Department of Health and Human Services Substance Abuse and Mental Health Services Administration Center for Substance Abuse Prevention

Collection Site Checklist

for the
Collection of Urine Specimens for
Federal Agency Workplace Drug Testing Programs

Effective January 2022

Note: This checklist applies to federal agency drug testing programs that come under Executive Order 12564 dated September 15, 1986, section 503 of Public Law 100-71, 5 U.S.C. section 7301 note dated July 11, 1987, and the Department of Health and Human Services Mandatory Guidelines for Federal Workplace Drug Testing Programs (82 FR 7920) dated January 23, 2017 (effective October 1, 2017). This checklist does not apply to specimens submitted for testing under U.S. Department of Transportation (DOT) Procedures for Transportation Workplace Drug and Alcohol Testing Programs (49 CFR Part 40).

Previous Versions of this Checklist are Obsolete

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Instructions

A federal agency must ensure that collectors and collection sites satisfy all requirements in subparts D, E, F, G and H of the *Mandatory Guidelines for Federal Workplace Drug Testing Programs* (UrMG) published on January 23, 2017 (effective October 1, 2017).

This Collection Site Checklist is designed to assist the Drug Program Coordinator or designee and Collection Site Personnel in evaluating collection site performance based on onsite inspections and self-evaluations. A federal agency is responsible for inspecting 5 percent (up to a maximum of 50) collection sites each year, selected randomly from those sites used to collect federal agency specimens. A federal agency must investigate reported collection site deficiencies (e.g., specimens reported as "rejected for testing" by a HHS-certified testing facility) and take appropriate action, which may include an onsite inspection, or virtual, or collection site self-evaluation using the *Collection Site Checklist for Collection of Urine Specimens for Federal Agency Workplace Drug Testing Programs* and the HHS Urine Specimen Collection Handbook.

Checklist

Each question in the *Collection Site Checklist for Collection of Urine Specimens for Federal Agency Workplace Drug Testing Programs* is designed to address the requirements in UrMG subparts D, E, F, G and H. Answer each question based on these requirements and your review of the collection site standard operating procedures, practice, and records.

- 1. Check the appropriate **YES** or **NO** answer for each checklist question.
- 2. If required for a **NO** answer, check the deficient area(s) for the checklist question.
- 3. Record comments in the space provided to explain the specific reason for each **NO** answer.

Section Evaluation

Each checklist section contains a section evaluation page. Use the section evaluation to summarize and classify the seriousness of identified deficiencies.

- 1. For each checklist question in the section with a **NO** answer, explain the potential problem or identified non-compliance.
- 2. Mark the overall section evaluation at the top of the page as appropriate:
 - Deficiencies require immediate corrective action by the collection site
 - Deficiencies were identified but do not require immediate correction action
 - No deficiencies were identified.

Collection Site Evaluation Form

- 1. In the Overall Section Summary, assign a numerical "score" for each checklist section, based on the section evaluation:
 - Record a "0" on the evaluation form for each section summary where serious deficiencies were identified.
 - Record a "1" for each section summary where deficiencies were identified but do not require immediate corrective action.
 - Record a "2" for each section summary where no deficiencies were identified.
- 2. In the appropriate "Inspector/Collection Site Reviewer" columns under "Overall Summary of Serious Deficiencies," list the sections identified as having serious deficiencies and those with no serious deficiencies.
- 3. Add the individual section scores to determine the rating and record the total in the "Rating" space for "Inspector/Collection Site Reviewer" under "Inspection Outcome."
- 4. Sign and date in the appropriate space at the bottom of the form. Inspectors sign the "Onsite Inspection by" line; Collection Site Reviewers sign the "Self-Evaluation by" line.

