Quantitation Report (Qedit)

Data Path : Z:\voasrv\HPCHEM1\MSVOA_W\Data\VW120821\

Data File : VW021231.D

Acq On : 08 Dec 2021 15:43

Operator : SY/VA

Sample : M4886-12REMS

Misc : 3.25g/10.0mL/MSVOA_W/SOIL
ALS Vial : 1 Sample Multiplier: 1

Quant Time: Dec 08 23:02:42 2021

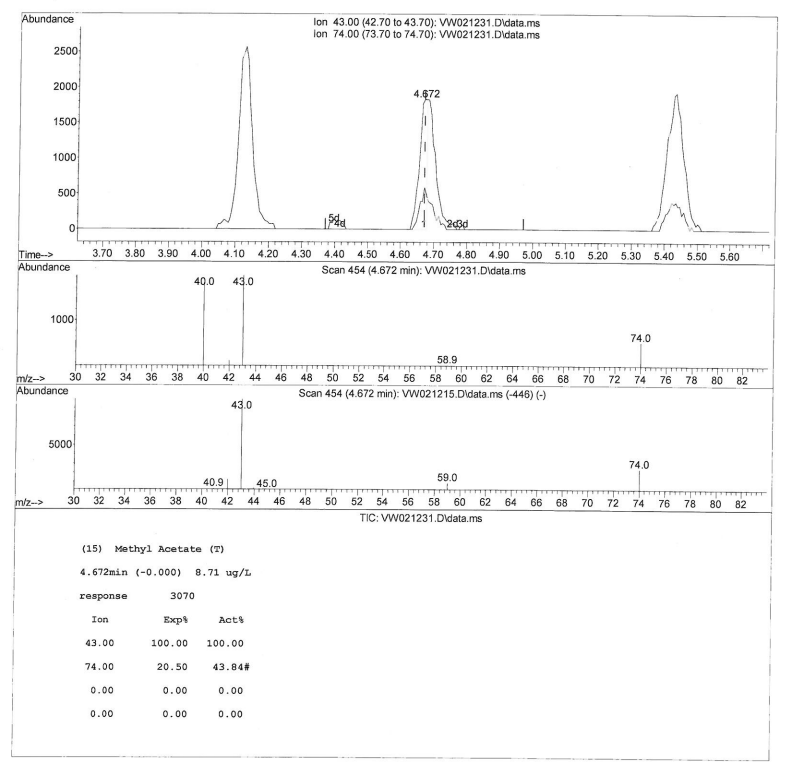
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_W\Method\SFAMWLM120321SMA.M

Quant Title : SFAM01.0

QLast Update: Wed Dec 08 22:58:53 2021 Response via: Initial Calibration



Manual IntegrationsAPPROVED



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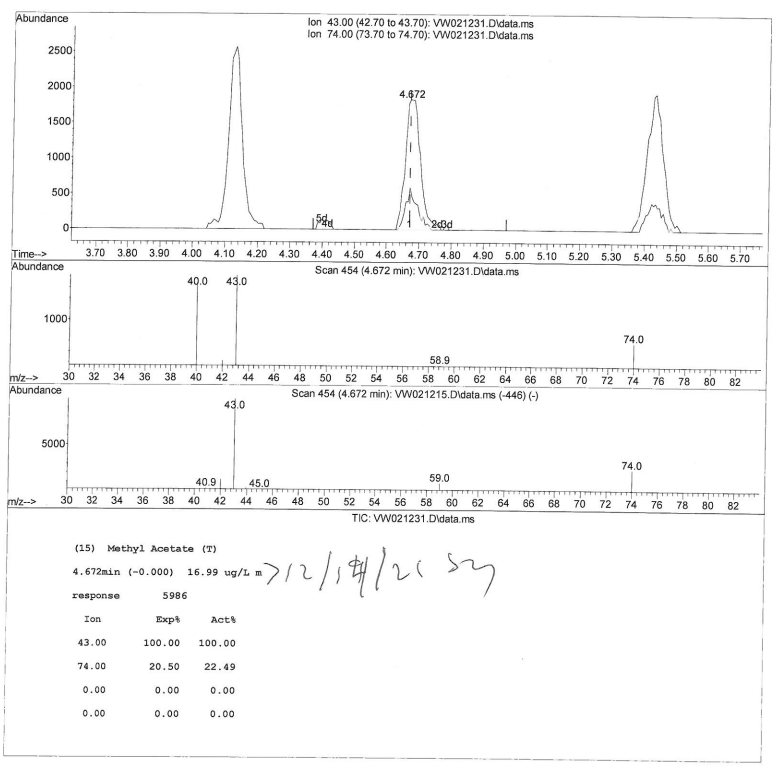
Quant Time: Dec 08 23:02:42 2021

