

Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
 Data File : BM039474.D
 Acq On : 19 Apr 2023 01:40
 Operator : CG/JU
 Sample : 02335-18
 Misc :
 ALS Vial : 25 Sample Multiplier: 1

Instrument :
 BNA_M
 ClientSampleId :

Integration Parameters: LSCINT.P

Integrator: RTE

Smoothing : OFF

Filtering: 5

Sampling : 1

Min Area: 0.1 % 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

Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
 Title : SVOA CALIBRATION

Signal : TIC: BM039474.D\data.ms

peak #	R.T. min	first scan	max scan	last scan	PK TY	peak height	corr. area	corr. % max.	% of total
1	3.652	77	79	95	rBV	127563	230720	0.27%	0.063%
2	5.128	321	330	335	rBV2	28121530	77981067	90.04%	21.368%
3	6.504	558	564	574	rBV	84164	156937	0.18%	0.043%
4	6.798	609	614	625	rVB	55039	96747	0.11%	0.027%
5	7.022	647	652	668	rBV	203860	385125	0.44%	0.106%
6	7.169	671	677	692	rBV	962087	1611295	1.86%	0.442%
7	7.369	704	711	735	rVB	867375	1517611	1.75%	0.416%
8	7.557	736	743	754	rVB	138836	251008	0.29%	0.069%
9	7.722	763	771	784	rBV	8135385	14308123	16.52%	3.921%
10	7.887	785	799	804	rBV2	26733836	86611059	100.00%	23.733%
11	8.210	843	854	858	rBV2	26741037	79457394	91.74%	21.772%
12	8.569	908	915	931	rBV	561816	1111755	1.28%	0.305%
13	9.004	982	989	992	rBV	1083690	1787815	2.06%	0.490%
14	9.051	992	997	1015	rVB	2465864	4313520	4.98%	1.182%
15	9.334	1040	1045	1053	rBV	53424	104922	0.12%	0.029%
16	9.457	1059	1066	1074	rBV	284182	466751	0.54%	0.128%
17	9.728	1105	1112	1134	rBV	822232	1505993	1.74%	0.413%
18	10.275	1198	1205	1221	rBV	1089594	2042785	2.36%	0.560%
19	10.516	1236	1246	1261	rBV	17129065	31724745	36.63%	8.693%
20	10.645	1261	1268	1275	rVV	1085281	1789994	2.07%	0.490%
21	10.792	1286	1293	1310	rBV	416222	843249	0.97%	0.231%
22	11.028	1325	1333	1346	rBV	1644182	2868397	3.31%	0.786%
23	11.198	1356	1362	1366	rBV	69020	119803	0.14%	0.033%
24	11.257	1366	1372	1377	rBV	238107	439303	0.51%	0.120%
25	11.475	1401	1409	1422	rBV	75535	200926	0.23%	0.055%
26	12.675	1602	1613	1624	rBV	992967	1724665	1.99%	0.473%
27	13.286	1710	1717	1730	rBV2	1609679	3338138	3.85%	0.915%
28	13.504	1748	1754	1765	rVB	331134	494938	0.57%	0.136%
29	13.904	1815	1822	1845	rBV	2651345	3837290	4.43%	1.051%
30	14.180	1861	1869	1877	rBV2	2937463	4440116	5.13%	1.217%
31	14.251	1877	1881	1888	rVB	80143	135327	0.16%	0.037%
32	14.486	1912	1921	1936	rVV	1743016	2610905	3.01%	0.715%
33	14.751	1960	1966	1971	rBV3	43621	115941	0.13%	0.032%
34	14.804	1971	1975	1986	rVB	89265	171422	0.20%	0.047%
35	15.480	2081	2090	2096	rBV	3604373	5323010	6.15%	1.459%
36	15.539	2096	2100	2107	rVV	304333	550915	0.64%	0.151%

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Peak separation: 5

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37	15.616	2107	2113	2127	rVV	1261598	1967048	2.27%	0.539%
38	15.851	2148	2153	2161	rVB	66321	97792	0.11%	0.027%
39	17.239	2381	2389	2399	rBV	2006909	2942141	3.40%	0.806%
40	17.339	2400	2406	2426	rVB	3976791	5834887	6.74%	1.599%
41	18.139	2537	2542	2551	rBV	116569	222011	0.26%	0.061%
42	18.233	2552	2558	2564	rVB4	61724	127985	0.15%	0.035%
43	19.421	2755	2760	2765	rBV2	81661	148349	0.17%	0.041%
44	19.639	2790	2797	2815	rBV	4751895	6785257	7.83%	1.859%
45	20.627	2962	2965	2969	rBV	161327	172835	0.20%	0.047%
46	21.145	3049	3053	3063	rBV4	77975	148792	0.17%	0.041%
47	21.427	3096	3101	3117	rBV	1905613	2812417	3.25%	0.771%
48	22.515	3283	3286	3291	rVB	120039	149528	0.17%	0.041%
49	23.056	3375	3378	3386	rVB	181566	275336	0.32%	0.075%
50	23.644	3471	3478	3496	rVV2	2653765	5653202	6.53%	1.549%
51	23.797	3498	3504	3517	rVB2	1169218	2495774	2.88%	0.684%
52	24.374	3599	3602	3612	rVB	232501	441718	0.51%	0.121%

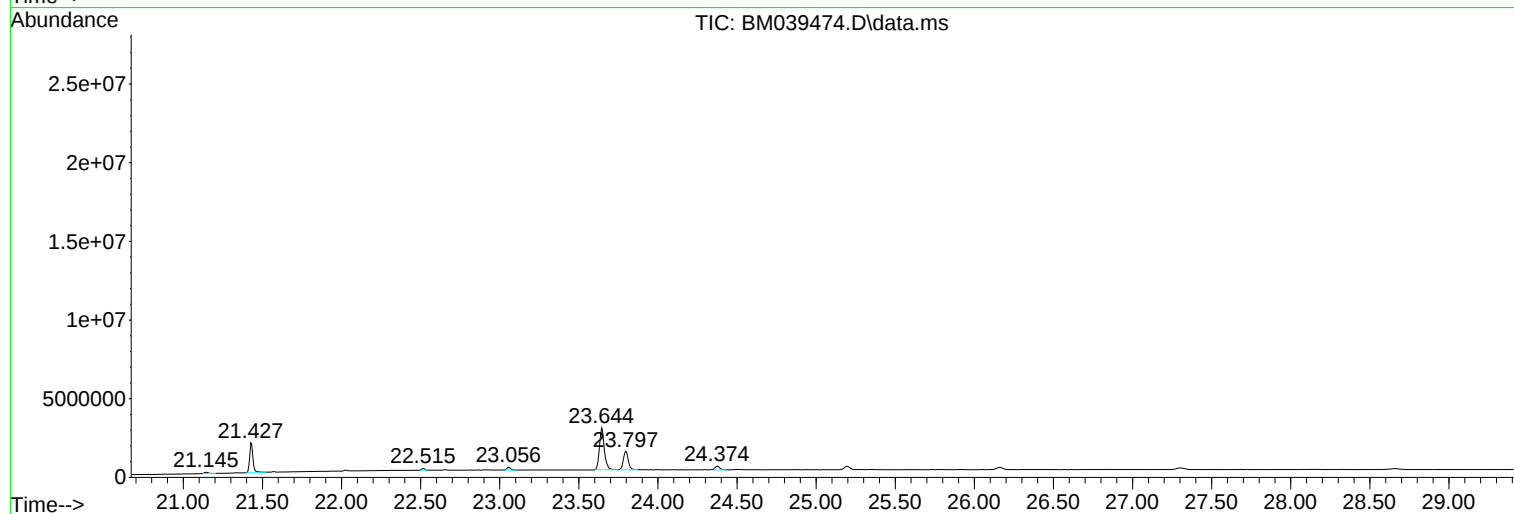
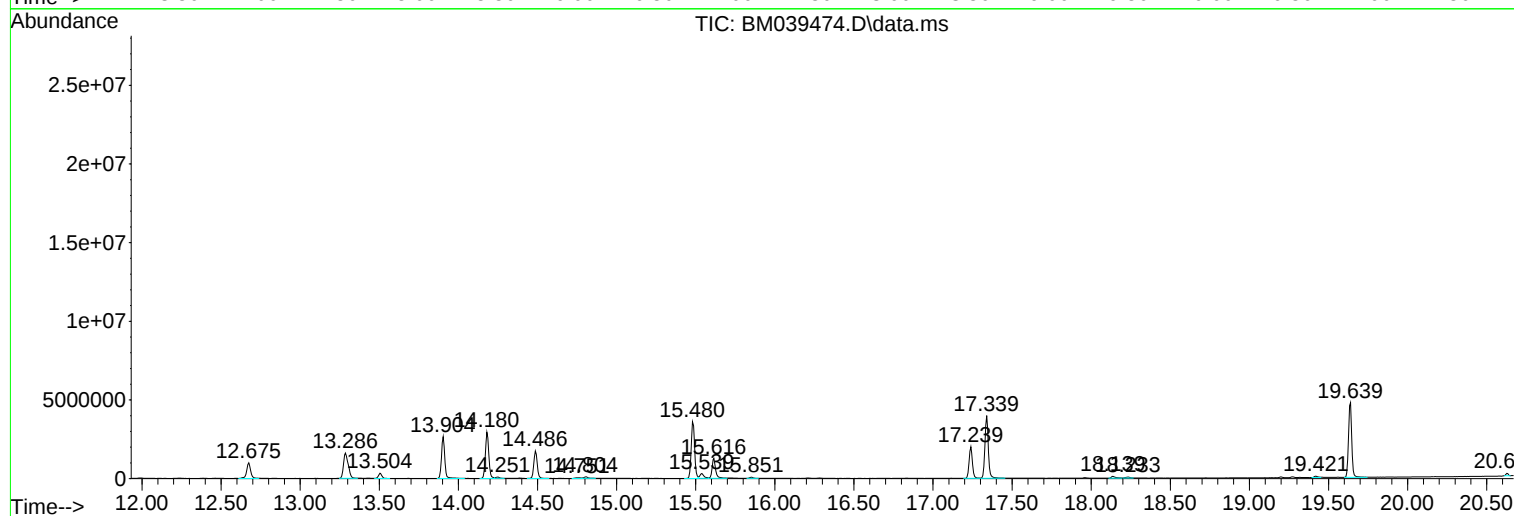
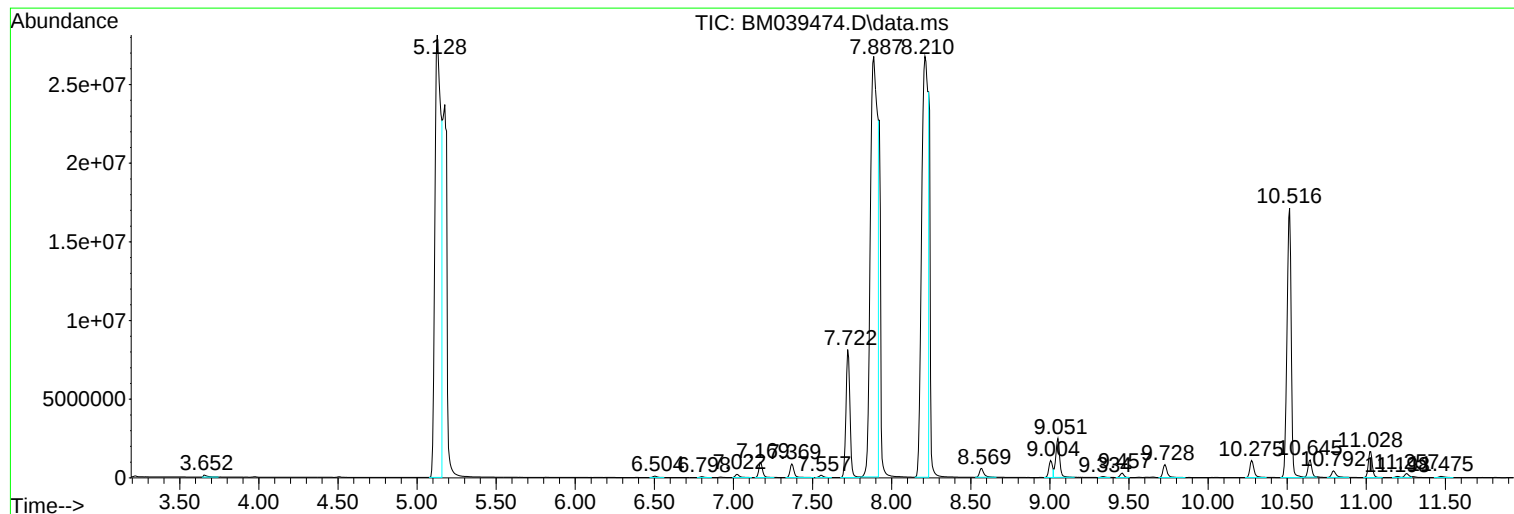
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P



