# 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 Oral Fluid 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 using Oral Fluid (84 FR 57554) dated October 25, 2019 (effective January 1, 2020). 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).

### **Table of Contents**

Instructions	Z
Checklist	
Section Evaluation	
Collection Site Evaluation Form	
A. Collection Site	
Section Evaluation	8
B. Personnel	
Collectors	
Collector Trainers	
Section Evaluation	
C. Specimen Collection Procedures	13
Completion of a Collection	
Section Evaluation	
D. Collection Problems	18
Insufficient Specimen	
Refusal to Test	
Collector Errors	21
Section Evaluation	
E. Collection Site Records	23
Section Evaluation	
Collection Site Evaluation Form	

#### References

(available at https://www.samhsa.gov/workplace

- 1. Federal Custody and Control Form (Federal CCF)
- 2. HHS Oral Fluid Specimen Collection Handbook for Federal Agency Workplace Drug Testing Programs (HHS Oral Fluid Collection Handbook)
- 3. Mandatory Guidelines for Federal Workplace Drug Testing Programs using Oral Fluid (OFMG). Published October 25, 2019 (84 FR 57554), effective January 1, 2020.

#### 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 using Oral Fluid* (OFMG) published on October 25, 2019 (effective January 1, 2020).

This Collection Site Checklist is designed to assist the Drug Program Coordinator (DPC) 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 an HHS-certified laboratory) and take appropriate action, which may include an onsite or virtual inspection or collection site self-evaluation using the *Collection Site Checklist for Collection of Oral Fluid Specimens for Federal Agency Workplace Drug Testing Programs* and the HHS Oral Fluid Specimen Collection Handbook for Federal Agency Workplace Drug Testing Programs.

#### Checklist

Each question in the Collection Site Checklist for Collection of Oral Fluid Specimens for Federal Agency Workplace Drug Testing Programs is designed to address the requirements in OFMG 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. Circle 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-1.	Does the collection site have provisions to ensure donor privacy during the specimen collection procedure?  ☐ YES ☐ NO
A-2.	Does the collection site have the following?  ☐ YES ☐ NO
	If <b>NO</b> , check the deficient area(s):
	a. A means for washing hands
	b. A suitable clean surface, inaccessible to the donor, for the collector to use as a work area
	c. A secure temporary storage area for maintaining specimens until they are transferred to an HHS-certified laboratory
A-3.	Does the collection site have procedures or restrictions to prevent the following?
	☐ YES ☐ NO
	If NO, check 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 or substitute the specimen, or otherwise adversely affect the oral fluid collection
A-4.	Does the collection site have the required supplies for federal agency oral fluid specimen collections?
	☐ YES ☐ NO
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 designated representative for each federal agency for which specimens are collected?
	☐ YES ☐ NO
	If <b>YES</b> , Is this information readily available to each collector, in the event that a problem or issue arises during a collection?
	☐ YES ☐ NO

**Collection Site** 

A.

A-7.	Does the collection site have procedures to prohibit the following individuals from serving as a specimen collector?							
	☐ YES ☐ NO							
	If <b>NO</b> , identify the deficient area(s):							
	a. Hiring official or donor's immediate supervisor unless there is no feasible alternative and 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 laboratory who can link the donor with the specimen drug test results							
	e. Relatives or close personal friends of the donor							

A-8. F	or th	e Collection Site Section:					
	Serious deficiencies were identified						
		Deficiencies were identified					
		No deficiencies were identified					
integri	ty of t	us deficiencies require immediate corrective action by the collection site to maintain the he collection process, to maintain the security and integrity of the specimens collected, and e privacy of the donors.					
	Des	cribe basis for the above selection:					

### B. Personnel

# Collectors

B-1.	During interview by the inspection team, did each collector demonstrate a working knowledge of the collection procedures described in the OFMG and any other guidance provided by the federal agency related to specimen collection procedures?  YES NO  If NO, identify the individual(s) and deficient area(s) of knowledge.
B-2.	Was documentation of training for each collector provided for review during the inspection?  YES  NO
	If <b>NO</b> , note the collector(s) with missing training documentation.
Answ	er questions B-3 through B-8 for the records provided.
B-3.	Does each collector maintain their training documentation?  ☐ YES ☐ NO
B-4.	Did each collector complete initial training for each collection device used before they began collecting specimens for a federal agency?  ☐ YES ☐ NO
B-5.	Has each collector (as applicable) completed refresher training at least every five years from the date of initial training?    NO  N/A

