Effective 1 October 2010 Revised November 2015

Urine Laboratory Application Form

National Laboratory Certification Program (NLCP)

RTI International Center for Forensic Sciences 3040 Cornwallis Road P.O. Box 12194 Research Triangle Park, North Carolina 27709

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NATIONAL LABORATORY CERTIFICATION PROGRAM URINE LABORATORY APPLICATION FORM

A. Applicant Laboratory

1.	Name of Laboratory:				
	Address:				
	City, State, ZIP:				
	Telephone: () FAX: ()				
	e-Mail:				
2.	Express delivery address (if different from above)				
	Address:				
	City, State, ZIP:				
3.	Designated Responsible Person (RP):				
	Title/Position:				
	Telephone: () Ext				
	e-Mail:				
	If applicable:				
	Designated Alternate RP (Alt-RP):				
	Title/Position:				
	Telephone: () Ext				
	e-Mail:				
4.	I understand that the answers provided in this application will be used to determine the applicant laboratory's potential eligibility				
	for the National Laboratory Cortification Brogram. To the best of				

used to determine the applicant laboratory's potential eligibility for the National Laboratory Certification Program. To the best of my knowledge and belief, the answers recorded herein are true and complete as of this date.

Signature, Designated RP

Date

NOTE: Any false, fictitious, or fraudulent statements or information presented in this application form could subject you to prosecution, monetary penalties, or both. See Sec. 18 U.S.C. 1001; 31 U.S.C. 3801-812.

B. General Laboratory Information

The following table is excerpted from Section 3.4 of the Mandatory Guidelines for Federal Workplace Drug Testing Programs (Federal Register, 73 FR 71858, 25 November 2008, effective 1 October 2010):

Initial Test Analyte	Initial Test Cutoff Concentration	Confirmatory Test Analyte	Confirmatory Test Cutoff Concentration		
Marijuana metabolites	50 ng/mL	THCA ¹	15 ng/mL		
Cocaine metabolites	150 ng/mL	Benzoylecgonine	100 ng/mL		
Opiate metabolites					
Codeine/Morphine ²	2000 ng/mL	Codeine	2000 ng/mL		
		Morphine	2000 ng/mL		
6-Acetylmorphine	10 ng/mL	6-Acetylmorphine	10 ng/mL		
Phencyclidine	25 ng/mL	Phencyclidine	25 ng/mL		
Amphetamines ³					
AMP/MAMP ⁴	500 ng/mL	Amphetamine	250 ng/mL		
		Methamphetamine ⁵	250 ng/mL		
MDMA ⁶	500 ng/mL	MDMA	250 ng/mL		
		MDA ⁷	250 ng/mL		
		MDEA ⁸	250 ng/mL		
¹ Delta-9-tetrahydrocanna	abinol-9-carboxylic acid (THCA).			
² Morphine is the target a	analyte for codeine/morpl	hine testing.			
		t kits may be used provided	the single test kit detects		
each target analyte independently at the specified cutoff.					
⁴ Methamphetamine is the target analyte for amphetamine/methamphetamine testing.					
		specimen must also contai	n amphetamine at a		
concentration equal to or greater than 100 ng/mL. ⁶ Methylenedioxymethamphetamine (MDMA).					
⁷ Methylenedioxyamphetamine (MDMA).					
⁸ Methylenedioxyethylamphetamine (MDA).					
meanylonoaloxyoarylam					

- To be eligible for certification, the laboratory must test for all drug analytes and specimen validity test measurands required by the Mandatory Guidelines for Federal Workplace Drug Testing Programs (*Federal Register*, 73 FR 71858, 25 November 2008, effective 1 October 2010). The laboratory must use the test methods specified by the Mandatory Guidelines for screening, differential, initial, and confirmatory tests (i.e., drug tests and specimen validity tests).
 - 1a. Does the laboratory have validated initial drug test assays for the drug classes required by the Mandatory Guidelines?

Yes No → LABORATORY NOT ELIGIBLE TO APPLY

1b. Does the laboratory use an immunoassay method approved, cleared, or otherwise recognized as accurate and reliable by the U.S. Food and Drug Administration (FDA) for the initial drug tests?



1c. Does the laboratory have validated confirmatory test assays for the drug analytes required by the Mandatory Guidelines? (*Note: testing for amphetamine and methamphetamine enantiomers is optional.*)

$$\begin{array}{cc} & \text{Yes} \\ \hline & \text{No} \end{array} \rightarrow \textbf{LABORATORY NOT ELIGIBLE TO APPLY} \end{array}$$

1d. Does the laboratory use methods combining chromatographic separation and mass spectrometric identification [e.g., gas chromatography/mass spectrometry (GC/MS), liquid chromatography/mass spectrometry (LC/MS), GC/MS/MS, LC/MS/MS)] for the confirmatory drug tests?

____ Yes No → LABORATORY NOT ELIGIBLE TO APPLY

1e. Does the laboratory have validated tests to assess specimen validity as required by the Mandatory Guidelines (i.e., at a minimum, tests for creatinine, pH, specific gravity, and one or more oxidizing adulterants)?

Yes No → LABORATORY NOT ELIGIBLE TO APPLY

1f. Does the laboratory perform testing for amphetamine and methamphetamine enantiomers?

 $\underbrace{\qquad Yes \rightarrow \textbf{COMMENT BELOW}}_{No}$

Briefly describe the procedure for analysis and reporting of the enantiomers:

2. Is the laboratory registered with the U.S. Drug Enforcement Agency (DEA)?

 $\begin{array}{ccc} & \text{Yes} \rightarrow \text{ATTACH PHOTOCOPY OF REGISTRATION CERTIFICATE} \\ & \text{No} \rightarrow \text{COMMENT BELOW} \end{array}$

If YES, which schedules are covered by the registration?

____1 ___2 ___2N ____3 ___3N ____4 ___5

If NO, explain how controlled reference materials are acquired:

3. Describe the State licensure requirements for urine forensic toxicology for the State in which the laboratory is located.

4. List laboratory certifications/licenses:

 States (List):
 CLIA/HCFA ¹ (List Specialties):
 CAP ² (List Specialties):
 Others (Specify):

¹Clinical Laboratory Improvement Amendments(CLIA)/Health Care Financing Administration (HCFA) ²College of American Pathologists (CAP)

4a. ATTACH PHOTOCOPIES OF ALL LICENSES AND CERTIFICATIONS INDICATED ABOVE.

C. Laboratory Standard Operating Procedures (SOP) Manual

1. For certification, the laboratory must have a complete SOP manual that will apply to testing of regulated specimens under the Mandatory Guidelines for Federal Workplace Drug Testing Programs (Federal Register, 73 FR 71858, 25 November 2008, effective 1 October 2010).

Note: Manufacturers' package inserts or instrument manuals are not considered formal procedures. A written SOP manual is required to be eligible to apply for certification and it must be completed before the laboratory is eligible to receive NLCP performance testing (PT) samples.

1a. Does the laboratory have a complete SOP manual for regulated drug testing?