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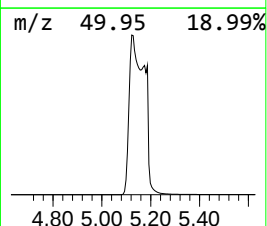
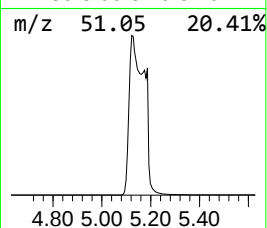
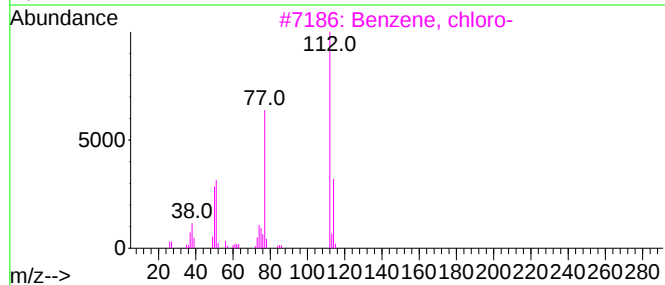
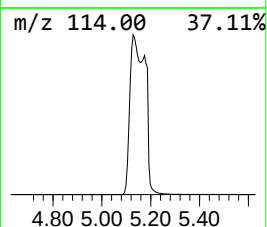
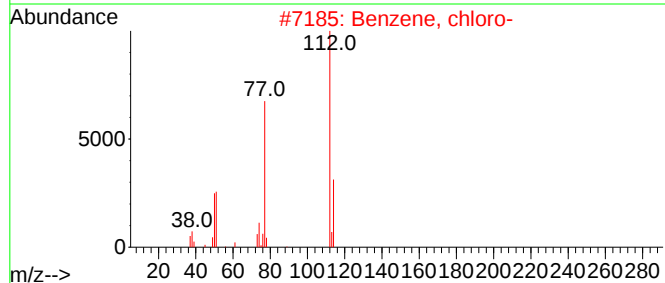
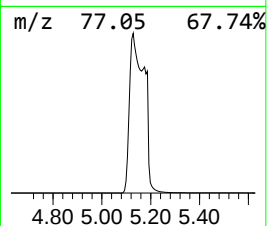
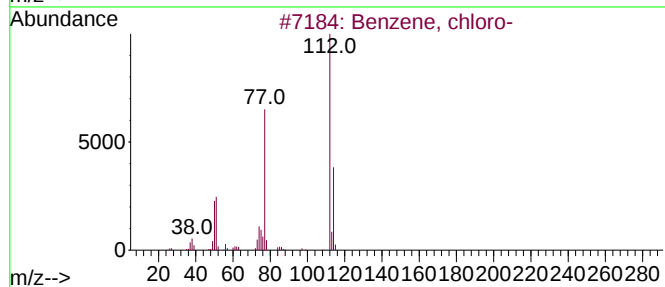
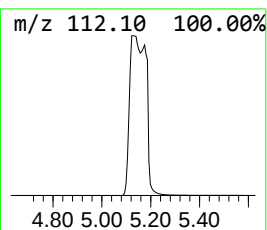
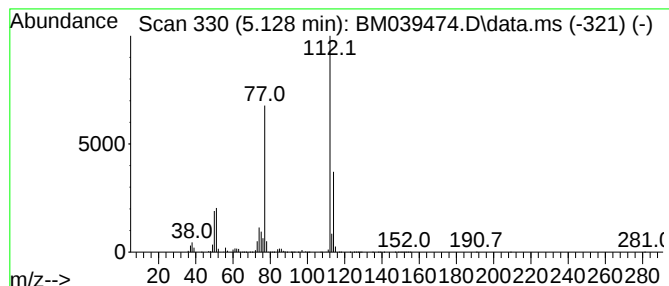
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 1 Benzene, chloro- Concentration Rank 5

R.T.	EstConc	Area	Relative to ISTD	R.T.
5.128	18.01 ng/ul	77981100	1,4-Dichlorobenzene-d4	7.851

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, chloro-	112	C6H5Cl	000108-90-7	96
2			Benzene, chloro-	112	C6H5Cl	000108-90-7	95
3			Benzene, chloro-	112	C6H5Cl	000108-90-7	94
4			Benzene, chloro-	112	C6H5Cl	000108-90-7	94
5			Benzene, chloro-	112	C6H5Cl	000108-90-7	94



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Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

