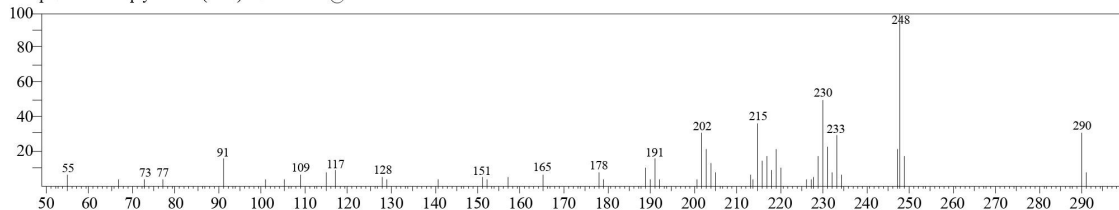


Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:26.350(Scan#:2336)
MassPeaks:27
RawMode:Single 26.350(2336) BasePeak:247.15(10000)
BG Mode:Averaged 26.220-26.430(2323-2344) Group 1 - Event 1

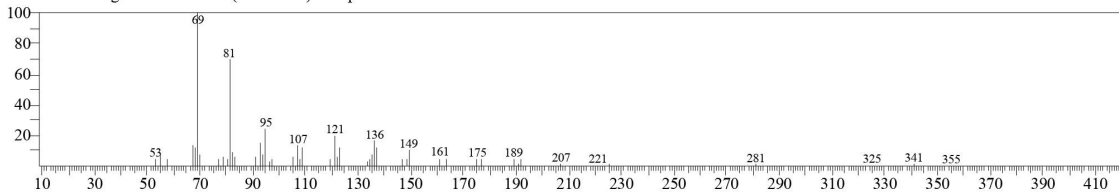


Spectrum2 #Library# MPW2011.lib Entry:3649 Formula:C20H18O2 CAS:0-00-0 MolWeight:290
MassPeaks:50 BasePeak:248.00(10000)
CompName:Amitriptyline-M -(CH3)2NOH AC @P699



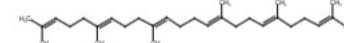
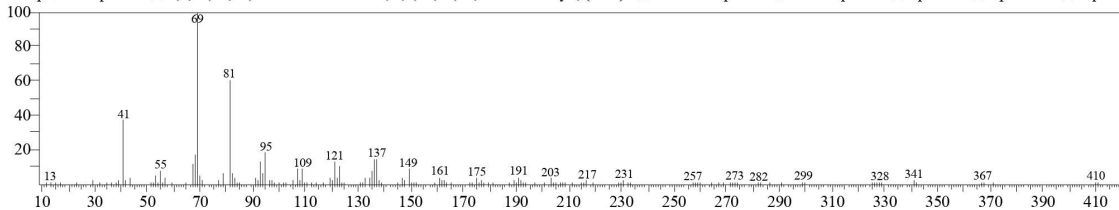
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:30.350(Scan#:2736)
MassPeaks:58
RawMode:Single 30.350(2736) BasePeak:69.10(10000)
BG Mode:Averaged 30.280-30.440(2729-2745) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:29859 Formula:C30H50 CAS:111-02-4 MolWeight:410
MassPeaks:146 BasePeak:69.00(10000)

CompName:Squalene \$\$ 2,6,10,14,18,22-Tetracosahexaene, 2,6,10,15,19,23-hexamethyl-, (all-E)- \$\$ all-trans-Squalene \$\$ trans-Squalene \$\$ Spinacen \$\$ Spinacene \$\$ Squalen



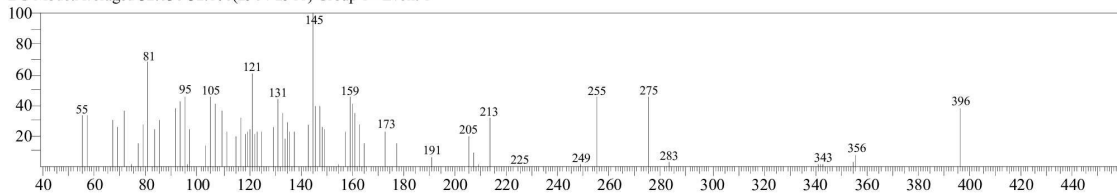
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:32.110(Scan#:2912)

MassPeaks:72

RawMode:Single 32.110(2912) BasePeak:145.15(10000)

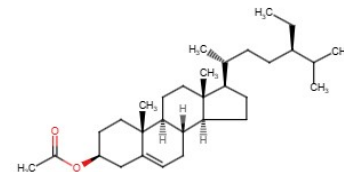
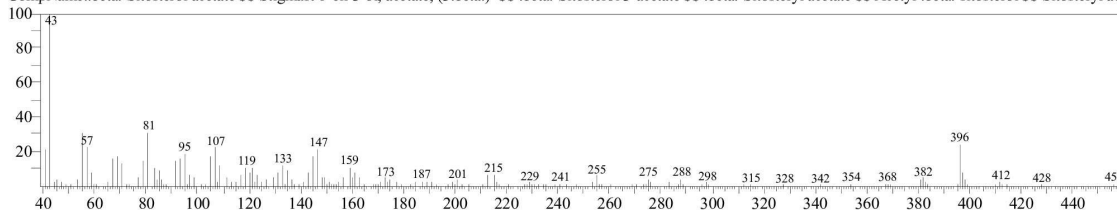
BG Mode:Averaged 32.030-32.170(2904-2918) Group 1 - Event 1



Spectrum2 #Library# NIST11s.lib Entry:30335 Formula:C31H52O2 CAS:915-05-9 MolWeight:456

MassPeaks:157 BasePeak:43.00(10000)

CompName:.beta.-Sitosterol acetate SS Stigmast-5-en-3-ol, acetate, (3.beta.)- SS .beta.-Sitosterol 3-acetate SS .beta.-Sitosteryl acetate SS Acetyl-.beta.-sitosterol SS Sitosteryl ace



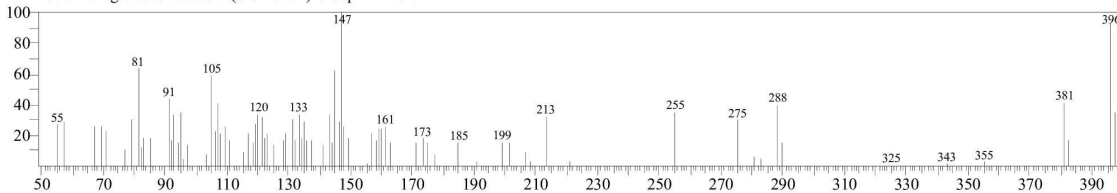
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:33.810(Scan#:3082)

MassPeaks:92

RawMode:Single 33.810(3082) BasePeak:147.15(10000)

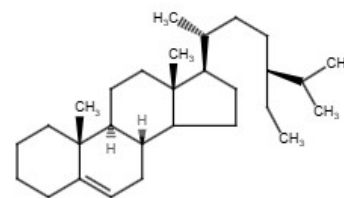
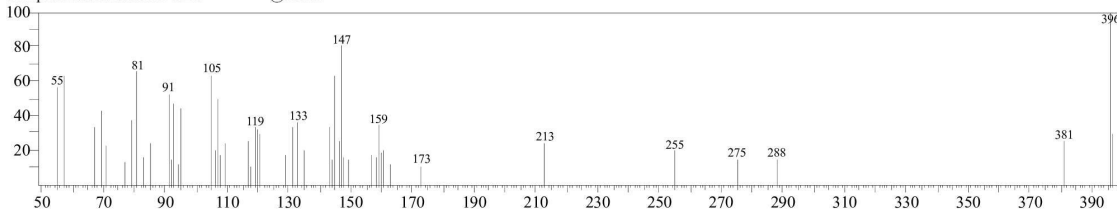
BG Mode:Averaged 33.700-33.900(3071-3091) Group 1 - Event 1



Spectrum2 #Library# MPW2011.lib Entry:7013 Formula:C29H48 CAS:0-00-0 MolWeight:396

MassPeaks:50 BasePeak:396.00(10000)

CompName:Clonasterol -H2O @P1182



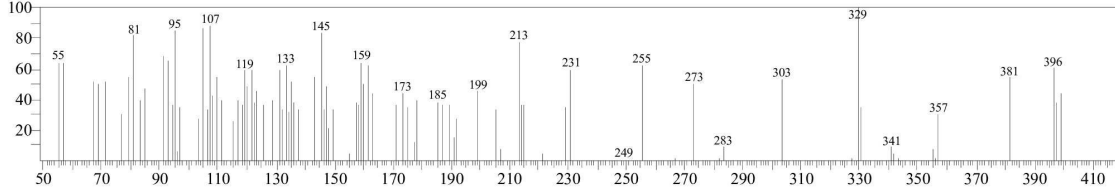
Spectrum Comparison

Spectrum1 #Data# UNK_2232024_3.qgd R.Time:37.420(Scan#:3443)

MassPeaks:95

RawMode:Single 37.420(3443) BasePeak:329.30(10000)

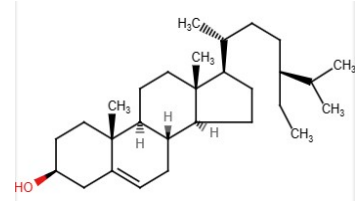
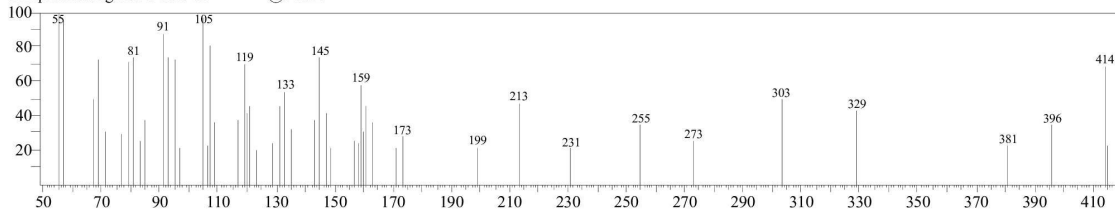
BG Mode:Averaged 37.330-37.510(3434-3452) Group 1 - Event 1