 $___ Yes$ $___ No → LABORATORY NOT ELIGIBLE TO APPLY$

LABORATORY SOP MANUAL INDEX

Indicate the location for each of these topics in the laboratory's SOP manual:

<u>TOPIC</u>	<u>SECTION</u>	<u>PAGE NO.</u>
Security Procedure for controlling access to the drug testing facility		
Procedure for controlling access to individual secured areas		
Procedure for documenting visitor access		
Accessioning (Specimen receipt) Procedure for receipt and processing of specimens		
Procedure for accessioning specimens received from another laboratory		
Procedure for problem/rejected specimens		
Chain-of-Custody Procedure for documenting all transfers of specimens		

<u>TOPIC</u>	<u>SECTION</u>	<u>PAGE NO.</u>
Procedure for documenting all transfers of aliquots		
Procedure for using an ECCF System (if applicable)		
Procedure for maintaining security of specimen bottles		
Procedure for maintaining security of specimen aliquots		
Procedure for sending a specimen to another laboratory		
Procedures for documenting all transfers of specimens received from another laboratory		
<i>Aliquot Preparation</i> Procedure for preparing initial drug test aliquots		
Procedure for preparing screening/differenti specimen validity test aliquots	al	
Procedure for preparing initial specimen validity test aliquots		
Procedure for preparing confirmatory specimen validity test aliquots		
Procedure for preparing confirmatory drug test aliquots		
Procedures for automated aliquotting equipment		
<i>Initial Drug Test</i> Principle of analysis		
Preparation of test materials, calibrators, and controls		

<u>TOPIC</u>	<u>SECTION</u>	<u>PAGE NO.</u>
Procedure for set-up and normal operation of instruments		
Procedure for instrument maintenance		
Procedure for assay calibration		
Procedure for calculating results		
Quality control (QC) procedure and criteria for acceptable results and corrective actions		
Procedure for validation of initial drug test methods		
References		
Second Initial Drug Test Criteria for use		
Principle of analysis		
Preparation of test materials, calibrators, and controls		
Procedure for set-up and normal operation of instruments		
Procedure for instrument maintenance		
Procedure for assay calibration		
Procedure for calculating results		
QC procedure and criteria for acceptable results and corrective actions		
Procedure for validation of second initial drug test methods		

<u>SECTION</u> PAGE NO.

<u>TOPIC</u>

References

Specimen Validity Tests

Note: Provide the following information for each specimen validity test (Initial, Confirmatory, Screening, Differential)

Creatinine Principle of analysis	
Preparation of test materials, calibrators, and controls	
Procedure for set-up and normal operation of instruments	
Procedure for instrument maintenance	
Procedure for assay calibration	
Procedures for conducting creatinine tests	
QC acceptance/rejection criteria and corrective action for creatinine tests	
Procedure for validation of creatinine test methods	
Procedure for periodic re-verification of creatinine test methods	
Special requirements, etc.	
References	
Specific Gravity Principle of analysis	
Preparation of calibrators and and controls	
Procedure for set-up and normal operation of instruments	
Procedure for instrument maintenance	

<u>TOPIC</u>	<u>SECTION</u>	<u>PAGE NO.</u>
Procedure for assay calibration		
Procedures for conducting specific gravity tests		
QC acceptance/rejection criteria and corrective action for specific gravity tests		
Procedure for validation of specific gravity test methods		
Special requirements, etc.		
References		
Criteria for identifying acceptable, dilute, invalid, and substituted specimens based on creatinine and specific gravity test results		
Procedure for designating reconfirmed results for split specimens as substituted		
pH Principle of analysis		
Preparation of test materials, calibrators, and controls		
Procedure for set-up and normal operation of instruments		
Procedure for instrument maintenance		
Procedure for assay calibration		
Procedures for conducting pH tests		
QC acceptance/rejection criteria and corrective action for pH tests		
Criteria for identifying acceptable, invalid, and adulterated specimens based on pH test results		

	<u>TOPIC</u>		<u>SECTION</u>	<u>PAGE NO.</u>	
	ure for designating reconfirm for split specimens as adulte on pH				
Procedu	ure for validation of pH test r	nethods _			
Special	requirements, etc.	-			
Referen	ces	-			
Oxidants Principle	e of analysis	-			
Prepara and con	tion of test materials, calibra trols	ators,			
	ure for set-up and normal on of instruments	-			
Procedu	ure for instrument maintenar	nce _			
Procedu	ure for assay calibration	-			
Procedu	ures for conducting oxidant t	ests _			
	eptance/rejection criteria rective action for oxidant tes	ts _			
and adu	for identifying acceptable, in Ilterated specimens based o test results				
results f	ure for designating reconfirm for split specimens as adulte pecific oxidant				
Procedu methods	ure for validation of oxidant t s	est			
	ure for periodic re-verification test methods	n of			
Special	requirements, etc.	-			
Referen	ces	-			
Urine, Labora	itory	10		October 2010, F	Rev. 1115

TOPIC SECTION PAGE NO. **Other Adulterants** Adulterant. Principle of analysis Preparation of test materials, calibrators, and controls Procedure for set-up and normal operation of instruments Procedure for instrument maintenance Procedure for assay calibration Procedures for conducting the test QC acceptance/rejection criteria and corrective action for the test Criteria for identifying acceptable, invalid, and adulterated specimens based on the adulterant test results Procedure for designating reconfirmed results for split specimens as adulterated Procedure for validation of the test methods Procedure for periodic re-verification of the test methods Special requirements, etc. References **Confirmatory Drug Tests** Principle of each analysis THCA Benzoylecgonine Codeine/Morphine

TOPIC	<u>SECTION</u>	<u>PAGE NO.</u>
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Preparation of test materials, calibrators, ar THCA	nd controls	
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Extraction procedures THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		<u> </u>
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Procedure for instrument maintenance		
Procedure for verifying the performance		
of the mass spectrometer(s)		
Procedure for instrument set-up and operat THCA	lion	
Benzoylecgonine		
Codeine/Morphine		
6-AcetyImorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Procedure for assay calibration THCA		
Benzoylecgonine		
Codeine/Morphine		
·		·

TOPIC	<u>SECTION</u>	<u>PAGE NO.</u>
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Procedure for calculating results THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Procedure when results exceed linearity THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Procedure for designating positive results THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		<u> </u>
Procedure for designating reconfirmed resu	ılts for split sp	pecimens
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		

<u>TOPIC</u>	<u>SECTION</u>	<u>PAGE NO.</u>
Amphetamines enantiomers		
QC procedure and QC acceptance criteria THCA		
Benzoylecgonine		
Codeine/Morphine		
6-AcetyImorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
Special requirements, etc.		
THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		
References		
THCA		
Benzoylecgonine		
Codeine/Morphine		
6-Acetylmorphine		
Phencyclidine		
Amphetamine/Methamphetamine		
MDMA/MDA/MDEA		
Amphetamines enantiomers		<u> </u>
Procedure for validation of confirmatory		
drug test methods		
Procedure for periodic re-verification		
of confirmatory drug test methods		
QC and Test Materials		
Procedures for preparing stock		
standards, etc.		
Procedures for preparing and verifying		
calibrators		

TOPIC	<u>SECTION</u>	<u>PAGE NO.</u>
Procedures for preparing and verifying controls		
Corrective procedure when QC verification results are out of control limits		
Procedures for preparing and verifying test materials		
Corrective procedure when test materials verification results are unacceptable		
<i>Quality Assurance (QA) Procedures</i> Procedures for monitoring control results		
Corrective procedure when QA review of control results shows problems or potential problems (e.g., trends, shifts, bias)		
<i>Equipment and Maintenance</i> Wash procedure for labware		
Procedure for determining accuracy and precision of pipetting devices		
Procedures for temperature-dependent equipment		
Procedures for centrifuges		
Procedures for analytical balances		
Safety procedures		
Administrative/Reporting Procedures Procedure for reviewing/certifying the test result(s) of a primary specimen		
Procedure for reporting the test result(s) of a primary specimen		
Procedure for reviewing/certifying the test result(s) of a split specimen		

<u>SECTION</u>	<u>PAGE NO.</u>
tems Proced	ures

D. Chain of Custody, Accessioning, and Security

The laboratory must have chain of custody, accessioning, and security procedures that ensure integrity is maintained for the original specimens and their aliquots. Procedures must address specimens received from collectors, Instrumented Initial Test Facilities (IITFs), and other laboratories. The chain of custody forms and procedures must account for all individuals who handle the specimens and aliquots. The chain of custody forms and procedures should provide a clear picture of the handling/transfers of specimens and aliquots from initial receipt to final disposition. The laboratory must ensure the security of specimens and aliquots during processing and placement in any storage locations.