B-6.	Do the initial and refresher training records for each collector document training on the following subjects?		
	☐ YES ☐ NO		
If NO,	identify the individual and records and check the deficient area(s):		
	<ul> <li>a. The steps to correctly perform a collection using each type of collection device to be used for federal agency specimens</li> </ul>		
	□ 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 collection process, to protect the privacy of donors, to ensure the security and integrity of specimens, and to maintain proper conduct		
B-7.	Do the initial and refresher training records for each collector document their proficiency in collections by successful completion of five (5) consecutive error-free mock collections?		
	☐ YES ☐ NO		
If NO,	identify the individual and records and check the deficient area(s):		
	a. Two uneventful scenarios		
	☐ b. One insufficient specimen quantity scenario		
	c. One scenario in which the donor refuses to sign the Federal CCF		
	<ul> <li>d. One scenario in which the donor refuses to initial the tamper-evident tube label/seal</li> </ul>		
B-8.	Do the initial and refresher training records for each collector include the following?  ☐ YES ☐ NO		
If NO,	identify the individual and records and check the deficient area(s):		
	a. Documentation that the training was conducted in person or by means allowing real-time observation and interaction between trainer and trainee.		
	☐ b. Written attestation by the trainer that the mock collections were error-free.		
	c. 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 demonstrate a working knowledge of the collection procedures described in the OFMG, the HHS Oral Fluid Collection Handbook, the manufacturer instructions for the specific collection device(s) and any other guidance provided by the federal agency related to the collection procedures?
	☐ YES ☐ NO
B-10.	Was documentation of training for each trainer provided for review during the inspection?
	☐ YES ☐ NO
	If <b>NO</b> , note the trainer(s) with missing training documentation.
Comp	blete 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> of the following qualifications?
	☐ YES ☐ NO
	<ul> <li>The trainer is qualified as a collector and has regularly conducted drug test collections for a period of at least one year using the specific collection device(s),</li> </ul>
	<ul> <li>The trainer successfully completed a "train the trainer" course given by an organization (e.g., manufacturer, private entity, contractor, or federal agency), including training on the specific collection device(s)</li> </ul>
B-13.	Has each trainer (as applicable) completed refresher training at least every five years from the date of initial training?
	☐ YES ☐ NO

B-14. For t	he Personnel Section:
	Serious deficiencies were identified
	Deficiencies were identified
	No deficiencies were identified
integrity of th	us deficiencies require immediate corrective action by the collection site to maintain the ne collection process, to maintain the security and integrity of the specimens collected, and privacy of the donors.
Describe ba	asis for the above selection:

C.	Specimen Collection Procedures				
C-1.	Does the collector pof a specimen?	orepa	re the collection site to deter the adulteration or substitution		
	YES		NO		
C-2.	Does the collector to collection site?	oegin	the collection without delay once the donor arrives at the		
	YES		NO		
C-3.		ctor c	arrive at the collection site at the assigned time for the drug ontact the federal agency representative to obtain guidance to be taken?		
	☐ YES		NO		
C-4.	Does the collector ptime?	perfor	m only one specimen collection at a		
	YES		NO		
C-5.	Does the collector p	orope	rly verify donor identity?		
	YES		NO		
	·	Proper forms of identification include:			
	Driver's license     Employee badg	o ioo	and by the employer		
	<ul> <li>Employee badge issued by the employer</li> <li>Photo identification issued by a federal, state, or local government agency</li> </ul>				
C-6.	Does the collector p	orovic	de identification to the donor when requested?		
C-7.			ibe the basic collection procedures to the donor and instruct read the instructions for completing the Federal CCF?		
	YES		NO		
C-8.	Does the collector a		er any reasonable and appropriate questions that the donor process?		
	YES		NO		
C-9.	Does the collector o	comp	lete the required information in Step 1 of the Federal CCF?		
	V   \ \	1 1	INIL 1		

C-10.	Does the collector t	take t	he following steps to deter specimen tampering?			
	If NO, check the de	If <b>NO</b> , check the deficient step(s):				
	a. Ask the dor	or to	open their mouth to allow inspection of the oral cavity			
	<ul> <li>b. Begin 10-minute wait period after inspection of the donor oral cavity prior to beginning specimen collection</li> </ul>					
	colored sali	va, o	oves an item from their mouth as instructed, has abnormally r indicates they have "dry mouth", provide water to the donor 0-minute wait period after donor has rinsed their mouth			
C-11.	Does the collector r Federal CCF?	note a	any unusual appearance or behavior of the donor on the			
	☐ YES		NO			
C-12.			w the procedures for a successful oral fluid specimen the device-specific manufacturer's instructions?			
	☐ YES		NO			
C-13.	Are unauthorized p collection procedure		nnel prohibited from entering the collection site during the			
	☐ YES		NO			
C-14.	Are only the collect	or an	d the donor allowed to handle the unsealed specimen?			
	YES		NO			
C-15.	Does the collector the entire collection		n present and maintain visual contact with the donor during ess?			
	YES		NO			
	If <b>NO</b> , check the deficient area(s):					
	a. Ensure the donor has positioned the specimen collection device properly for collection					
	b. Ensure the	colle	ction is performed correctly			
	c. Ensure collection device is working properly					