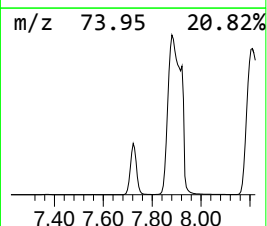
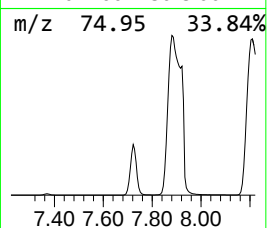
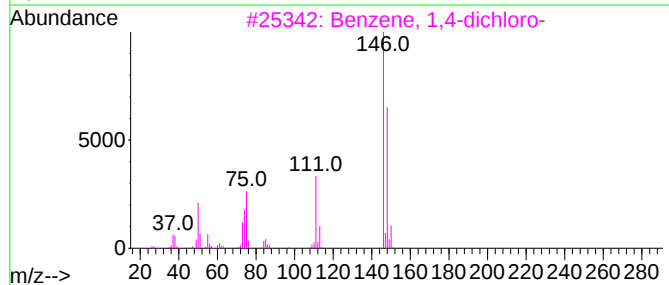
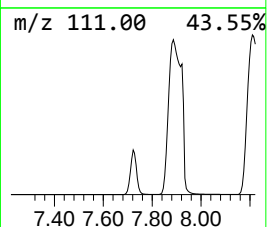
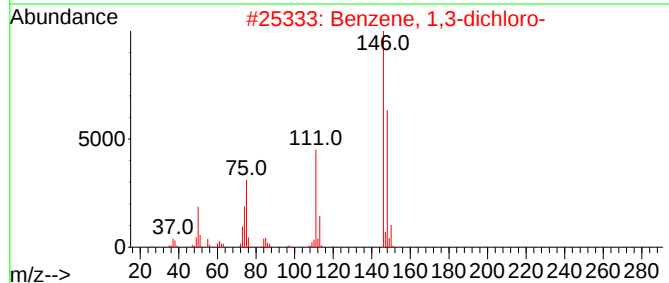
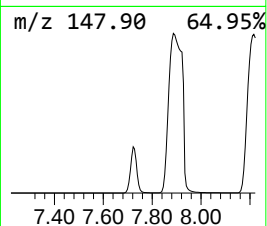
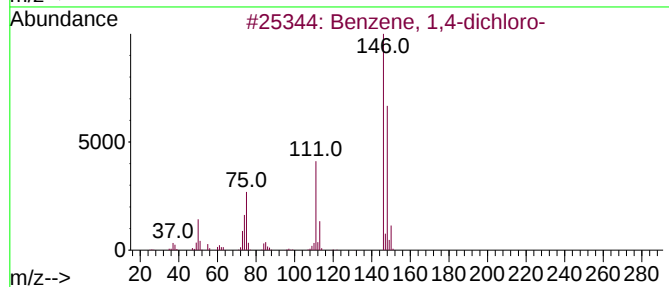
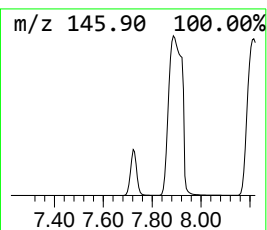
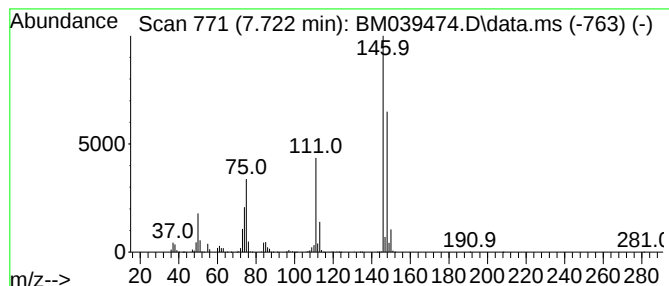
Quant Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 2 Benzene, 1,4-dichloro- Concentration Rank 11

R.T.	EstConc	Area	Relative to ISTD	R.T.
7.722	3.30 ng/ul	14308100	1,4-Dichlorobenzene-d4	7.851

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1,4-dichloro-	146	C6H4Cl2	000106-46-7	98
2			Benzene, 1,3-dichloro-	146	C6H4Cl2	000541-73-1	98
3			Benzene, 1,4-dichloro-	146	C6H4Cl2	000106-46-7	97
4			Benzene, 1,4-dichloro-	146	C6H4Cl2	000106-46-7	96
5			Benzene, 1,2-dichloro-	146	C6H4Cl2	000095-50-1	96



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Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

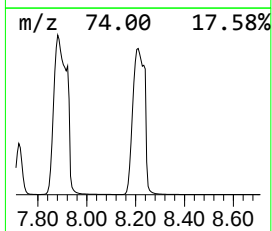
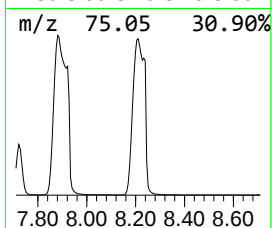
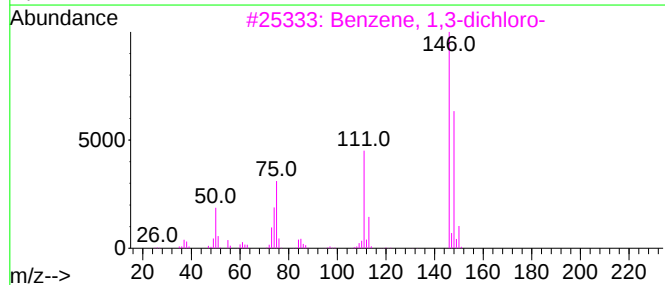
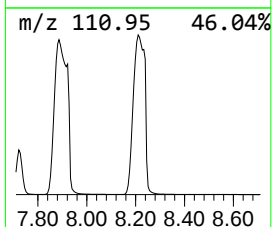
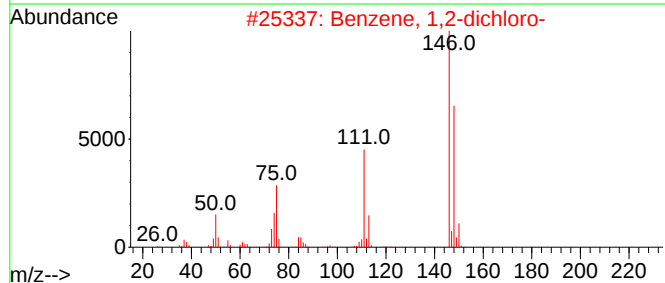
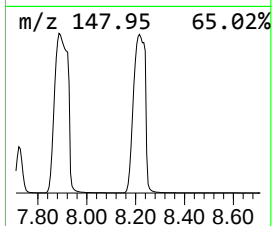
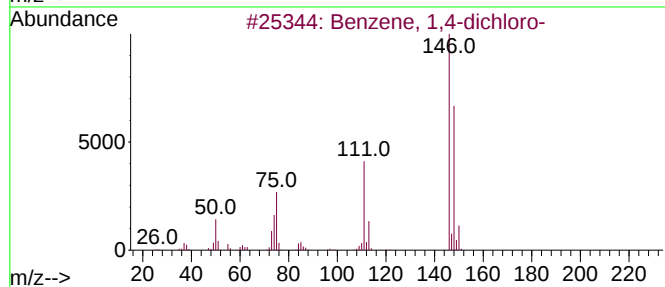
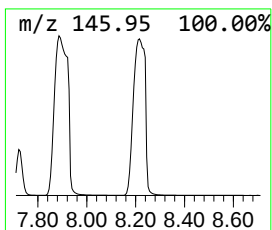
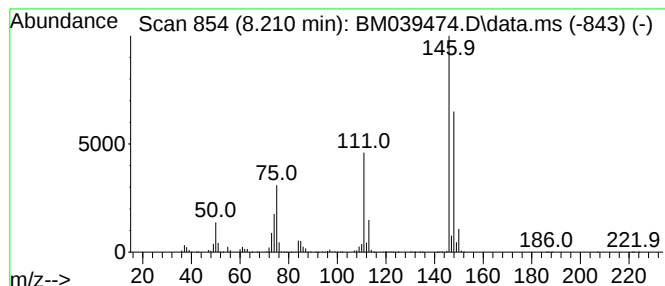
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 3 Benzene, 1,2-dichloro- Concentration Rank 4

R.T.	EstConc	Area	Relative to ISTD	R.T.
8.210	18.35 ng/ul	79457400	1,4-Dichlorobenzene-d4	7.851

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1,4-dichloro-	146	C6H4Cl2	000106-46-7	98
2			Benzene, 1,2-dichloro-	146	C6H4Cl2	000095-50-1	98
3			Benzene, 1,3-dichloro-	146	C6H4Cl2	000541-73-1	97
4			Benzene, 1,2-dichloro-	146	C6H4Cl2	000095-50-1	97
5			Benzene, 1,2-dichloro-	146	C6H4Cl2	000095-50-1	96