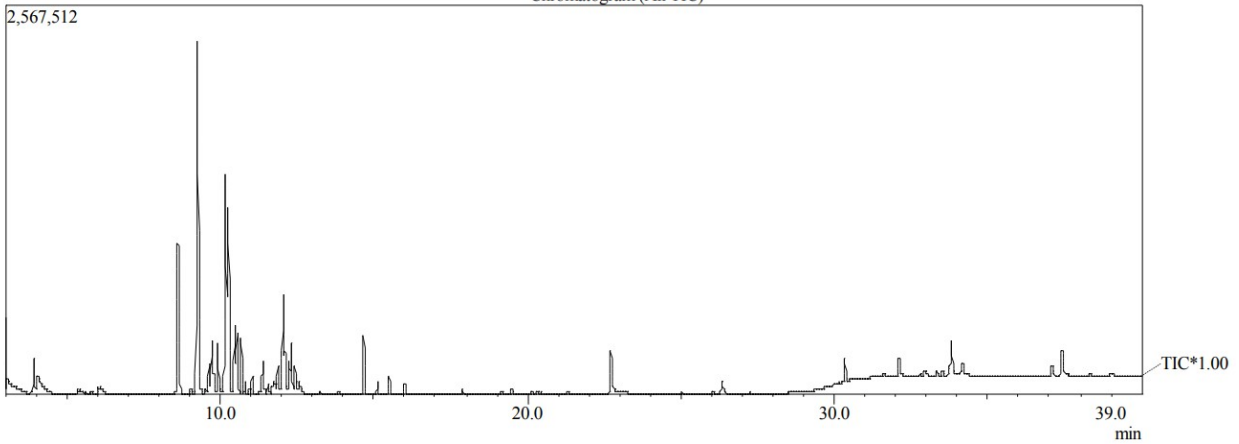
Spectrum2 #Library# MPW2011.lib Entry:7358 Formula:C29H50O CAS:83-47-6 MolWeight:414

MassPeaks:50 BasePeak:55.00(10000)

CompName:Stigmast-5-en-3-ol @P1232



Chromatogram (All TIC)



Chromatogram (Zoom)

2,598,965

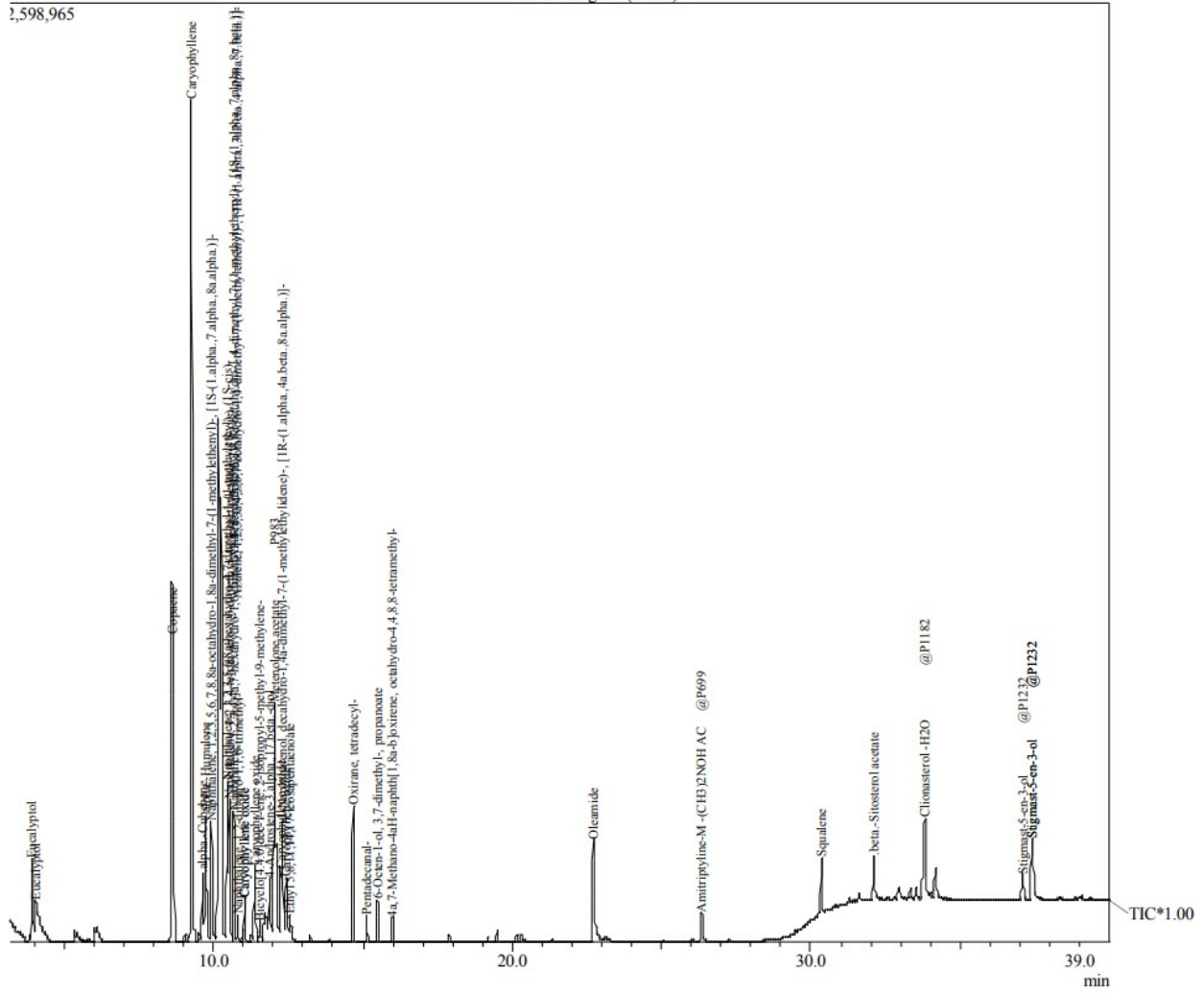

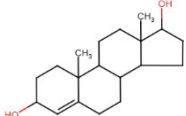




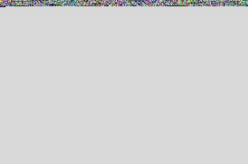

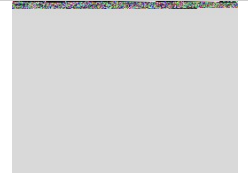
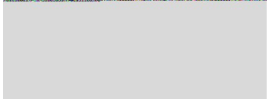

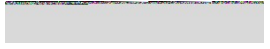
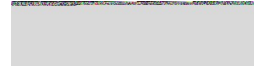


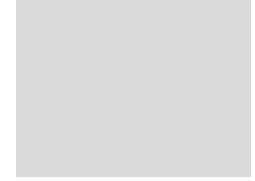
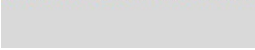
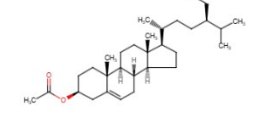
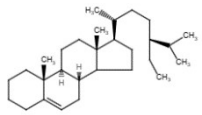
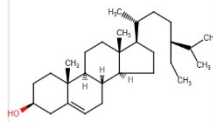


Table S4. Identified Bioactive Compounds Structures and Description

CID	Name	Structure	Type of Phytochemical	SMILES	CID	Name	Structure	Type of Phytochemical	SMILES
1	Eucalyptol		Terpenoid	<chem>CC12CCC(CC1)C(C)(C)O2</chem>	15	4-Androstene-3. α .,17. β .-diol		Steroid	<chem>CC12CCC3C(CCC4=CC(O)CCC34C)C1CCC2O</chem>
2	α -Copaene		Terpenoid	<chem>CC(C)C1CCC2(C)C3CC=C(C)C2C13</chem>	16	Methenolone acetate		Steroid	<chem>CC(=O)OC1CCC2C3CCC4CC(=O)C=C(C)C4(C)C3CCC12C</chem>
3	Caryophyllene		Terpenoid	<chem>[H][C@]12CC(C)(C)[C@]1([H])CC\C(C)=C\CCC2=C</chem>	17	Juniper camphor		Terpenoid	<chem>CC(C)=C1CC[C@]2(C)CCC[C@](C)(O)C2C1</chem>
4	α -Cubebene		Terpenoid	<chem>CC(C)C1CCC(C)C23CC=C(C)C2C13</chem>	18	Ethyl 5,8,11,14,17-icosapentaenoate		Fatty acid	<chem>CCOC(=O)CCC\C=C\C\C=C\C\C=C\C\C=C\C\C=C\C\C=C\C</chem>
5	Humulene		Terpenoid	<chem>C\C1=C/CC(C)(C)\C=C\C\C(C)=C\CC1</chem>	19	Linoleic Acid		Fatty Acid	<chem>CCCCCCCCCCCCCCC1CO1</chem>
6	Valencene		Terpenoid	<chem>C[C@@H]1CCC=C2CC[C@@H](C[C@@]1]2C)C(C)=C</chem>	20	1-Pentadecanal		Fatty Aldehyde	<chem>CCCCCCCCCCCCCCC=O</chem>

7	α -Bulnesene	Terpenoid	<chem>C[C@@H]1CCC2=C(C)CC[C@@H](CC12)C(C)=C</chem>	21	6-Octen-1-ol, 3,7-dimethyl-, propanoate		Terpenoid/terpene alcohol	<chem>CCC(=O)OCCC(C)CCC=C(C)C</chem>
8	γ -Gurjunene	Terpenoid	<chem>CC1CCC2C(C)CCC(C=C12)C(C)=C</chem>	22	4a,7-Methano-4aH-naphth[1,8a-b]oxirene, octahydro-4,4,8,8-tetramethyl-		Terpenoid	<chem>CC1(CCC2C3(C14)CCC(C4)C3(C)O2)C</chem>
9	δ -Cadinene, (+)-	Terpenoid	<chem>[H][C@]1(CCC(C)=C2CCC(C)=C[C@]12[H])C(C)C</chem>	23	Oleamide		Fatty Acid amide	<chem>CCCCCCCC=CCCCCCCC(=O)N</chem>
10	(-)-Calamenene	Terpenoid	<chem>CC(C)[C@@H]1CC[C@H](C)C2=C1C=C(C)C=C2</chem>	24	Amitriptyline-M-(CH ₃) ₂ NOH AC		Amine	<chem>C=1C2CCCCC2\C(=C\C=C)C2CC(CC2C1)OC(C)=O</chem>
11	Cadinadiene-1,4	Terpenoid	<chem>CC(C)C1CCC(C)C2=CCC(C)=CC12</chem>	25	Squalene		Terpenoid	<chem>CC(=CCCC(=CCC(C)=CCCC=C(C)CC(C)=C(C)CCC=C(C)C)C)C</chem>
12	Naphthalene, 1,2-dihydro-1,1,6-trimethyl-	Benzenoids, Aromatic	<chem>CC1=CC2=C(C=C1)C(C)(C)CC=C2</chem>	26	β -Sitosterol acetate		Terpenoid	<chem>CCC(CCC(C)C1CC2C1(CCC3C2CC=C4C3(CCC(C4)OC(=O)C)C)C(C)C</chem>