1. Provide a description of the laboratory's chain of custody procedures for the following:

Specimen Receiving/Accessioning

-Receipt of specimen packages, how they are handled, who reviews the accuracy of the information on the custody and control forms and how discrepancies are documented -Assignment of laboratory accession numbers

-Handling and resolution of problems with specimen bottles and/or custody and control forms

-Description of collection kit to be used

- -Location of temporary storage area(s)
- Procedures for electronic (digital) or combination (electronic and paper) Federal CCF (if applicable)

Aliquotting Procedures

-Aliquotting from the original specimen bottles (i.e., who and where)

-The aliquotting procedure (pouring or pipetting and amounts) used for preparing aliquots for initial drug tests, screening/differential specimen validity tests, initial specimen validity tests, confirmatory drug tests, and confirmatory specimen validity tests

-Transfer of aliquots from the individuals performing the aliquotting to those who will be testing the aliquots

Initial Drug Tests (First and Second Tests)

-Handling and testing of aliquots by laboratory personnel

-Maintenance of chain of custody and aliquot identity during the testing

Specimen Validity Tests (Initial, Confirmatory, Screening, Differential)

-Handling and testing of aliquots by laboratory personnel -Maintenance of chain of custody and aliquot identity during the testing

Confirmatory Drug Tests

-Handling and testing of aliquots by laboratory personnel -Maintenance of chain of custody and aliquot identity during the testing

Disposition of Specimens and Aliquots

-Handling of original specimen bottles and aliquots after testing is completed

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-Procedure for transferring positive, adulterated, substituted, and invalid specimens to long-term frozen storage

Note: (1)Insert here. (2) Do not exceed a total of 4 pages.

- 2. Will the laboratory use an electronic (digital) or combination (electronic and paper) Federal CCF?
 - Yes → Provide the items on the Electronic CCF System Submission List (attached) No
- 3. Attach a flowchart and/or examples of chain of custody documents showing how regulated specimens and aliquots will be processed and their custody documented (chain of custody documents may be referenced and/or provided as examples for clarification).
- 4. Will regulated specimens be accessioned in a limited access, secure area?
 - $\underbrace{ Yes } \\ No \rightarrow \textbf{LABORATORY NOT ELIGIBLE TO APPLY}$
- 5. Will regulated specimens be tested in a limited access, secure area?

___ Yes

No \rightarrow LABORATORY NOT ELIGIBLE TO APPLY

- 6. Attach a floorplan of the laboratory indicating the areas to be used for accessioning, testing of specimens, and storage of specimens, aliquots, and records. Include information to describe how the areas are secured and what security devices are utilized (e.g., which walls are outside walls; which are secured up to the ceiling; the location and type of security devices such as magnetic key cards, cipher locks, padlocks; location of secured storage areas such as refrigerators or freezers and how they are secured).
- 7. Will the original specimens be maintained in a limited access, secured area at all times?
 - $\begin{array}{cc} & \text{Yes} \\ \hline & \text{No} \end{array} \rightarrow \textbf{LABORATORY NOT ELIGIBLE TO APPLY} \end{array}$
 - 7a. Where will the original specimens be stored?

	Before testing?
	During testing?
	After testing is complete?
7b.	Who will have access to the specimen storage areas?
	Before testing?
	During testing?
	After testing is complete?
	en testing is complete, will all positive, adulterated, substituted, and invalid specimens (

8. When testing is complete, will all positive, adulterated, substituted, and invalid specimens (A and B Bottles) and split specimens be retained in long-term frozen storage in their original containers?

____ Yes \rightarrow # of days to be stored:_____

____ No → LABORATORY NOT ELIGIBLE TO APPLY

8a. How will specimens (A and B Bottles) and split specimens be stored?

E. Records

The laboratory must maintain records to support test results (i.e., including but not limited to all associated QC results, analytical data, chain of custody documents and associated administrative records) for at least two years. The laboratory must also maintain method validation records for past and current procedures, instrument validation records, records documenting the standard operating procedures used at any given time period, and records of the education, training, and certification of all employees associated with regulated testing. The laboratory must have security measures in place to limit access to electronic and hardcopy records to essential authorized personnel.

1. Will the laboratory maintain records supporting specimen test results for at least two years?

$$\begin{array}{cc} & \text{Yes} \\ \hline & \text{No} \end{array} \rightarrow \textbf{LABORATORY NOT ELIGIBLE TO APPLY} \end{array}$$

1a. Will there be a secured area for the storage of records supporting specimen test results?

2. Will the laboratory limit records access to authorized personnel?

- 3. Attach data packages using the format described in Section R of the NLCP Manual for Urine Laboratories to support (1) a positive drug test result and (2) an adulterated, substituted, or invalid result based on specimen validity testing.
- 4. In addition to the data packages described above: if the laboratory will use more than one technology (e.g., GC/MS, GC/MS/MS, LC/MS/MS) for confirmatory drug tests, attach data and documentation from a confirmatory drug test batch using each additional technology.

F. Personnel

Qualifications for a Responsible Person Candidate

1. RP Candidate's Name:

LAST FIRST MI

MIDDLE

The candidate must provide the following for review of his/her eligibility:

- (a) A detailed description of the experience and qualifications specifically addressing the RP requirements as stated in the Mandatory Guidelines (Section 11.3);
- (b) A current résumé or curriculum vitae; and
- (c) Official copies with raised seal of all academic undergraduate and graduate transcripts.
- 2. To be eligible for review as an RP, at least one of the following questions must be answered "yes":
 - 2a. Is the candidate certified/licensed by the State in which the laboratory is located and any other State requiring personnel licensure as a Laboratory Director in forensic or clinical laboratory toxicology?
 - $___ Yes → In which State(s)? _____ No$
 - 2b. Does the candidate have a Ph.D. in one of the natural sciences?

_ Yes → In which field? ____ GO TO QUESTION 3.

- No \rightarrow GO TO QUESTION 2C.
- 2c. Does the candidate have training and experience comparable to a Ph.D. in one of the natural sciences, such as a medical or scientific degree with additional training and laboratory/research experience in biology, chemistry, and pharmacology or toxicology?

 Yes \rightarrow Describe :	
 No	

- 3. An RP must have extensive experience in forensic toxicology with emphasis on the collection and analysis of biological specimens for drugs of abuse. To be eligible for review as an RP, both of the following questions must be answered "yes":
 - 3a. Does the candidate have two years or more of postdoctoral experience or at least six years of experience in forensic toxicology beyond any other degree?

 $Yes \rightarrow$	Describe:
 $\rm No \rightarrow$	CANDIDATE NOT ELIGIBLE AS RP

3b. Does the candidate have appropriate experience in forensic applications of analytical toxicology (e.g., publications, court testimony, conducting research on the toxicology of drugs of abuse) or qualify as an expert witness in forensic toxicology?

 Yes→ Describe:	
 No → CANDIDATE NOT ELIGIBLE AS RP	

4. In the table below, enter the candidate's education.

Education	Name of School	Major and Minor Fields of Study	Diploma, Certificate or Degree Received
College or University			
Other Schools Attended			

- 5. Is the candidate a full-time or part-time employee of the laboratory?
 - ____ Full-time (at least 40 hours per week)
 - ____ Part-time _____ hours per week

If not a full- or part-time employee, what is the relationship between the candidate and the laboratory?

