# URINE LABORATORY INFORMATION CHECKLIST

# NATIONAL LABORATORY CERTIFICATION PROGRAM (NLCP)

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# NATIONAL LABORATORY CERTIFICATION PROGRAM URINE LABORATORY CHECKLIST

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Urine, Laboratory October 2010

#### I. URINE LABORATORY INFORMATION CHECKLIST

#### A. Instructions for the Laboratory

#### **Pre-inspection Materials**

Before each scheduled inspection, the NLCP sends instructions to the laboratory listing the required pre-inspection materials with due dates for submission. The required materials depend on the inspection type (e.g., initial inspection, maintenance inspection, records audit, special inspection). The following describes some items that may be required.

#### 1. NLCP Urine Laboratory Information Checklist (Sections B and C)

The laboratory provides up-to-date information to the NLCP on its drug testing operation (i.e., staffing, facility, and procedures) using the NLCP Urine Laboratory Information Checklist (Sections B and C). The information is maintained in NLCP records and is verified by the inspection team (i.e., inspectors, records auditors) at each NLCP inspection.

#### 2. <u>Laboratory Operation Schedule/Inspection Schedule</u>

The laboratory provides a schedule of its operations to the NLCP, listing the days and hours for various processes (e.g., receiving, accessioning, initial testing, confirmation aliquotting, confirmatory drug test extractions, certification). Using this schedule, NLCP staff prepare a tentative schedule for the inspection team. The lead inspector determines the final schedule for the inspection team at most NLCP inspections. The lead auditor determines the final schedule for a records audit.

#### 3. Key Staff Interview List

The laboratory provides a Staff Interviews List to the NLCP, listing key staff, their job titles, and work schedules. NLCP staff select individuals from the list to be interviewed at the inspection and return the list to the laboratory, instructing the laboratory to ensure that the selected individuals are available for interview during the inspection. In addition to interacting with laboratory staff in the course of the inspection, the inspection team conducts formal interviews (i.e., 10-15 minutes each) with the selected staff members to evaluate their knowledge and ability to fulfill job duties.

#### 4. Laboratory Computer Systems (Section P)

To facilitate the inspection of the laboratory's computer system, the NLCP directs the laboratory to perform a self-assessment using Section P, Laboratory Computer Systems. The laboratory provides the completed Section P to the inspection team at the beginning of the inspection.

#### 5. Floor plan of the laboratory

#### 6. <u>Laboratory data packages</u>

The laboratory provides two data packages to the NLCP: one for a positive specimen and one for a specimen that was reported as adulterated, substituted, or invalid based on specimen validity testing (i.e., invalid-abnormal pH, invalid-inconsistent creatinine and specific gravity results, or invalid-possible <adulterant>activity). These data packages should contain all chain of custody forms, worksheets, initial drug test data, screening/differential specimen validity test data, initial specimen validity test data, confirmatory specimen validity test data, confirmatory drug test data, and reports pertaining to the specimen. The program-required format for data packages is described in Section R of the NLCP Manual for Urine Laboratories. These must be recent specimens, processed since the last NLCP inspection using the laboratory's current procedures.

#### 7. Hotel list

The laboratory provides a list of several hotels/motels located in close proximity to the laboratory and to the airport. Hotels selected should ensure the safety and welfare of the inspectors during the inspection.

#### 8. Directions

The laboratory provides a clear, precise map with directions describing the routes from the airport to the hotels and from the hotels to the laboratory.

#### **Non-Negative Specimen List (NNSL)**

Prior to each NLCP inspection that includes a records audit, the NLCP notifies the laboratory of the specified audit period (e.g., the six-month period ending one month prior to the month of the inspection). The laboratory is required to identify all regulated specimens reported during that time period as positive, adulterated, substituted, invalid, rejected, reconfirmed, or failed to reconfirm. In addition, the laboratory must identify all specimens received for testing from an Instrumented Initial Test Facility (IITF), *including specimens reported as negative*. The laboratory must submit to the NLCP a list of these specimens, with specific information for each specimen. The laboratory also provides a monthly summary for the records audit period listing the numbers of regulated specimens reported as positive, adulterated, substituted, invalid, negative, rejected, reconfirmed, or failed to reconfirm.

