Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV111021\

Data File : VV023416.D

Acq On : 12 Nov 2021 02:59

Operator : SY/MD Sample : VSTDCCC005EC

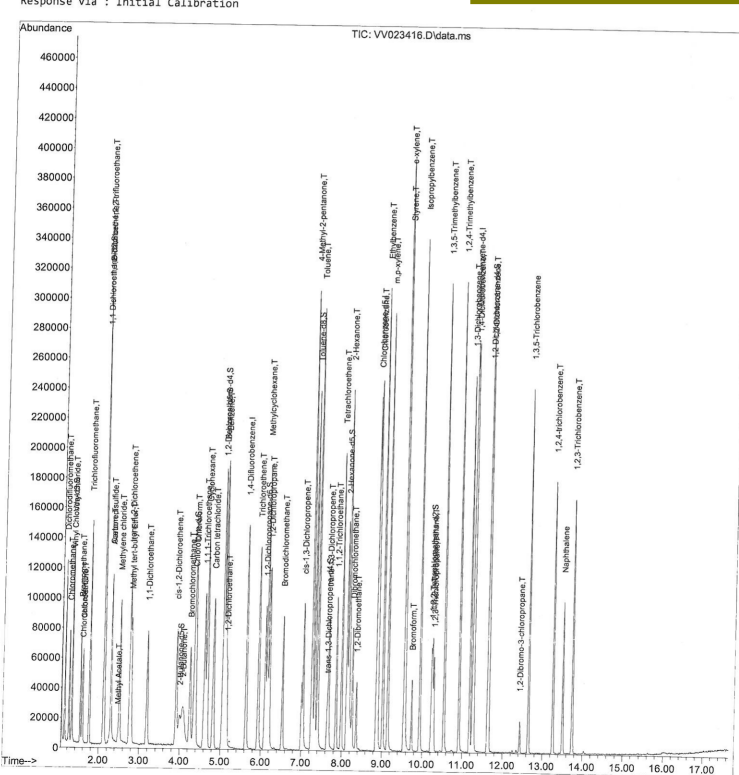
Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 25 Sample Multiplier: 1

Quant Time: Nov 12 04:10:10 2021

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

Quant Title : TRACE VOA SFAM1.0 QLast Update : Fri Nov 12 02:02:21 2021 Response via : Initial Calibration Instrument: MSVOA_V LabSampleId: VSTDCCC005EC

Manual Integrations APPROVED



Quantitation Report (Qedit)

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV111021\

Data File: VV023416.D

Acq On : 12 Nov 2021 02:59

Operator : SY/MD Sample : VSTDCCC005EC

Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 25 Sample Multiplier: 1

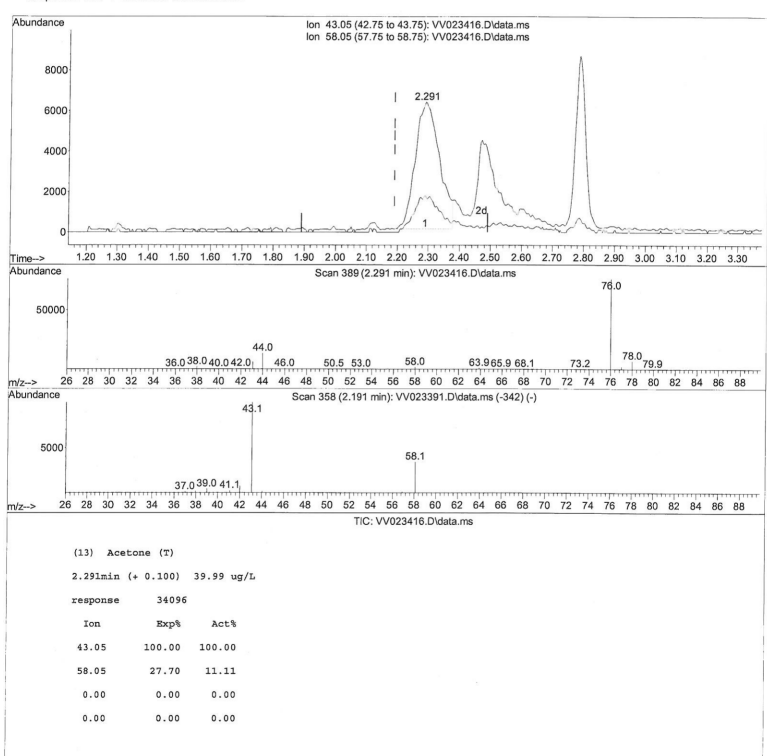
Quant Time: Nov 12 04:10:10 2021

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

Quant Title : TRACE VOA SFAM1.0

QLast Update : Fri Nov 12 02:02:21 2021 Response via : Initial Calibration Instrument : MSVOA_V LabSampleId : VSTDCCC005EC

Manual IntegrationsAPPROVED



Quantitation Report (Qedit)