# **Completion of a Collection**

C-16.	(e.g., 1 mL of undilu	uted [	e the donor has provided a sufficient volume of oral fluid neat] oral fluid for tube A and 1 mL of undiluted [neat] oral ollowing types of split specimen collections?			
	YES		NO			
	If <b>NO</b> , check the deficient area(s):					
	a. Two specim	iens (	collected simultaneously using two separate collection			
	☐ b. Two specim	ens o	collected serially with two separate collection devices			
			collected simultaneously using a single collection device that en into two separate collection tubes			
			en collected using a single collection device that is wo specimen tubes.			
C-17.	Does the collector o	comp	lete the required information in Step 2 of the Federal CCF?			
C-18.	Does the collector r	eport	a refusal if the donor refuses to complete the collection?			
C-19.	evident label/seal fr	om th	onor, does the collector place the appropriate tamper- ne Federal CCF over the lid/cap of each tube to ensure that noved without destroying the label/seal?			
C-20.	collector apply the	unaco	el/seal does not adhere to the tube or is damaged, does the ceptable label/seal to the tube, and apply a second, separate eal the specimen tube?			
C-21.	Does the collector r them on the tubes?		d the date of the collection on the tube seals after placing			
	YES		NO			
C-22.	Does the collector a on the tubes?	ask th	ne donor to initial the specimen tube seals after placing them			
	YES		NO			

C-23.	Does the collector instruct the donor to read and sign the donor certification statement and to fill out the donor portion in Step 5 on Copy 2 of the Federal CCF?				
	YES		NO		
C-24.			lete the collector chain of custody section and document (as applicable) in Step 4 on Copy 1 of the Federal CCF?		
C-25.	container, seal the	conta	the sealed specimen tubes inside the leak-resistant ainer, and include Copy 1 in the package with the specimen eparate from the specimen tubes)?  NO		
C-26	Does the collector r	orovio	de Copy 5 of the Federal CCF to the donor?		
0 20.	YES		NO		
C-27.	-	•	are the sealed tamper-resistant package containing the deral CCF for transport to the HHS-certified laboratory?		
	YES		NO		
C-28.			and Federal CCF appropriately safeguarded until they are the HHS-certified laboratory?		
	YES		NO		
C-29.		of th	Copy 2 of the Federal CCF to the Medical Review Officer e Federal CCF to the agency's designated representative		
	YES		NO		
C-30.	Are specimens sub collection or during  YES		d to an HHS-certified laboratory within 24 hours after the next business day?  NO		

C-31. For the Specimen Collection Procedures Section:					
Serious deficiencies were identified					
☐ Deficiencies were identified					
☐ No deficiencies were identified					
<u>Note</u> : Serious deficiencies require immediate corrective action by the collection site to maintain the integrity of the collection process, to maintain the security and integrity of the specimens collected, and to ensure the privacy of the donors.					
Describe basis for the above selection:					

#### D. Collection Problems

Insufficient Specimen – Questions D-1 through D-7 pertain to collections where the donor states they are unable to provide an oral fluid specimen.

D-1.	Does the collector attempt an oral fluid collection in the event the donor states they are unable to provide a specimen?						
	☐ YES ☐ NO						
D-2.	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., up to 8 ounces of water)?						
	☐ YES ☐ NO						
D-3.	Does the collector allow the donor up to one hour to provide a sufficient specimen?  YES  NO						
D-4.	Does the collector direct the donor to remain in a designated area at the collection site during the wait period?  YES  NO						
D-5.	Does the collector record each failed collection attempt on the Remarks line of the Federal CCF?  YES  NO						
D-6.	Does the collector discontinue the collection procedure in the following situations?  YES NO  The donor states that they are unable to provide a specimen						
	The donor has not provided sufficient volume of specimen in one hour from the time of the donor's first attempt						

D-7.	When the donor has not provided a sufficient specimen, does the collector end the collection procedure and take the following steps?						
	☐ YES ☐ NO						
	If <b>NO</b> , check the deficient area(s):						
	a. Mark the "None Provided" checkbox in Step 2 of the Federal CCF						
	<ul> <li>b. Record the reason for not collecting the specimen on the Remarks line in Step 2 of the Federal CCF</li> </ul>						
	c. Notify the agency's designated representative for authorization to collect an alternate specimen or follow the standard protocol from the federal agency						
	☐ d. Discard the oral fluid collected (if any)						
	<ul> <li>e. Discard Copy 1 of the Federal CCF (no valid specimen was collected) and maintain Copy 3 in the collection records</li> </ul>						
	f. Distribute the remaining Federal CCF copies within 24 hours or the next business day:						
	Send Copy 2 to the MRO						