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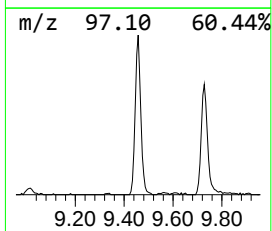
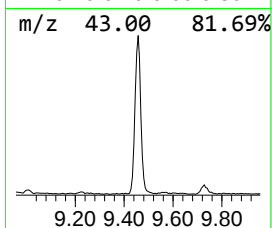
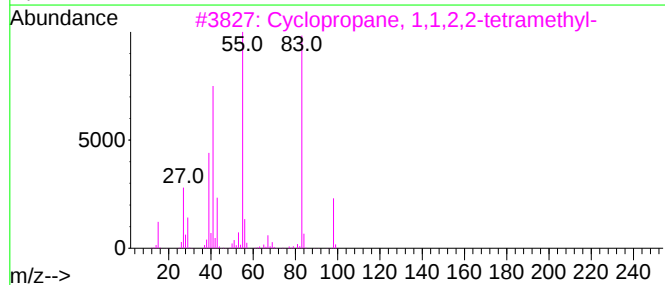
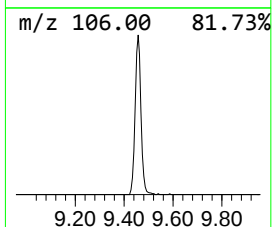
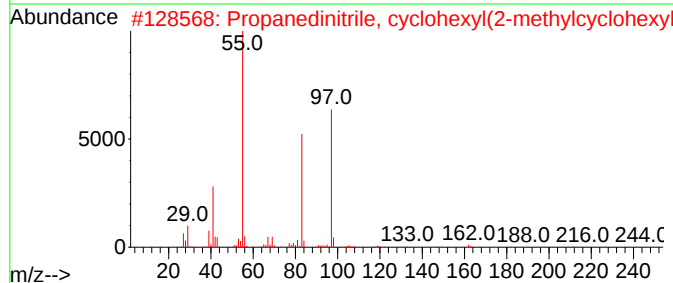
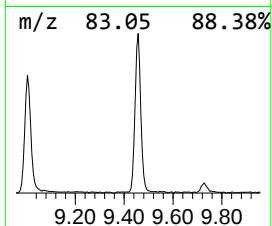
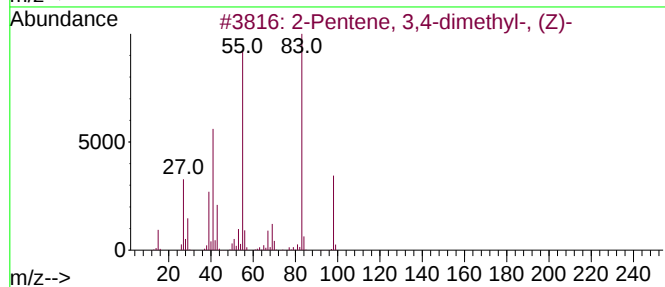
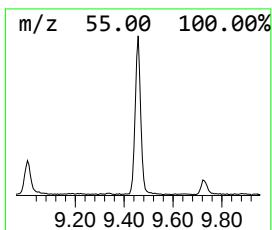
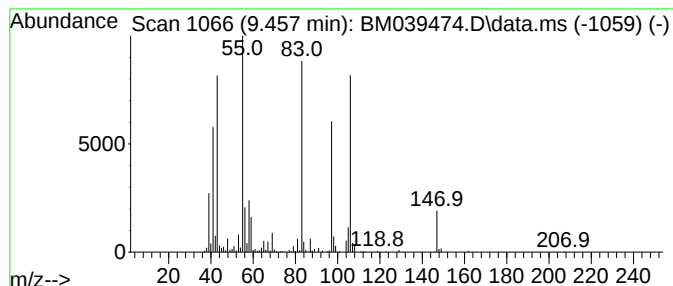
Quant Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
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Peak Number 4 (DEL) Alkane: Straight-Chai... Concentration Rank 6

R.T.	EstConc	Area	Relative to ISTD	R.T.
9.457	5.22 ng/ul	466751	Naphthalene-d8	10.645

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			2-Pentene, 3,4-dimethyl-, (Z)-	98	C7H14	004914-91-4	22
2			Propanedinitrile, cyclohexyl(2-m...	244	C16H24N2	074764-55-9	22
3			Cyclopropane, 1,1,2,2-tetramethyl-	98	C7H14	004127-47-3	22
4			2-Pentene, 4,4-dimethyl-, (E)-	98	C7H14	000690-08-4	22
5			2-Pentene, 2,3-dimethyl-	98	C7H14	010574-37-5	16



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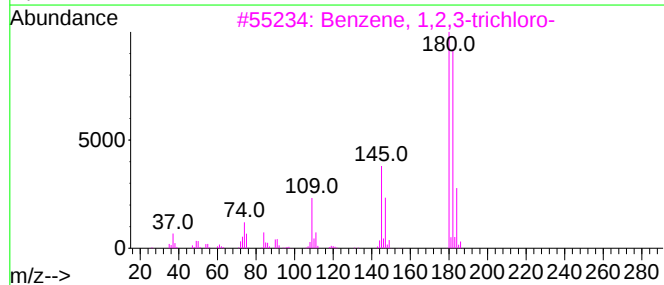
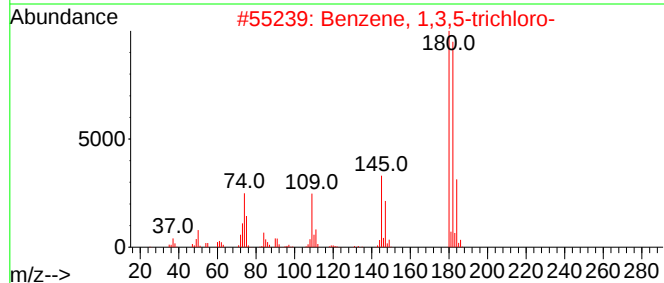
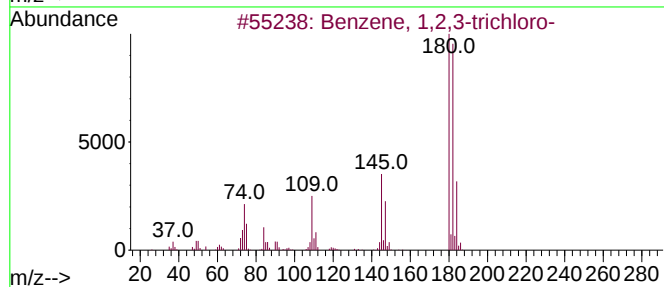
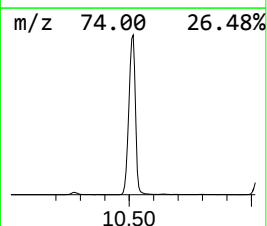
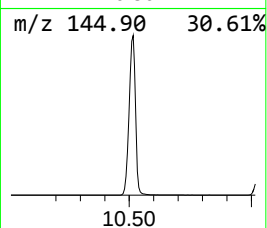
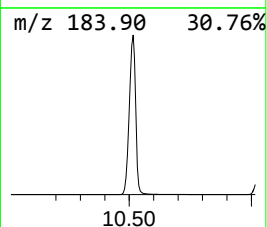
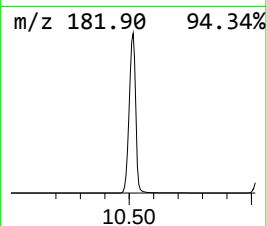
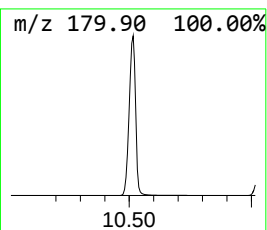
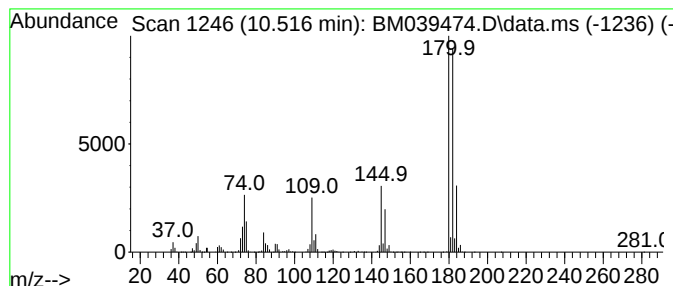
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 5 Benzene, 1,2,3-trichloro- Concentration Rank 1

R.T.	EstConc	Area	Relative to ISTD	R.T.
10.516	354.47 ng/ul	31724700	Naphthalene-d8	10.645

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1,2,3-trichloro-	180	C6H3Cl3	000087-61-6	98
2			Benzene, 1,3,5-trichloro-	180	C6H3Cl3	000108-70-3	98
3			Benzene, 1,2,3-trichloro-	180	C6H3Cl3	000087-61-6	98
4			Benzene, 1,2,3-trichloro-	180	C6H3Cl3	000087-61-6	97
5			Benzene, 1,3,5-trichloro-	180	C6H3Cl3	000108-70-3	97