13	Caryophyllene oxide		Terpenoid	<chem>[H][C@]12CCC(=C)[C@]3([H])CC(C)(C)[C@]3([H])CC[C@@]1(C)O2</chem>	27	Clionasterol - H2O		Phytosterol	<chem>[H][C@@]12CC=C3CCCC[C@]3(C)[C@@]1([H])CC[C@]1(C)[C@H](CCC21)[C@H](C)CC[C@@H](CC)C(C)C</chem>
14	Bicyclo[4.4.0]dec-1-ene, 2-isopropyl-5-methyl-9-methylene-		Terpenoid	<chem>CC(C)C1=C2CC(=C)CCC2C(C)CC1</chem>	28	Stigmast-5-en-3-ol		Sterol lipid	<chem>CCC(CCC(C)C1CC2C1(CCC3C2CC=C4C3(CCC(C4)O)C)C(C)C.O</chem>

Note: Compound Identification (CID), Simplified Molecular-Input Line-Entry System (SMILES)

Table S5. SwissADME Analysis Report of Top Bioactive Compounds

CLASS	Parameters	Compound #							Control		
		1	2	3	4	5	6	7	Rivaroxaban	Apixaban	Betrixaban
Physico-chemical Properties	MW (g/mol)	220.35	290.44	344.49	222.37	212.33	220.35	288.34	435.88	459.50	451.91
	# Heavy Atoms	16	21	25	16	15	16	22	29	34	32
	# Aromatic Heavy Atoms	0	0	0	0	0	0	12	11	17	18
	Fraction Csp3	0.87	0.89	0.82	0.87	0.77	1	0.05	0.32	0.28	0.13
	# Rotatable Bonds	0	0	2	0	8	0	3	6	5	9
	# H-bond acceptors	1	2	3	1	2	1	2	5	5	5
	# H-bond donors	0	2	0	1	0	0	0	1	1	3
	Molar Refractivity	68.27	86.32	99.9	70.46	65.42	66.11	90.59	114.09	132.70	125.23
Lipophilicity	TPSA (Å ²)	12.53	40.46	43.37	20.23	26.3	12.53	26.3	116.42	110.76	107.41
	iLOGP	3.11	2.92	3.19	3.1	3.57	3.06	3.23	2.95	3.62	2.72
	XLOGP3	3.56	3.41	4.45	3.89	4.24	3.85	4.84	2.49	2.24	3.56
	WLOGP	3.94	3.67	4.7	4.06	3.71	3.77	4.5	1.76	1.94	3.75
	MLOGP	3.67	3.7	4.1	3.67	3.32	3.81	4.24	1.41	1.76	2.83
	Silicos-IT Log P	4.07	2.99	4.13	3.75	3.58	4.18	4.87	2.84	1.96	3.24
	Consensus Log P	3.67	3.34	4.11	3.69	3.68	3.73	4.34	2.29	2.30	3.22
Water Solubility	ESOL Class	Soluble	Soluble	Moderately soluble	Soluble	Soluble	Soluble	Moderately soluble	Soluble	Moderately Soluble	Moderately Soluble
	Ali Class	Soluble	Soluble	Moderately soluble	Moderately soluble	Moderately soluble	Soluble	Moderately soluble	Moderately Soluble	Moderately Soluble	Moderately Soluble
	Silicos-IT Class	Soluble	Soluble	Moderately soluble	Soluble	Soluble	Soluble	Moderately soluble	Moderately Soluble	Moderately Soluble	Poorly Soluble
Pharmacokinetics	GI Absorption	High	High	High	High	High	High	High	High	High	High
	BBB Permeant	Yes	Yes	Yes	Yes	Yes	Yes	Yes	No	No	No
	Pgp Substrate	No	Yes	No	No	No	No	No	Yes	Yes	No
	CYP1A2 Inhibitor	No	No	No	No	No	No	No	No	No	No
	CYP2C19 Inhibitor	Yes	No	No	No	No	No	Yes	Yes	Yes	Yes
	CYP2C9 Inhibitor	Yes	No	Yes	Yes	No	No	Yes	Yes	Yes	Yes
	CYP2D6 Inhibitor	No	No	No	No	No	No	No	No	Yes	Yes
	CYP3A4 Inhibitor	No	No	No	No	No	No	No	Yes	Yes	Yes
	Log Kp (cm/s)	-5.12	-5.65	-5.24	-4.89	-4.58	-4.91	-4.62	-7.19	-7.51	-6.53
Drug-likeness	Lipinski #	0	0	0	0	0	0	1	0	0	0
	Ghose #	0	0	0	0	0	0	0	0	1	0
	Veber #	0	0	0	0	0	0	0	0	0	0

	Egan #	0	0	0	0	0	0	0	0	0	0
	Muegge #	1	0	0	1	0	1	0	0	0	0
	Bioavailability Score	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55
Medicinal Chemistry	PAINS # Alerts	0	0	0	0	0	0	0	0	0	0
	Brenk # Alerts	2	1	0	1	1	1	2	0	0	2
	Leadlikeness # Violations	2	0	1	2	3	2	1	1	1	3
	Synthetic Accessibility	4.35	4.96	4.87	3.69	2.84	4.87	3.33	3.63	3.48	3.05