A.	Collection Site		
A-1.		e collection site have provisions to ensure donor privacy during n procedure?	the specimen YES NC
A-2.	Does the	e collection site have the following?	☐ YES ☐ NC
	If NO , ch	neck the deficient area(s):	
	☐ a.	A means for washing hands	
	□ b.	A suitable clean surface, inaccessible to the donor, for the collwork area	lector to use as a
	☐ c.	A secure temporary storage area for maintaining specimens utransferred to an HHS-certified test facility	ntil they are
A-3.	Does the	e collection site have procedures or restrictions to prevent the fo	ollowing?
	If NO , ch	neck the deficient area(s):	
	□ a.	Unauthorized access to the site during the collection	
	□ b.	Unauthorized access to the collection materials/supplies	
	□ c.	Unauthorized access to collection site records	
	☐ d.	Donor access to items that could be used to adulterate, substi specimen (e.g., soap, disinfectants, cleaning agents, water)	tute, or dilute the

A-4.	Does the collection site have the required supplies for federally regular specimen collections?	ated urine
A-5.	Is access to collection supplies restricted to authorized personnel?	☐ YES ☐ NO
A-6.	Does the collection site have the name and telephone number of the crepresentative for each federal agency for which specimens are collected.	•

lf	Υ	ES.
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		is information readily available to each collector, in the event that a problem or earises during a collection?
A-7.		e collection site have procedures to prohibit the following individuals from as a specimen collector?
	If NO , ic	lentify the deficient area(s):
	☐ a.	Hiring official or donor's immediate supervisor <u>unless</u> there is no feasible alternative <u>and</u> the individual is a trained collector
	□ b.	Co-worker in the same testing pool or who works with the donor on a daily basis
	□ c.	The applicant or employee (i.e., the specimen donor)
	☐ d.	Employee of an HHS-certified Instrumented Initial Test Facility (IITF) or HHS-certified laboratory who can link the donor with the specimen drug test results
	□ е.	Relatives or close personal friends of the donor

A-8.	For the Collection Site Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
integri	Serious deficiencies require immediate corrective action by the collection site to maintain the ty of the collection process, to maintain the security and integrity of the specimens collected, and ure the privacy of the donors.
Descri	be basis for the above selection:

B. Personnel

Collectors

B-1.	knowled Specime	nterview by the inspection team, did each collector demonstr dge of the collection procedures described in the UrMG, the F en Collection Handbook, and any other guidance provided by to specimen collection procedures?	HHS Urine
	If NO , io	dentify the individual(s) and deficient area(s) of knowledge.	
B-2.	Was do	cumentation of training for each collector provided for review on?	during the
	If NO , n	ote the collector(s) with missing training documentation.	
Ancu	or alloca	tions B 2 through B 9 for the records provided	
AIISW	er quest	tions B-3 through B-8 for the records provided.	
B-3.	Does ea	ach collector maintain their training documentation?	☐ YES ☐ NO
B-4.		h collector complete initial training before they began collection agency?	ng specimens for a
B-5.		ch collector (as applicable) completed refresher training at lea e date of initial training?	ast every five years YES NO
B-6.		nitial and refresher training records for each collector docume g subjects?	ent training on the
	If NO , io	dentify the individual and records and check the deficient area	a(s):
	Па.	The steps to correctly perform a collection for federal agence	cy specimens
	b.	The proper completion and distribution of the Federal CCF	•
	c.	Problem collections	
	d.	Fatal and correctable flaws and how to correct problems in	collections
	e.	Collector responsibilities to maintain the integrity of the colle protect the privacy of donors, to ensure the security and integrated and to maintain proper conduct	ection process, to

B-7.	. Do the initial and refresher training records for each collector document their proficie in collections by successful completion of five (5) consecutive error-free mock collections?		
	If NO , id	entify the individual and records and check the deficient area(s):	
	□ a. □ b. □ c. □ d. □ e.	One uneventful scenario One insufficient specimen quantity scenario One temperature is out of range scenario One scenario in which the donor refuses to sign the Federal CCF One scenario in which the donor refuses to initial the tamper-evident bottle label/seal	
B-8.	Do the in	nitial and refresher training records for each collector include the following? ☐ YES ☐ NO	
	If NO , id	entify the individual and records and check the deficient area(s):	
	☐ a. ☐ b. ☐ c.	Documentation that the training was conducted in person or by means allowing real-time observation and interaction between trainer and trainee. Written attestation by the trainer that the mock collections were error-free. Documentation of the trainer's qualifications at the time of the training.	