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

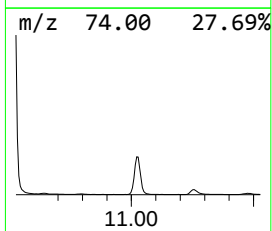
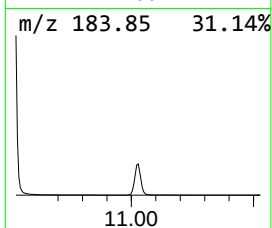
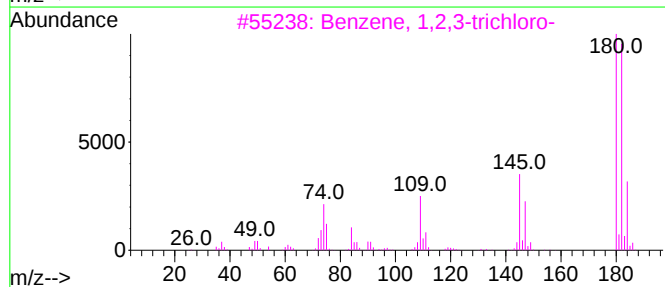
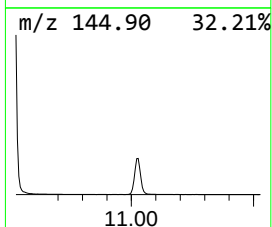
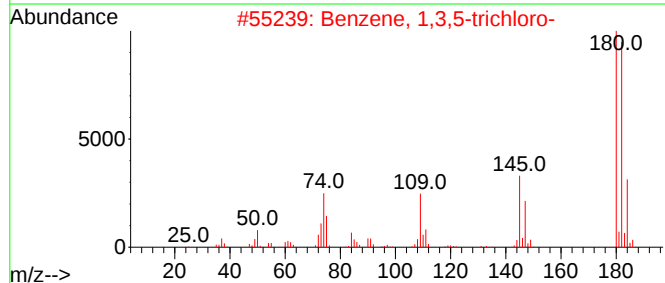
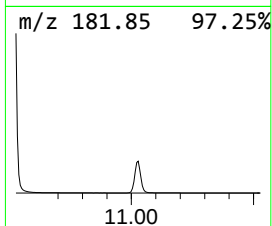
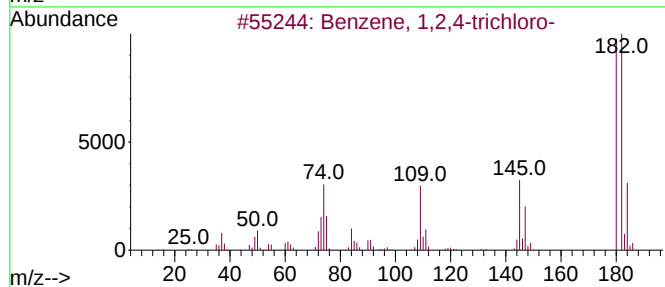
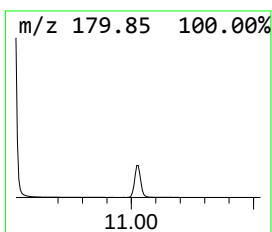
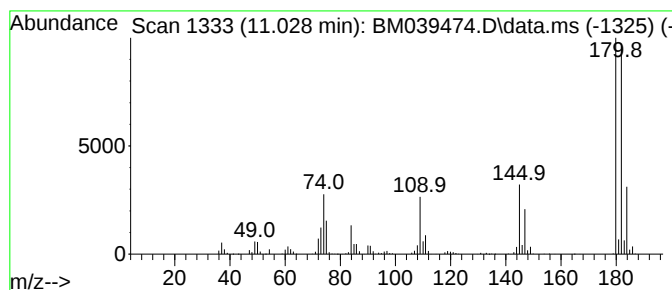
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 6 Benzene, 1,2,4-trichloro- Concentration Rank 2

R.T.	EstConc	Area	Relative to ISTD	R.T.
11.028	32.05 ng/ul	2868400	Naphthalene-d8	10.645

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1,2,4-trichloro-	180	C6H3Cl3	000120-82-1	98
2			Benzene, 1,3,5-trichloro-	180	C6H3Cl3	000108-70-3	98
3			Benzene, 1,2,3-trichloro-	180	C6H3Cl3	000087-61-6	98
4			Benzene, 1,2,3-trichloro-	180	C6H3Cl3	000087-61-6	98
5			Benzene, 1,3,5-trichloro-	180	C6H3Cl3	000108-70-3	96



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

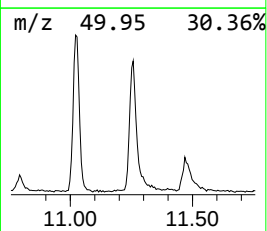
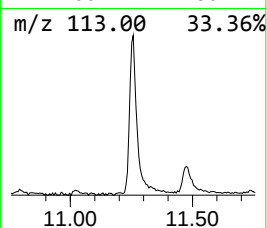
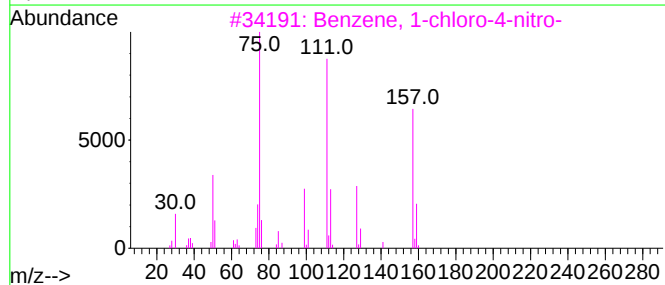
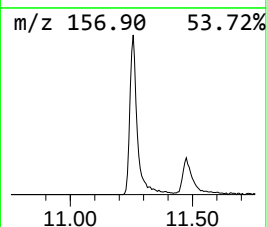
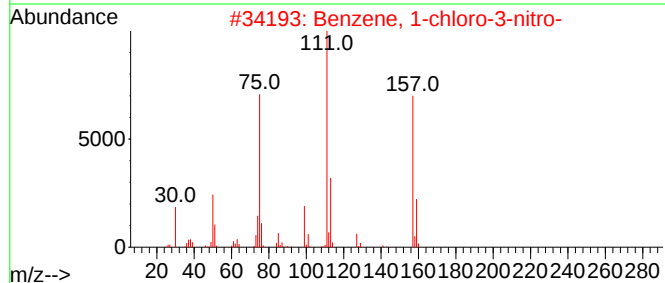
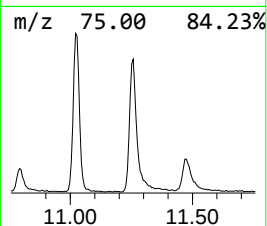
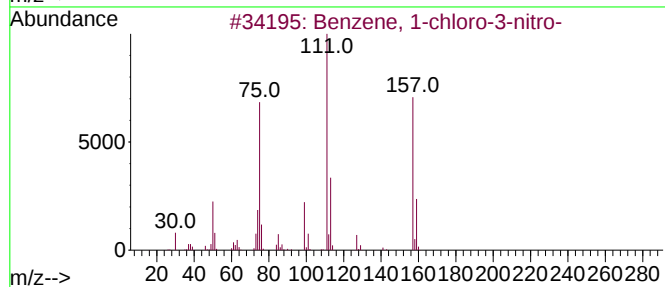
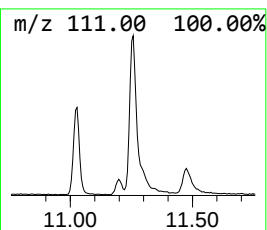
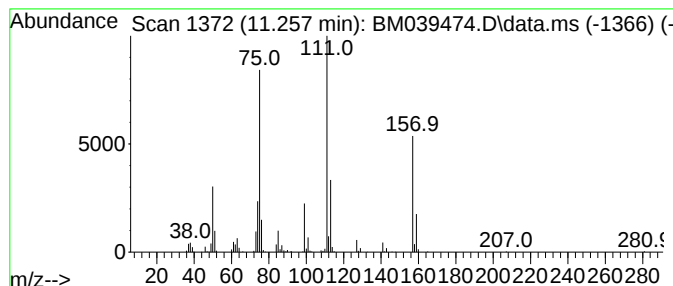
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 7 Benzene, 1-chloro-3-nitro- Concentration Rank 7

R.T.	EstConc	Area	Relative to ISTD	R.T.
11.257	4.91 ng/ul	439303	Naphthalene-d8	10.645

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1-chloro-3-nitro-	157	C6H4ClNO2	000121-73-3	96
2			Benzene, 1-chloro-3-nitro-	157	C6H4ClNO2	000121-73-3	94
3			Benzene, 1-chloro-4-nitro-	157	C6H4ClNO2	000100-00-5	94
4			Benzene, 1-chloro-3-nitro-	157	C6H4ClNO2	000121-73-3	93
5			Benzene, 1-chloro-4-nitro-	157	C6H4ClNO2	000100-00-5	91



