

0.00

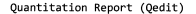
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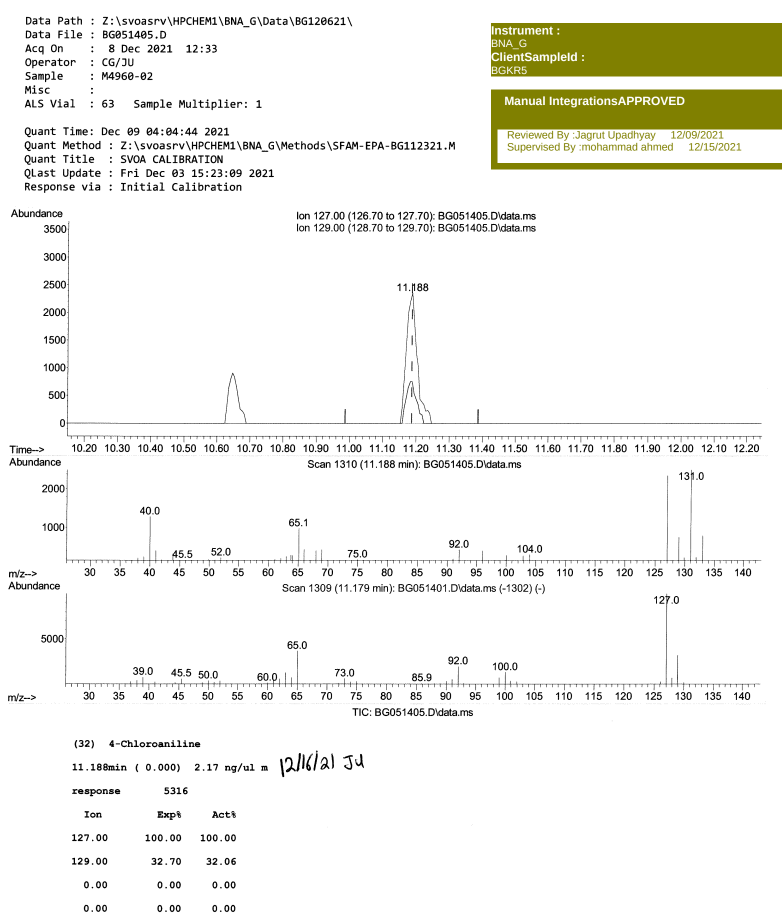
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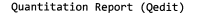
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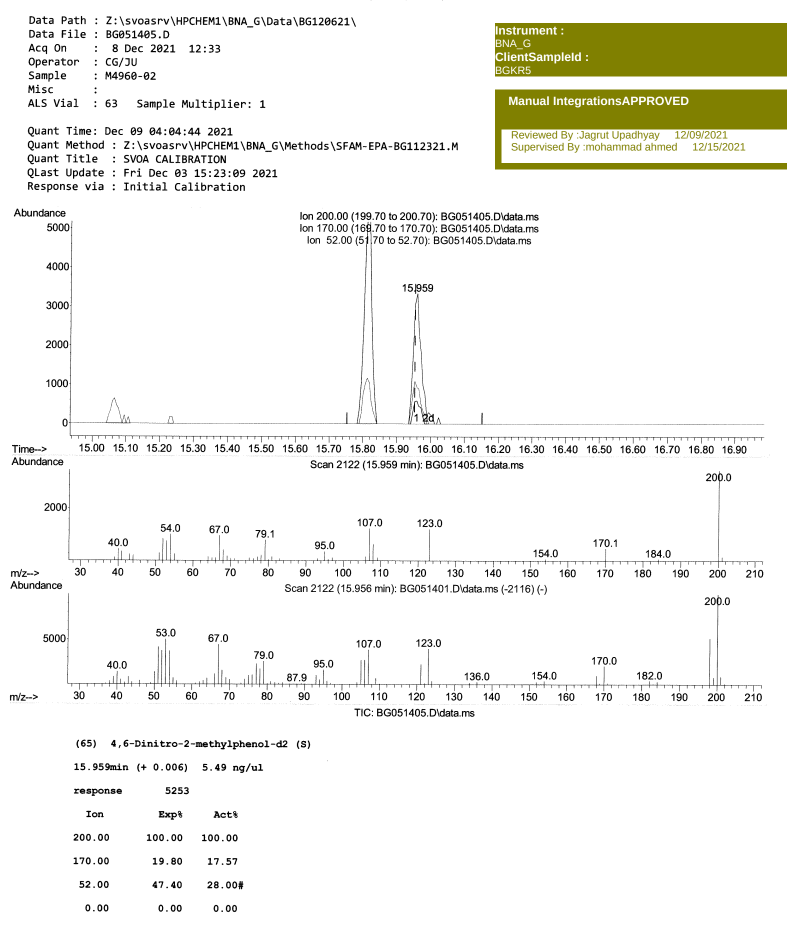
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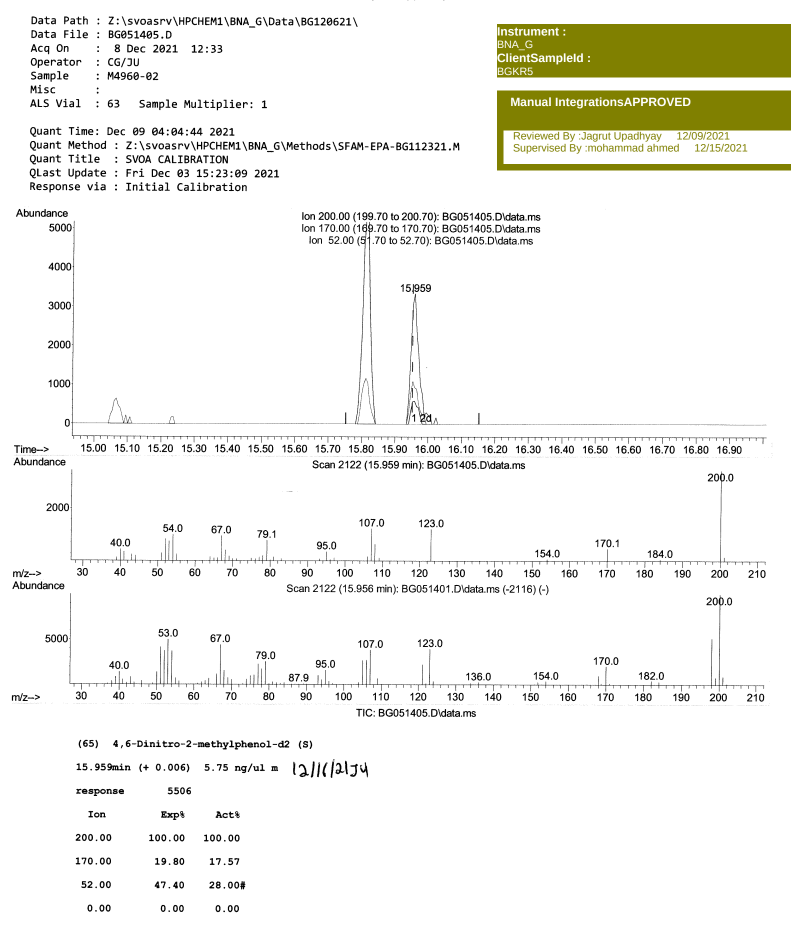