The NLCP provides instructions for the NNSL to the laboratory prior to the inspection. These instructions include, but are not limited to, the following:

#### 1. Format for NNSL spreadsheet

#### 2. NNSL categories:

- The laboratory will provide information concerning results reported for the following NNSL categories: amphetamine/methamphetamine/enantiomers, methylenedioxymethamphetamine (MDMA)/methylenedioxyamphetamine (MDA)/ methylenedioxyethylamphetamine (MDEA), benzoylecgonine, opiates, phencyclidine, cannabinoids, adulterated, invalid, substituted, and rejected.
- If the laboratory has tested a regulated specimen for an additional Schedule I or II drug upon request of a Federal Agency and reported the specimen as positive (i.e., drug present at or above the cutoff used for the test), the laboratory must submit a separate NNSL sheet for that drug.
- If no specimen is identified for a specific category, the laboratory must submit that sheet indicating "None."

#### 3. Specimens to be included on the NNSL:

- Specimens reported positive, adulterated, substituted, invalid, rejected, reconfirmed, and failed to reconfirm.
- Specimens received for testing from an IITF, including those reported negative.

The laboratory must remove all known NLCP performance testing (PT) samples.

#### 4. Requirements for records assembly

The NLCP selects specimens from the submitted NNSL for review during the inspection and provides the selected list to the laboratory and to the lead auditor. The laboratory must organize and assemble records for each of the selected specimens to facilitate their review by the audit team during the inspection. At a minimum, records must be assembled by NNSL category and in chronological order, to facilitate their location within labeled storage folders/boxes. Auditors must be able to retrieve all records (excluding failed batches) pertaining to a specimen on the selected NNSL with a minimum of assistance from the laboratory staff.

During the inspection, the lead auditor and the Responsible Person (RP) will prepare an inventory of records for the selected specimens on the NNSL that were not available for review. The RP must forward the missing records to the NLCP for subsequent review and follow-up.

#### **Laboratory Preparation Criteria List**

Prior to each inspection, the NLCP sends a Laboratory Preparation Criteria List to the laboratory, listing materials that must be available for the inspection team upon their arrival at the laboratory. Materials include a copy of the standard operating procedures (SOP) manual for each inspection team member, NLCP PT records, personnel files, quality assurance (QA)/quality control (QC) records, reagent records, validation records, a timeline of any changes in QC criteria and control acceptance limits during the records audit period, and documentation of security procedures (e.g., access rosters and visitor logs for each secured area). Other items may be requested for review prior to or during the inspection.

B.	<b>Laboratory Information</b> (completed by the laboratory)	
B-1.	Name of Laboratory:Address:	
	City, State, ZIP:	
	Telephone: ( ) FAX: ( ) _ e-Mail:	
B-2.	Responsible Person(s) RP's name: RP's title:	
	RP's name:RP's title:	
	RP's name:RP's title:	
	Alternate Responsible Person(s) Alt-RP's name: Alt-RP's title:	
	Alt-RP's name:Alt-RP's title:	
B-3.	I certify that the statements and information present and C are true and correct as of this date. I affirm to read and are familiar with the current version of the Urine Laboratories. I also recognize my responsible amended Sections B and C to the inspectors at the inspection if changes are made between the date of the inspection.	hat the key staff have NLCP Manual for lity for providing beginning of the
Note:	Any false, fictitious, or fraudulent statements or information pasections B and C or misrepresentations relative thereto may be Law and could subject you to prosecution, monetary penaltie U.S.C. 1001; 31 U.S.C. 3801-812).	<u>riolate Federal</u>
	Signature, Responsible Person	Date
	Signature, Responsible Person	Date
	Signature, Responsible Person	Date

-	<del></del>
-	
-	
D	ays/hours of operation of the forensic urine drug testing laboratory:
_	days per week;hours per day
lf	≤ 6 days, indicate the day(s) that the laboratory is routinely <b>not</b> operational:
	Poes the laboratory have a U.S. Drug Enforcement Agency (DEA) egistration?  YES
lf	YES, for which schedules?
	122N33N45
lf	NO, explain how controlled reference materials are acquired:
-	
	escribe the State licensure requirements for urine forensic toxicology or the State in which the laboratory is located:
-	
Li	ist laboratory certifications/licenses:
_	States (List):
_	CLIA/HCFA <sup>1</sup> (List Specialties):
	CAP² (List Specialties):
<sup>1</sup> C	Others (Specify): Clinical Laboratory Improvement Amendments(CLIA)/Health Care Financing Administration (H College of American Pathologists (CAP)

