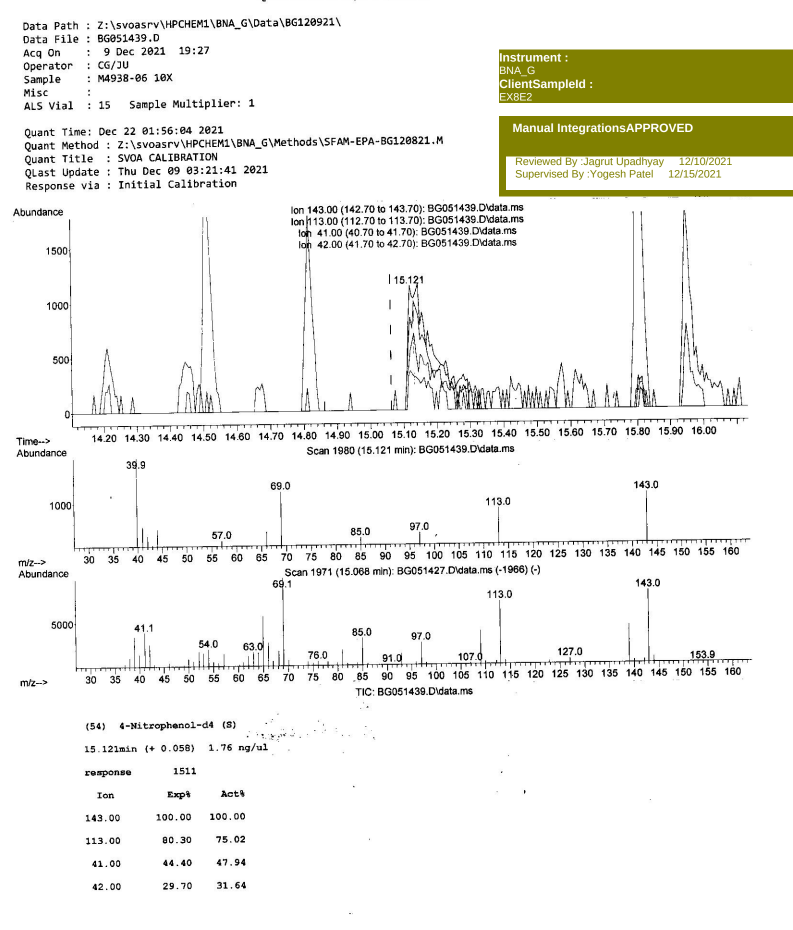


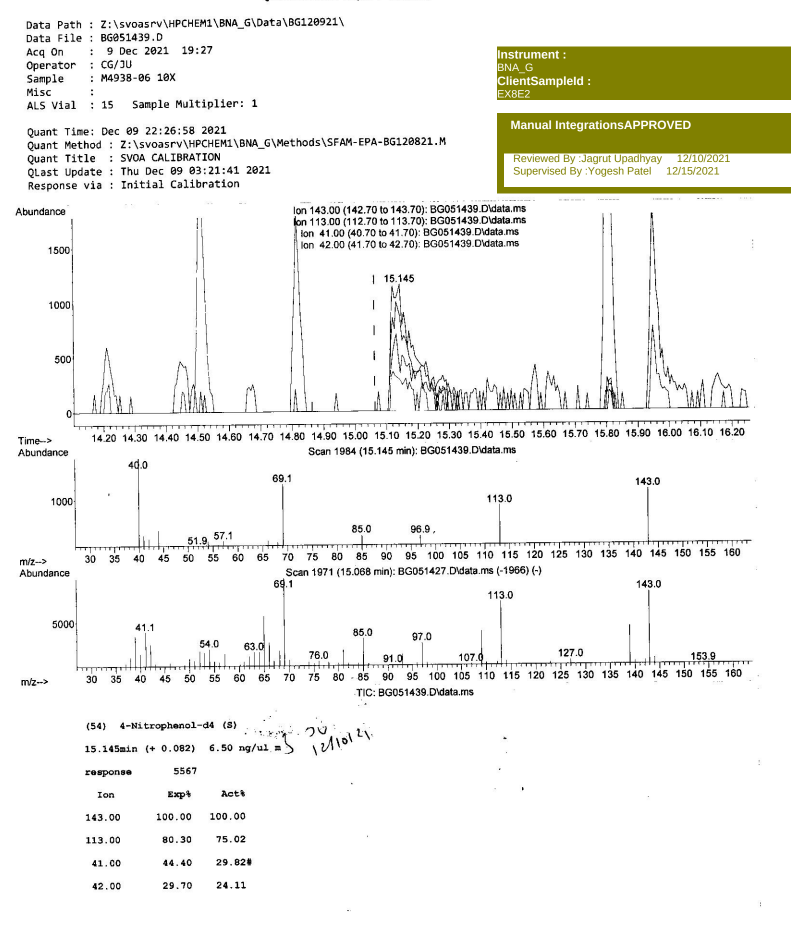
Quantitation Report (Qedit)



