

Method Path : Z:\HPCHEM1\BNA_M\METHODS\
 Method File : SOM01.2-EPA-SIM-BM061115.M
 Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION
 Last Update : Thu Jun 11 07:52:35 2015
 Response Via : Initial Calibration

Calibration Files

0.1 =BM001636.D 0.2 =BM001637.D 0.4 =BM001638.D
 0.8 =BM001639.D 1 =BM001640.D

Compound		0.1	0.2	0.4	0.8	1	Avg	%RSD

1) I	1,4-Dichlorobenzene-d	-----ISTD-----						
2) SURR1	1,4-Dioxane-d8						0.000	-1.00
3)	1,4-Dioxane						0.000	-1.00
4) I	Naphthalene-d8	-----ISTD-----						
5)	Naphthalene	1.073	1.033	1.069	1.053	1.039	1.053	1.68
6) SURR2	Methylnaphthalene	0.479	0.479	0.484	0.483	0.489	0.483	0.90
7)	2-Methylnaphthalene	0.659	0.651	0.662	0.657	0.671	0.660	1.16
8) I	Acenaphthene-d10	-----ISTD-----						
9)	Acenaphthylene	1.929	1.861	1.898	1.958	1.904	1.910	1.88
10) C	Acenaphthene	1.484	1.421	1.450	1.445	1.426	1.445	1.72
11)	Fluorene	1.552	1.539	1.544	1.575	1.547	1.551	0.89
12) I	Phenanthrene-d10	-----ISTD-----						
13)	Pentachlorophenol		0.074	0.080	0.087	0.079	0.080	6.89
14)	Phenanthrene	1.167	1.131	1.161	1.173	1.141	1.155	1.57
15)	Anthracene	1.134	1.081	1.087	1.136	1.102	1.108	2.32
16) SURR	Fluoranthene-d10	0.847	0.781	0.808	0.819	0.780	0.807	3.49
17) C	Fluoranthene	1.318	1.185	1.244	1.255	1.196	1.240	4.29
18) I	Chrysene-d12	-----ISTD-----						
19)	Pyrene	1.649	1.647	1.610	1.576	1.582	1.613	2.13
20)	Benzo(a)anthracene	1.338	1.289	1.302	1.275	1.239	1.289	2.81
21)	Chrysene	1.500	1.383	1.414	1.334	1.290	1.384	5.80
22) I	Perylene-d12	-----ISTD-----						
23)	Benzo(b)fluoranthene	1.677	1.499	1.654	1.471	1.455	1.551	6.82
24)	Benzo(k)fluoranthene	1.870	1.558	1.372	1.518	1.468	1.557	12.09
25) C	Benzo(a)pyrene	1.618	1.397	1.402	1.366	1.332	1.423	7.91
26)	Indeno(1,2,3-cd)pyr	2.048	1.631	1.530	1.469	1.442	1.624	15.26
27)	Dibenzo(a,h)anthrac	1.695	1.319	1.217	1.172	1.145	1.309	17.20
28)	Benzo(g,h,i)perylene	1.996	1.555	1.395	1.318	1.278	1.509	19.38

(#) = Out of Range