Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV110521\

Data File: VV023239.D

Acq On : 05 Nov 2021 17:57

Operator : SY/MD Sample : M4535-05MSD

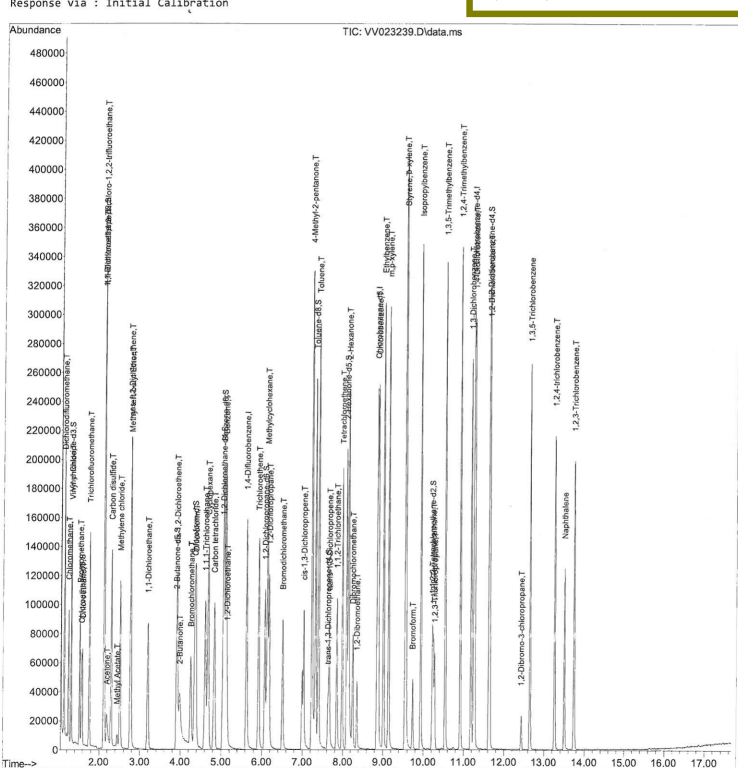
Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 22 Sample Multiplier: 1

Quant Time: Nov 09 03:27:19 2021

Quant Method : Z:\voasrv\HPCHEM1\MSVOA V\Method\SFAMVTR110421WMA.M

Quant Title : TRACE VOA SFAM1.0 QLast Update : Tue Nov 09 02:04:24 2021 Response via : Initial Calibration Instrument :
MSVOA_V
ClientSampleId :

Manual Integrations APPROVED



Quantitation Report (Qedit)

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV110521\

Data File: VV023239.D

Acq On : 05 Nov 2021 17:57

Operator : SY/MD Sample : M4535-05MSD

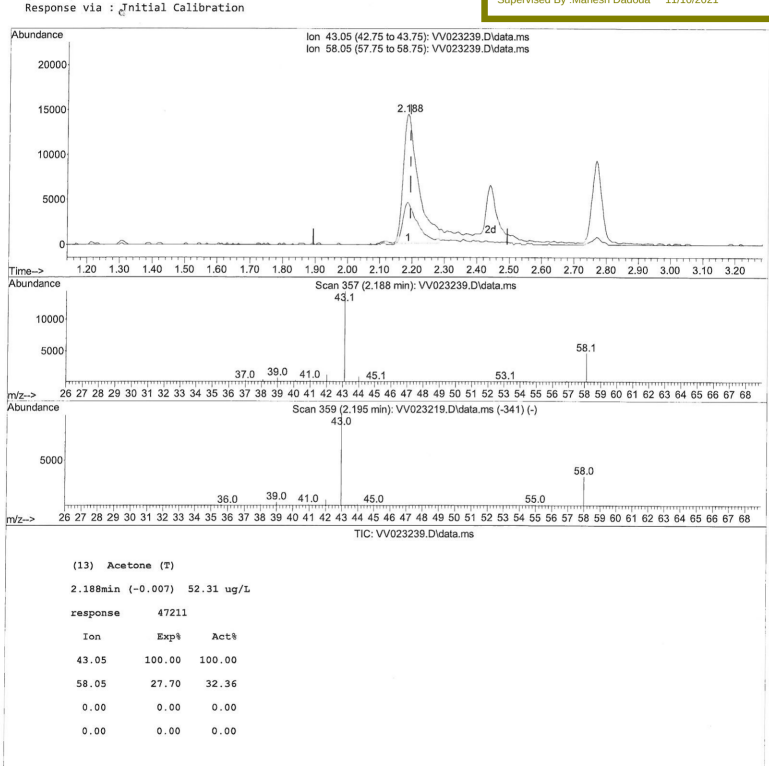
Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 22 Sample Multiplier: 1