Collector Trainers

Answer the remaining Section B questions if collection site employees serve as collector trainers.

B-9.	During interview by the inspection team, did each collector trainer demonstrated working knowledge of the collection procedures described in the UrMG Collection Handbook, and any other guidance provided by the federal at the collection procedures?	, the HHS Urine
B-10.	Was documentation of training for each trainer provided for review durin inspection?	ng the
	If NO , note the trainer(s) with missing training documentation.	
Comp	plete the remaining Section B questions for the records provided.	
B-11.	Does each trainer maintain their training documentation?	☐ YES ☐ NO
B-12.	Do the training records for each collector trainer document <u>at least one</u> qualifications?	of the following
	The trainer is qualified as a collector and has regularly conducted ditest collections for a period of at least one year,	rug
	The trainer successfully completed a "train the trainer" course given an organization (e.g., manufacturer, private entity, contractor, or federal agency)	by
B-13.	Has each trainer (as applicable) completed refresher training at least en from the date of initial training?	very five years

B-14.	For the Personnel Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
the inte	Serious deficiencies require immediate corrective action by the collection site to maintain egrity of the collection process, to maintain the security and integrity of the specimens ed, and to ensure the privacy of the donors.
Descril	pe basis for the above selection:

C.	Specimen Collection Procedures			
C-1.	Does the collector prepare the restroom to deter the dilution or substituti specimen?		fa ES [] NO
	Required steps:			
	 Placing bluing agent in the toilet or turning off the water supply and flushing the toilet 			
	 Securing any other water source in the enclosure where urination occurs 			
C-2.	Does the collector begin the collection without delay once the donor arricollection site?		it the ES [] NO
C-3.	When a donor does not arrive at the collection site at the assigned time test, does the collector contact the federal agency representative to obtathe appropriate action to be taken?	ain gu		_
C-4.	Does the collector perform only one specimen collection at a time?	☐ Y	ES [] NO
C-5.	Does the collector properly verify donor identity?	□ Y	ES [] NO
	Proper forms of identification include:			
	Driver's license			
	Employee badge issued by the employer			
	 Photo identification issued by a federal, state, or local government agency 			
C-6.	Does the collector provide identification to the donor when requested?	□ Y	ES [] NO
C-7.	Does the collector describe the basic collection procedures to the donor donor that they may read the instructions for completing the Federal CC	F?	instru	ct the

C-0.		e collection process?	YES NO
C-9.	Does the	e collector complete the required information in Step 1 of the F	Federal CCF?
C-10.	Does the	e collector take the following steps to deter specimen tamperir	ng?
	If NO , ch	neck the deficient step(s):	
	☐ a.	Ask the donor to remove any unnecessary outer clothing (e.g. hat, etc.)	յ., coat, jacket,
	□ b.	Ask the donor to leave all other personal belongings (e.g., briwith the outer clothing or in another secured location	efcase, purse)
	c.	Direct the donor to empty their pockets and display the items	for inspection
	☐ d.	Secure any items that could be used to adulterate a specime have been inadvertently brought by the donor to the collection	
	□ e.	Direct the donor to wash and dry their hands under the collect	tor's supervision
C-11.	Does the Federal	e collector note any unusual appearance or behavior of the do CCF?	nor on the
C-12.	Does the	e collector give the donor the following collection instructions?	☐ YES ☐ NO
	If NO , ch	neck the deficient area(s):	
	☐ a.	Provide at least 45 mL of urine	
	☐ b.	Do not flush the toilet	
	∐ c.	Provide the specimen in a reasonable time (set by the collect	•
	∐ d.	Return with the specimen as soon as they have finished prov specimen	iding the
C-13.		uthorized personnel prohibited from entering the collection sitent n procedure?	e during the
C-14.	Are only	the collector and the donor allowed to handle the unsealed sp	oecimen?
C-15.	time the	the collector and the donor maintain visual contact with the specimen is transferred to the collector until specimen bottles or shipment?	