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Data File: VV023416.D

Acq On : 12 Nov 2021 02:59

Operator : SY/MD

Sample : VSTDCCC005EC

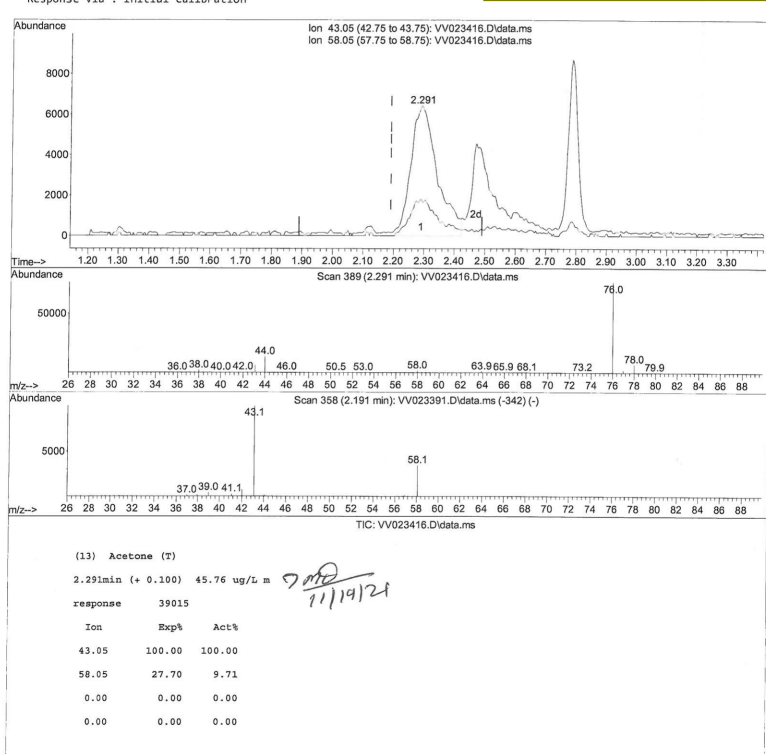
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ALS Vial : 25 Sample Multiplier: 1

Quant Time: Nov 12 04:10:10 2021

Quant Title : TRACE VOA SFAM1.0

QLast Update : Fri Nov 12 02:02:21 2021 Response via : Initial Calibration Instrument:
MSVOA_V
LabSampleId:
VSTDCCC005EC

Manual IntegrationsAPPROVED



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Operator : SY/MD Sample : VSTDCCC005EC

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Quant Title : TRACE VOA SFAM1.0 QLast Update : Fri Nov 12 02:02:21 2021 Response via : Initial Calibration Instrument: MSVOA_V **LabSampleld:** VSTDCCC005EC

Manual IntegrationsAPPROVED

Compound			Response		1.00	53	
Internal Standards							
 1,4-Difluorobenzene 	5.625	114	129283	5.000	ug/L	0.00	
28) Chlorobenzene-d5	8.857	117	126461	5.000	ug/L	0.00	
58) 1,4-Dichlorobenzene-d4	11.249	152	67974	5.000	ug/L	0.00	
System Monitoring Compounds							
4) Vinyl Chloride-d3	1.304	65	34402	4.248	ug/L	0.00	
Spiked Amount 5.000	Range 40	- 130	Recover	ry =	85.000%		
7) Chloroethane-d5	1.574	69	28163	4.267	ug/L	0.00	
Spiked Amount 5.000	Range 65	- 130	Recover		85.400%		
11) 1,1-Dichloroethene-d2	2.111	63	67594	4.458	ug/L	0.00	
Spiked Amount 5.000	Range 60	- 125	Recover	ry =	89.200%		
20) 2-Butanone-d5	3.995	46	58718	42.082	ug/L	0.09	
Spiked Amount 50.000	Range 40	- 130	Recover	'y =	84.160%		
24) Chloroform-d	4.352	84	82743	4.794	ug/L	0.00	
Spiked Amount 5.000	Range 70		Recover	'y =	95.800%		
26) 1,2-Dichloroethane-d4	5.047	65	36162	4.659	ug/L	0.00	
Spiked Amount 5.000	Range 70	- 130	Recover	-	93.200%		
32) Benzene-d6	5.053	84	153025	4.716	ug/L	0.00	
Spiked Amount 5.000	Range 70	- 125	Recover	'y =	94.400%		
36) 1,2-Dichloropropane-d6	6.092	67	44366	4.645	ug/L	0.02	
Spiked Amount 5.000	Range 60	- 140	Recover	·y =	92.800%		
41) Toluene-d8	7.326	98	146313	4.812	ug/L	0.00	
	Range 70		Recover		96.200%		
43) trans-1,3-Dichloroprop		79		4.312	Managara Vincensia Indiana	0.00	
55.00	Range 55		Recover		86.200%		
46) 2-Hexanone-d5	8.108	63	52479	39.382		0.02	
	Range 45		Recover		78.760%		
56) 1,1,2,2-Tetrachloroeth			30938	4.504		0.00	
	Range 65		Recover		90.000%	na cana	
66) 1,2-Dichlorobenzene-d4				4.350		0.00	
Spiked Amount 5.000	Range 80	- 120	Recover	y =	87.000%		
Target Compounds	12 (2022	200			Qval		
2) Dichlorodifluoromethane	1.127	85	61595	4.886		99	
3) Chloromethane	1.240	50	51414	4.797		94	
5) Vinyl chloride	1.310	62	53293	4.979		98	
6) Bromomethane	1.529	94	31060	4.539		100	
8) Chloroethane	1.590	64	30463	4.931		94	
9) Trichlorofluoromethane	1.757	101	85310	5.304		97	
10) 1,1,2-Trichloro-1,2,2		101	38532	4.759		96	
12) 1,1-Dichloroethene	2.121	96	37376	4.848		92	Met
13) Acetone	2.291	43	39015m	45.761		~~	1110/21
14) Carbon disulfide	2.294	76	112299	3.860		99	11/17/01
15) Methyl Acetate	2.471	43	12783	5.298		89	27
16) Methylene chloride	2.510	84	37229	3.309	-	95	
17) Methyl tert-butyl Ether	2.789	73	84098	4.955	-	99	
18) trans-1,2-Dichloroethene	2.754	96	40177	4.239	100	97	
19) 1,1-Dichloroethane	3.188	63	70001	4.375		100	
21) 2-Butanone	4.072	43	71336	51.752	_	92	
22) cis-1,2-Dichloroethene	3.908	96	43011	4.716		95 96	
23) Bromochloromethane	4.249	128	22741	5.407	ng/r #	86	