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

# **Refusal to Test**

D-8.	Does the collector report a "refusal to test" in the following situations?						
	☐ YES ☐ NO						
	If <b>NO</b> , 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						
	<ul> <li>b. The donor leaves the collection site before completion of the collection (except for leaving before the collection has begun for a pre-employment test)</li> </ul>						
	c. The donor fails to provide a specimen (e.g., oral fluid or another authorized alternate specimen type)						
	d. The donor fails to provide a sufficient amount of oral fluid when directed and it has been determined through a required medical evaluation that there is no legitimate medical explanation for the failure						
	<ul><li>e. The donor fails or declines to participate in an alternate specimen collection (e.g., urine)</li></ul>						
	f. The donor fails to cooperate with any part of the testing process (e.g., refuses to empty pockets, disrupts the collection process, fails to rinse mouth when directed by the collector, refuses to provide a split specimen)						
	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 admits to the collector that they have adulterated or substituted their specimen						
D-9.	When reporting a "refusal to test," does the collector take the following steps?						
	☐ YES ☐ NO						
	If <b>NO</b> , check the deficient step(s):						
	a. Discard the oral fluid collected (if any)						
	b. Immediately notify the agency's designated representative of the refusal (e.g., by telephone, secure fax machine, e-mail)						
	<ul> <li>c. Document the refusal to test with appropriate comments, signature, and date in the Remarks line of Step 2 of the Federal CCF</li> </ul>						
	<ul> <li>d. Send all copies of the Federal CCF to the federal agency's designated representative</li> </ul>						

### Collector Errors

D-10.	to packaging the sp	collector realizes that an incorrect or expired Federal CCF was used prior ling the specimen tubes, does the collector document on the form that the is a federal agency specimen and provide the reason for the incorrect					
	☐ YES		NO				
D-11.	Does the collector puthe HHS-certified la		e a memorandum for the record (MFR) when requested by ory or MRO?				
	YES		NO				

D-12. For the Collection Problems Section:
Serious deficiencies were identified
☐ Deficiencies were identified
☐ No deficiencies were identified
<u>Note</u> : Serious deficiencies require immediate corrective action by the collection site to maintain the integrity of the collection process, to maintain the security and integrity of the specimens collected, and to ensure the privacy of the donors.
Describe basis for the above selection:

### E. Collection Site Records

E-1.	Are collection site records including Copy 3 of the Federal Custody and Control Form (Federal CCF) stored for a minimum of two years?					
	☐ YES	□ NO				
E-2.	Are collection site confidentiality?	cords stored	and disposed of in a manner that ensures donor			
	☐ YES	☐ NO				
E-3.	Have collectors pro	erly complete	d the Federal CCF?			
	☐ YES	□ NO				
E-4.	Are edits to the Fe	eral CCF prop	erly made, initialed, and dated?			
	☐ YES					

E-5. For the Collection Site Records Section:					
Serious deficiencies were identified					
☐ Deficiencies were identified					
☐ No deficiencies were identified					
<u>Note</u> : Serious deficiencies require immediate corrective action by the collection site to maintain the integrity of the collection process, to maintain the security and integrity of the specimens collected, and to ensure the privacy of the donors.					
Describe basis for the above selection:					

#### **Collection Site Evaluation Form**

Overall Section Summary							
Checklist Sections	Defici	ious encies ied (0)		ciencies tified (1)	No Deficiencies Identified (2)		
A. Collection Site							
B. Personnel							
C. Specimen Collection Procedures							
D. Collection Problems							
E. Collection Site Records							
Overall Summary of Serious Deficiencies (List Sections)							
Section		s Deficier e identifie		No Serious Deficiencies were identified			
Inspector / Collection Site Reviewer							
Federal Agency/ Designee							
Inspection Outcome							
Rating (out of 10)  Inspector / Collection Site Reviewer	/10	Acceptable: rating ≥ 5 and no more than o section with serious deficiencies  /10 Unacceptable: rating < 5 or more than one section with serious deficiencies			cies more than one		
Federal Agency/ Designee	/10	Outcome:					
Additional Comments:							
Acceptable Outcome for Inspection:							
Self-Evaluation by: Date							
Onsite Inspection by:				Date			
Approved by:		Date					
Position/Title:							