Quant Time: Nov 09 03:27:19 2021

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\SFAMVTR110421WMA.M

Quant Title : TRACE VOA SFAM1.0 QLast Update : Tue Nov 09 02:04:24 2021 Response via : Initial Calibration Instrument: MSVOA_V ClientSampleId: H4621MSD

Manual IntegrationsAPPROVED



Quantitation Report (Qedit)

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV110521\

Data File: VV023239.D

Acg On : 05 Nov 2021 17:57

Operator : SY/MD Sample : M4535-05MSD

Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 22 Sample Multiplier: 1

Quant Time: Nov 09 03:27:19 2021

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\SFAMVTR110421WMA.M

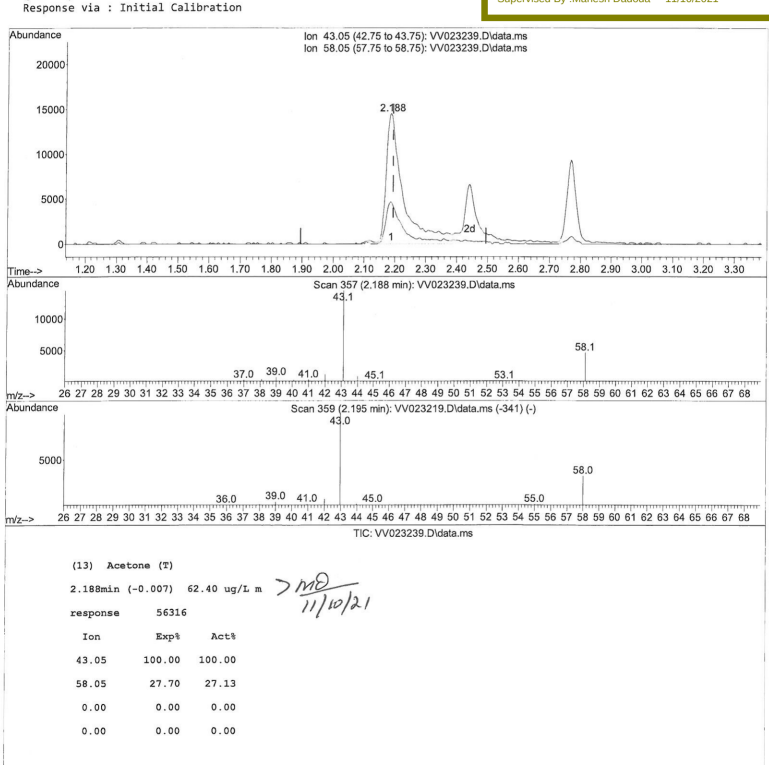
Quant Title : TRACE VOA SFAM1.0

QLast Update : Tue Nov 09 02:04:24 2021

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Operator : SY/MD

Sample : M4535-05MSD
Misc : 25.0mL/MSVOA_V/WATER Misc ALS Vial : 22 Sample Multiplier: 1

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Quant Title : TRACE VOA SFAM1.0 QLast Update : Tue Nov 09 02:04:24 2021 Response via : Initial Calibration