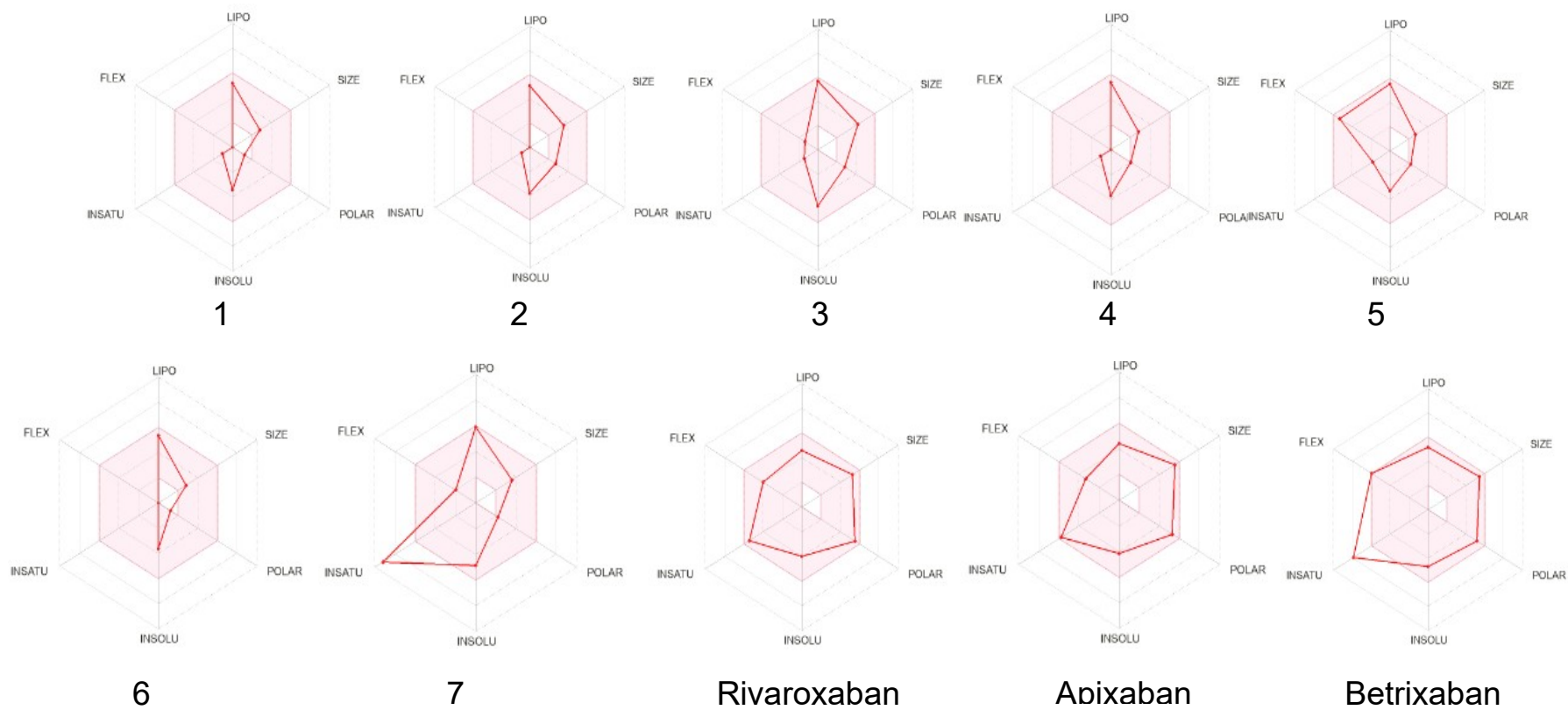


Figure S3. Radar Plot of the Bioactive Compound

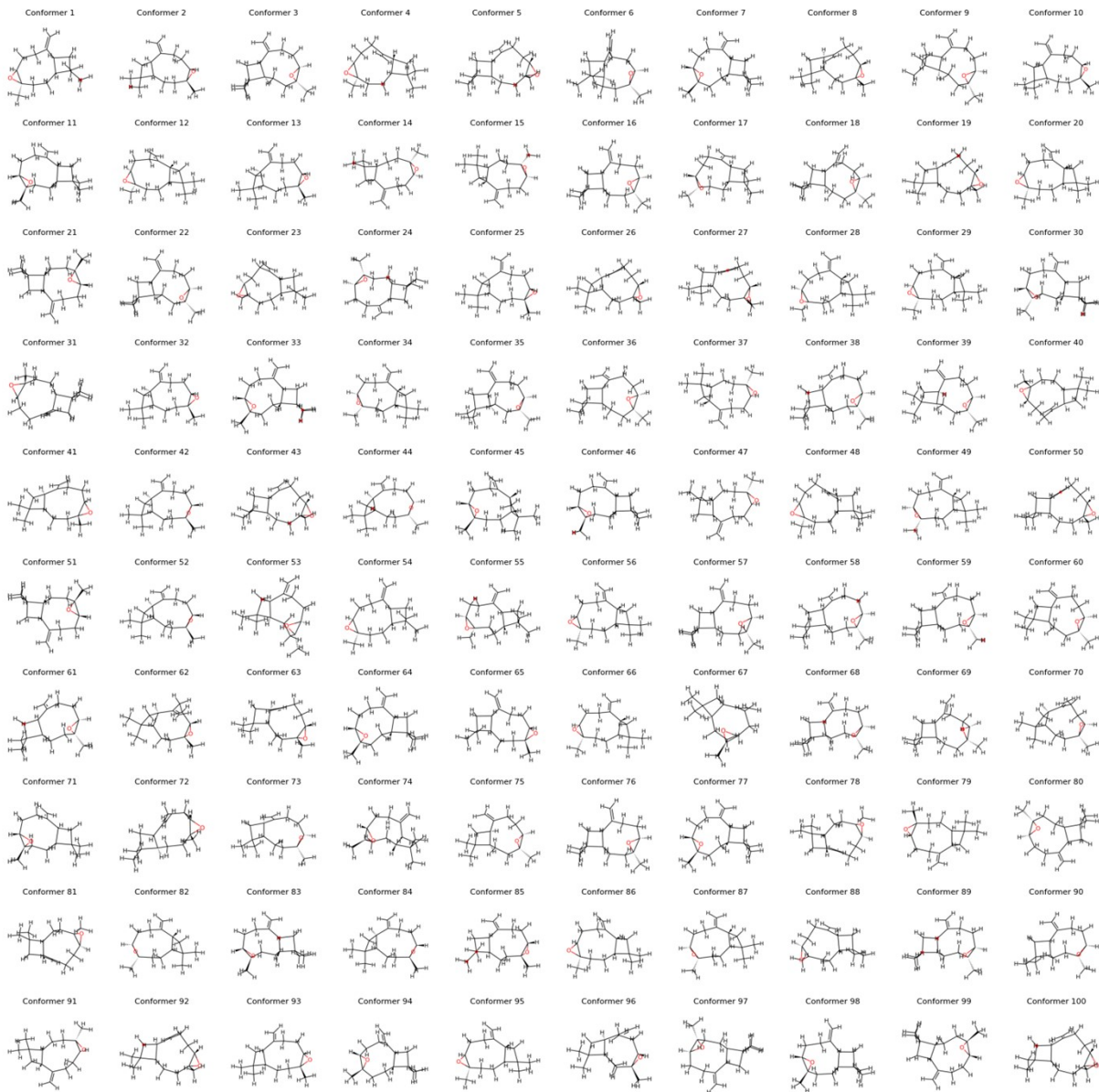
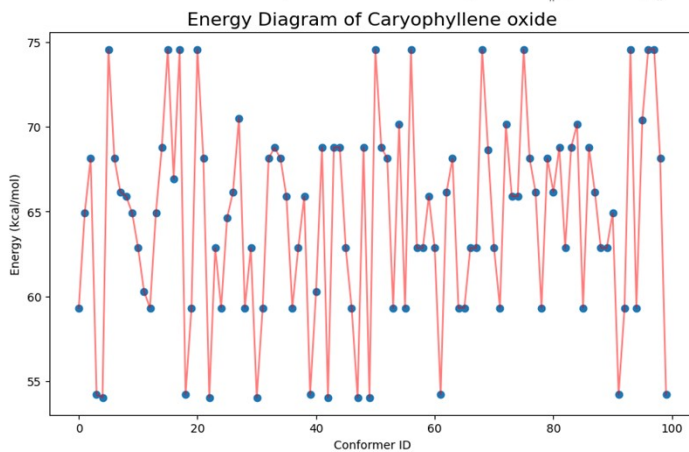


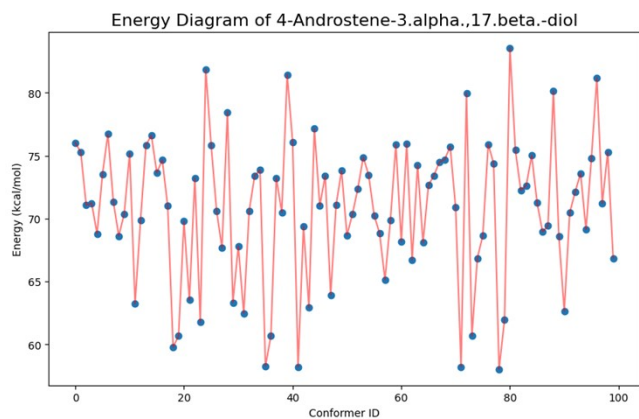
Figure S4.