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B-9. List name, job title, education, and licenses/certifications for the following key staff:

Note: (1) May attach separate sheet listing additional key staff

(2) Indicate (\*) individuals new to the positions in the last 6 months

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

	a.	Is licensure and/or certification required for any of the above positions in the State in which the laboratory is located?	YES	NO
	If YE	ES, describe requirements:		
B-10.		ere is more than one RP, briefly describe how the RPs share the onsibilities for the various laboratory operations and procedures.		
B-11.		cribe the administrative relationships that exist for the <b>key staff</b> of orensic drug testing laboratory (see B-9 above):		
	a.	To whom does the RP(s) report?		
	b.	Who evaluates the performance of the RP(s)?		
	C.	What staff administratively report <i>directly</i> to the RP(s)?		
	d.	The RP(s) evaluates the performance of which staff members?		
	e.	Which staff members do not report to the RP(s)?		
B-12.	Does	s the laboratory test any Federal agency specimens for drugs		
	othe	r than those specified in the HHS Guidelines?  ES, list the drug(s) and answer a and b below:	YES	NO

	a.	Does the laboratory have a copy of the HHS waiver for a Federal agency to test the additional drug(s) on a routine basis	s? YES	NC
	b.	Does the laboratory maintain written authorization from Federa agencies to test the additional drug(s) on a case-by-case basis		NC
B-13.	drugs	ge number of specimens analyzed by the laboratory each day to of abuse during the six months preceding submission of cons B and C (both regulated and non-regulated specimens)		
		Specify the months		
		Total specimens/day		
	How v	vas this number derived?		
B-14.		otal number of staff who have authorized access to the secure sic drug testing laboratory facility:		
		individuals		

B-15. List the total numbers of staff who are trained and routinely perform the following activities *for regulated specimens*:

Activity	No. of Individuals
Accessioning	
Initial drug testing	
Screening/initial specimen validity testing	
Confirmatory specimen validity testing	
Extraction	
Confirmatory drug testing	
Certification	

#### **C.** Laboratory Procedures (completed by the laboratory)

**NOTE:** Any computer interface communicating any form of data from an HHS-certified IITF to an HHS-certified laboratory must be approved by the NLCP prior to implementation. The IITF and/or laboratories must submit a detailed plan to the NLCP for review. Affected test facilities will be subject to inspection to verify compliance with NLCP requirements. HHS-certified laboratories are prohibited from transmitting data to an HHS-certified IITF through a computer interface.

C-1. Provide a description of the laboratory's procedures for the following:

#### Security

- Building
- Department
- Specimens
- Records
  - Note: (1) Insert here.
    - (2) Do not exceed a total of one page.
- C-2. Provide a description of the laboratory's 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.
- Handling problems with specimen bottles and/or custody and control forms.
- Assignment of laboratory accession numbers.
- Location of temporary storage area(s).
  - Note: (1) Insert here.
    - (2) Do not exceed a total of one page.
- C-3. Provide a description of the laboratory's procedures for the following:

#### **Aliquotting Procedures**

- Aliquotting of the original specimen bottles (i.e., who and where).
- The actual aliquotting procedure (pouring or pipetting and amounts) used for preparing aliquots for initial drug tests, specimen validity tests, and confirmatory drug tests.
- Transfer of aliquots from the individuals performing the aliquotting to those who will be testing the aliquots.
  - Note: (1) Insert here.
    - (2) Do not exceed a total of one page.

C-4.	Provide a description of the laboratory's procedures for the following:
	<ul> <li>Specimen Accessioning</li> <li>Introduction and/or aliquotting of blind controls into the test batches by accessioning personnel.</li> <li>If applicable, preparation and submission of blind samples as donor specimens from external sources.</li> </ul>
	Note: (1) <u>Insert here.</u> (2) <u>Do not exceed a total of one page.</u>
C-5.	Provide a description of the laboratory's procedures for the following:
	<ul> <li>First and Second Initial Drug Tests</li> <li>Handling and testing of aliquots by laboratory personnel.</li> <li>Maintenance of chain of custody during the testing.</li> </ul>
	Note: (1) <u>Insert here.</u> (2) <u>Do not exceed a total of one page.</u>
C-6.	Provide a description of the laboratory's procedures for the following:
	<ul> <li>First and Second Initial Drug Tests</li> <li>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, are regulated and non-regulated specimens tested in the same batch?).</li> <li>The distribution of specimens and QC samples within each batch.</li> </ul>
	<ul> <li>The acceptance criteria for each control (open and blind) in each batch and when and by whom these are evaluated and documented.</li> <li>The criteria for accepting all donor specimen results or only a partial number of donor specimens in a batch.</li> </ul>
	Note: (1) <u>Insert here.</u> (2) <u>Do not exceed a total of one page.</u>
C-7.	Provide the following information for the first and second Initial Drug Tests:
	Describe the procedure(s) and acceptance criteria for calibration:

-	nalytes:		
-			
-			
Р	rovide a de	escripti	on of the laboratory's procedures for the following:
s •	Handling	and te	y Tests (Initial, Confirmatory and Screening/Differer esting of aliquots by laboratory personnel. for chain of custody during the testing.
	Note:	(1) (2)	Insert here.  Do not exceed a total of one page.
			or a legible flowchart that comprehensively describes t nen Validity Testing.
	Note:	(1) (2)	Insert here.  Do not exceed a total of one page.
а	outlin	e/flow@	inges to the specimen validity testing chart during the time period of the NNSL audit, with a date of each change.
-			
_			

The distribution of specimens and QC samples within each batch.
 The acceptance criteria for each control (open and blind) in each batch.

How batches are constituted.

• The acceptance criteria for each control (open and blind) in each batch and when and by whom these are evaluated and documented.

	Note:		Insert here.  Do not exceed a total of one page.
C-11.			ng information for the Specimen Validity Tests (i.e., initial, screening/differential tests):
	Describe the	proce	dures and acceptance criteria for calibration:
	Describe the measurands		od used to calculate the concentrations/responses of
C-12.	Provide a de	scription	on of the laboratory's procedures for the following:
	Confirmator	y Dru	g Tests

• The criteria for accepting all donor specimen results or only a partial number

of donor specimens in a batch.

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(2) Do not exceed a total of one page.

• Handling and testing of aliquots by laboratory personnel.

• Maintenance of chain of custody during the testing.

Note: (1) <u>Insert here.</u>

C-13. Provide a description of the laboratory's procedures for the following:

#### **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, are regulated and non-regulated specimens tested in the same batch?).
- The distribution of the donor specimens and QC samples within each batch.
- 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 a donor specimen result, reextracting a specimen, or reinjecting a specimen.
  - Note: (1) Insert here.
    - (2) Do not exceed a total of one page.
- C-14. Provide the following information for the Confirmatory Drug Tests:

Describe the requirements for calibration including criteria for exclusion	sion
of unsatisfactory calibrators:	
Describe the method used to calculate the concentrations of analyte for each calibration procedure used by the laboratory:	es:

C-15. Provide a description of the laboratory's procedures for the following:

#### **Certification/Reporting Procedures**

- Review of all calibration data and control data.
- Review of chain of custody forms.
- Review of specimen data.
- Documentation and certification of results.
- Release/reporting of results.
- Verification of information (e.g., CCF and computer resident result).

Note: (1) Insert here.

(2) Do not exceed a total of one page.

C-16. Provide a description of the laboratory's procedures for the following: **Electronic Reporting Procedures**  Release of computer-generated electronic reports. Note: (1) Insert here. (2) Do not exceed a total of one page. C-17. Provide 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 C-18. Does the laboratory use an off-site computer information system?

YES NO If YES.

If YES,
Address:
City, State, ZIP:

C-19. Provide a description of the laboratory's procedures for the following:

#### **Disposition of Specimens and Aliquots**

- Handling of original specimen bottles and aliquots after testing is completed.
- Procedure for transferring positive, adulterated, substituted, and invalid specimens to long-term frozen storage.
  - Note: (1) Insert here.
    - (2) <u>Do not exceed a total of one page.</u>