Completion of a Collection

C-16.	After receiving the specimen from the donor, whenever practical, does allow the donor to wash their hands and to flush the toilet?		collector YES	NO
C-17.	Does the collector check the specimen temperature within four minutes the specimen from the donor and check the appropriate box in Step 2 c CCF?	of the		_
C-18.	Does the collector inspect the specimen for adulteration or substitution physical characteristics of the urine?	•	xamining YES □	the NO
C-19.	Does the collector check the specimen volume to ensure that the specileast 45 mL of urine?		contains YES 🗌	at NO
C-20.	In the presence of the donor, does the collector pour at least 30 mL into at least 15 mL into "Bottle B"?		ottle A" aı YES ☐	nd NO
C-21.	Does the collector discard excess urine (unless it is used for a clinical to physical examination required by a federal agency)?		s part of YES 🔲	a NO
C-22.	In the presence of the donor, does the collector place the appropriate to label/seal from the Federal CCF over the lid/cap of each bottle to ensur cannot be removed without destroying the label/seal?	re tha		
C-23.	If the tamper-evident label/seal does not adhere to the bottle or is dama collector apply the unacceptable label/seal to the bottle, and apply a set tamper-evident seal to seal the specimen bottle?	cond		
C-24.	Does the collector record the date of the collection on the bottle seals a on the bottles?	:	placing tl YES 🔲	nem NO
C-25.	Does the collector ask the donor to initial the specimen bottle seals after on the bottles?	<u> </u>	icing thei	m NO

C-26.	Does the collector instruct the donor to read and sign the donor certificand to fill out the donor portion in Step 5 on Copy 2 of the Federal CCF		state	men	ıt
			YES	<u> </u>	NO
C-27.	Does the collector complete the collector chain of custody section in St of the Federal CCF?	<u> </u>	on Co YES	<u> </u>	
C-28.	Does the collector place the sealed specimen bottles inside the leak-re seal the container, and include Copy 1 in the package with the specime compartment separate from the specimen bottles)?	en (i.		a	er, NO
C-29.	Does the collector provide Copy 5 of the Federal CCF to the donor?		YES	□ I	NO
C-30.	Does the collector prepare the sealed tamper-resistant package contains specimen bottles and Federal CCF for transport to the HHS-certified tellITF or laboratory)?	st fac			NO
C-31.	Are the specimen bottles and Federal CCF appropriately safeguarded retrieved for transport to the HHS-certified test facility (i.e., IITF or labo	rator	•		NO
C-32.	Does the collector send Copy 2 of the Federal CCF to the Medical Rev (MRO) and Copy 4 of the Federal CCF to the agency's designated representation and the collection or during the next business day?	reser		with	hin NO
C-33.	Are specimens submitted to an HHS-certified test facility (i.e., IITF or la 24 hours after the collection or during the next business day?		itory) v YES		in NO

C-34.	For the Specimen Collection Procedures Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
the inte	Serious deficiencies require immediate corrective action by the collection site to maintain egrity of the collection process, to maintain the security and integrity of the specimens ed, and to ensure the privacy of the donors.
Descri	be basis for the above selection:

D. Collection Problems

Direct Observed Collections

D-1.	Does the collector initiate a direct observed collection in the following situations? ☐ YES ☐ NO				
	If NO , check the deficient area(s):				
	☐ a.	Specimen temperature is outside the acceptable range			
	b.	Specimen appearance indicative of tampering (abnormal physical characteristic such as unusual color, excessive foaming when shaken, unusual odor)			
	☐ c.	Donor conduct indicates an attempt to adulterate or substitute the specimen, and the donor has already provided a specimen			
D-2.	Does th	e collector take the following steps before conducting a direct observed on?			
	If NO , c	heck the deficient step(s):			
	☐ a.	Contact a collection site supervisor for concurrence with the collector's decision for a direct observed collection			
	□ b.	Explain to the donor why a direct observed collection is being conducted			
	☐ c.	Inform the donor that the gender of the observer will match the donor's gender, which is determined by the donor's gender identity			
D-3.		an individual is allowed to serve as the observer for a direct observed collection, collector/collection site supervisor provide training on the following subjects? ☐ YES ☐ NO			
	If NO , identify the individual and records and check the deficient area(s):				
	□ a.	The steps necessary to perform a direct observed collection correctly			
	□ b.	Maintaining visual contact with the collection container throughout the collection process, to maintain the integrity and security of the specimen			
	□ c.	Ensuring the privacy of the donor			
	☐ d.	Observing the collection in a professional manner, to minimize discomfort of the donor			
	☐ e.	Avoiding conduct that could be interpreted as offensive or inappropriate			
D-4.		e collector ensure that the observer for each direct observed collection meets wing requirements?			

Does the collector properly document the direct observed collection in Step 2 of the Federal CCF?
If NO , check the deficient step(s):
 a. Mark the checkbox for an observed collection b. Record the name and gender of the observer on the Remarks line c. Record the reason for the observed collection on the Remarks line
ored Collections
Does the collector initiate a monitored collection in the following situations?
The collection is being conducted in a public restroom
The restroom used for the collection has a water source that cannot be disabled or secured
Does the collector ensure that the monitor for each monitored collection meets <u>at least</u> <u>one</u> of the following requirements?
Same gender as the donor
 A trained medical professional (e.g., nurse, doctor, physician's assistant, technologist, or technician) who is licensed or certified to practice where the collection occurs
Does the collector record the name of the monitor (if not the collector) on the Remarks line in Step 2 on Copy 1 of the Federal CCF?
icient Specimen
When the donor has demonstrated that they are unable to provide a sufficient specimen, does the collector offer the donor a reasonable amount of fluid to drink (e.g., an 8 ounce glass of water every 30 minutes, not to exceed 40 ounces over a period of 3 hours)?

• Trained in direct observed specimen collection procedures

• Same gender as the donor

D-10.	Does the	e collector allow the donor up to three hours to provide a suffici	ent specimen YES	i? NO
D-11.		ction procedures prohibit combining urine collected from separa specimen of sufficient volume?	ate voids to	NO
D-12.		e donor remain under the direct observation of the collector to passibly compromising the collection process?	orevent the do	onor NO
D-13.		e collector record the time of each attempt to provide a sufficier en (e.g., on the Remarks line of the Federal CCF)?	nt volume of	NO
D-14.	Does the	e collector discontinue the collection procedure in the following	situations?	NO
	• The	donor states that they are unable to provide a specimen		
	• The	donor has not provided sufficient volume of specimen in three s from the time of the donor's first attempt		
D-15.		e donor has not provided a sufficient specimen, does the collect n procedure and take the following steps?	ctor end the	NO
	If NO , ch	neck the deficient area(s):		
	☐ a.	Mark the "None Provided" checkbox in Step 2 of the Federal C	CCF	
	□ b.	Record the reason for not collecting the specimen on the Rem 2 of the Federal CCF	arks line in S	tep
	☐ C.	Notify the agency's designated representative for authorization alternate specimen or follow the standard protocol from the fee		
	☐ d.	Discard the urine collected (if any)		
	☐ e.	Discard Copy 1 of the Federal CCF (no valid specimen was comaintain Copy 3 in the collection records	ollected) and	
	☐ f.	Distribute the remaining Federal CCF copies within 24 hours of business day:	or the next	
	• Seno	1 Conv 2 to the MRO		