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

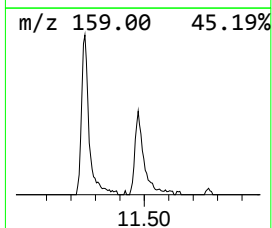
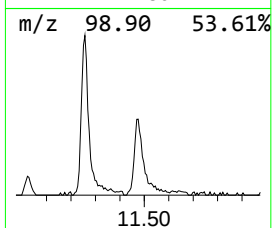
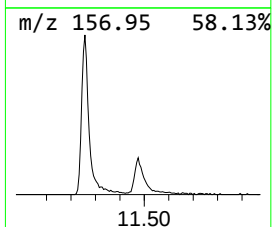
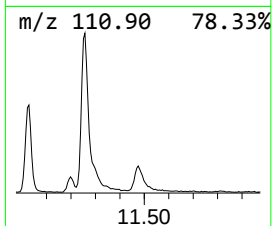
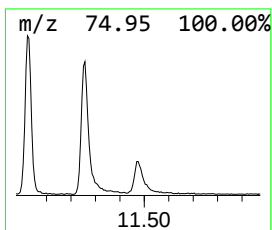
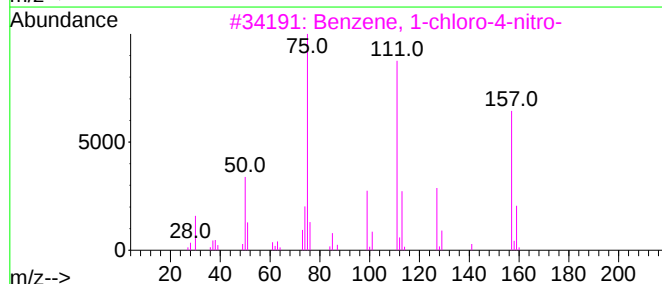
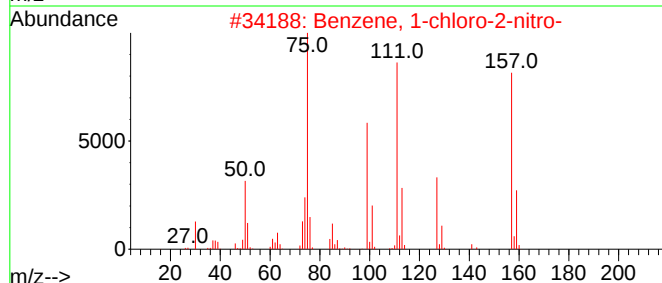
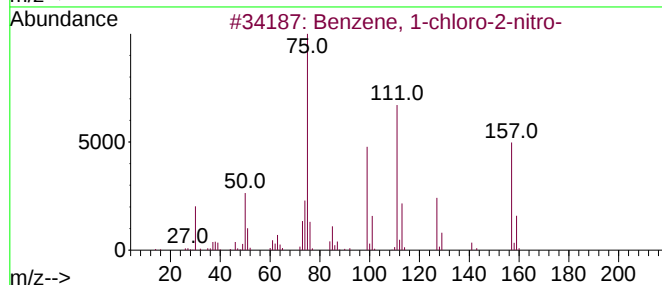
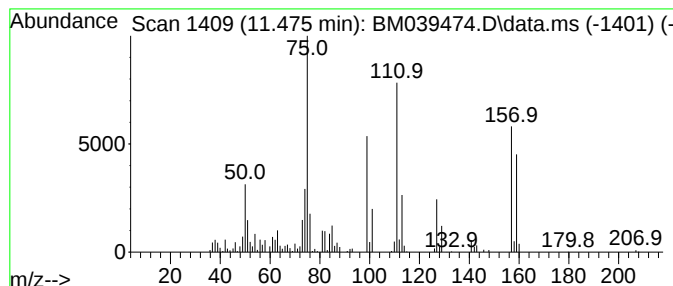
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 8 Benzene, 1-chloro-2-nitro- Concentration Rank 12

R.T.	EstConc	Area	Relative to ISTD	R.T.
11.475	2.24 ng/ul	200926	Naphthalene-d8	10.645

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1-chloro-2-nitro-	157	C6H4ClNO2	000088-73-3	98
2			Benzene, 1-chloro-2-nitro-	157	C6H4ClNO2	000088-73-3	97
3			Benzene, 1-chloro-4-nitro-	157	C6H4ClNO2	000100-00-5	93
4			Benzene, 1-chloro-2-nitro-	157	C6H4ClNO2	000088-73-3	90
5			Benzene, 1-chloro-4-nitro-	157	C6H4ClNO2	000100-00-5	81



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

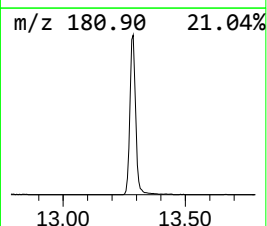
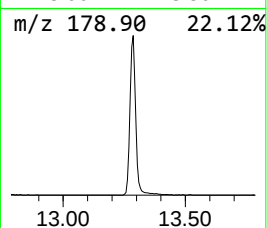
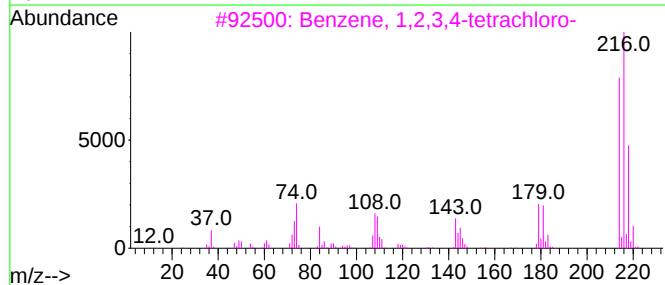
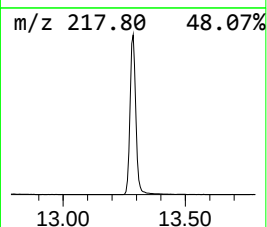
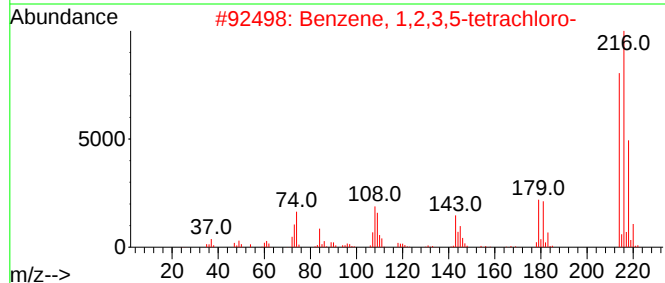
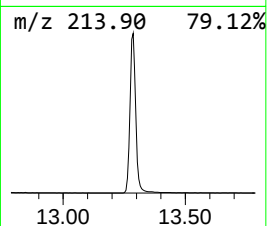
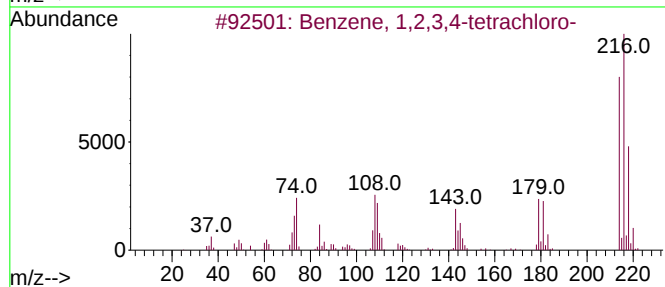
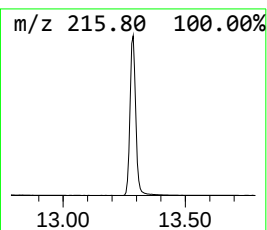
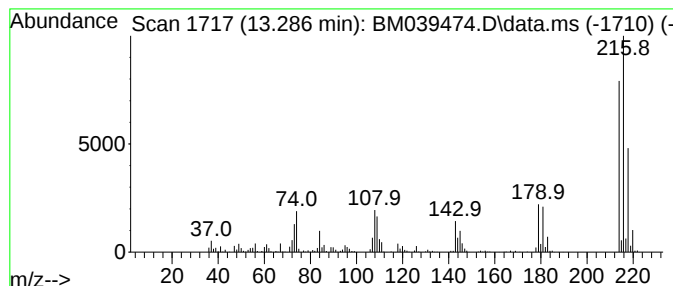
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 9 Benzene, 1,2,3,4-tetrachloro- Concentration Rank 3

R.T.	EstConc	Area	Relative to ISTD	R.T.
13.286	25.57 ng/ul	3338140	Acenaphthene-d10	14.486

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Benzene, 1,2,3,4-tetrachloro-	214	C6H2Cl4	000634-66-2	99
2			Benzene, 1,2,3,5-tetrachloro-	214	C6H2Cl4	000634-90-2	99
3			Benzene, 1,2,3,4-tetrachloro-	214	C6H2Cl4	000634-66-2	99
4			Benzene, 1,2,3,5-tetrachloro-	214	C6H2Cl4	000634-90-2	99
5			Benzene, 1,2,4,5-tetrachloro-	214	C6H2Cl4	000095-94-3	99



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
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ClientSampleId :