#### **Complete the C Tables:**

Table C-1-a.	First and Second Initial Drug Test Methods and Instruments
Table C-1-b.	First Initial Drug Test QC samples
Table C-1-c.	Second Initial Drug Test QC samples
Table C-2-a-1.	Initial Specimen Validity Test Methods and Instruments (continued on <b>Table C-2-a-2</b> as needed)
Table C-2-b-1.	Confirmatory Specimen Validity Test Methods and Instruments (continued on <b>Table C-2-b-2</b> as needed)
Table C-2-c-1.	Screening/Differential Specimen Validity Test Methods and Instruments (continued on <b>Table C-2-c-2</b> as needed)
Table C-2-d-1.	Initial Specimen Validity Test QC samples (continued on <b>Table C-2-d-2</b> as needed)
Table C-2-d-3.	Confirmatory Specimen Validity Test QC samples (continued on <b>Table C-2-d-4</b> as needed)
Table C-2-d-5.	Screening/Differential Specimen Validity Test QC samples
Table C-3-a.	Primary and Alternate Confirmatory Drug Test Methods
Table C-3-b-1.	Primary Confirmatory Drug Test Methods and Instruments – Gas Chromatography (GC)
Table C-3-b-2.	Alternate Confirmatory Drug Test Methods and Instruments – GC
Table C-3-b-3.	Primary Confirmatory Drug Test Methods and Instruments – Liquid Chromatography (LC)
Table C-3-b-4.	Alternate Confirmatory Drug Test Methods and Instruments – LC
Table C-3-c-1.	Primary Confirmatory Drug Test Methods and Instruments – Mass Spectrometry (MS)
Table C-3-c-2.	Alternate Confirmatory Drug Test Methods and Instruments –MS
Table C-3-c-3.	Primary Confirmatory Drug Test Methods and Instruments – Tandem Mass Spectrometry
Table C-3-c-4.	Alternate Confirmatory Drug Test Methods and Instruments – Tandem Mass Spectrometry
Table C-3-d-1.	Primary Confirmatory Drug Test QC Samples
Table C-3-d-2.	Alternate Confirmatory Drug Test QC Samples

#### Initial Drug Test Methods and Instruments

		First Initia	I Drug Test Me	thods and Ins	truments		
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							
Average Number of federally regulated specimens tested daily							
Average Number of Batches with federally regulated specimens tested daily							
*If "Other" is	s selected, pleas						
			ial Drug Test M	lethods and Ir	nstruments		
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							
	s selected, pleas		6-AM = 6-acetylmorphine		MDMA = methylenedioxy		

THCA =  $\Delta$ 9-tetrahydrocannabinol-9-carboxylic acid MOR = morphine BZE = benzoylecgonine

PCP = phencyclidine

6-AM = 6-acetylmorphine MAMP = methamphetamine MDMA = methylenedioxymethamphetamine

### First Initial Drug Test QC Samples

1st initi		Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	BQC 1	BQC 2
	Conc										
THCA	Matrix										
	Source										
D.7.E	Conc										
BZE	Matrix										
	Source										
MOR	Conc										
WOR	Matrix										
	Source										
6-AM	Conc										
O-Aivi	Matrix										
	Source										
PCP	Conc										
1 01	Matrix Source										
	Conc										
MAMP	Matrix										
	Source										
	Conc										
MDMA	Matrix										
	Source										
	*If "(	Other" is select	ted, please spe	cify:	•		•	•	•	•	

BQC = blind quality control sample

# Second Initial Drug Test QC Samples

2nd initi		Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	BQC 1	BQC 2
test											
	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										
	*If "Other" is selected, please specify:										

#### Table C-2-a-1

# Initial Specimen Validity Test Methods and Instruments

Initial SVT	Creatinine	SG	рН	Nitrite	Gen.Oxid.	Other:	Other:
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 se	l <mark>lected, pleas</mark>	e specify:					

SG = specific gravity
Gen. Oxid. = general oxidant

LOD = limit of detection

ULOL= upper limit of linearity

LOQ = limit of quantitation

### Table C-2-a-2

# 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	selected, pleas	se specify:					

# Table C-2-b-1 Confirmatory Specimen Validity Test Methods and Instruments

Confirmatory SVT	Creatinine	SG	рН	Nitrite	Other:	Other:	Other:
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 se	lected, please	specify:					

### Table C-2-b-2

# Confirmatory Specimen Validity Test Methods and Instruments

Confirmatory SVT	Other:	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:					

### Table C-2-c-1

### 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 selecte	d, please specify:				

# Table C-2-c-2 Screening/Differential Specimen Validity Test Methods and Instruments

Screening/Differential SVT cont.	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 selected	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 selected, please specify:										