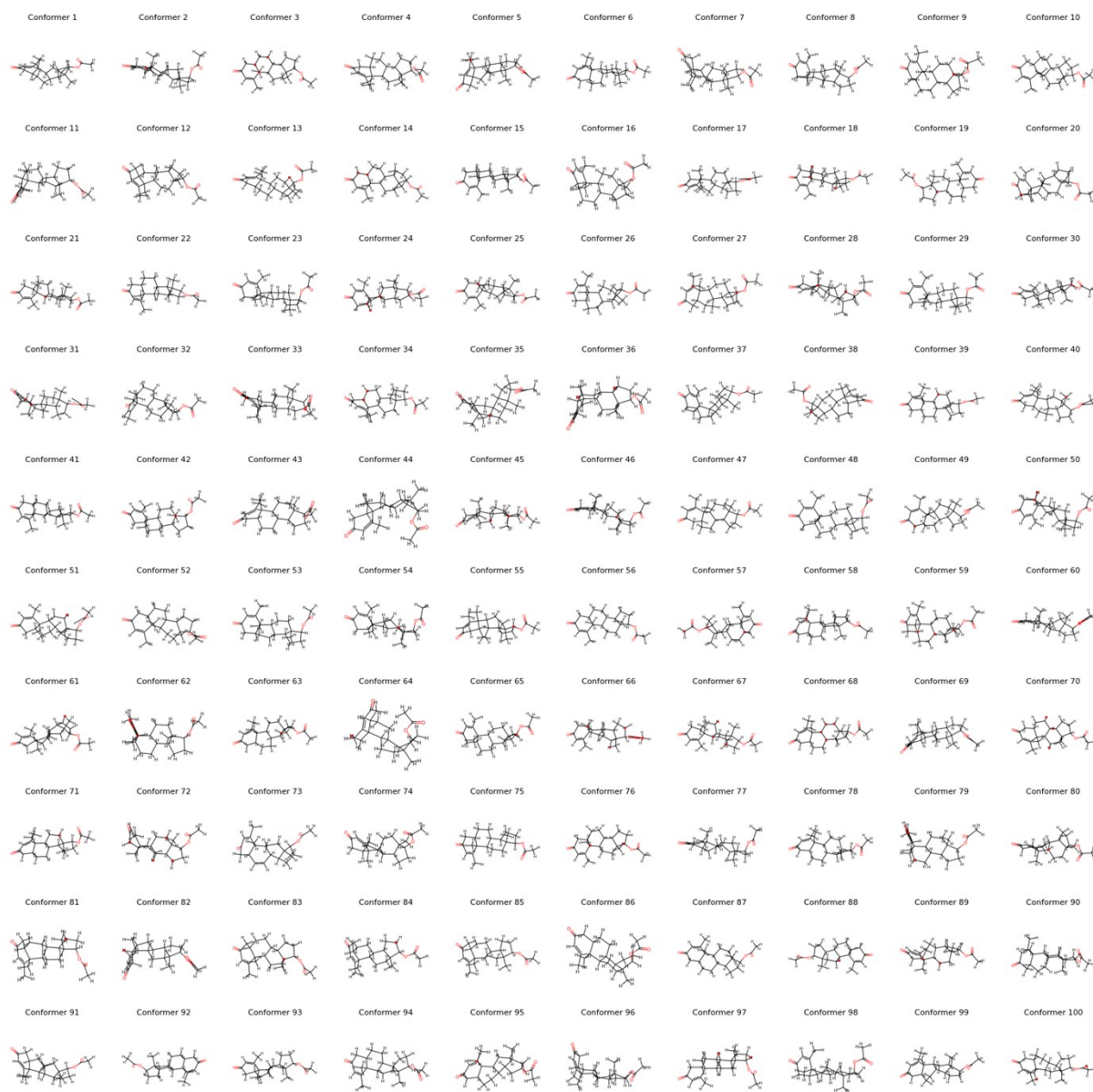
Analysis of



*Conformational
Compound 1*

Figure S5. Conformational Analysis of Compound 2





Energy Diagram of Methenolone Acetate

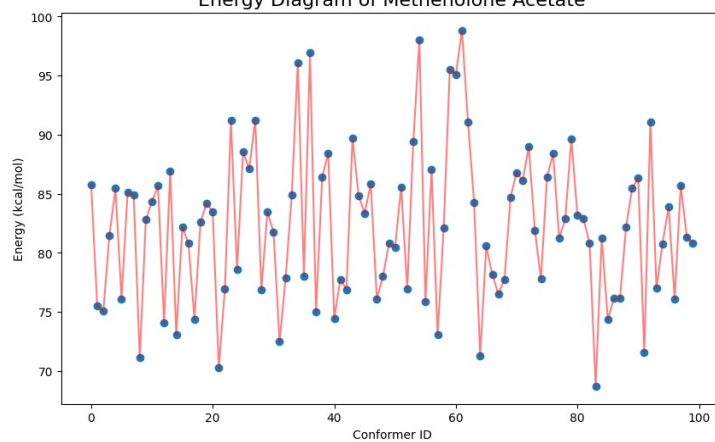


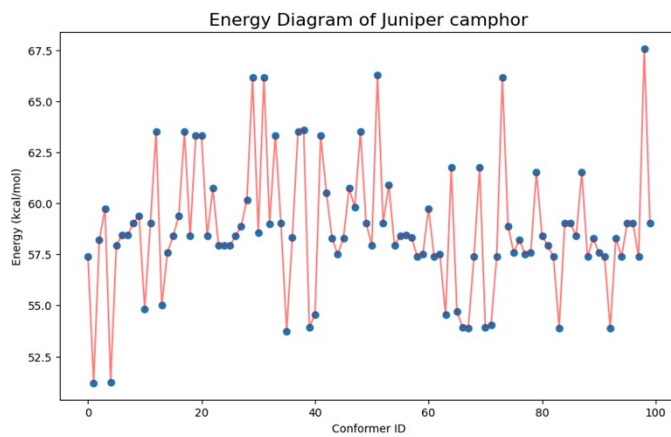
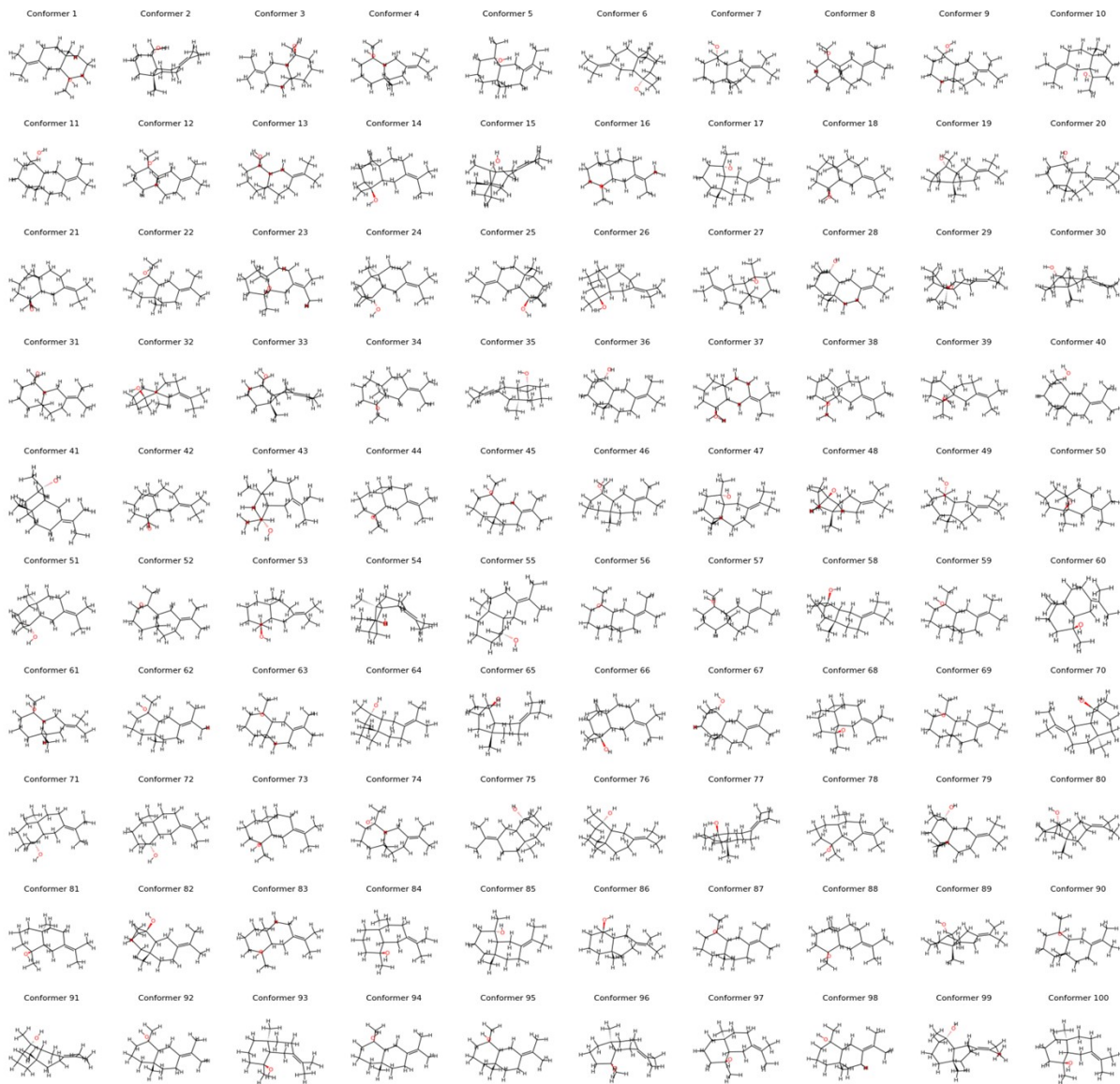
Figure S6.

Analysis of

Conformational

Compound 3

Figure S7. Conformational Analysis of Compound 4



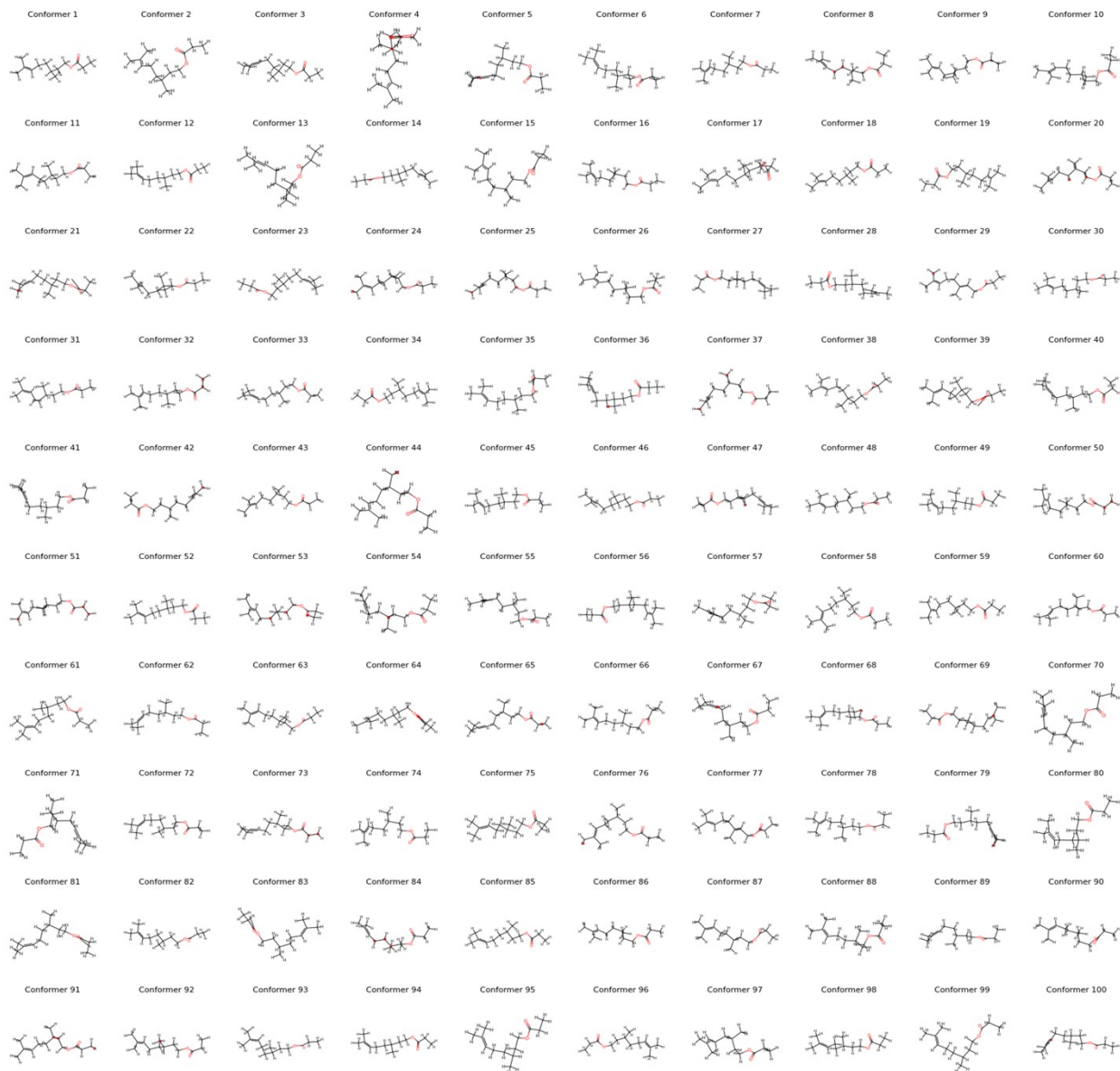


Figure S8. Conformational Analysis of Compound 5

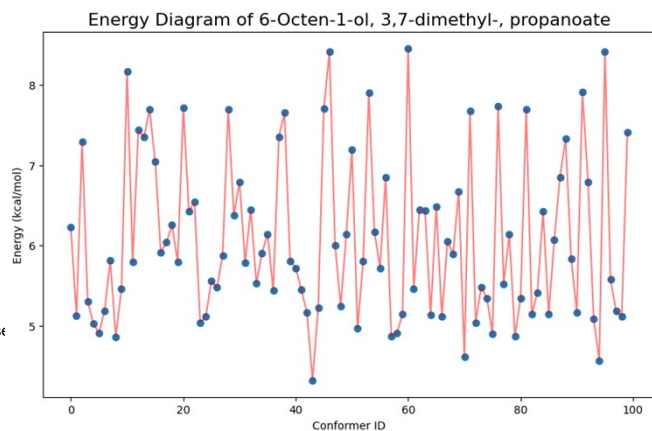
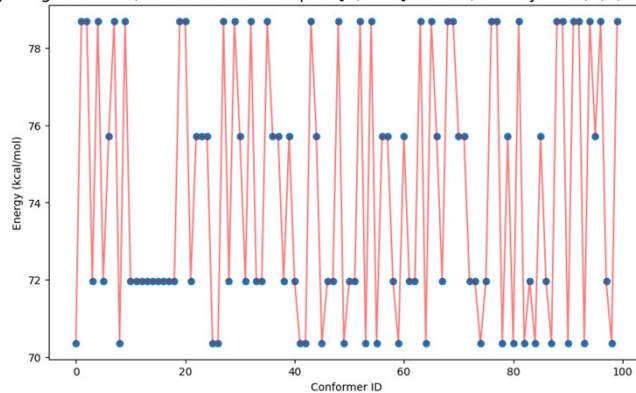




Figure S9. Conformational Analysis of Compound 6

Energy Diagram of 4a,7-Methano-4aH-naphth[1,8a-b]oxirene, octahydro-4,4,8,8-tetramethyl-