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\Data\VV111021\

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Sample : VSTDCCC005EC
Misc : 25.0mL/MSVOA_V/WATER ALS Vial : 25 Sample Multiplier: 1

Quant Time: Nov 12 04:10:10 2021

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

Quant Title : TRACE VOA SFAM1.0 QLast Update : Fri Nov 12 02:02:21 2021

Response via : Initial Calibration

Instrument : MSVOA_V **LabSampleld**: VSTDCCC005EC

Manual IntegrationsAPPROVED

Compound	R.T.	QIon	Response	Conc Units Dev	(Min)
25) Chloroform	4.378	83	94503	5.541 ug/L	95
27) 1,2-Dichloroethane	5.146	62	47661	5.253 ug/L	98
29) 1,1,1-Trichloroethane	4.606	. 97	83687	5.449 ug/L	99
30) Cyclohexane	4.667	56	70969	5.157 ug/L	96
31) Carbon tetrachloride	4.825	117	75357	5.463 ug/L	98
33) Benzene	5.101	78	190415	5.387 ug/L	100
34) Trichloroethene	5.928	95	49789	5.297 ug/L	96
35) Methylcyclohexane	6.143	83	72231	4.869 ug/L	97
37) 1,2-Dichloropropane	6.194	63	45287	5.488 ug/L	99
38) Bromodichloromethane	6.529	83	59827	5.410 ug/L	96
39) cis-1,3-Dichloropropene	7.040	75	59650	5.026 ug/L	98
40) 4-Methyl-2-pentanone	7.255	43	210298	54.950 ug/L	98
42) Toluene	7.397	91	209879	5.552 ug/L	98
44) trans-1,3-Dichloropropene	7.664	75	49787	5.056 ug/L	97
45) 1,1,2-Trichloroethane	7.850	97	31424	5.300 ug/L	98
47) Tetrachloroethene	7.982	164	43280	5.313 ug/L	99
48) 2-Hexanone	8.159	43	164492	61.339 ug/L	97
49) Dibromochloromethane	8.255	129	39930	5.315 ug/L	98
50) 1,2-Dibromoethane	8.362	107	29821	5.427 ug/L	100
51) Chlorobenzene	8.886	112	129228	5.143 ug/L	99
52) Ethylbenzene	9.017	91	204643	5.132 ug/L	99
53) m,p-xylene	9.143	106	82129	5.248 ug/L	99
54) o-xylene	9.548	106	76300	5.197 ug/L	99
55) Styrene	9.564	104	135064	5.370 ug/L	97
57) 1,1,2,2-Tetrachloroethane	10.246	83	33766	5.198 ug/L	98
59) Bromoform	9.735	173	22471	5.535 ug/L #	98
60) Isopropylbenzene	9.934	105	206596	5.296 ug/L	99
61) 1,2,3-Trichloropropane	10.278	75	25471	5.641 ug/L	99
62) 1,3,5-Trimethylbenzene	10.542	105	163233	5.047 ug/L	98
63) 1,2,4-Trimethylbenzene	10.915	105	168667	5.240 ug/L	98
64) 1,3-Dichlorobenzene	11.181	146	103350	5.186 ug/L	98
65) 1,4-Dichlorobenzene	11.275	146	102565	5.039 ug/L	99
67) 1,2-Dichlorobenzene	11.644	146	94020	5.272 ug/L	99
68) 1,2-Dibromo-3-chloropr	12.429	75	4965	5.161 ug/L	84
69) 1,3,5-Trichlorobenzene	12.644	180	73740	4.725 ug/L	98
70) 1,2,4-trichlorobenzene	13.262	180	54505	4.362 ug/L	99
71) Naphthalene	13.503	128	81095	4.401 ug/L	99
72) 1,2,3-Trichlorobenzene	13.744	180	51662	4.725 ug/L	98

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