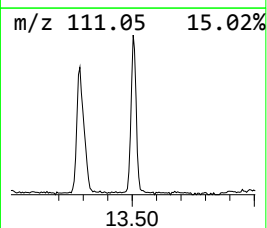
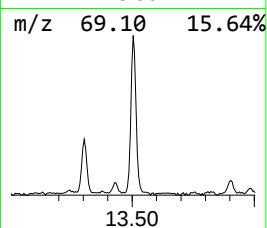
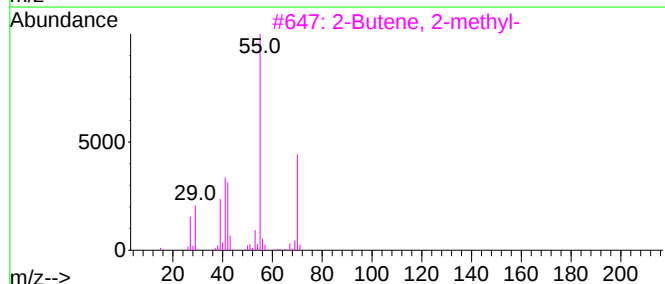
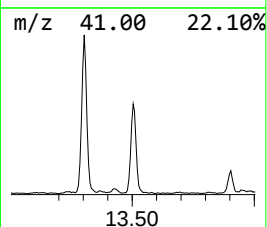
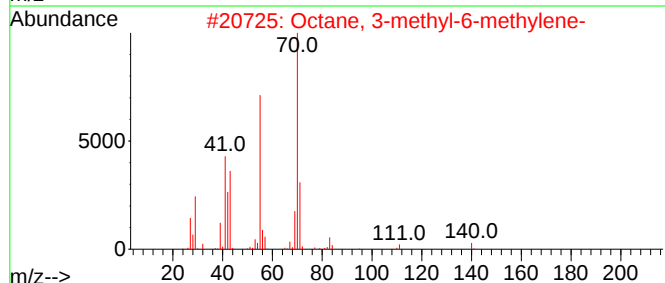
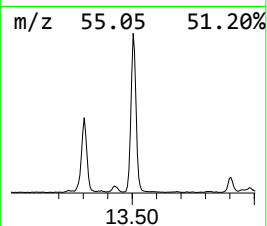
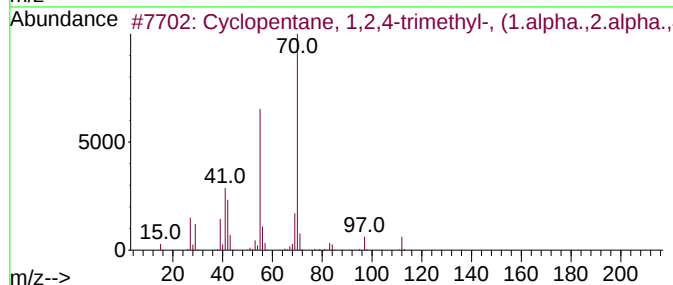
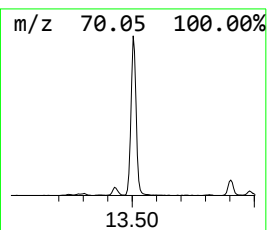
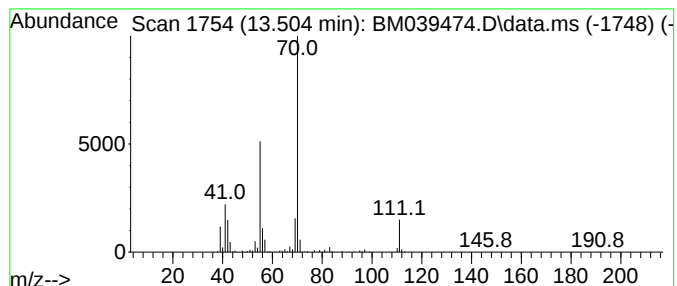
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 10 (DEL) Alkane: Cyclic13.504 Concentration Rank 9

R.T.	EstConc	Area	Relative to ISTD	R.T.
13.504	3.79 ng/ul	494938	Acenaphthene-d10	14.486

Hit#	of	5	Tentative ID	MW	MolForm	CAS#	Qual
1			Cyclopentane, 1,2,4-trimethyl-, ...	112	C8H16	004850-28-6	80
2			Octane, 3-methyl-6-methylene-	140	C10H20	074630-07-2	72
3			2-Butene, 2-methyl-	70	C5H10	000513-35-9	64
4			Heptane, 3-methylene-	112	C8H16	001632-16-2	64
5			Cyclopropane, 1,2-dimethyl-, cis-	70	C5H10	000930-18-7	59



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
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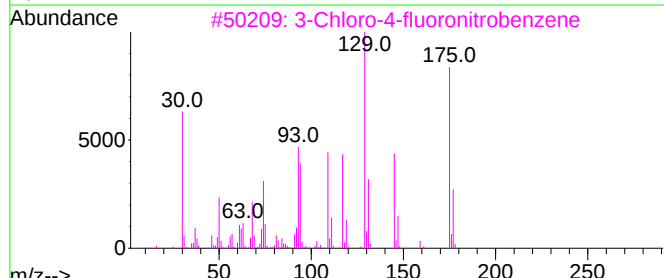
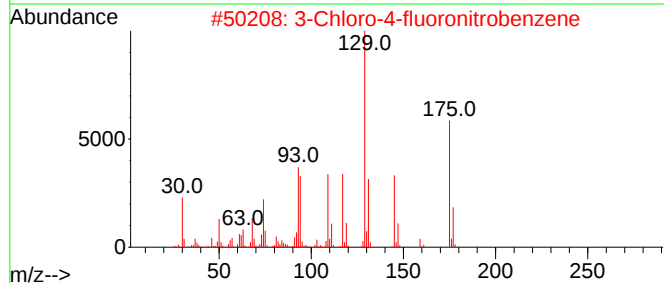
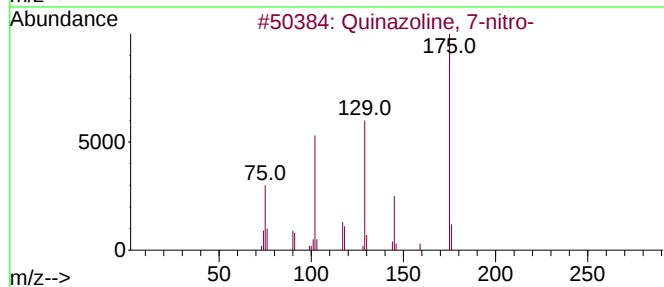
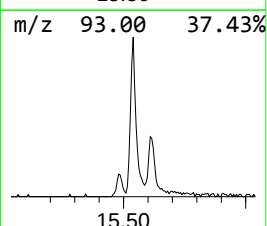
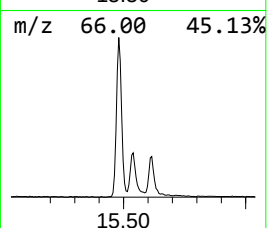
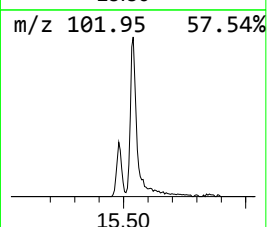
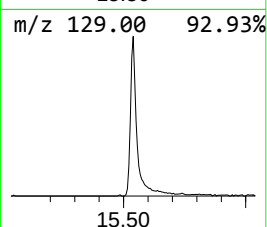
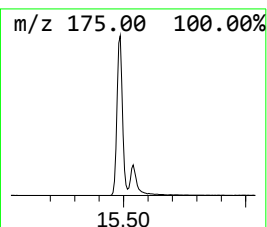
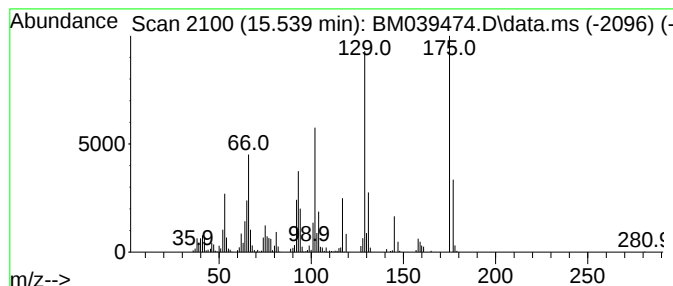
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ClientSampleId :

Quant Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 11 Quinazoline, 7-nitro- Concentration Rank 8

R.T.	EstConc	Area	Relative to ISTD		R.T.	
15.539	4.22 ng/ul	550915	Acenaphthene-d10		14.486	
Hit# of 5	Tentative ID		MW	MolForm	CAS#	Qual
1	Quinazoline, 7-nitro-		175	C8H5N3O2	007557-00-8	50
2	3-Chloro-4-fluoronitrobenzene		175	C6H3ClFN02	000350-30-1	43
3	3-Chloro-4-fluoronitrobenzene		175	C6H3ClFN02	000350-30-1	38
4	3-Chloro-4-fluoronitrobenzene		175	C6H3ClFN02	000350-30-1	38
5	2-Amino-6-chloropyrazine		129	C4H4ClN3	033332-28-4	35