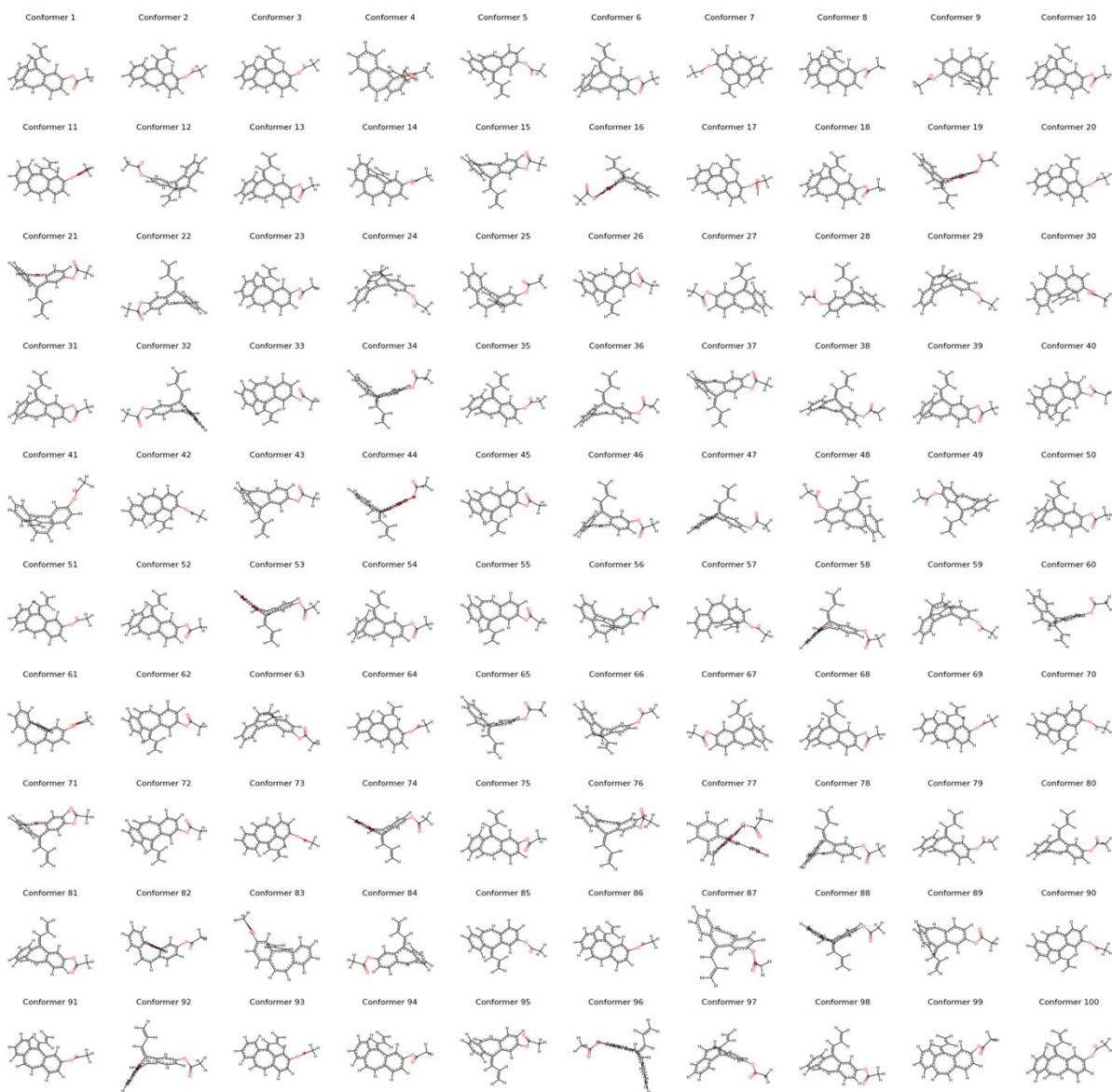
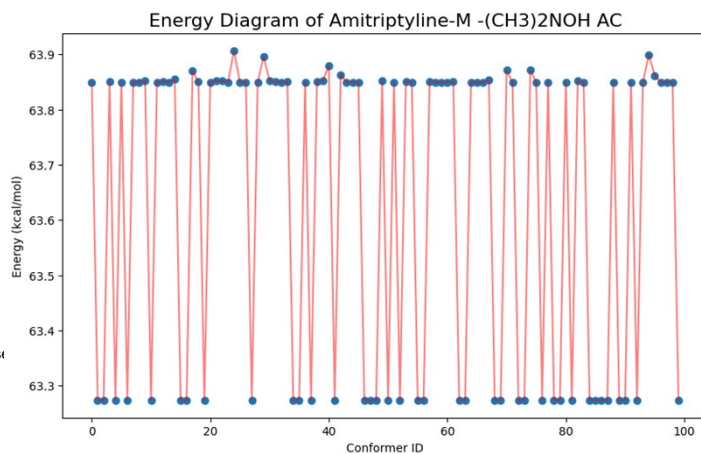


Figure S10. Conformational Analysis of Compound 7



Active Site	Mode	Compound #													
		1		2		3		4		5		6		7	
		Vina	Vinardo	Vina	Vinardo	Vina	Vinardo	Vina	Vinardo	Vina	Vinardo	Vina	Vinardo	Vina	Vinardo
II	1	-6.5	-6.1	-6.8	-6.0	-7.6	-6.8	-6.0	-5.3	-4.7	-5.6	-5.7	-4.8	-7.2	-6.6
	2	-6.4	-6.0	-6.6	-5.7	-7.2	-5.9	-5.8	-5.0	-4.7	-5.0	-5.6	-4.6	-6.8	-6.6
	3	-5.9	-5.9	-6.5	-5.6	-6.4	-4.6	-5.7	-4.8	-4.6	-4.8	-5.5	-4.4	-6.7	-6.3
	4	-5.8	-5.7	-6.4	-5.2	-6.3	-3.1	-5.6	-4.7	-4.6	-4.8	-5.3	-3.5	-6.8	-6.2
	5	-5.8	-5.6	-6.4	-5.1	-6.3	-3.1	-5.6	-4.7	-4.4	-4.6	-5.1	-3.3	-6.1	-5.6
	6	-5.6	-5.4	-6.4	-5.0	-6.2	-2.9	-5.6	-4.0	-4.4	-4.4	-4.8	-3.2	-	-
	7	-5.6	-4.4	-6.3	-5.0	-6.2	-	-5.4	-3.6	-4.3	-4.4	-4.6	-3.2	-	-
	8	-5.5	-4.2	-6.2	-4.5	-6.1	-	-5.2	-3.5	-4.3	-4.3	-	-3.1	-	-
	9	-4.5	-	-5.7	-3.8	-	-	-	-3.3	-4.2	-4.2	-	-3.0	-	-
	10	-3.7	-	-4.4	-3.0	-	-	-	-3.1	-4.2	-	-	-2.8	-	-
III	1	-5.5	-4.8	-6.6	-4.8	-7.9	-5.7	-5.2	-4.1	-4.9	-3.5	-5.3	-3.9	-6.9	-4.4
	2	-5.4	-4.6	-6.1	-4.5	-6.6	-4.6	-5.2	-4.0	-4.8	-3.0	-5.2	-3.3	-6.5	-4.3
	3	-5.3	-4.3	-5.6	-4.3	-6.2	-4.5	-5.0	-3.9	-4.7	-3.0	-5.0	-3.3	-6.4	-4.3
	4	-5.2	-4.2	-5.6	-4.3	-6.1	-4.4	-4.9	-3.7	-4.7	-2.9	-4.9	-3.2	-6.3	-4.1
	5	-5.1	-3.9	-5.5	-4.2	-5.5	-4.4	-4.8	-3.6	-4.4	-2.8	-4.8	-3.2	-6.0	-4.1
	6	-5.1	-3.8	-5.5	-4.0	-5.5	-4.4	-4.7	-3.6	-4.2	-2.8	-4.8	-3.1	-5.9	-4.1
	7	-4.8	-3.6	-5.3	-3.9	-5.2	-4.3	-4.7	-3.5	-4.2	-2.8	-4.8	-2.9	-5.7	-4.1
	8	-4.6	-3.6	-5.2	-3.9	-5.1	-4.3	-4.6	-3.4	-4.1	-2.8	-4.8	-2.8	-5.6	-4.0
	9	-4.5	-3.5	-5.2	-3.8	-5.0	-4.2	-4.5	-3.3	-3.9	-2.8	-4.6	-2.8	-5.5	-4.0
	10	-4.4	-3.4	-5.2	-3.6	-4.9	-4.0	-4.5	-3.2	-3.9	-2.5	-4.6	-2.8	-5.3	-3.9
V	1	-5.6	-4.4	-7.0	-6.1	-7.0	-5.8	-6.4	-5.3	-4.7	-4.3	-5.0	-3.8	-6.8	-6.1
	2	-5.5	-4.4	-6.7	-5.7	-6.9	-5.6	-6.0	-5.0	-4.6	-4.0	-4.9	-3.5	-6.6	-6.0
	3	-5.3	-4.3	-6.7	-5.6	-6.6	-5.3	-5.3	-4.5	-4.5	-4.0	-4.8	-3.5	-6.5	-6.0
	4	-5.2	-4.3	-6.6	-5.5	-6.6	-5.1	-5.2	-4.4	-4.3	-4.0	-4.7	-3.4	-6.3	-5.9
	5	-5.2	-4.2	-6.6	-5.5	-6.3	-5.0	-5.2	-4.3	-4.3	-3.9	-4.7	-3.4	-6.2	-5.7
	6	-5.1	-4.2	-6.5	-5.5	-6.3	-5.0	-5.0	-4.2	-4.3	-3.9	-4.6	-3.3	-6.0	-5.7
	7	-5.1	-4.1	-6.5	-5.4	-6.2	-4.9	-5.0	-4.1	-4.3	-3.8	-4.5	-3.3	-5.9	-5.6
	8	-5.1	-4.1	-6.4	-5.2	-6.4	-4.9	-4.9	-4.0	-4.3	-3.8	-4.4	-3.3	-5.8	-5.6
	9	-5.1	-4.0	-6.3	-5.1	-6.0	-4.7	-4.9	-4.0	-4.2	-3.7	-4.4	-3.1	-5.7	-5.5
	10	-5.0	-4.0	-6.2	-5.0	-6.0	-4.6	-4.6	-4.0	-4.2	-3.7	-4.4	-3.0	-5.6	-5.3

Table S6. CFIXa Docking Scores

Table S7. Conformational Changes on Best Ligand-Complex per Bioactive Compound in CFIXa

