

Data Path : Z:\VOASRV\HPCHEM1\MSVOA N\DATA\VN092320\
 Data File : VN063599.D
 Acq On : 23 Sep 2020 19:18
 Operator : JC/MD
 Sample : L4083-05
 Misc : 5.00mL/MSVOA N/WATER
 ALS Vial : 21 Sample Multiplier: 1

Instrument :
 MSVOA_N
 ClientSampleId :
 RE135D1-20200917

Manual Integrations
 APPROVED

MMDadoda
 9/24/2020 1:38:06 PM

Quant Time: Sep 24 04:05:15 2020
 Quant Method : Z:\VOASRV\HPCHEM1\MSVOA_N\METHODS\82N091620W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 16 13:05:20 2020
 Response via : Initial Calibration

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Pentafluorobenzene	7.63	168	204663	50.00	ug/l	0.00
34) 1,4-Difluorobenzene	8.55	114	379893	50.00	ug/l	0.00
63) Chlorobenzene-d5	11.38	117	386449	50.00	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	13.32	152	160345	50.00	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	7.99	65	175507	50.38	ug/l	0.00
Spiked Amount 50.000			Recovery	=	100.76%	
35) Dibromofluoromethane	7.55	113	125821	47.84	ug/l	0.00
Spiked Amount 50.000			Recovery	=	95.68%	
50) Toluene-d8	10.06	98	496639	50.22	ug/l	0.00
Spiked Amount 50.000			Recovery	=	100.44%	
62) 4-Bromofluorobenzene	12.38	95	193392	53.35	ug/l	0.00
Spiked Amount 50.000			Recovery	=	106.70%	
Target Compounds						
3) Chloromethane	2.04	50	1691	0.551	ug/l	96
16) Acetone	3.78	43	3699m	3.304	ug/l	
30) Chloroform	7.33	83	8126	1.592	ug/l	97
44) Trichloroethene	8.81	130	4681m	1.670	ug/l	

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

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