Data Path : Z:\svoasrv\HPCHEM1\BNA_G\Data\BG120621\ Data File : BG051405.D Acq On : 8 Dec 2021 12:33 Operator : CG/JU Sample : M4960-02 Misc : ALS Vial : 63 Sample Multiplier: 1					Instrument : BNA_G ClientSampleId : BGKR5 Manual IntegrationsAPPROVED	
Quant Time: Dec 09 04:04:44 2021 Quant Method : Z:\svoasrv\HPCHEM1\BNA_G\Methods\SFAM-EPA-BG112321.M Quant Title : SVOA CALIBRATION QLast Update : Fri Dec 03 15:23:09 2021 Response via : Initial Calibration						
	Compound	R.T.	QIon	Response	Conc Units Dev(Min)
Inte	rnal Standards					
1)	1,4-Dichlorobenzene-d4	8.191	152	23001	20.000 ng/ul	-0.01
20)	Naphthalene-d8	11.017	136	103095	20.000 ng/ul	-0.01
38)	Acenaphthene-d10	14.819	164	69955	20.000 ng/ul	-0.01
	Phenanthrene-d10	17.569	188	155086	20.000 ng/ul	-0.01
	Chrysene-d12	21.875	240	133645	20.000 ng/ul	0.00
88)	Perylene-d12	25.277	264	133878	20.000 ng/ul	0.00
System Monitoring Compounds						
	1,4-Dioxane-d8	3.526	96	1418	2.142 ng/uL	-0.02
4)	Pyridine-d5	3.967	84	8216	4.230 ng/ul	
	Phenol-d5	7.357		29416	12.940 ng/ul	0.00
9)	<pre>Bis-(2-Chloroethyl)eth</pre>	7.504	67	18396	12.885 ng/ul	
11)	2-Chlorophenol-d4	7.721	132	22036	13.461 ng/ul	
15)	4-Methylphenol-d8	8.908	113	23105	12.595 ng/ul	0.00
21)	Nitrobenzene-d5	9.366	128	11686	13.428 ng/ul	-0.01
24)	2-Nitrophenol-d4	10.095	143	12790	13.028 ng/ul	0.00
28)	2,4-Dichlorophenol-d3	10.647	165	21384	12.838 ng/ul	0.00
31)	4-Chloroaniline-d4	11.158	131	29708	12.190 ng/ul	0.00
46)	Dimethylphthalate-d6	14.214	166	75997	14.119 ng/ul	-0.01
49)	Acenaphthylene-d8	14.519	160	97649	14.387 ng/ul	-0.01
54)	4-Nitrophenol-d4	15.066	143	9956	11.427 ng/ul	0.02
	Fluorene-d10	15.812		70032	14.448 ng/ul	-0.01
65)	4,6-Dinitro-2-methylph	15.959	200	5506m 🗲	5.753 ng/ul>	0.00 12/16/2174
73)	Anthracene-d10	17.674	188	127394	17.175 ng/ul	0.00
81)	Pyrene-d10	19.954	212	146571	18.125 ng/ul	0.00
92)	Benzo(a)pyrene-d12	25.042	264	126882	17.746 ng/ul	0.00
Targe	Target Compounds Ovalue					
-	4-Chloroaniline	11.188	127	5316m ·	> 2.173 ng/ul >	
•	Bis(2-ethylhexyl)phtha			15937	2.742 ng/ul#	98

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