Compound #	Vina Score	Vinardo Score	Active Site	Bond Length (Å)			Bond Angle (Å)					
				Atom	l_i	l_r	Δl	Atom	\angle_i	\angle_r	$\Delta \angle$	
1	-6.5	-6.1	II	H:TYR99								
				CG - CD1	1.39556	1.39549	-0.00007	α (CD1-CG-CD2)	118.05	118.05	0	
				CG - CD2	1.38966	1.38993	0.00027	α (CG-CD2-CE2)	121.11	121.12	0.01	
				CD1 - CE1	1.39409	1.39403	-0.00006	α (CD2-CE2-CZ)	119.59	119.57	-0.02	
				CD2 - CE2	1.39517	1.39501	-0.00016	α (CE2-CZ-CE1)	120.23	120.23	0	
				CE1 - CZ	1.37854	1.37844	-0.00010	α (CZ-CE1-CD1)	119.64	119.67	0.03	
				CE2 - CZ	1.38939	1.38949	0.00010	α (CE1-CD1-CG)	121.24	121.21	-0.03	
				H:TRP215								
				CD2 - CE2	1.41412	1.41425	0.00013	α (CE3-CD2-CE2)	118.63	118.63	0	
				CD2 - CE3	1.39553	1.39532	-0.00021	α (CD2-CE2-CZ2)	122.34	122.36	0.02	
				CE2 - CZ2	1.411	1.41021	-0.00079	α (CE2-CZ2-CH2)	117.54	117.55	0.01	
				CE3 - CZ3	1.39216	1.39257	0.00041	α (CZ2-CH2-CZ3)	120.93	120.92	-0.01	
				CZ2 - CH2	1.3775	1.37761	0.00011	α (CH2-CZ3-CE3)	121.18	121.20	0.02	
CZ3 - CH2	1.40946	1.40908	-0.00038	α (CZ3-CE3-CD2)	119.30	119.27	-0.03					
2	-7.0	-6.1	V	L:PHE98								
				CG - CD1	1.39561	1.39508	-0.000530	α (CD1-CG-CD2)	118.35	118.36	0.01	
				CG - CD2	1.38125	1.38094	-0.000310	α (CG-CD2-CE2)	120.93	120.94	0.01	
				CD1 - CE1	1.40154	1.40109	-0.000450	α (CD2-CE2-CZ)	120.20	120.19	-0.01	
				CD2 - CE2	1.39071	1.39084	0.000130	α (CE2-CZ-CE1)	119.86	119.84	-0.02	
				CE1 - CZ	1.37139	1.37223	0.000840	α (CZ-CE1-CD1)	119.83	119.83	0	
				CE2 - CZ	1.38619	1.38583	-0.000360	α (CE1-CD1-CG)	120.79	120.80	0.01	
				L:LYS100								
				CB - CG	1.51808	1.51855	0.000470	α (CB-CG-CD)	110.42	110.41	-0.01	
				CG - CD	1.51948	1.51874	-0.000740					
CE - NZ	1.49714	1.49794	0.000800	α (CE-NZ-HZ1)	109.30	109.30	0					
HZ1 - NZ	1.01014	1.00985	-0.000290									
3	-7.6	-6.8	II	H:TYR99								
				CG - CD1	1.39556	1.39531	-0.000250	α (CD1-CG-CD2)	118.05	118.05	0	
CG - CD2	1.38966	1.39003	0.000370	α (CG-CD2-CE2)	121.11	121.11	0					

				CD1 - CE1	1.39409	1.39417	0.000080	α (CD2-CE2-CZ)	119.59	119.53	-0.06
				CD2 - CE2	1.39517	1.39534	0.000170	α (CE2-CZ-CE1)	120.23	120.27	0.04
				CE1 - CZ	1.37854	1.37766	-0.000880	α (CZ-CE1-CD1)	119.64	119.66	0.02
				CE2 - CZ	1.38939	1.38953	0.000140	α (CE1-CD1-CG)	121.24	121.22	-0.02
				H:TRP215							
				CG - CD1	1.36247	1.3615	-0.000970	α (CE3-CD2-CE2)	118.63	118.67	0.04
				CG - CD2	1.44127	1.44155	0.000280	α (CD2-CE2-CZ2)	122.34	122.33	-0.01
				CD1 - NE1	1.37831	1.37861	0.000300	α (CE2-CZ2-CH2)	117.54	117.55	0.01
				CD2 - CE2	1.41412	1.41369	-0.000430	α (CZ2-CH2-CZ3)	120.93	120.94	0.01
				NE1 - CE2	1.37471	1.37452	-0.000190	α (CH2-CZ3-CE3)	121.18	121.17	-0.01
				CD2 - CE3	1.39553	1.3958	0.000270	α (CZ3-CE3-CD2)	119.30	119.26	-0.04
				CE2 - CZ2	1.411	1.41062	-0.000380	α (CE2-CD2-CG)	107.19	107.18	-0.01
				CE3 - CZ3	1.39216	1.39215	-0.000010	α (CD2-CG-CD1)	106.27	106.26	-0.01
				CZ2 - CH2	1.3775	1.37728	-0.000220	α (CG-CD1-NE1)	110.25	110.29	0.04
				CZ3 - CH2	1.40946	1.4098	0.000340	α (CD1-NE1-CE2)	109.01	108.97	-0.04
				HE1 - NE1	1.00952	1.00894	-0.000580	α (NE1-CE2-CD2)	107.23	107.25	0.02
				H:TYR128							
				CG - CD1	1.39175	1.39183	0.000080	α (CD1-CG-CD2)	118.01	118	-0.01
				CG - CD2	1.3885	1.38838	-0.000120	α (CG-CD2-CE2)	121.21	121.19	-0.02
				CD1 - CE1	1.39351	1.39339	-0.000120	α (CD2-CE2-CZ)	119.57	119.63	0.06
				CD2 - CE2	1.39727	1.39715	-0.000120	α (CE2-CZ-CE1)	120.29	120.25	-0.04
4	-6.4	-5.3	V	CE1 - CZ	1.38569	1.38633	0.000640	α (CZ-CE1-CD1)	119.38	119.36	-0.02
				CE2 - CZ	1.38262	1.3824	-0.000220	α (CE1-CD1-CG)	121.43	121.47	0.04
				H:ILE129B							
				CB - CG1	1.53349	1.53338	-0.000110	α (CD1-CG1-CB)	114.11	114.10	-0.01
				CB - CG2	1.54926	1.54956	0.000300	α (CG1-CB-CG2)	112.14	112.09	-0.05
				CG1 - CD1	1.57649	1.57657	0.000080				
				H:TYR99							
				CG - CD1	1.39556	1.39527	-0.000290	α (CD1-CG-CD2)	118.05	118.08	0.03
				CG - CD2	1.38966	1.38882	-0.000840	α (CG-CD2-CE2)	121.11	121.12	0.01
				CD1 - CE1	1.39409	1.39521	0.001120	α (CD2-CE2-CZ)	119.59	119.54	-0.05
5	-4.7	-5.6	II	CD2 - CE2	1.39517	1.39589	0.000720	α (CE2-CZ-CE1)	120.23	120.27	0.04
				CE1 - CZ	1.37854	1.37795	-0.000590	α (CZ-CE1-CD1)	119.64	119.64	0
				CE2 - CZ	1.38939	1.38904	-0.000350	α (CE1-CD1-CG)	121.24	121.19	-0.05
				H:TRP215							

				CD2 - CE2	1.41412	1.41369	-0.000430	α (CE3-CD2-CE2)	118.63	118.65	0.02
				CD2 - CE3	1.39553	1.39571	0.000180	α (CD2-CE2-CZ2)	122.34	122.29	-0.05
				CE2 - CZ2	1.411	1.41166	0.000660	α (CE2-CZ2-CH2)	117.54	117.57	0.03
				CE3 - CZ3	1.39216	1.393	0.000840	α (CZ2-CH2-CZ3)	120.93	120.95	0.02
				CZ2 - CH2	1.3775	1.37717	-0.000330	α (CH2-CZ3-CE3)	121.18	121.15	-0.03
				CZ3 - CH2	1.40946	1.40887	-0.000590	α (CZ3-CE3-CD2)	119.30	119.30	0
				H:TYR99							
				CG - CD1	1.39556	1.39634	0.000780	α (CD1-CG-CD2)	118.05	117.99	-0.06
				CG - CD2	1.38966	1.38971	0.000050	α (CG-CD2-CE2)	121.11	121.14	0.03
				CD1 - CE1	1.39409	1.39403	-0.000060	α (CD2-CE2-CZ)	119.59	119.30	-0.29
				CD2 - CE2	1.39517	1.39488	-0.000290	α (CE2-CZ-CE1)	120.23	120.23	0
				CE1 - CZ	1.37854	1.37858	0.000040	α (CZ-CE1-CD1)	119.64	119.63	-0.01
				CE2 - CZ	1.38939	1.38915	-0.000240	α (CE1-CD1-CG)	121.24	121.26	0.02
6	-5.7	-4.8	II	H:TRP215							
				CD2 - CE2	1.41412	1.41467	0.000550	α (CE3-CD2-CE2)	118.63	118.61	-0.02
				CD2 - CE3	1.39553	1.39476	-0.000770	α (CD2-CE2-CZ2)	122.34	122.31	-0.03
				CE2 - CZ2	1.411	1.41127	0.000270	α (CE2-CZ2-CH2)	117.54	117.54	0
				CE3 - CZ3	1.39216	1.39205	-0.000110	α (CZ2-CH2-CZ3)	120.93	120.95	0.02
				CZ2 - CH2	1.3775	1.37681	-0.000690	α (CH2-CZ3-CE3)	121.18	121.16	-0.02
				CZ3 - CH2	1.40946	1.40937	-0.000090	α (CZ3-CE3-CD2)	119.30	119.33	0.03
				H:TYR99							
				CG - CD1	1.39556	1.39527	-0.000290	α (CD1-CG-CD2)	118.05	118.06	0.01
				CG - CD2	1.38966	1.3897	0.000040	α (CG-CD2-CE2)	121.11	121.08	-0.03
				CD1 - CE1	1.39409	1.39499	0.000900	α (CD2-CE2-CZ)	119.59	119.60	0.01
				CD2 - CE2	1.39517	1.39539	0.000220	α (CE2-CZ-CE1)	120.23	120.25	0.02
				CE1 - CZ	1.37854	1.3782	-0.000340	α (CZ-CE1-CD1)	119.64	119.59	-0.05
				CE2 - CZ	1.38939	1.38977	0.000380	α (CE1-CD1-CG)	121.24	121.27	0.03
				CZ - OH	1.37181	1.3714	-0.000410	α (CE1-CZ-OH)	118.89	118.88	-0.01
7	-7.2	-6.6	II	HH - OH	0.959771	0.959907	0.000136	α (CZ-OH-HH)	106.98	107.04	0.06

