

Data Path : Z:\SVOASRV\HPCHEM1\BNA G\DATA\BG040820\
 Data File : BG044975.D
 Acq On : 8 Apr 2020 13:59
 Operator : CG/JU
 Sample : L2129-11DL 5X
 Misc :
 ALS Vial : 5 Sample Multiplier: 1

Instrument :
 BNA_G
 ClientSampleId :

Quant Time: Apr 08 14:38:59 2020
 Quant Method : Z:\SVOASRV\HPCHEM1\BNA G\METHODS\8270-BG031020.M
 Quant Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION
 QLast Update : Wed Apr 08 13:01:06 2020
 Response via : Initial Calibration

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	8.04	152	99882	20.00	ng	0.00
21) Naphthalene-d8	10.84	136	407552	20.00	ng	0.00
39) Acenaphthene-d10	14.67	164	280080	20.00	ng	0.00
64) Phenanthrene-d10	17.42	188	547314	20.00	ng	0.00
76) Chrysene-d12	21.69	240	452592	20.00	ng	0.00
87) Perylene-d12	24.90	264	503494	20.00	ng	0.00

System Monitoring Compounds

5) 2-Fluorophenol	5.61	112	203350	31.69	ng	0.00
7) Phenol-d6	7.20	99	305353	32.75	ng	-0.01
23) Nitrobenzene-d5	9.19	82	199545	21.49	ng	0.00
42) 2,4,6-Tribromophenol	16.16	330	118172	32.22	ng	0.00
45) 2-Fluorobiphenyl	13.29	172	456373	23.33	ng	0.00
79) Terphenyl-d14	20.03	244	512710	21.19	ng	0.00

Target Compounds

						Qvalue
9) Benzaldehyde	7.18	77	1307	Below Cal		81
19) n-Nitroso-di-n-propylamine	9.19	70	25917	3.937 ng	#	67
26) 2-Nitrophenol	9.75	139	89	5.539 ng	#	62
32) Benzoic acid	10.25	122	75	9.683 ng	#	38
50) Dimethylphthalate	14.12	163	53673	2.406 ng		99
51) 2,6-Dinitrotoluene	14.67	165	41499	9.229 ng	#	22
52) Acenaphthene	14.73	154	81098	4.623 ng		100
54) 2,4-Dinitrophenol	14.82	184	60	8.346 ng	#	1
55) Dibenzofuran	15.07	168	116127	4.565 ng	#	95
56) 4-Nitrophenol	15.07	139	42796	10.847 ng	#	2
58) Fluorene	15.72	166	97302	4.650 ng		96
65) 4,6-Dinitro-2-methylphenol	15.86	198	62	7.748 ng	#	1
71) Phenanthrene	17.46	178	1602309	54.784 ng		98
72) Anthracene	17.55	178	311329	10.795 ng		97
73) Carbazole	17.81	167	179990	6.497 ng		98
75) Fluoranthene	19.47	202	1651834	47.434 ng		98
78) Pyrene	19.83	202	1381084	41.592 ng		98
81) Benzo(a)anthracene	21.67	228	820980	25.191 ng		99
83) Chrysene	21.74	228	779800	25.049 ng		97
86) Indeno(1,2,3-cd)pyrene	28.53	276	354643	8.689 ng	#	96
88) Benzo(b)fluoranthene	23.89	252	870710	26.195 ng		99
89) Benzo(k)fluoranthene	23.89	252	870710	27.680 ng		99
90) Benzo(a)pyrene	24.74	252	600342	19.302 ng		98
92) Benzo(g,h,i)perylene	29.67	276	308952	9.966 ng		96

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