# Initial Specimen Validity Test QC Samples

Initial SVT	QC cont.	Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
Other (enter name):	Target Value Matrix										
ŕ	Source										
Other (enter	Target Value										
name):	Matrix										
	Source										
Other (enter	Target Value										
name):	Matrix										
	Source										
Other (enter name):	Target Value										
namej.	Matrix Source										
Other (enter											
name):	Target Value Matrix										
	Source										
Other (enter	Target Value										
name):	Matrix										
	Source										
Other (enter	Target Value										
name):	Matrix										
	Source										
Other (enter	Target Value										
name):	Matrix										
<b>417</b> ()	Source										
*11 "	Otner" is sel	ected, please sp	pecity:								

# Confirmatory Specimen Validity Test QC Samples

Confirmator	ry SVT QC	Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
Creatinine	Target Value Matrix Source										
SG	Target Value Matrix Source										
рН	Target Value Matrix Source										
Nitrite	Target Value Matrix Source										
Gen Oxid	Target Value Matrix Source										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix Source										
*If "Other" i		lease specify:									

# Confirmatory Specimen Validity Test QC Samples

Confirmato	ry SVT QC nt.	Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
Other (enter name):	Target Value Matrix										
	Source										
Other (enter name):	Target Value Matrix										
	Source										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix Source										
Other (enter name):	Target Value Matrix										
*If "Other" i	Source is selected, p	lease specify:									

# Screening/Differential Specimen Validity Test QC Samples

Screening/Diffe QC		Cal 1	Cal 2	Cal 3	Cal 4	Cal 5	Control 1	Control 2	Control 3	Control 4	Control 5
	Target Value										
Specific Gravity											
	Source										
	Target Value										
	Matrix										
	Source Target Value										
Other (enter name):	Matrix										
	Source										
	Target Value										
	Matrix										
	Source										
Other (enter name):	Matrix										
	Source										
Other (enter name):	Matrix										
	Source										
	Target Value										
Other (enter name).	Matrix										
	Source										
Other (enter name):	Target Value										
	Matrix										
	Source										
*If "Other" is	*If "Other" is selected, please specify:										

Primary Confirmatory Drug Test Methods													
Primary Confirmatory Drug Test	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MDMA	MDA	MDEA		
Method													
Internal Standard													
Int. Std. Isotope													
Type and Number													
Int. Std. Conc.*													
LOD*													
LOQ*													
ULOL*													
Carryover Limit*													
Maximum Batch Size													
Average Number of Batches with HHS specimens tested daily													
*If "Other" is sele	cted, pleas	e specify:											
			Alter	nate Confi	rmatory D	rug Test N	Methods						
Alternate													
Confirmatory Drug Test	THCA	BZE	COD	MOR	6-AM	PCP	AMP	MAMP	MDMA	MDA	MDEA		
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:											

\* ng/mL

COD = codeine MDA = methylenedioxyamphetamine
AMP = amphetamine MDEA = methylenedioxyethylamphetamine

Table C-3-b-1
Primary Confirmatory Drug Test Methods and Instruments - Gas Chromatography

	Prim	nary Confirmat	ory Drug Test	Methods and	Instruments -	Gas Chromatogra	aphy	
Primary Confirmatory Drug	THCA	BZE	COD/MOR	6-AM	PCP	AMPHETAMINE	S (select analytes from dr	opdown lists below)
Test - Gas Chromatography			002/111011	• 1				
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 addition	al information bel	ow		
Cryotrapping (Y/N)								
2nd GC Column Type								
2nd GC Column Length								
(m)								
*If "Other" is se	elected, please s	specify:						

Table C-3-b-2
Alternate Confirmatory Drug Test Methods and Instruments - Gas Chromatography

	Alteri	nate Confirma	tory Drug Test	t Methods and	Instruments -	Gas Chromatog	raphy	
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 Methods	s: provide addition	al information belo	DW .		
Cryotrapping (Y/N)								
2nd GC Column Type								
2nd GC Column Length (m)								
	elected, please	specify:						

Table C-3-b-3
Primary Confirmatory Drug Test Methods and Instruments - Liquid Chromatography

Primary C	onfirmatory I	Drug Test Me	ethods and In	struments- L	iquid Chroma	atography	
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 select	ed, please spe	ecify:			<u> </u>		l

Table C-3-b-4
Alternate Confirmatory Drug Test Methods and Instruments- Liquid Chromatography