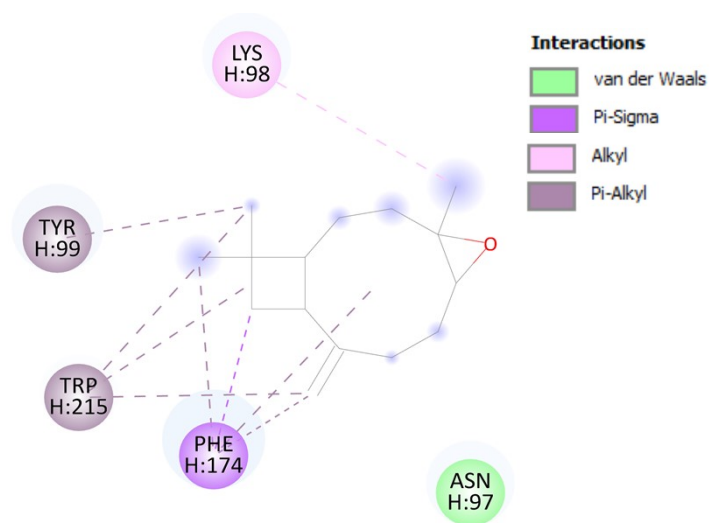
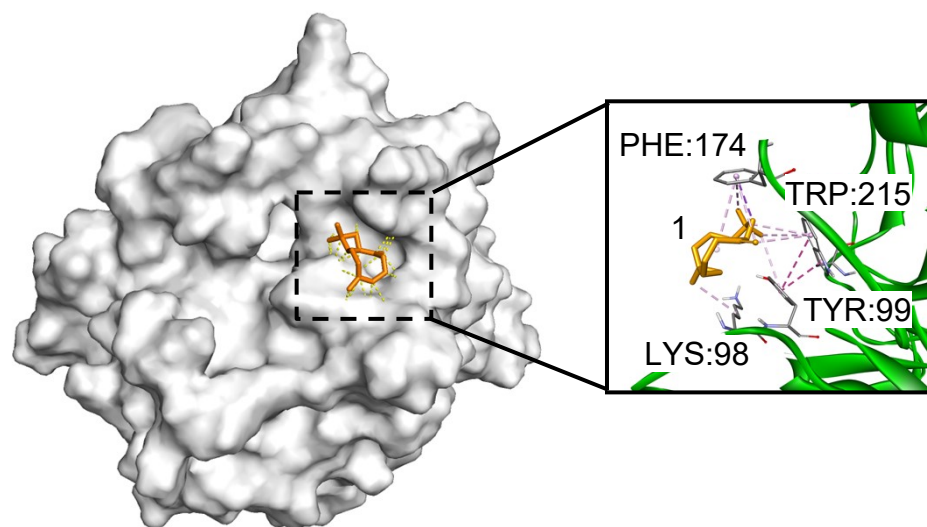
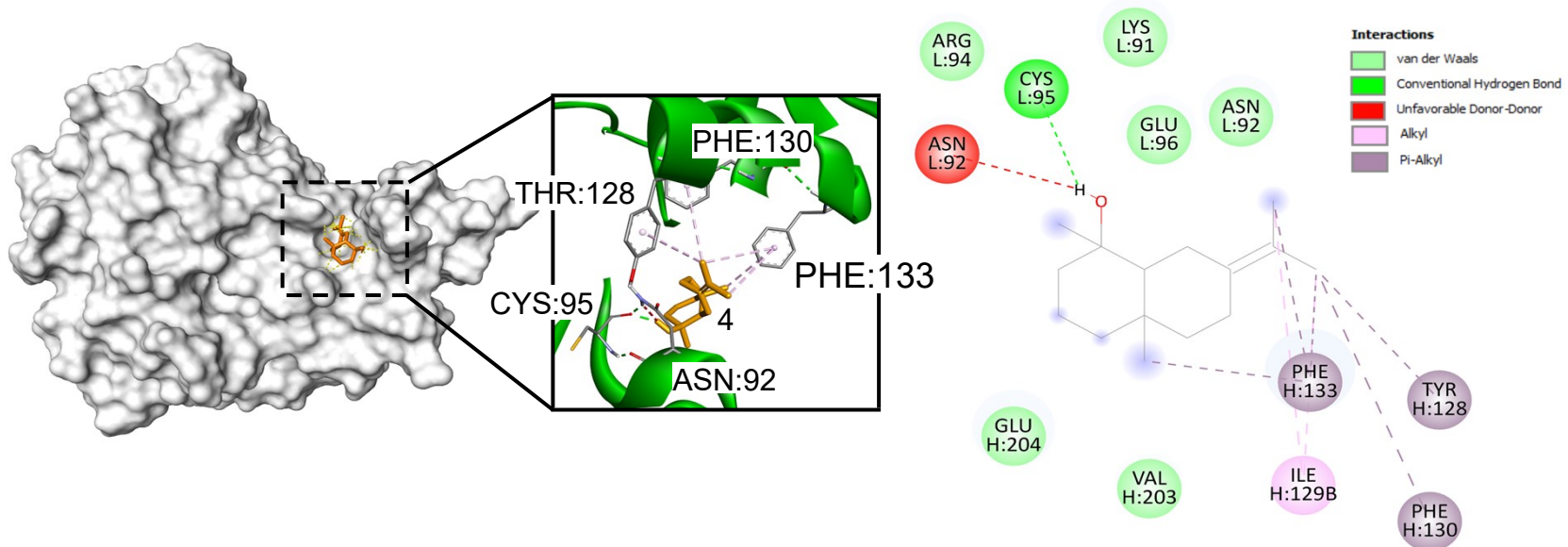


Figure S11.
Compound 1 Best Binding Mode on CFIXa at Active Site II

Figure S12. *Compound 4 Best Binding Mode on CFIXa at Active Site V*



S13. Compound 5 Best Binding Mode on CFIXa at Active Site II

Figure

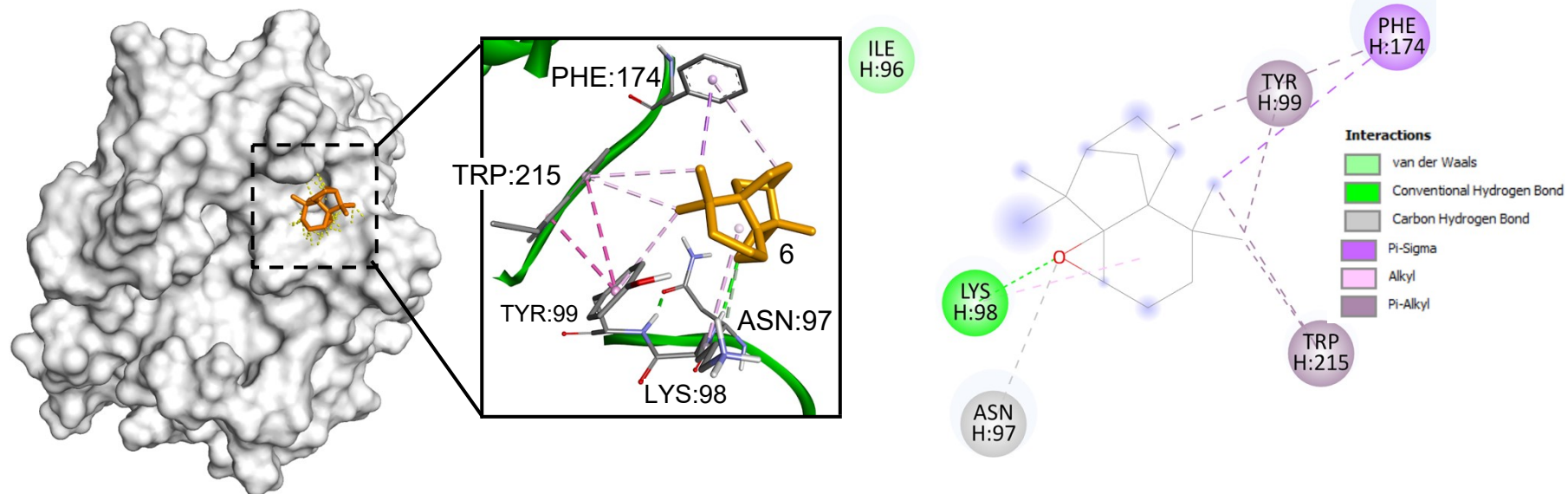
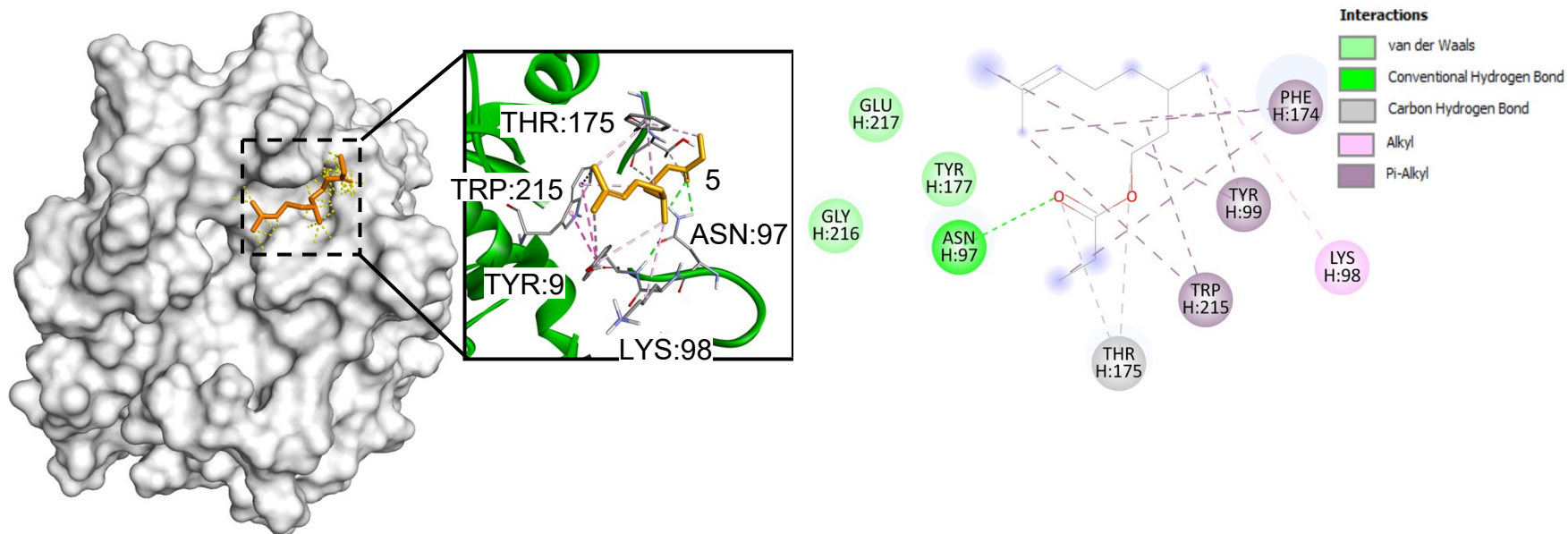


Figure S14. Compound 6 Best Binding Mode on CFXa at Active Site V