Instrument : MSVOA_V ClientSampleId: H4621MSD

Manual IntegrationsAPPROVED

Compound	R.T.			Conc Units Dev(
Internal Standards						
1) 1,4-Difluorobenzene	5.619	114	136855	5.000 ug/L	0.00	
28) Chlorobenzene-d5	8.853		133507	5.000 ug/L	0.00	
58) 1,4-Dichlorobenzene-d4	11.249	152	73606	5.000 ug/L	0.00	
System Monitoring Compounds						
4) Vinyl Chloride-d3	1.307	65	41164	4.801 ug/L	0.00	
Spiked Amount 5.000	Range 40	- 130	Recove	ry = 96.000%		
7) Chloroethane-d5	1.568	69	35216	5.040 ug/L	0.00	
Spiked Amount 5.000	Range 65	- 130		ry = 100.800%		
11) 1,1-Dichloroethene-d2	2.111	63		4.924 ug/L	0.00	
Spiked Amount 5.000	Range 60			ry = 98.400%		
20) 2-Butanone-d5	3.899		80028	54.181 ug/L	0.00	
Spiked Amount 50.000	Range 40			ry = 108.360% 5.102 ug/L	100100000	
24) Chloroform-d Spiked Amount 5.000	4.352 Range 70		93224 Recover	ry = 102.000%	0.00	
26) 1,2-Dichloroethane-d4	5.037		41629		0.00	
Spiked Amount 5.000	Range 70			ry = 101.400%	0.00	
32) Benzene-d6	5.053	84		5.110 ug/L	0.00	
	Range 70			y = 102.200%		
36) 1,2-Dichloropropane-d6	6.072	67	51943		0.00	
Spiked Amount 5.000	Range 60	- 140	Recover	y = 103.000%		
41) Toluene-d8	7.317	98		5.163 ug/L	0.00	
•	Range 70	- 130		y = 103.200%		
43) trans-1,3-Dichloroprop		79		4.767 ug/L	0.00	
ACCOUNTS OF THE PROPERTY OF TH	Range 55			ry = 95.400%		
46) 2-Hexanone-d5	8.091	63		55.111 ug/L	0.00	
	Range 45			ry = 110.220%	0.00	
56) 1,1,2,2-Tetrachloroeth Spiked Amount 5.000			37699	5.199 ug/L ry = 104.000%	0.00	
66) 1,2-Dichlorobenzene-d4	Range 65 11.625			5.026 ug/L	0.00	
	Range 80			y = 100.600%	0.00	
Sparked / mount	mange or		11000101	, 1001000%		
Target Compounds				Qva:		
2) Dichlorodifluoromethane		85	66437	4.979 ug/L	99	
3) Chloromethane	1.240	50	58681	5.172 ug/L	97	
5) Vinyl chloride6) Bromomethane	1.310 1.523	62 94	58004 36012	5.119 ug/L 4.972 ug/L	100 97	
8) Chloroethane	1.584	64	32910	5.033 ug/L	96	
9) Trichlorofluoromethane	1.754	101	84141	4.942 ug/L	100	
10) 1,1,2-Trichloro-1,2,2		101	43126	5.031 ug/L	99	
12) 1,1-Dichloroethene	2.121	96	41020	5.026 ug/L	88	
13) Acetone	2.188	43	56316m	62.399 ug/L	> MO	
14) Carbon disulfide	2.294	76	151093	4.906 ug/L	100 1//10/21	
15) Methyl Acetate	2.442	43	11575	4.532 ug/L	96 / /	A ₁
16) Methylene chloride	2.506	84	49043	4.118 ug/L	95	
17) Methyl tert-butyl Ether	2.770	73	97128	5.407 ug/L	96	
18) trans-1,2-Dichloroethene	2.760	96	50142	4.998 ug/L	97	
19) 1,1-Dichloroethane	3.191	63	88325	5.214 ug/L	98	
21) 2-Butanone	3.982	43	72628	49.774 ug/L	97	
22) cis-1,2-Dichloroethene	3.915	96 120	51227	5.306 ug/L #	91	
23) Bromochloromethane	4.252	128	22834	5.129 ug/L	83	