#### Alternate Confirmatory Drug Test Methods and Instruments- Liquid Chromatography Alternate Confirmatory Drug Test -MDMA/MDA/ **THCA** AMP/MAMP BZE COD/MOR 6-AM **PCP** Liquid Chromatography 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 Hq B Solvent (Organic) Component 1 Component 2 Component 3 Component Ratio (1:2:3) Instrument Manufacturer Number of Units \*If "Other" is selected, please specify:

### Table C-3-c-1 Primary Confirmatory Drug Test Methods and Instruments - Mass Spectrometry (MS)

		Prima	ary Confirmato	ory Drug Test	Methods and	Instruments -	- Mass Spectr	ometry (MS)			
Primary 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 sel	lected, please	specify:			•	•			•	•	•

<sup>\*</sup>Minimum required

		Alterna	ate Confirmat	ory Drug Test	Methods and	d Instruments	- Mass Spect	rometry (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 sel	ected, please	specify:									

<sup>\*</sup>Minimum required

Table C-3-c-3
Primary Confirmatory Drug Test Methods and Instruments - Tandem Mass Spectrometry

	Primary	y Confirma	atory Drug	Test Met	hods and	Instrumen	ts - Tande	em Mass S	Spectrome	etry	
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*	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$
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*	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	→	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$
Int. Std. Qualifier Transition 1*	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$	$\rightarrow$
Int. Std. Qualifier Transition 2	$\rightarrow$	$\rightarrow$	$\rightarrow$	<b>→</b>	<b>→</b>	$\rightarrow$	<b>→</b>	<b>→</b>	$\rightarrow$	<b>→</b>	$\rightarrow$
Int. Std. Qualifier Transition 3	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>
*If "Other"	is selected	<mark>, please spe</mark>	ecify:								

<sup>\*</sup>Minimum required

Table C-3-c-4
Alternate Confirmatory Drug Test Methods and Instruments - Tandem Mass Spectrometry

	Alternat	te Confirm	atory Drug	g Test Met	thods and	Instrumer	nts - Tande	em Mass S	Spectrome	etry							
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*	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Qualifier Transition 1*	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Qualifier Transition 2	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Qualifier Transition 3	$\rightarrow$	→	÷	→	<del>)</del>	÷	<del>)</del>	<del>)</del>	<del>)</del>	÷	<del>)</del>						
Int. Std. Quantifier Transition*	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Int. Std. Qualifier Transition 1*	<b>&gt;</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Int. Std. Qualifier Transition 2	$\rightarrow$	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
Int. Std. Qualifier Transition 3	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>	<b>→</b>						
*If "Other"	is selected	, please sp	ecify:			*If "Other" is selected, please specify:											

<sup>\*</sup>Minimum required

# Primary Confirmatory Drug Test QC Samples

Primary Confir Test	matory Drug QC	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
<b></b>	Concentration									
THCA	Matrix									
	Source									
BZE	Concentration									
	Matrix Source									
	Concentration									
COD	Matrix									
	Source									
MOR	Concentration									
WOR	Matrix Source									
	Concentration									
6-AM	Matrix									
	Source									
505	Concentration									
PCP	Matrix									
	Source									
AMP	Concentration									
	Matrix Source									
	Concentration									
MAMP	Matrix									
	Source									
MDMA	Concentration									
WIDWIA	Matrix Source									
	Concentration									
MDA	Matrix									
	Source									
MDEA	Concentration									
MDEA	Matrix									
*If "Other" is	Source s selected, ple	ase specify:								
i Other R	s selected, ple	ase specify.								

# Alternate Confirmatory Drug Test QC Samples

Alternate Confi Test	irmatory Drug QC	Cal 1	Cal 2	Cal 3	Cal 4	Control 1	Control 2	Control 3	Control 4	Control 5
	Concentration									
THCA	Matrix									
	Source									
	Concentration									
BZE	Matrix Source									
	Concentration									
COD	Matrix									
	Source									
MOR	Concentration									
WOT	Matrix Source									
	Concentration									
6-AM	Matrix									
	Source									
PCP	Concentration									
1 01	Matrix Source									
	Concentration									
AMP	Matrix									
	Source									
MAMP	Concentration									
IVIAIVII	Matrix Source									
	Concentration									
MDMA	Matrix									
	Source									
MDA	Concentration									
MDA	Matrix Source									
	Concentration									
MDEA	Matrix									
	Source									
*If "Other" is	s selected, ple	ase specify:								