6. How many hours per week will the candidate work in the forensic urine drug testing laboratory?

_____ HOURS PER WEEK

7. How long has the candidate been associated with the laboratory?

_____ YEARS

Qualifications for an Alternate Responsible Person Candidate

1. Alt-RP Candidate's Name: ____

LAST FIRST MIDDLE

The candidate must provide the following for review of his/her eligibility:

- (a) A detailed description of the experience and qualifications specifically addressing the RP requirements as stated in the Mandatory Guidelines (Section 11.3);
- (b) A current résumé or curriculum vitae; and
- (c) Official copies with raised seal of all academic undergraduate and graduate transcripts.
- 2. An alt-RP must be capable of fulfilling RP duties for a limited time (i.e., up to 180 days). An alt-RP candidate's qualifications are compared to RP requirements as follows:
 - 2a. Is the candidate certified/licensed by the State in which the laboratory is located and any other State requiring personnel licensure as a Laboratory Director in forensic or clinical laboratory toxicology?
 - $\underline{\qquad} Yes \rightarrow \text{In which State(s)?} \underline{\qquad} No$
 - 2b. Does the candidate have a Ph.D. in one of the natural sciences?
 - - No \rightarrow GO TO QUESTION 2C.
 - 2c. Does the candidate have training and experience comparable to a Ph.D. in one of the natural sciences, such as a medical or scientific degree with additional training and laboratory/research experience in biology, chemistry, and pharmacology or toxicology?

 Yes \rightarrow Describe:	
 No	

- 3. An alt-RP candidate must have appropriate experience in forensic toxicology.
 - 3a. How many years of experience does the candidate have in forensic toxicology (including experience with the collection and analysis of biological specimens for drugs of abuse) beyond any degree?

_____YEARS

3b. Does the candidate have appropriate training and/or experience in all operations of the forensic drug testing laboratory (i.e., including training and experience as a certifying scientist)?

___ Yes

____ No \rightarrow CANDIDATE NOT ELIGIBLE AS AN ALT-RP

Urine, Laboratory

4. In the table below, enter the candidate's education.

Education	Name of School	Major and Minor Fields of Study	Diploma, Certificate or Degree Received
College or University			
Other Schools Attended			

- 5. Is the candidate a full-time or part-time employee of the laboratory?
 - Full-time (at least 40 hours per week)Part-time _____ hours per week

If not a full- or part-time employee, what is the relationship between the candidate and the laboratory?

6. How many hours per week will the candidate work in the forensic urine drug testing laboratory?

_____ HOURS PER WEEK

7. How long has the candidate been associated with the laboratory?

_____YEARS

Personnel Certifications and Licenses

1. List the name, job title, education, and licenses/certifications for the following key staff:

Note: (1) Attach a résumé for each individual listed below.

(2) Attach a separate sheet as needed to list all individuals in these positions.

	Name	Job Title	Education	License/ Certification
Certifying Technician(s)				
Certifying Scientist(s)				
Supervisor(s)				
Other Key Staff				

2. Is licensure and/or certification required for any of the above positions in the State in which the laboratory is located?

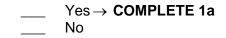
 $\underbrace{ Yes }_{No \rightarrow \textbf{GO TO SECTION G} }$

If YES, describe requirements:

G. Quality Control

For certification, the laboratory must have clearly defined QC procedures that are consistently applied, subject to review, and prompt appropriate corrective action upon failure to meet established acceptance criteria.

1. Are instrument function checks reviewed prior to batch analysis?



1a. What is the title and/or position of the person responsible for these checks?

Title/Position:

2. Are corrective actions documented when controls, instrument responses, etc., fail defined acceptance criteria?

Yes

No → LABORATORY NOT ELIGIBLE TO APPLY

3. Are all QC results reviewed by the Certifying Technician/Scientist prior to the release of the results?

___ Yes

No \rightarrow LABORATORY NOT ELIGIBLE TO APPLY

4. Is the QA/QC program under the direct supervision of a Quality Control Supervisor?

___ Yes No → COMPLETE 4a

4a. What is the title/position of the person responsible for the QA/QC program?

Title/Position:

5. Is the QA/QC program reviewed periodically by the Responsible Person Candidate?



5a. What is the title/position of the person responsible for the periodic review?

Title/Position:

6. Are there written procedures that are employed to routinely detect clerical and analytical errors prior to reporting results?

 $\begin{array}{cc} & \text{Yes} \\ \hline & \text{No} \end{array} \rightarrow \textbf{LABORATORY NOT ELIGIBLE TO APPLY} \end{array}$

7. For certification, the laboratory must have a QC program that includes both blind and open QC samples. At a minimum, these must include the number and type of QC samples described in the Mandatory Guidelines for drug and specimen validity tests.

Urine, Laboratory

Provide a description of the laboratory's procedures for the following:

Specimen Accessioning

- Introduction and/or aliquotting of blind samples into the test batches by accessioners
- Content and concentration of each blind sample
- If applicable, preparation and submission of blind samples as donor specimens from external sources

Initial Drug Tests (First and Second)

- How batches are constituted (e.g., how many specimens are in a batch, is it constituted in one session or are specimens added to the batch throughout the day?)
- The distribution of the donor specimens and QC samples within each batch
- The procedure(s) and acceptance criteria for calibration and when and by whom the calibration data are evaluated and documented
- The acceptance criteria for each control (open and blind) in each batch and when and by whom these are evaluated and documented
- The criteria for accepting all donor specimen results or only a partial number of donor specimens in a batch

Specimen Validity Tests (Initial, Confirmatory, Screening, Differential)

- How batches are constituted (e.g., how many specimens are in a batch, is it constituted in one session or are specimens added to the batch throughout the day?)
- The distribution of the donor specimens and QC samples within each batch
- The procedure(s) and acceptance criteria for calibration and when and by whom the calibration data are evaluated and documented
- The acceptance criteria for each control (open and blind) in each batch and when and by whom these are evaluated and documented
- The criteria for accepting all donor specimen results or only a partial number of donor specimens in a batch
- Include an outline or a legible flowchart that comprehensively describes the laboratory's specimen validity testing. The laboratory's submission must identify any "reflex" testing, the use of two separate aliquots, the initial and confirmatory methods for each analytical parameter, and any screening or differential tests.

Confirmatory Drug Tests

- How batches are constituted (e.g., how many specimens are in a batch, is it constituted in one session or are specimens added to the batch throughout the day?)
- The distribution of the donor specimens and QC samples within each batch
- The procedure and acceptance criteria for calibration, including criteria for exclusion of unsatisfactory calibrators
- The acceptance criteria for each control (open and blind) in each batch and when and by whom these are evaluated and documented
- The criteria for accepting, re-extracting, or reinjecting a specimen

Note: (1) Insert here.

(2) Do not exceed a total of 3 pages.

H. Review and Reporting

The laboratory must have adequate procedures to ensure the thorough review and accurate reporting of results.

1. Briefly describe the procedures for reviewing initial drug test data and certifying negative results (i.e., title/position of reviewers, electronic/hardcopy documents reviewed, QC review):

2. Briefly describe the procedures for reviewing specimen validity test data/results (i.e., screening, differential, initial and confirmatory tests): _____

3. Briefly describe the procedures for reviewing confirmatory drug test data and certifying results (i.e., title/position of reviewers, electronic/hardcopy documents reviewed, QC review):

4. Briefly describe the procedures for the reporting of results. If the laboratory will use electronic reporting for any regulated specimens, describe procedures to ensure confidentiality:

5. Is the laboratory's custody and control form (CCF) identical to the OMB-approved Federal CCF to be used for all specimens submitted for testing under the Mandatory Guidelines?

Yes→ ATTACH EXAMPLE OF LABORATORY'S CUSTODY AND CONTROL FORM
No→LABORATORY NOT ELIGIBLE TO APPLY

6. Does the laboratory's report form for split specimens contain all required elements as described in Section U of the NLCP Manual for Urine Laboratories?

___ Yes→ ATTACH EXAMPLE OF LABORATORY'S SPLIT SPECIMEN REPORT FORM

- ___ No
- 7. Will the laboratory use computer-generated electronic reports for specimens submitted for testing under the Mandatory Guidelines?

