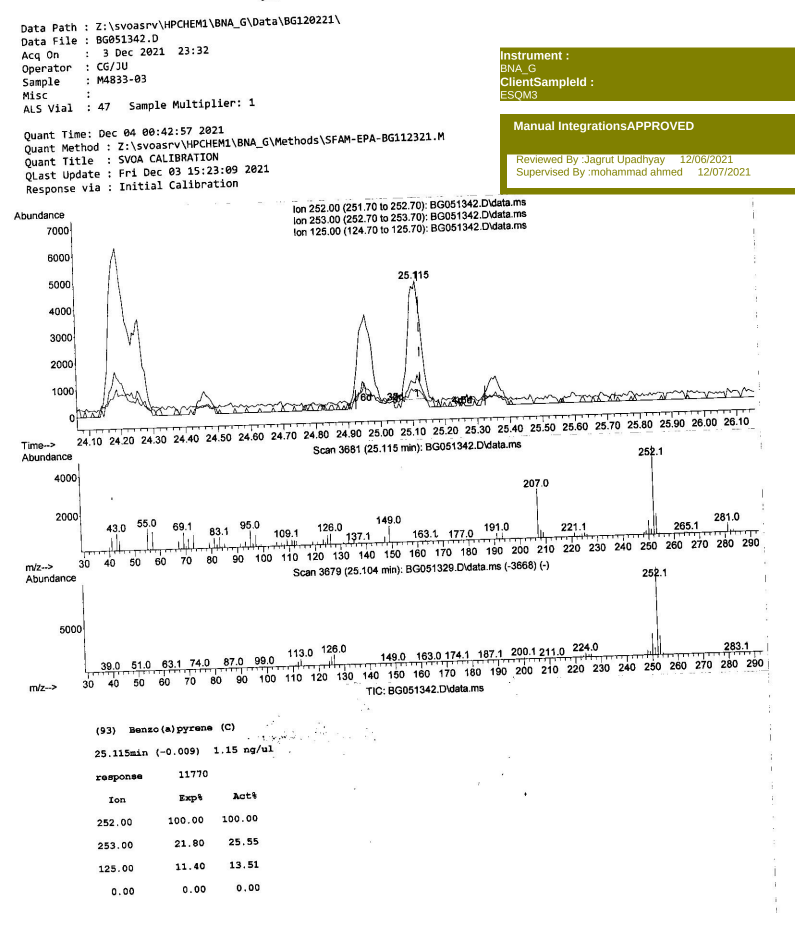
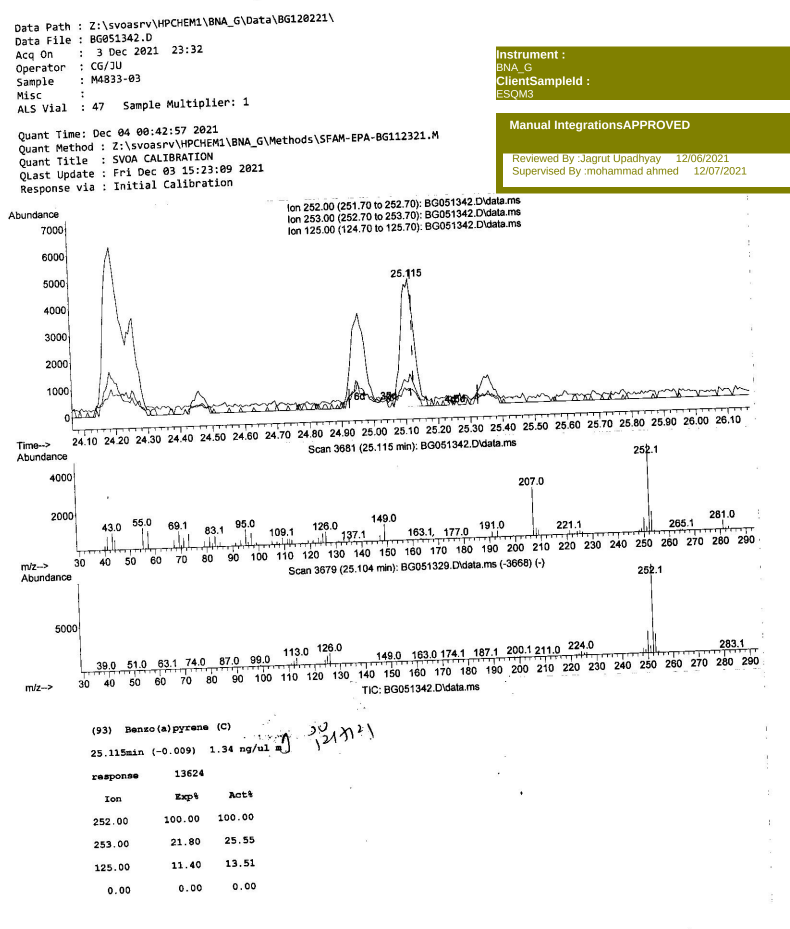


