

Data Path : Z:\HPCHEM1\BNA_M\DATA\BM052216\
 Data File : BM005594.D
 Acq On : 22 May 2016 19:38
 Operator : UM/SJ
 Sample : H3087-08
 Misc :
 ALS Vial : 13 Sample Multiplier: 1

Instrument :
 BNA_M
 ClientSampleId :
 JHGK3

Manual Integrations
 APPROVED

umangi
 5/23/2016 8:35:45 PM

Quant Time: May 23 06:57:47 2016
 Quant Method : Z:\HPCHEM1\BNA_M\METHODS\SOM02.2-EPA-BM052016.M
 Quant Title : SVOA CALIBRATION
 QLast Update : Mon May 23 05:32:02 2016
 Response via : Initial Calibration

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) 1,4-Dichlorobenzene-d4	7.81	152	41921	20.00	ng/ul	0.00
18) Naphthalene-d8	10.60	136	207966	20.00	ng/ul	0.00
35) Acenaphthene-d10	14.44	164	132618	20.00	ng/ul	0.00
61) Phenanthrene-d10	17.19	188	338045	20.00	ng/ul	0.00
75) Chrysene-d12	21.36	240	494631	20.00	ng/ul	0.00
83) Perylene-d12	23.64	264	580166	20.00	ng/ul	0.00

System Monitoring Compounds

3) 1,4-Dioxane-d8	3.29	96	3983	4.16	ng/uL	0.00
5) Phenol-d5	6.98	99	69264	20.16	ng/ul	0.00
7) Bis-(2-Chloroethyl)ether-d	7.15	67	41321	20.75	ng/ul	0.00
9) 2-Chlorophenol-d4	7.35	132	55255	19.27	ng/ul	0.00
13) 4-Methylphenol-d8	8.52	113	54371	19.31	ng/ul	0.00
19) Nitrobenzene-d5	8.96	128	29372	18.47	ng/ul	0.00
22) 2-Nitrophenol-d4	9.69	143	32898	18.80	ng/ul	0.00
26) 2,4-Dichlorophenol-d3	10.22	165	62075	18.83	ng/ul	0.00
29) 4-Chloroaniline-d4	10.74	131	74738	22.64	ng/ul	0.00
43) Dimethylphthalate-d6	13.85	166	223034	20.60	ng/ul	0.00
46) Acenaphthylene-d8	14.13	160	265207	19.60	ng/ul	0.00
51) 4-Nitrophenol-d4	14.64	143	31013	15.15	ng/ul	0.00
57) Fluorene-d10	15.43	176	198733	21.10	ng/ul	0.00
62) 4,6-Dinitro-2-methylphenol	15.55	200	19061	9.80	ng/ul	0.00
70) Anthracene-d10	17.28	188	325041	20.19	ng/ul	0.00
76) Pyrene-d10	19.57	212	426487	19.04	ng/ul	0.00
87) Benzo(a)pyrene-d12	23.49	264	563365	20.80	ng/ul	0.00

Target Compounds

						Qvalue
14) Acetophenone	8.63	105	5812	1.37	ng/ul	96
44) Dimethylphthalate	13.90	163	128896	12.05	ng/ul	99
47) Acenaphthylene	14.16	152	18198	1.32	ng/ul	99
58) Fluorene	15.49	166	18848	1.82	ng/ul	95
69) Phenanthrene	17.23	178	258939	13.71	ng/ul	99
71) Anthracene	17.32	178	58666	3.05	ng/ul	99
74) Fluoranthene	19.24	202	392045	18.44	ng/ul	99
77) Pyrene	19.60	202	441289	15.52	ng/ul	97
80) Benzo(a)anthracene	21.34	228	180363	6.20	ng/ul	96
82) Chrysene	21.40	228	216316	7.82	ng/ul	98
85) Benzo(b)fluoranthene	22.95	252	230848	6.75	ng/ul	98
86) Benzo(k)fluoranthene	22.99	252	59472m	1.84	ng/ul	
88) Benzo(a)pyrene	23.54	252	199192	6.00	ng/ul	98
89) Indeno(1,2,3-cd)pyrene	25.93	276	129406	3.28	ng/ul	96
91) Benzo(g,h,i)perylene	26.64	276	126681	3.82	ng/ul	98

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

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