

Data Path : Z:\svoasrv\HPCHEM1\BNA_P\Data\BP123021\
 Data File : BP008618.D
 Acq On : 30 Dec 2021 21:00
 Operator : CG/JU
 Sample : M5244-22
 Misc :
 ALS Vial : 7 Sample Multiplier: 1

Instrument :
 BNA_P
 ClientSampleId :
 CONC-9B

Quant Time: Dec 31 08:15:04 2021
 Quant Method : Z:\svoasrv\HPCHEM1\BNA_P\Methods\8270E-BP122821.M
 Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION
 QLast Update : Thu Dec 30 08:44:33 2021
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) 1,4-Dichlorobenzene-d4	7.893	152	807618	20.000	ng	0.00
21) Naphthalene-d8	10.693	136	3266401	20.000	ng	0.00
39) Acenaphthene-d10	14.522	164	2275153	20.000	ng	0.00
64) Phenanthrene-d10	17.269	188	4924586	20.000	ng	0.00
76) Chrysene-d12	21.351	240	4883438	20.000	ng	-0.01
86) Perylene-d12	23.774	264	4902391	20.000	ng	-0.01
System Monitoring Compounds						
5) 2-Fluorophenol	5.469	112	3468199	78.384	ng	0.00
7) Phenol-d6	7.057	99	5022340	81.549	ng	0.00
23) Nitrobenzene-d5	9.051	82	3039084	55.349	ng	0.00
42) 2,4,6-Tribromophenol	16.010	330	2038647	66.229	ng	0.00
45) 2-Fluorobiphenyl	13.151	172	8952890	57.012	ng	0.00
79) Terphenyl-d14	19.827	244	12528672	52.383	ng	0.00
Target Compounds						
						Qvalue
9) Benzaldehyde	7.028	77	374603	10.687	ng	99
15) Benzyl Alcohol	8.163	79	14457898	327.742	ng	99
84) Bis(2-ethylhexyl)phtha...	21.268	149	1283020	6.703	ng	98

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

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