**Biomonitoring Quality Assurance Support Program** 

Analytical Method Report UNIVERSAL PESTICIDES

## **Laboratory Information**

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State Report:

Results for Method:

## Sample Preparation Information:

3 Does your method use automation:

4 Does your sample preparation method include:

5 Solid phase extraction platform:

Additional information on sample prep procedure:

6 What is the method sample volume size:

7 Name of enzyme:

Enzyme Vendor Enzyme Concentration Amount How long do you incubate the samples Temperature during incubations

### **HPLC Configuration:**

- 8 Instrument manufacturer
- 9 What is the flow rate:
- 10 What is the method run time:
- 11 What is the sample injection volume:
- 12 Column name and Manufacturer
- 13 Column dimensions
- 14 Elution Type:
- 15 Mobile Phase A Composition
- 16 Mobile Phase B Composition

### **Mass Spectrometer Configuration**

Have you optimized the MS Parameters for your method? (Analytes, 17 Precursor and Product Ions, Collision Energy) 18 What is the ionization mode:

<sup>20</sup> Please complete the table for each analytes LOD, precursor and product ion transitions:

Analytical and Internal Standards Please complete the table for metabolite standards:

# **Additional Method Questions**

30 Which proficiency testing programs do you participate in?

what is the average number of samples analyzed per month for this 31 method?

- 32 Have you checked the accuracy of the method using NIST SRMs?
- 33 What volume of sample is required for BQASP Analysis?

Please provide a screenshot of your results chromatography:

Select State	
Select Method	
Yes or No	
Select SPE	
N/A	
Type description here	
	(please include units)
	Degrees Celsius
Select	Type Other Here
Select	
	 Describe composition for Mobile Phase A
	Describe composition for Mobile Phase B

Yes or No Select Mode

Analyte	LOD
Example: MCPP	0.4 ng/ml
13C4-MCPP	

Analytical and Internal Standard	Vendor
Phthalate Metabolites	Cambridge Isotope Laboratory
Select	- 
	_ 
Select	_Please type other programs here

Right click in the textbox Click format shape Click Fill Options Select picture from your saved file

Calibration Range	Precursor Ion (mass)	Product Ion (mass)
0.035 - 350 ng/ml	251	103
	225	103

Purity		

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#### **UNIVERSAL PESTICIDES**

PT Event ID: 20190FUPSU Participant: Analyst: Reviewer: Units of Result:

Sample ID	<u>Analyte</u>	Reported Value
20190F001UPSU	3,5,6-trichloro-2-pyridinol	
20190F001UPSU	3-phenoxybenzoic acid	
20190F001UPSU	2-Isopropyl-4-methyl-6-hydroxypyrimidine	
20190F001UPSU	3-ph2-diethylamino-6-methyl pyrimidin-4-olenoxybenzoic acid	
20190F001UPSU	Malathion dicarboxylic acid	
20190F001UPSU	Trans-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane-1- carboxylic acid	
20190F001UPSU	Cis-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane-1- carboxylic acid	
20190F001UPSU	4-fluoro-3-phenoxy-benzoic acid	
20190F001UPSU	Para-nitrophenol	
20190F001UPSU	2,4-dichlorophenoxyacetic acid	

Sample ID	<u>Analyte</u>	Reported Value
20190F002UPSU	3,5,6-trichloro-2-pyridinol	
20190F002UPSU	3-phenoxybenzoic acid	
20190F002UPSU	2-Isopropyl-4-methyl-6-hydroxypyrimidine	
20190F002UPSU	3-ph2-diethylamino-6-methyl pyrimidin-4-olenoxybenzoic acid	
20190F002UPSU	Malathion dicarboxylic acid	
20190F002UPSU	Trans-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane-1- carboxylic acid	
20190F002UPSU	Cis-3-(2,2-dichlorovinyl)-2,2-dimethylcyclopropane-1- carboxylic acid	
20190F002UPSU	4-fluoro-3-phenoxy-benzoic acid	
20190F002UPSU	Para-nitrophenol	
20190F002UPSU	2,4-dichlorophenoxyacetic acid	

By submitting this form, we attest that the results reported were produced in this laboratory from the analysis of proficiency testing samples that were introduced into the routine workflow of the laboratory and analyzed using protocols and procedures with the same frequency routinely applied to patient specimens.

We further attest that the laboratory did not discuss or engage in any communications with anyone outside of our laboratory regarding the proficiency test or the results obtained.

CDC estimates the average public reporting burden for this collection of information as 45 minutes per response, including the time for reviewing instructions, searching existing data/information sources, gathering and maintaining the data/information needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB Control Number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC/ATSDR Information Collection Review Office, 1600 Clifton Road NE, MS D-74, Atlanta, Georgia 30333; ATTN: PRA (0920-xxxx).

Form Approved OMB No. 0920-xxxx Exp. Date xx/xx/20xx