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Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\SFAMVTR110421WMA.M

Quant Title : TRACE VOA SFAM1.0 QLast Update : Tue Nov 09 02:04:24 2021 Response via : Initial Calibration

Instrument : MSVOA_V ClientSampleId: H4621MSD

Manual IntegrationsAPPROVED

Compound	R.T.	QIon	Response	Conc Units Dev	(Min)
25) Chloroform	4.378	83	92869	5.144 ug/L	99
27) 1,2-Dichloroethane	5.137	62	48011	4.999 ug/L	99
29) 1,1,1-Trichloroethane	4.609	97	83660	5.159 ug/L	99
30) Cyclohexane	4.680	56	72563	4.994 ug/L	98
31) Carbon tetrachloride	4.828	117	74829	5.138 ug/L	97
33) Benzene	5.101	78	195251	5.232 ug/L	100
34) Trichloroethene	5.915	95	50372	5.076 ug/L	98
35) Methylcyclohexane	6.133	83	76328	4.873 ug/L	96
37) 1,2-Dichloropropane	6.175	63	46637	5.354 ug/L	100
38) Bromodichloromethane	6.513	83	60075	5.146 ug/L	96
39) cis-1,3-Dichloropropene	7.030	75	59331	4.735 ug/L	98
40) 4-Methyl-2-pentanone	7.226	43	231207	57.225 ug/L	99
42) Toluene	7.387	91	216659	5.428 ug/L	98
44) trans-1,3-Dichloropropene	7.654	75	50690	4.876 ug/L	99
45) 1,1,2-Trichloroethane	7.841	97	31799	5.080 ug/L	97
47) Tetrachloroethene	7.979	164	43540	5.063 ug/L	97
48) 2-Hexanone	8.140	43	166275	58.732 ug/L	98
49) Dibromochloromethane	8.246	129	40888	5.155 ug/L	97
50) 1,2-Dibromoethane	8.355	107	30171	5.201 ug/L	94
51) Chlorobenzene	8.882	112	133650	5.038 ug/L	99
52) Ethylbenzene	9.014	91	215996	5.131 ug/L	98
53) m,p-xylene	9.140	106	86333	5.226 ug/L	95
54) o-xylene	9.545	106	81012	5.227 ug/L	98
55) Styrene	9.561	104	137276	5.170 ug/L	97
57) 1,1,2,2-Tetrachloroethane	10.242	83	36072	5.260 ug/L	97
59) Bromoform	9.731	173	22058	5.017 ug/L	98
60) Isopropylbenzene	9.934	105	217992	5.161 ug/L	99
61) 1,2,3-Trichloropropane	10.275	75	25862	5.289 ug/L	98
62) 1,3,5-Trimethylbenzene	10.541	105	176516	5.040 ug/L	99
63) 1,2,4-Trimethylbenzene	10.914	105	181973	5.220 ug/L	99
64) 1,3-Dichlorobenzene	11.181	146	110034	5.098 ug/L	97
65) 1,4-Dichlorobenzene	11.271	146	110242	5.002 ug/L	99
67) 1,2-Dichlorobenzene	11.644	146	99613	5.158 ug/L	99
68) 1,2-Dibromo-3-chloropr	12.429	75	5303	5.091 ug/L	96
69) 1,3,5-Trichlorobenzene	12.648	180	81914	4.848 ug/L	99
70) 1,2,4-trichlorobenzene	13.262	180	64528	4.769 ug/L	99
71) Naphthalene	13.503	128	97752	4.899 ug/L	99
72) 1,2,3-Trichlorobenzene	13.744	180	59151	4.996 ug/L	100

^{(#) =} qualifier out of range (m) = manual integration (+) = signals summed