SFAM-EPA-BG120821.M Wed Dec 22 01:56:16 2021

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Quantitation Report (Qedit)



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Data Path : Z:\svoasrv\HPCHEM1\BNA	A_G\Data\BC	512092	21\			
Data File : BG051439.D Aco On : 9 Dec 2021 19:27						
					Inst	trument :
Operator : CG/JU Sample : M4938-06 10X					BNA	_G
					Clie	entSampleId :
Misc : ALS Vial : 15 Sample Multiplie	r: 1				EX8	
ALS Vial : 15 Sample Multiplie						
Quant Time: Dec 09 22:26:58 2021 Quant Method : Z:\svoasrv\HPCHEM1 Quant Title : SVOA CALIBRATION QLast Update : Thu Dec 09 03:21:4 Response via : Initial Calibratio	1 2021	hods'	SFAM-EPA-	BG120821.M	F	Ianual IntegrationsAPPROVED Reviewed By :Jagrut Upadhyay 12/10/2021 Supervised By :Yogesh Patel 12/15/2021
		on R	esponse	Conc Units Dev(M	in) 	
Internal Standards	8.188 1	52	24205	20.000 ng/ul	0.00	
1) 1,4-Dichlorobenzene-d4	11.008 1	36	108048	20.000 ng/ul	0.00	
20) Naphthalene-d8	14.816 1	.64	73532	20.000 ng/ul	0.00	
38) Acenaphthene-d10	17.565 1	88	166147	20.000 ng/ul	0.00	
64) Phenanthrene-d10		40	150250	20.000 ng/ul	0.00	
79) Chrysene-d12		264	140914	20.000 ng/ul	0.00	
88) Perylene-d12	231200					
System Monitoring Compounds				- 171 (ul	0.00	
3) 1,4-Dioxane-d8	3.529	96	1573	2.134 ng/uL	0.02	
4) Pyridine-d5	3.981	84	17257	8.153 ng/ul 7.153 ng/ul	0.02	
7) Phenol-d5	7.372	99	17626	33,762 ng/ul	0.00	
9) Bis-(2-Chloroethyl)eth	7.501	67	53350	26.088 ng/ul	0.00	1
11) 2-Chlorophenol-d4		132	45735	17.493 ng/ul	0.00	
15) 4-Methylphenol-d8		113	33862	33.975 ng/ul	0.00	
21) Nitrobenzene-d5		128	31844	33.374 ng/ul	0.00	
24) 2-Nitrophenol-d4	10.0	143	35397	31.627 ng/ul	0.00	
28) 2,4-Dichlorophenol-d3		165	54570	26.586 ng/ul	0.00	
31) 4-Chloroaniline-d4		131	67094 216860	38.114 ng/ul	0.00	A 11
46) Dimethylphthalate-d6	14.216	166			0.00	20/10/21
49) Acenaphthylene-d8	14.516	160	256890 5567m) 6.499 ng/ul	0.08	1211
54) 4-Nitrophenol-d4	15.145	143	188967	37.308 ng/ul	0.00	
cal Eluorene-d10	15.809		33618	34.050,ng/ul	0.00	
65) 4.6-Dinitro-2-methylph	. 15.956		325473	41.867 ng/ul	0.00	
73) Anthracene-d10	17.000		382400	42.343 ng/ul	0.00	
81) Pyrene-d10	19.951 25.027		320062	44.036 ng/ul	0.00	
92) Benzo(a)pyrene-d12	25.02/	204	520002			
					/alue	
Target Compounds	7.401	94	5327	2.112 ng/ul	97	
8) Phenol	11.061		6590	1.111 ng/ul	98	
20) Nanhthalene	11.001					
	• * * * * * * * * * * * * * * * * * * *		10	() stepping	cummed	

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

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