Yes →ATTACH EXAMPLE REPORTS (SEE BELOW) No

If YES, attach an example of the laboratory's computer-generated electronic report for each of the following laboratory results:

- Negative
- Negative, Dilute
- Rejected
- Cocaine Metabolite Positive
- 6-AM/Morphine/Codeine Positive
- Amphetamine/Methamphetamine Positive
- d-Methamphetamine (if applicable)
- MDMA/MDA/MDEA Positive
- Substituted
- Invalid Result
- Specimen Adulterated: pH
- Specimen Adulterated: Others as Pertinent
- Split Specimen: Reconfirmed
- Split Specimen: One or More Primary Specimen Results Not Reconfirmed
- 8. Will the laboratory send a data file report in lieu of a formatted electronic report?

__ Yes → ATTACH EXAMPLE DATA FILE REPORTS (reflecting what will be sent) No

9. Does the laboratory plan to use an electronic (digital) or combination (electronic and paper) Federal CCF for reporting? Note: Section D of the NLCP Manual for Urine Laboratories describes the allowable formats for the Federal CCF. ___ Yes ___ No

If YES, specify the CCF type(s) and supplier(s):

I. Laboratory Computers and Information Systems

Laboratory computer systems include any computer system used in processing regulated specimens. Such systems are typically used for accessioning specimens, batch assignment and scheduling, capturing test results, tabulating QC data, and reporting final results. HHS-certified laboratories are prohibited from transmitting data to an IITF through a computer interface. Any computer interface communicating any form of data from an HHS-certified IITF to a laboratory must be approved by the NLCP prior to implementation. The applicant IITF and/or laboratories must submit a detailed plan to the NLCP for review.

1. Give a brief description of the computer system to be utilized by the laboratory. Is it a "stand alone" system used solely by the laboratory, part of a local system (e.g., a hospital system), or part of a multi-laboratory corporate system? (If not on-site, provide information on its location and organizational control of the system.)

2. Give a brief description of how the laboratory plans to use the computer system in regulated specimen processing: _____

3. Is the laboratory computer system maintained in a secure area?



Attach a floorplan identifying the laboratory computer system location. Include information to describe how the area is secured and what security devices are utilized (e.g., which walls are outside walls; which are secured up to the ceiling; the location and type of security devices such as magnetic key cards, cipher locks, padlocks).

- 4. Does the laboratory limit functional access to the laboratory computer system?
 - ____ Yes ____ No

Urine, Laboratory

5. Does the laboratory have a System Security Plan (SSP) for each information system used for regulated drug testing, including corporate systems and external service provider systems?

Yes No → LABORATORY NOT ELIGIBLE TO APPLY

6. Will the laboratory use an external service provider (e.g., LIMS provider, software service provider, ECCF provider, report provider) to perform services on the laboratory's behalf related to regulated drug testing?

6a. Does the laboratory have a signed contract/agreement with each external service provider that includes the priority elements listed in the Priority Elements for Contracts/Agreements with External Service Providers?

Complete the NLCP Application Tables

Table 1-a.	First and Second Initial Drug Test Methods and Instruments
Table 1-b.	First Initial Drug Test QC Samples
Table 1-c.	Second Initial Drug Test QC Samples
Table 2-a-1.	Initial Specimen Validity Test Methods and Instruments (continued on Table 2-a-2 as needed)
Table 2-b-1.	Confirmatory Specimen Validity Test Methods and Instruments (continued on Table 2-b-2 as needed)
Table 2-c-1.	Screening/Differential Specimen Validity Test Methods and Instruments (continued on Table 2-c-2 as needed)
Table 2-d-1.	Initial Specimen Validity Test QC Samples (continued on Table 2-d-2 as needed)
Table 2-d-3.	Confirmatory Specimen Validity Test QC Samples (continued on Table 2-d-4 as needed)
Table 2-d-5.	Screening/Differential Specimen Validity Test QC Samples
Table 3-a.	Primary and Alternate Confirmatory Drug Test Methods

Table 3-b-1.	Primary Confirmatory Drug Test Methods and Instruments – Gas Chromatography (GC)
Table 3-b-2.	Alternate Confirmatory Drug Test Methods and Instruments – GC
Table 3-b-3.	Primary Confirmatory Drug Test Methods and Instruments – Liquid Chromatography (LC)
Table 3-b-4.	Alternate Confirmatory Drug Test Methods and Instruments – LC
Table 3-c-1.	Primary Confirmatory Drug Test Methods and Instruments – Mass Spectrometry (MS)
Table 3-c-2.	Alternate Confirmatory Drug Test Methods and Instruments –MS
Table 3-c-3.	Primary Confirmatory Drug Test Methods and Instruments – Tandem Mass Spectrometry
Table 3-c-4.	Alternate Confirmatory Drug Test Methods and Instruments – Tandem Mass Spectrometry
Table 3-d-1.	Primary Confirmatory Drug Test QC Samples
Table 3-d-2. Table 4-a.	Alternate Confirmatory Drug Test QC Samples Amphetamines Enantiomer Test Methods
Table 4-b.	Amphetamines Enantiomer QC Samples
Table 4-c.	Description of Enantiomer Calculations

Priority Elements for Contracts/Agreements with External Service Providers

- 1. Limiting access to regulated specimen information
- 2. Implementing appropriate safeguards to prevent unauthorized use or disclosure of the information, including implementing applicable federal requirements with regard to regulated specimen and drug test information
- 3. Reporting to the HHS-certified test facility any use or disclosure of the information not provided for by the contract, including incidents that constitute incidents that constitute data breaches of unsecured regulated specimen and drug test information
- 4. Disclosing information to HHS related to regulated specimens and drug tests
- 5. Arranging for disposition of regulated specimen data (i.e., disposal in accordance with specified record retention periods; transfer of records to the HHS-certified test facility upon termination of the agreement)
- 6. Notifying the HHS-certified test facility prior to allowing any subcontractors to have access to regulated specimen and drug test information
- 7. Ensuring that any subcontractors agree to the same restrictions and conditions that apply to the external service provider with respect to regulated specimen and drug test information.

Electronic CCF System Submission List

Items to be submitted for review:

- 1. <u>**Process Overview**</u>. A detailed overview of all processes involving the Federal ECCF from initiation until final disposition, including:
 - Assigning unique specimen identification numbers
 - Initiation of the ECCF
 - o Collection
 - Specimen shipment (labels/seals for specimen bottles and bags)
 - CCF distribution at the end of collection
 - Collector/collection site records storage and disposal
 - o Specimen tracking
 - Test facility accessioning
 - Test facility reporting
 - Test facility records storage and disposal
 - Medical Review Officer review and completion of the CCF
 - MRO reporting
 - MRO records storage and disposal
- 2. Topic Outline of Proposed SOPs An outline of topics to be addressed in:
 - HHS-certified test facility standard operating procedures (SOPs) for accessioning, certification, reporting
 - Procedures/Instructions for other Federal ECCF users including collectors, MROs, and MRO staff

<u>Note</u>: Proposed Federal ECCF instructions or proposed SOP Table of Contents may be submitted <u>Examples</u>: Screenshots, tables of contents

- 3. <u>Training Plans</u> Training for Federal ECCF system users, including:
 - Federal ECCF system users (IITF staff, laboratory staff, collectors, MROs, MRO staff as applicable)
 - Other individuals given access to regulated specimen data (e.g., IT staff)
 - Security awareness training must address forensic records and regulated specimen donor PII

Note: RP must document review and approval of training plans and materials

- 4. **System/Network Diagram** Logical network diagram including, at a minimum:
 - o Firewalls
 - Network security devices
 - o Servers
 - o Workstations

Electronic CCF System Submission List

- o Primary routers/switches
- Remote access devices
- Internet connection(s)
- System Security Plan (SSP) Plan that reflects NIST 800-53 or other recognized security standard, and provides an overview of the security requirements of the system, describes the controls in place or planned for meeting those requirements, and delineates responsibilities and expected behavior of all individuals who access the system.
 - The ability to generate accurate and complete copies of records in both human readable and electronic form suitable for inspection, review, and copying upon request of authorized parties (e.g., the MRO, federal agency, or SAMHSA)
 - Protection of records to enable accurate and ready retrieval through the records retention period
 - o Limiting system access to authorized individuals
 - Secure, computer-generated, time-stamped audit trails to independently record the date and time of operator entries and actions that create, modify, or delete records from the time of initiation of the Federal CCF (changes should be evident when reviewing the original record, and any electronic or paper copy of the original record)
 - Use of authority checks to ensure that only authorized individuals can use the system, electronically sign a record, access the operation or computer system input or output device, alter a record, or perform the operation at hand
- 6. <u>System Validation Plan</u> Plan for testing and evaluating information system security controls to ensure effective implementation.