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_W\Method\SFAMWLM120321SMA.M

Quant Title : SFAM01.0

QLast Update : Wed Dec 08 22:58:53 2021 Response via : Initial Calibration Instrument : MSVOA_W ClientSampleld : EX892REMS

Manual Integrations APPROVED



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Quant Title : SFAM01.0

QLast Update : Wed Dec 08 22:58:53 2021 Response via : Initial Calibration

Instrument : MSVOA_W ClientSampleId : EX892REMS

Manual IntegrationsAPPROVED

Compound	R.T	. QIon	Response	Conc Un	its Dev((Min)	
Internal Standards							
1) 1,4-Difluorobenzen	e 8.84	8 114	67134	25.000	ug/L	# 0.00	
28) Chlorobenzene-d5	11.63	4 117	58753	25.000		0.00	
58) 1,4-Dichlorobenzen	e-d4 13.55	4 152	27884	25.000		0.00	
System Monitoring Compo	unds						
4) Vinyl Chloride-d3	2.35	5 65	18832	20.134	ug/L	0.00	
	000 Range 3	0 - 150	Recove		80.520%		
7) Chloroethane-d5	2.88	5 69	15612	24.933	ug/L	0.00	
Spiked Amount 25.	000 Range 3	0 - 150	Recover	ry =	99.720%		
11) 1,1-Dichloroethene			31058	20.785	ug/L	0.00	
Spiked Amount 25.	000 Range 4	5 - 110	Recover	ry =	83.120%		
21) 2-Butanone-d5	7.08	0 46	7535	35.318	ug/L	0.00	
Spiked Amount 50.	000 Range 20	0 - 135	Recover	History and the second of the	70.640%		
24) Chloroform-d	7.64	7 84	33986	20.214	ug/L	0.00	
Spiked Amount 25.			Recover		80.840%		
26) 1,2-Dichloroethane			17146	18.641	100 C	0.00	
Spiked Amount 25.0			Recover		74.560%		
32) Benzene-d6	8.27		62868	20.927		0.00	
Spiked Amount 25.0			Recover		83.720%		
36) 1,2-Dichloropropand			17193	20.595		0.00	
Spiked Amount 25.0 41) Toluene-d8	000 Range 70 10.323		Recover		82.360%	0.00	
Spiked Amount 25.0			59893	19.444		0.00	
43) trans-1,3-Dichlorop			Recover 6226	`y = 16.424	77.760%	0 00	
Spiked Amount 25.6			Recover		65.680%	0.00	
47) 2-Hexanone-d5	10.921		5792	36.262		0.00	
Spiked Amount 50.0			Recover		72.520%	0.00	
56) 1,1,2,2-Tetrachloro			14876	19.261		0.00	
Spiked Amount 25.0			Recover		77.040%	0,00	
66) 1,2-Dichlorobenzene	e-d4 13.847	152	19123	19.141		0.00	
Spiked Amount 25.0	000 Range 75	- 120	Recover		76.560%		
Farget Compounds					Qva1	Lue	
Dichlorodifluoromet	hane 2.007	85	3312	10.100		99	
Chloromethane	2.221	. 50	16514	22.495		100	
Vinyl chloride	2.361	62	26347	20.659		99	
6) Bromomethane	2.782	94	16940	19.977	ug/L	98	
Chloroethane	2.922	64	14027	26.391		95	
Trichlorofluorometh	ane 3.257	101	19706	32.001		99	
10) 1,1,2-Trichloro-1,2	,2 4.068	101	19049	19.763	ug/L	91	
12) 1,1-Dichloroethene	4.044		16408	18.830	ug/L	81	
13) Acetone	4.129		7201	37.053		82	A
14) Carbon disulfide	4.385		37358	15.445		99 - î	11/2/12
15) Methyl Acetate	4.672		5986m	16.987	10.00	(2)	19/21
16) Methylene chloride	4.922		20121	19.024		83	11
17) trans-1,2-Dichloroe			16775	18.080	_	85	
18) Methyl tert-butyl E			30385	21.698	-	94	
19) 1,1-Dichloroethane	6.220		28792	19.570	100 miles	98	
20) cis-1,2-Dichloroeth 22) 2-Butanone	ene 7.171 7.177		18454 8383	18.748		76	
23) Bromochloromethane	7.519		8383	31.170 19.175		89	
23) DI OMOCITTOI OMECHATIE	7.519	120	0031	19.1/3	ug/L #	66	

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Misc : 3.25g/10.0mL/MSVOA_W/SOIL ALS Vial : 1 Sample Multiplier: 1

Quant Time: Dec 08 23:02:42 2021

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_W\Method\SFAMWLM120321SMA.M

Quant Title : SFAM01.0

QLast Update : Wed Dec 08 22:58:53 2021 Response via : Initial Calibration

Instrument : MSVOA_W ClientSampleId : EX892REMS

Manual IntegrationsAPPROVED

Compound	R.T.	QIon	Response	Conc Units Dev	(Min)
25) Chloroform	7.677	7 83	32477	19.665 ug/L	97
27) 1,2-Dichloroethane	8.403	62	20542	19.015 ug/L	96
29) Cyclohexane	7.958	56	25231	19.466 ug/L #	81
30) 1,1,1-Trichloroethane	7.878	97	30214	22.056 ug/L	95
31) Carbon tetrachloride	8.073	117	26780	21.038 ug/L	98
33) Benzene	8.323	78	68473	20.494 ug/L	100
34) Trichloroethene	9.092	95	18558	20.134 ug/L	93
35) Methylcyclohexane	9.335	83	27334	17.618 ug/L #	84
37) 1,2-Dichloropropane	9.372	63	15456	20.468 ug/L #	95
38) Bromodichloromethane	9.646	83	21486	20.241 ug/L	99
39) cis-1,3-Dichloropropene	10.073	75	22732	18.611 ug/L	94
40) 4-Methyl-2-pentanone	10.213	43	18026	35.041 ug/L #	91
42) Toluene	10.390	91	76561	20.062 ug/L	97
44) trans-1,3-Dichloropropene	10.603	75	19056	17.244 ug/L	100
45) 1,1,2-Trichloroethane	10.786	97	12876	20.338 ug/L	90
46) Tetrachloroethene	10.860	164	16941	20.071 ug/L	94
48) 2-Hexanone	10.969	43	13473	36.305 ug/L #	97
49) Dibromochloromethane	11.128	129	14728	18.613 ug/L	97
50) 1,2-Dibromoethane	11.231	107	12146	18.813 ug/L	97
51) Chlorobenzene	11.658	112	47674	18.696 ug/L #	87
52) Ethylbenzene	11.731	91	80093	18.643 ug/L	94
53) m,p-Xylene	11.841	106	33228	18.697 ug/L	95
54) o-Xylene	12.164	106	31871	19.020 ug/L	94
55) Styrene	12.182	104	50442	17.669 ug/L	89
57) 1,1,2,2-Tetrachloroethane	12.713	83	14090	18.503 ug/L #	98
59) Bromoform	12.347	173	7462	18.765 ug/L #	97
60) Isopropylbenzene	12.463	105	85843	22.152 ug/L	96
61) 1,2,3-Trichloropropane	12.768	75	10544	22.573 ug/L #	92
62) 1,3,5-Trimethylbenzene	12.945	105	69789	21.505 ug/L	98
63) 1,2,4-Trimethylbenzene	13.249	105	67245	20.602 ug/L	95
64) 1,3-Dichlorobenzene	13.493	146	34334	18.816 ug/L	92
65) 1,4-Dichlorobenzene	13.572	146	32809	17.791 ug/L	93
67) 1,2-Dichlorobenzene	13.865	146	31794	19.570 ug/L	94
68) 1,2-Dibromo-3-chloropr	14.481	75	1873	17.758 ug/L #	62
69) 1,3,5-Trichlorobenzene	14.627	180	20353	15.177 ug/L	98
70) 1,2,4-trichlorobenzene	15.127	180	13468	12.222 ug/L	98
71) Naphthalene	15.365	128	31375	15.316 ug/L	100
72) 1,2,3-Trichlorobenzene	15.554	180	12459	12.955 ug/L	99

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

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 $\label{lem:quant_Method} \mbox{Quant Method} : \mbox{Z:\normaliant} \mbox{VOA_W\mbox{Method}\scalebox{SFAMWLM120321SMA.M}} \mbox{$M$$.} \mbox{Method} : \mbox{Z:\normaliant} \mbox{$M$$.} \mbox{$M$$.} \mbox{M} \mbox{$M$$.} \mbox{M} \mbox{$M$$.} \mbox{M} \mbox{$M$$.} \mbox{M} \m$

Quant Title : SFAM01.0

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ClientSampleId:
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