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

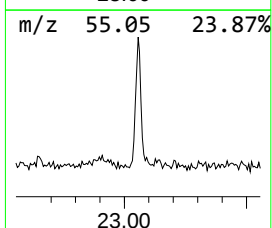
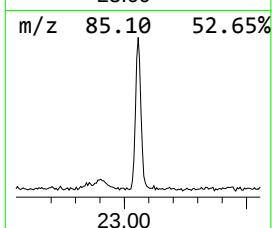
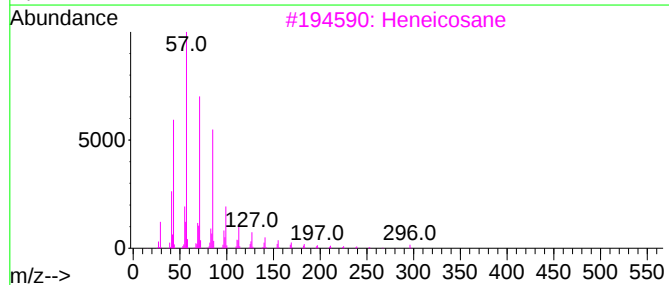
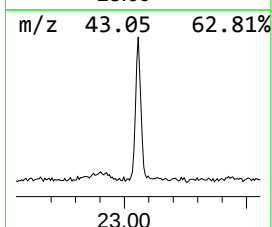
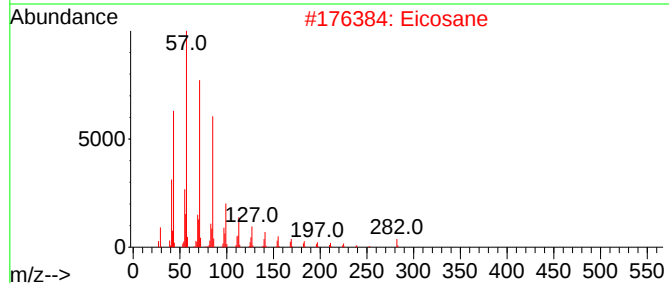
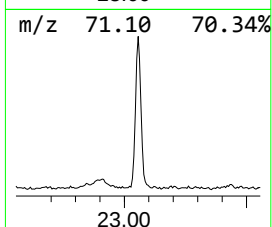
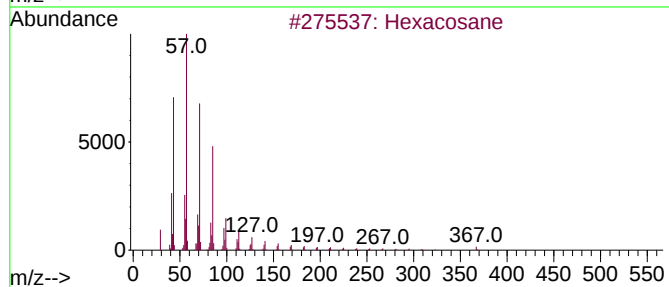
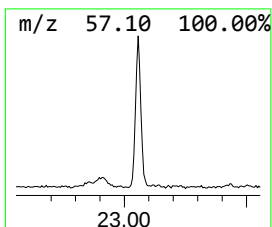
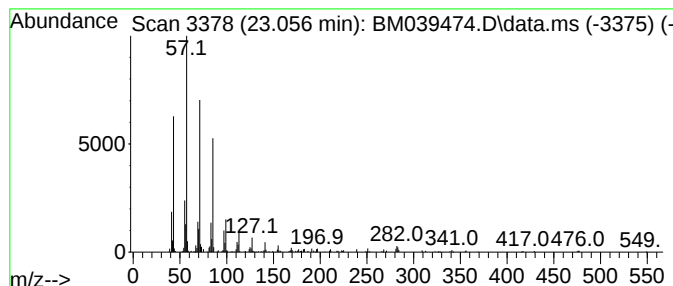
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Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 12 (DEL) Alkane: Straight-Chai... Concentration Rank 13

R.T.	EstConc	Area	Relative to ISTD	R.T.
23.056	2.21 ng/ul	275336	Perylene-d12	23.797

Hit#	of 5	Tentative ID	MW	MolForm	CAS#	Qual
1		Hexacosane	366	C26H54	000630-01-3	93
2		Eicosane	282	C20H42	000112-95-8	93
3		Heneicosane	296	C21H44	000629-94-7	90
4		Nonadecane	268	C19H40	000629-92-5	90
5		Heneicosane	296	C21H44	000629-94-7	87



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

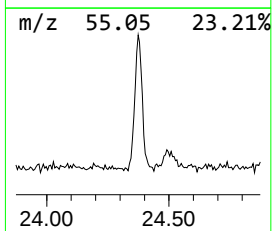
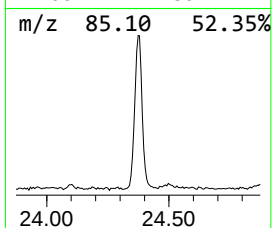
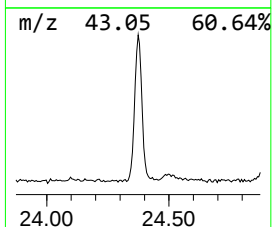
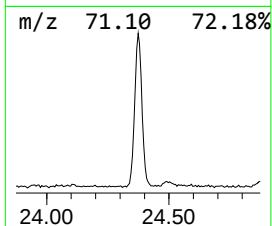
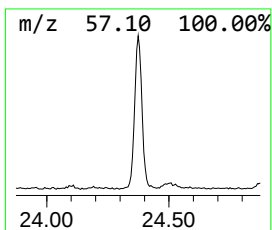
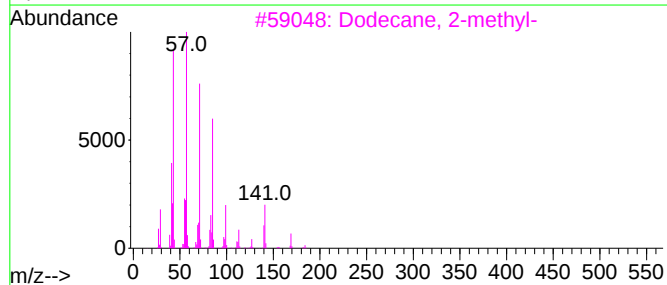
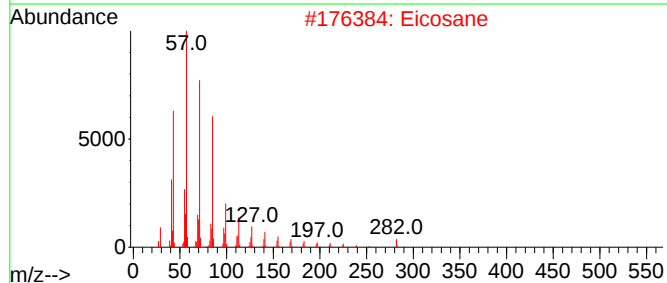
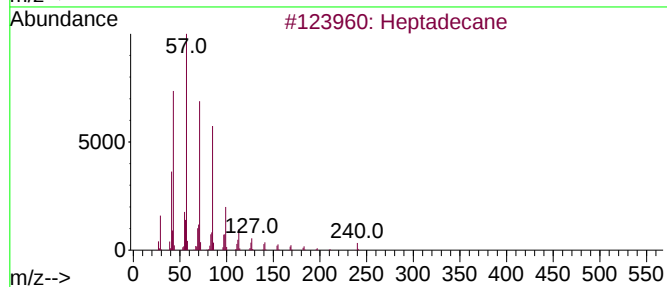
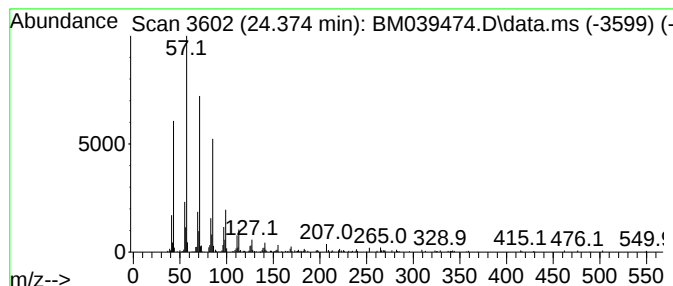
Quant Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST20.L
TIC Integration Parameters: LSCINT.P

Peak Number 13 (DEL) Alkane: Straight-Chai... Concentration Rank 10

R.T.	EstConc	Area	Relative to ISTD	R.T.
24.374	3.54 ng/ul	441718	Perylene-d12	23.797

Hit#	of 5	Tentative ID	MW	MolForm	CAS#	Qual
1		Heptadecane	240	C17H36	000629-78-7	93
2		Eicosane	282	C20H42	000112-95-8	91
3		Dodecane, 2-methyl-	184	C13H28	001560-97-0	90
4		Octacosane, 2-methyl-	408	C29H60	001560-98-1	87
5		Hexatriacontane	507	C36H74	000630-06-8	83



Data Path : Z:\svoasrv\HPCHEM1\BNA_M\Data\BM041823\
Data File : BM039474.D
Acq On : 19 Apr 2023 01:40
Operator : CG/JU
Sample : 02335-18
Misc :
ALS Vial : 25 Sample Multiplier: 1

Instrument :
BNA_M
ClientSampleId :

Quant Method : Z:\svoasrv\HPCHEM1\BNA_M\Methods\SFAM-EPA-BM041823.MA.M
Quant Title : SVOA CALIBRATION

TIC Library : C:\Database\NIST0.L
TIC Integration Parameters: LSCINT.P

TIC Top Hit name	RT	EstConc	Units	Response	--Internal Standard--			
					#	RT	Resp	Conc
Benzene, chloro-	5.128	18.0	ng/ul	77981100	1	7.851	86611100	20.0
Benzene, 1,4-di...	7.722	3.3	ng/ul	14308100	1	7.851	86611100	20.0
Benzene, 1,2-di...	8.210	18.4	ng/ul	79457400	1	7.851	86611100	20.0
(DEL) Alkane: S...	9.457	5.2	ng/ul	466751	2	10.645	1789990	20.0
Benzene, 1,2,3-...	10.516	354.5	ng/ul	31724700	2	10.645	1789990	20.0
Benzene, 1,2,4-...	11.028	32.0	ng/ul	2868400	2	10.645	1789990	20.0
Benzene, 1-chlo...	11.257	4.9	ng/ul	439303	2	10.645	1789990	20.0
Benzene, 1-chlo...	11.475	2.2	ng/ul	200926	2	10.645	1789990	20.0
Benzene, 1,2,3,...	13.286	25.6	ng/ul	3338140	3	14.486	2610910	20.0
(DEL) Alkane: C...	13.504	3.8	ng/ul	494938	3	14.486	2610910	20.0
Quinazoline, 7-...	15.539	4.2	ng/ul	550915	3	14.486	2610910	20.0
(DEL) Alkane: S...	23.056	2.2	ng/ul	275336	6	23.797	2495770	20.0
(DEL) Alkane: S...	24.374	3.5	ng/ul	441718	6	23.797	2495770	20.0