<u>Note:</u> The HHS-certified test facility must provide documentation of security control testing and evaluation at NLCP inspections.

Examples of records to be provided include

- Periodic records checks
- o Independent security monitoring by IITF/laboratory IT staff
- A report from an independent auditor regarding compliance with relevant industry standards

7. External ECCF Provider Agreement with HHS-Certified Test Facility An

HHS-certified test facility that plans to use an external ECCF system must have a contract/ agreement signed by each laboratory Responsible Person (RP)/IITF Responsible Technician (RT) and an authorized representative of the ECCF provider that:

 Specifies the responsibilities of the ECCF provider and states restrictions and conditions that apply to the ECCF provider with respect to regulated specimen and drug test information

Electronic CCF System Submission List

- Establishes the permitted and required uses and disclosures of regulated specimen and drug test information by the ECCF provider
- o Addresses, at a minimum, these priority elements:
 - Limiting access to regulated specimen information
 - Implementing appropriate safeguards to prevent unauthorized use or disclosure of the information, including implementing applicable federal requirements with regard to regulated specimen and drug test information
 - Reporting to the HHS-certified test facility any use or disclosure of the information not provided for by the contract, including incidents that constitute incidents that constitute data breaches of unsecured regulated specimen and drug test information
 - Disclosing information to HHS related to regulated specimens and drug tests
 - Arranging for disposition of regulated specimen data (i.e., disposal in accordance with specified record retention periods; transfer of records to the HHS-certified test facility upon termination of the agreement)
 - Notifying the HHS-certified test facility prior to allowing any subcontractors to have access to regulated specimen and drug test information
 - Ensuring that any subcontractors agree to the same restrictions and conditions that apply to the ECCF provider with respect to regulated specimen and drug test information.

Note: The agreement/contract must be provided for NLCP review with the initial ECCF submission and with other ECCF system documentation at each inspection.

Table 1-a

Initial Drug Test Methods and Instruments

		First Ini	itial Drug Test	Methods and I	nstruments		
First Initial Drug Test	THCA (marijuana metabolites)	BZE (cocaine metabolites)	MOR (opiate metabolites)	6-AM	PCP	MAMP (amphetamines)	MDMA
Kit and							
Manufacturer							
Analyzer and							
Manufacturer							
Number of							
Analyzer Units							
Calibration Method							
Maximum Batch							
Size							
*If "Other" is	s selected, pleas	se specify:					
		Second I	nitial Drug Tes	st Methods and	I Instruments		
Second Initial Drug Test	THCA (marijuana metabolites)	BZE (cocaine metabolites)	MOR (opiate metabolites)	6-AM	PCP	MAMP (amphetamines)	MDMA
Kit and		,					
Manufacturer							
Analyzer and							
Manufacturer							
Number of							
Analyzer Units							
Calibration Method							
Maximum Batch							
Size							
*If "Other" is	s selected, pleas	se specify:					

THCA = Δ 9-tetrahydrocannabinol-9-carboxylic acid BZE = benzoylecgonine

6-AM = 6-acetylmorphine MOR = morphine PCP = phencyclidine

MAMP = methamphetamine

MDMA = methylenedioxymethamphetamine

First Initial Drug Test QC Samples

1st initi test	al drug QC	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	BQC 1	BQC 2
T 110.4	Conc.										
THCA	Matrix Source										
	Conc.										
BZE	Matrix										
	Source										
	Conc.										
MOR	Matrix										
	Source										
6-AM	Conc. Matrix										
0 / 11	Source										
	Conc.										
PCP	Matrix										
	Source										
	Conc.										
MAMP	Matrix										
	Source Conc.										
MDMA	Matrix										
	Source										
* f "(*If "Other" is selected, please specify:										
	ROC - blind quality control comple										

BQC = blind quality control sample

Table 1-c

Second Initial Drug Test QC Samples

2nd initia test QC		Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	BQC 1	BQC 2
	Conc.										
THCA	Matrix Source										
	Conc.										
BZE	Matrix										
	Source Conc.										
MOR	Matrix										
	Source										
	Conc.										
6-AM	Matrix Source										
	Conc.										
PCP	Matrix										
	Source Conc.										
MAMP	Matrix										
	Source										
	Conc.										
MDMA	Matrix Source										
*lf "(*If "Other" is selected, please specify:										

Initial Specimen Validity Test Methods and Instruments

Initial SVT	Creatinine	SG	лЦ	Nitrite	Gen.Oxid.	Other:	Other:
	Creatinine	30	рН	Millite	Gen.Oxid.		
Method		4 dec. place refractometer					
Kit Manufacturer							
Analyzer and							
Manufacturer							
Number of							
Analyzer Units							
Unit of	mg/dL			mcg/mL			
Measurement	IIIg/dE			mcg/me			
Target Analyte of							
Assay							
Target Analyte of Calibrator							
Calibration Method							
LOD							
LOQ							
ULOL							
Carryover Limit							
Maximum Batch							
Size							
*If "Other" i	s selected, please	specify:					
SG = specific gravity		LOD = limit of detection	on	ULOL= upper limit of linearity			

Gen. Oxid. = general oxidant

LOQ = limit of quantitation

Initial Specimen Validity Test Methods and Instruments

Initial SVT cont.	Other:	Other:	Other:	Other:	Other:	Other:	Other:				
Method											
Kit Manufacturer											
Analyzer and											
Manufacturer											
Number of											
Analyzer Units											
Unit of											
Measurement											
Target Analyte of											
Assay											
Target Analyte of											
Calibrator											
Calibration Method											
LOD											
LOQ											
ULOL											
Carryover Limit											
Maximum Batch											
Size											
*If "Other" is	*If "Other" is selected, please specify:										

Confirmatory Specimen Validity Test Methods and Instruments

Confirmatory SVT	Creatinine	SG	pН	Nitrite	Other:	Other:	Other:					
· · · · · · · · · · · · · · · · · · ·			I.									
Method		4 dec. place refractometer										
Kit Manufacturer												
Analyzer and Manufacturer												
Number of Analyzer Units												
Unit of Measurement	mg/dL			mcg/mL								
Target Analyte of												
Assay												
Target Analyte of Calibrator												
Calibration Method												
LOD												
LOQ												
ULOL												
Carryover Limit												
Maximum Batch Size												
*If "Other" is s	*If "Other" is selected, please specify:											

Table 2-b-2

Confirmatory Specimen Validity Test Methods and Instruments

Confirmatory SVT	Other:	Other:	Other:	Other:	Other:	Other:
cont.						
Method						
Kit Manufacturer						
Analyzer and						
Manufacturer						
Number of Analyzer						
Units						
Unit of						
Measurement						
Target Analyte of						
Assay						
Target Analyte of						
Calibrator						
Calibration Method						
LOD						
LOQ						
ULOL						
Carryover Limit						
Maximum Batch						
Size						
*If "Other" is s	selected, pleas	e specify:				