- Send Copy 4 to the federal agency's designated representative

Refusal to Test

D-16.	6. Does the collector report a "refusal to test" in the following situations? YES NO				
	If NO , check the deficient area(s):				
	☐ a.	The donor fails to appear for any test (except a pre-employment test) within a reasonable time as determined by the federal agency			
	□ b.	The donor fails to provide a specimen (e.g., urine or another authorized alternate specimen type)			
	☐ C.	The donor fails to cooperate with any part of the testing process (e.g., refuses to empty pockets, disrupts the collection process, fails to wash hands when directed by the collector)			
	☐ d.	The donor fails to allow a direct observed collection when required			
	☐ e.	The donor fails to follow the observer's instructions related to the direct observed collection			
	☐ f.	The donor fails to allow a monitored collection when required			
	g.	The donor brings materials to the collection site for the purpose of adulterating, substituting, or diluting the specimen			
	☐ h.	The donor attempts to adulterate, substitute, or dilute the specimen			
	☐ i.	The donor leaves the collection site before completion of the collection (except for leaving before the collection has begun for a pre-employment test)			
	☐ j.	The donor possesses or wears a prosthetic or other device that could be used to interfere with the collection process			
	☐ k.	The donor admits to the collector that they have adulterated or substituted their specimen			
D-17.	When re	eporting a "refusal to test," does the collector take the following steps? ☐ YES ☐ NO			
	If NO, check the deficient step(s):				
	☐ a.	Discard the urine collected (if any)			
	□ b.	Immediately notify the agency's designated representative of the refusal (e.g., by telephone, secure fax machine, e-mail)			
	☐ c.	Document the refusal to test with appropriate comments, signature, and date in the Remarks line of Step 2 of the Federal CCF			
	☐ d.	Send all copies of the Federal CCF to the federal agency's designated representative			

Collector Errors

D-18.	When the collector realizes that an incorrect or expired Federal CCF is used prior to packaging the specimen bottles, does the collector document on the form that the specimen is a federal agency specimen and provide the reason for the incorrect form? YES NO
D-19.	Does the collector provide a memorandum for the record (MFR) when requested by the HHS-certified test facility (i.e., laboratory, IITF), or MRO?

D-20.	For the Collection Problems Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
the inte	Serious deficiencies require immediate corrective action by the collection site to maintain egrity of the collection process, to maintain the security and integrity of the specimens ed, and to ensure the privacy of the donors.
Descri	be basis for the above selection:

E.	Collection Site Records	
E-1.	Are collection site records including Copy 3 of the Federal Custody and (Federal CCF) stored for a minimum of two years?	d Control Form
E-2.	Are collection site records stored and disposed of in a manner that ensconfidentiality?	sures donor
E-3.	Have collectors properly completed the Federal CCF?	☐ YES ☐ NO
E-4.	Are edits to the Federal CCF properly made, initialed and dated?	☐ YES ☐ NO

E-5.	For the Collection Site Records Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
the inte	Serious deficiencies require immediate corrective action by the collection site to maintain egrity of the collection process, to maintain the security and integrity of the specimens ed, and to ensure the privacy of the donors.
Descri	be basis for the above selection:

Collection Site Evaluation Form

Overall Section Summary						
Checklist Sections		Serious Deficiencies Identified (0)	Deficiencies Identified (1)	No Deficiencies Identified (2)		
A. Collection Site						
B. Personnel						
C. Specimen Collection Procedures						
D. Collection Problems						
E. Collection Site Record	S					
Overa	all Summ	nary of Serious	Deficiencies			
		s Deficiencies e identified		eficiencies were ntified		
Inspector / Collection Site Reviewer			[
Federal Agency/ Designee						
	Ins	pection Outcon	ne			
Rating (out of 1	0)	Acceptable: rating serious deficienci	g ≥ 5 <u>and</u> no more tha ies	n one section with		
Inspector / Collection Site Reviewer	/10	Unacceptable: rai	ting < 5 <u>or</u> more than ies	one section with		
Federal Agency/ Designee	/10	Outcome:				
Additional Comments:						
Acceptable Outcome for Inspection:						
Self-Evaluation by: Date:						
Onsite Inspection by: Date:						
Approved by:	Approved by: Date:					
Position/Title:						