

Recipient Name (Select)

State

Federal Award Identification Number

Report Frequency

Date Completed

Project Period Start Date

Project End Date

Budget Period Start Date

Budget Period End Date

Principal Investigator (PI)

PI Email

PI Phone

Select "Yes" for all applicable award tracks for which reporting information is included:

The information below applies only to requirements of the Paperwork Reduction Act of 1995.

The burden time for this collection of information is estimated to average 36.67 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden to the address to the right:

"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 number."

Department of Health and Human Services
Food and Drug Administration
Office of Operations
Paperwork Reduction Act (PRA) Staff
PRASStaff@fda.hhs.gov

DO NOT SEND YOUR COMPLETED FORM TO THIS PRA STAFF EMAIL ADDRESS.

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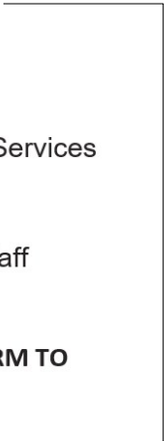


PRA statement

Human Services
Division

(PRA) Staff

ATTACHED FORM TO
PROCESS.



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ESS.



TO BE FILLED OUT BY FDA

Date:	
Matrices:	
Analyte(s):	
Methods:	
Comments:	

Lab Name

Are you able to analyze any or all the matrices listed using the method(s) cited?

Please list what matrices you could test.

Has your lab analyzed these matrices using these methods in the past?

Which methods do you use? Please list all methods available.

Are the methods
validated for the
matrices?

Please list matrix, method and
validation status.

Do you have trained staff
proficient for this analysis?

Are any or all of methods
on your scope of
accreditation?

Please list which methods
are on your accreditation
scope.

Are you willing to pivot
current approved
sampling plan to this
activity?

How many samples can
you do in a week?
(Estimate only)

How many samples
would you be willing
test in total? (Estimate
only)?

Are you able to
arrange collection of
these samples in your
state?

Please Provide Any Additional Information To
Explain Your Labs Capabilities