Screening/Differential Specimen Validity Test Methods and Instruments

Screening/Differential SVT	SG	рН	Other:	Other:	Other:				
Method									
Kit Manufacturer									
Analyzer and									
Manufacturer									
Number of Analyzer									
Units									
Unit of Measurement									
Target Analyte of Assay									
Target Analyte of									
Calibrator									
Calibration Method									
LOD									
LOQ									
ULOL									
Carryover Limit									
Maximum Batch Size									
*If "Other" is selected, please specify:									

Table 2-c-2	Screeni	ng/Differential	Specimen Valid	lity Test	
Screening/Differential	Other:	Other:	Other:	Other:	Other:
SVT cont.					
Method					
Kit Manufacturer					
Analyzer and					
Manufacturer					
Number of Analyzer					
Units					
Unit of Measurement					
Target Analyte of Assay					
Target Analyte of					
Calibrator					
Calibration Method					
LOD					
LOQ					
ULOL					
Carryover Limit					
Maximum Batch Size					
*If "Other" is selecte	ed, please specify:				

Initial Specimen Validity Test QC Samples

Initial	SVT QC	Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
	Target value										
Creatinine											
	Source										
	Target value										
SG	Matrix										
	Source										
	Target value										
pН	Matrix										
-	Source										
	Target value										
Nitrite	Matrix										
	Source										
	Target value										
Gen Oxid	Matrix										
	Source										
*	If "Other" is a	selected, please	specify:								

Initial SVT C		Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5		
Other (enter name):	Target Value Matrix											
Other (enter name):	Source Target Value Matrix											
Other (enter name):	Source Target Value											
Other (enter hame).	Matrix Source											
Other (enter name):	Target Value Matrix											
Other (enter name):	Source Target Value Matrix											
Other (enter name):	Source Target Value											
Other (enter name):	Matrix Source											
Other (enter name):	Target Value Matrix											
Other (enter name):	Source Target Value Matrix											
*If "Other" is	Matrix Image: Source Image: Source Image: Source *If "Other" is selected, please specify: Image: Source Image: Source											

Confirmatory Specimen Validity Test QC Samples

Confirm	atory SVT	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
	Target Value									
Creatinine	Matrix									
	Source									
	Target Value									
SG	Matrix									
	Source									
	Target Value									
pН	Matrix									
	Source									
	Target Value									
Nitrite	Matrix									
	Source									
	Target Value									
Gen Oxid	Matrix									
	Source									
*lf	⁻ "Other" is s	selected, please	specify:							

Confirmatory SV	T QC cont.	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
Other (enter name):	Target Value									
	Matrix Source									
Other (enter neme)	Target Value									
	Matrix									
	Source									
Other (enter name):	Target Value Matrix									
	Source									
	Target Value									
Other (enter name):	Matrix									
	Source									
Other (enter name):	Target Value									
	Matrix									
	Source									
Other (enter name):	Target Value									
	Matrix									
	Source									
Other (enter name):	Target Value Matrix									
	Source									
	Target Value									
Other (enter name):	Matrix									
	Source									
*If "Other" is s		se specify:		l			ı	ļ	1	

Screening/Differential Specimen Validity Test QC Samples

Screening/Differ QC	rential SVT	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
	Target Value									
Specific Gravity	Matrix									
	Source									
	Target Value									
рН	Matrix									
	Source									
Other (enter name):	Target Value Matrix									
	Source									
	Target Value									
Other (enter name):	Matrix									
	Source								Control 4	
	Target Value									
Other (enter name):	Matrix									
	Source									
	Target Value									
Other (enter name):	Matrix									
	Source									
Other (enter neme)	Target Value									
Other (enter name):	Matrix									
	Source									
Other (enter name):	Target Value									
	Matrix									
	Source									
*If "Other" is a	selected, ple	ase specify:								

			Prim	ary Confir	matory Dr	ug Test M	lethods		
Primary Confirmatory Drug Test	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Method									
Internal Standard									
Int. Std. Isotope									
Type and Number									
Int. Std. Conc.*									
LOD*									
LOQ*									
ULOL*									
Carryover Limit*									
Maximum Batch									
Size									
*If "Other" is sele	cted, pleas	e specify:							
			Alter	nate Conf	irmatory D	orug Test I	Methods		
Alternate Confirmatory Drug Test	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.*	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD*	THCA	BZE		MOR	6-AM	PCP	AMP	MAMP	MD
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD* LOQ*	THCA	BZE		MOR	6-AM	PCP	AMP	MAMP	
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD* LOQ* ULOL*		BZE		MOR	6-AM	PCP	AMP	MAMP	
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD* LOQ* ULOL* Carryover Limit*		BZE		MOR	6-AM	PCP	AMP		
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD* LOQ* ULOL* Carryover Limit* Maximum Batch		BZE		MOR	6-AM				
Confirmatory Drug Test Method Internal Standard Int. Std. Isotope Type and Number Int. Std. Conc.* LOD* LOQ* ULOL* Carryover Limit*				MOR	6-AM				

* ng/mL

COD = codeine AMP = amphetamine MDA = methylenedioxyamphetamine

MDEA = methylenedioxyethylamphetamine

DMA	MDA	MDEA

DMA	MDA	MDEA

Primary Confirmatory Drug Test Methods and Instruments - Gas Chromatography

	Prin	nary Confirma	tory Drug Test	Methods and	Instruments -	Gas Chromatogra	aphy	
Primary Confirmatory Drug						AMPHETAMINE	S (select analytes from dr	opdown lists below)
Test - Gas Chromatography	THCA	BZE	COD/MOR	6-AM	PCP			
Extraction Method								
Volume Used (mL)								
Hydrolysis Method								
Derivatizing Reagent								
Split/Splitless Injection								
Inj. Port Temp (°C)								
Isothermal or Gradient								
Column Type								
Column Length (m)								
Instrument Manufacturer								
Number of Units								
			GC/GC Method	s: provide additior	al information be	low		•
Cryotrapping (Y/N)								
2nd GC Column Type								
2nd GC Column Length								
(m)								
*If "Other" is se	elected, please	specify						

Alternate Confirmatory Drug Test Methods and Instruments - Gas Chromatography

	Alternate Confirmatory Drug Test Methods and Instruments - Gas Chromatography											
Primary Confirmatory Drug Test - Gas Chromatography	THCA	BZE	COD/MOR	6-AM	PCP	AMPHETAMINE	S (select analytes from dr	opdown lists below)				
Extraction Method												
Volume Used (mL)												
Hydrolysis Method												
Derivatizing Reagent												
Split/Splitless Injection												
Inj. Port Temp (°C)												
Isothermal or Gradient												
Column Type												
Column Length (m)												
Instrument Manufacturer												
Number of Units												
			GC/GC Method	s: provide additior	al information belo	W						
Cryotrapping (Y/N)												
2nd GC Column Type												
2nd GC Column Length												
(m)												
*If "Other" is s	elected, please	specify										

Table 3-b-3

Primary Confirmatory Drug Test Methods and Instruments - Liquid Chromatography

Primar	Primary Confirmatory Drug Test Methods and Instruments - Liquid Chromatography											
Primary Confirmatory Drug Test - Liquid Chromatography	THCA	BZE	COD/MOR	6-AM	PCP	AMP/MAMP	MDMA/MDA/ MDEA					
Extraction Method												
Volume Used (mL)												
Hydrolysis Method												
Injection Volume												
Isocratic or Gradient												
Guard Column (Y/N)												
Flow Rate (mL/min)												
Temperature (°C)												
Column Type												
Column Length (cm)												
Column Diameter												
Column Particle Size												
A Solvent (Buffer)												
Buffer Type												
Molarity												
pH B Solvent (Organic)												
Component 1												
Component 2												
Component 3												
Component Ratio (1:2:3)												
Instrument Manufacturer												
Number of Units												
*If "Other" is selected, ple	ease specify											