Quantitation Report (Qedit)



Quantitation Report (Qedit)



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Data Path : Z:\svoasrv\HPCHEM1\BN Data File : BG051342.D Acq On : 3 Dec 2021 23:32 Operator : CG/JU Sample : M4833-03 Misc : ALS Vial : 47 Sample Multiplie		\BG12	0221\			BN Cli	strument : A_G entSampleld : QM3
Quant Time: Dec 04 00:42:57 2021 Quant Method : Z:\svoasrv\HPCHEM1 Quant Title : SVOA CALIBRATION QLast Update : Fri Dec 03 15:23:0 Response via : Initial Calibratic	9 2021	lethod	Is\SFAM-EPA·	-BG112321	.M		Manual IntegrationsAPPROVED Reviewed By :Jagrut Upadhyay 12/06/2021 Supervised By :mohammad ahmed 12/07/2021
Compound		lon	Response				
Internal Standards		450	20622	20.000 n	a/u1	0.00	
1) 1,4-Dichlorobenzene-d4	8.194		30623	20.000 n	8896	0.00	
20) Naphthalene-d8	11.020	136	137919 90742	20.000 m		0.00	
38) Acenaphthene-d10	14.821	164	189939	20.000 п	1450 M	0.00	
64) Phenanthrene-d10	17.577	188	169939	20.000 n		0.00	
79) Chrysene-d12	21.878	240 264	158305	20.000 m	All and a second s	0.00	
88) Perylene-d12	25.274	204	138303	201000 1	·B/ 41		
Custom Monitoning Compounds							
System Monitoring Compounds	3.535	96	2542	2.885 r	ng/uL	0.00	
3) 1,4-Dioxane-d8	3.964	84	16636	6.434 r		-0.02	
4) Pyridine-d5	7.354	99	55229	18.248 r		0.00	
7) Phenol-d5	7.506	67	39324	20.688		0.00	I.
9) Bis-(2-Chloroethyl)eth	7.724	132	42418	19.463 r	-	0.00	2 <u>/</u>
11) 2-Chlorophenol-d4	8.911	113	29253	11.977		0.00	
15) 4-Methylphenol-d8	9.369	128	23957	20.577 1		0.00	
21) Nitrobenzene-d5	10.098	143	26262	19.997 1		0.00	
24) 2-Nitrophenol-d4		165	41826	18.771		0.00	
28) 2,4-Dichlorophenol-d3	10.644	131	37639	11.544		0.00	
31) 4-Chloroaniline-d4	14.216	166	151377	21.681		0.00	
46) Dimethylphthalate-d6		160	199297	22.636		0.00	
49) Acenaphthylene-d8	14.522	143	16651	14.733		0.00	
54) 4-Nitrophenol-d4	15.051		143955	22.896		0.00	
60) Fluorene-d10	15.814	176	12011	10.248		0.00	
65) 4,6-Dinitro-2-methylph	15.949	200 188	216663	23.851		0.00	
73) Anthracene-d10	17.671	212	259378	26,713		0.00	
81) Pyrene-d10	19.957 25.039	264	217725	25.752	a second s	0.00	
92) Benzo(a)pyrene-d12	23.039	204	21//23	201100			
Terret Compounds					Qv	alue	
Target Compounds	19.622	202	32515	2.726	ng/ul	97	
80) Fluoranthene	19.980		27619	2.368	1000 N	98	
82) Pyrene	21.854		12374	1.137		96	20
85) Benzo(a)anthracene	21,925		14656	1.402	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	98	JAN L
87) Chrysene 90) Benzo(b)fluoranthene	24.193		21856 1	2.046		98	
93) Benzo(a)pyrene	25.115		13624m	1.337	ng/ul		
55) Denzo(a)pyrene							
(#) = qualifier out of range (r	n) = man	ual i	ntegration	(+) = się	gnals s	summed	

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SFAM-EPA-BG112321.M Sat Dec 04 00:51:24 2021

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