Table 3-b-4

Alternate Confirmatory Drug Test Methods and Instruments - Liquid Chromatography

			-				
Alternate Confirmatory Drug Test - Liquid Chromatography	THCA	BZE	COD/MOR	6-AM	PCP	AMP/MAMP	MDMA/MDA/ MDEA
Extraction Method							
Volume Used (mL)							
Hydrolysis Method							
njection Volume							
socratic or Gradient							
Guard Column (Y/N)							
Flow Rate (mL/min)							
Temperature (°C)							
Column Type							
Column Length (cm)							
Column Diameter							
Column Particle Size							
A Solvent (Buffer)							
Buffer Type							
Molarity							
pH							
B Solvent (Organic)							
Component 1							
Component 2 Component 3							
Component Ratio (1:2:3)							
nstrument Manufacturer							
Number of Units							

		Prir	mary Confirma	atory Drug Te	st Methods a	nd Instrument	s - Mass Spec	ctrometry (MS	5)		
Primary Confirmatory Drug Test - Mass Spectrometry	THCA	BZE	COD	MOR	6-AM	РСР	AMP	MAMP	MDMA	MDA	MDEA
Instrument											
Manufacturer											
Number of Units											
Inlet System											
Ionization											
Ion Focus											
Full Scan Mass											
Range											
Calibration Type											
Analyte Quantifier Ion											
Analyte Qualifier Ion 1*											
Analyte Qualifier Ion 2*											
Analyte Qualifier Ion 3											
Int. Std. Quantifier Ion											
Int. Std. Qualifier Ion 1*											
Int. Std. Qualifier Ion 2											
*If "Other" is se	elected, pleas	e specify									

		Alter	nate Confirma	atory Drug Te	st Methods ar	nd Instruments	s - Mass Spec	ctrometry (MS)		
Alternate Confirmatory Drug Test - Mass Spectrometry	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MDMA	MDA	MDEA
Instrument Manufacturer											
Number of Units											
Inlet System											
Ionization											
Ion Focus											
Full Scan Mass Range											
Calibration Type											
Analyte Quantifier Ion											
Analyte Qualifier Ion 1*											
Analyte Qualifier Ion 2*											
Analyte Qualifier Ion 3											
Int. Std. Quantifier Ion											
Int. Std. Qualifier Ion 1*											
Int. Std. Qualifier ion 2											
*If "Other" is se	elected, pleas	e specify									

Primary Confirmatory Drug Test Methods and Instruments -Tandem Mass Spectrometry

	Р	rimary Conf	irmatory Dru	ig Test Meth	nods and Ins	struments -	Tandem Ma	ss Spectron	netry		
Primary Confirmatory Drug Test - Tandem Mass Spectrometry	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MDMA	MDA	MDEA
Instrument Manufacturer											
Number of Units Ionization											
Configuration											
Calibration Type											
Quantifier Transition*	\rightarrow										
Qualifier Transition 1*	\rightarrow										
Qualifier Transition 2	\rightarrow										
Qualifier Transition 3	\rightarrow										
Int. Std. Quantifier Transition*	→	<i>→</i>	÷	\rightarrow	\rightarrow	÷	÷	÷	\rightarrow	\rightarrow	\rightarrow
Int. Std. Qualifier Transition 1*	→	÷	→	→	\rightarrow	→	<i>→</i>	\rightarrow	→	→	\rightarrow
Int. Std. Qualifier Transition 2	÷	÷	÷	÷	\rightarrow	÷	÷	÷	>	>	\rightarrow
Int. Std. Qualifier Transition 3	→	÷	÷	<i>→</i>	\rightarrow	÷	÷	÷	→	→	\rightarrow
*If "Other" is sele	cted, please	specify									

Alternate Confirmatory Drug Test Methods and Instruments -Tandem Mass Spectrometry

Alternate Confirmatory Drug Test Methods and Instruments - Tandem Mass Spectrometry											
Alternate Confirmatory Drug Test - Tandem Mass Spectrometry	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MDMA	MDA	MDEA
Instrument Manufacturer											
Number of Units											
Ionization											
Configuration											
Calibration Type											
Quantifier Transition*	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow
Qualifier Transition 1*	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow
Qualifier Transition 2	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow
Qualifier Transition 3	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow	\rightarrow
Int. Std. Quantifier Transition*	\rightarrow	÷	÷	\rightarrow	\rightarrow	\rightarrow	<i>→</i>	÷	\rightarrow	\rightarrow	→
Int. Std. Qualifier Transition 1*	\rightarrow	÷	→	<i>→</i>	\rightarrow	\rightarrow	→	→	<i>→</i>	\rightarrow	→
Int. Std. Qualifier Transition 2	<i>→</i>	<i>→</i>	→	<i>→</i>	<i>→</i>	→	<i>→</i>	<i>→</i>	<i>→</i>	<i>→</i>	<i>→</i>
Int. Std. Qualifier Transition 3	→	<i>→</i>	<i>→</i>	→	→	→	→	<i>></i>	→	→	→
*If "Other" is sele	*If "Other" is selected, please specify										

Table 3-d-1

Primary Confirmatory Drug Test QC Samples

Primary Conf	irmatory Drug Test QC	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
ТНСА	Concentration Matrix Source									
BZE	Concentration Matrix Source									
COD	Concentration Matrix Source									
MOR	Concentration Matrix Source									
6-AM	Concentration Matrix Source									
РСР	Concentration Matrix Source									
AMP	Concentration Matrix Source									
MAMP	Concentration Matrix Source									
MDMA	Concentration Matrix Source									
MDA	Concentration Matrix Source									
MDEA	Concentration Matrix Source									
*If "Othe	Source Source *If "Other" is selected, please specify									

Table 3-d-2

Alternate Confirmatory Drug Test QC Samples

Alternate Confirr	natory Drug Test QC	Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
THCA	Concentration Matrix Source										
BZE	Concentration Matrix Source										
COD	Concentration Matrix Source										
MOR	Concentration Matrix Source										
6-AM	Concentration Matrix Source										
PCP	Concentration Matrix Source										
AMP	Concentration Matrix										
MAMP	Source Concentration Matrix										
MDMA	Source Concentration Matrix										
MDA	Source Concentration Matrix										
MDEA	Source Concentration Matrix										
*If "Othe	Source Source Image: Control of the second										

Table C-4-aAMPS Enantiomer Test Methods

Amphetamines Enantiomer Drug Test	D-AMP	L-AMP	D-MAMP	L-MAMP
Method				
Internal Standard				
Int. Std. Isotope Type and				
Number				
Int. Std. Conc.*				
LOD*				
LOQ*				
ULOL*				
Carryover Limit*				
*If "Other" is selected, p	please specify:			
Extraction Method				
Volume Used (mL)				
Derivatizing Reagent				
Enantiomer Calculation				
*If "Other" is selected, p	please specify:			
Reflex ALL AMPs positive specimens for DL testing?				

* ng/mL

AMPS Ena	intiomer QCs	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
	Concentration									
D-AMP	Matrix									
	Source									
	Concentration									
L-AMP	Matrix									
	Source									
	Concentration									
D-MAMP	Matrix									
	Source									
	Concentration									
L-MAMP	Matrix									
	Source									
*If "Other	is selected, pleas	se specify:								

* ng/mL

Describe in detail the method used to calculate the percentages of Dand L-enantiomers (e.g., use of peak abundances, concentrations, use of an internal standard). Indicate if concentrations are calculated that can be compared to the total methamphetamine result obtained in the amphetamines method.

Type description below: