

Kim Reynolds, Governor Chris Cournoyer, Lieutenant Governor

Reporting HIV and AIDS in Iowa

What's reportable AIDS has been a reportable disease in Iowa since February 1983. HIV became reportable by name in Iowa on July 1, 1998. Iowa Administrative Code 641.11.6., below, details reporting.

641—11.6(141A) Reporting of diagnoses and HIV-related tests, events, and conditions to the department.

11.6(1) The following constitute reportable events related to HIV infection:

a. A test result indicating HIV infection, including:

(1) Confirmed positive results on any HIV-related test or combination of tests, including antibody tests, antigen tests, cultures, and nucleic acid amplification tests.

(2) A positive result or report of a detectable quantity on any other HIV detection (non-antibody) tests, and results of all viral loads, including nondetectable levels.

b. AIDS and AIDS-related conditions, including all levels of CD4+ T-lymphocyte counts.

c. Birth of an infant to an HIV-infected mother (perinatal exposure) or any (positive, negative, or undetectable) non-antibody detection test (antigen test, viral culture, viral load, or qualitative nucleic acid amplification test) on an infant 18 months of age or younger.

d. Death resulting from an AIDS-related condition, or death of a person with HIV infection.

11.6(2) Within seven days of the receipt of a person's confirmed positive test result indicating HIV infection, the director of a plasma center, blood bank, clinical laboratory or public health laboratory that performed the test or that requested the confirmatory test shall make a report to the department on a form provided by the department.

11.6(3) Within seven days of the receipt of a test result indicating HIV infection, which has been confirmed as positive according to prevailing medical technology, or immediately after the initial examination or treatment of a person infected with HIV, the physician or other health care provider at whose request the test was performed or who performed the initial examination or treatment shall make a report to the department on a form provided by the department.

11.6(4) Within seven days of diagnosing a person as having AIDS or an AIDS-related condition, the diagnosing physician shall make a report to the department on a form provided by the department. **11.6(5)** Within seven days of the death of a person with HIV infection, the attending physician shall make a report to the department on a form provided by the department.

11.6(6) Within seven days of the birth of an infant to an HIV-infected mother or a receipt of a laboratory result (positive, negative, or undetectable) of a non-antibody detection test (antigen test, viral culture, viral load, or qualitative nucleic acid amplification test) on an infant 18 months of age or younger, the attending physician shall make a report to the department on a form provided by the department. **11.6(7)** The report shall include:

a. The person's name, address, date of birth, gender, race and ethnicity, marital status, and telephone number.

b. The name, address and telephone number of the plasma center, blood bank, clinical laboratory or public health laboratory that performed or requested the test, if a test was performed.

c. The address of the physician or other health care provider who requested the test.

d. If the person is female, whether the person is pregnant.

11.6(8) All people who experience a reportable event while receiving services in the state, regardless of state of residence, shall be reported.

Need reporting forms? Want to call in a report? Have questions? Need surveillance data? Alagie "Al" Jatta, HIV Surveillance Coordinator: 515-281-6918 or 515-322-8819 <u>Alagie.Jatta@hhs.iowa.gov</u>, Nabila Adamu, HIV Surveillance Epidemiologist: 515-721-8486 <u>Nabila.Adamu@hhs.iowa.gov</u>; COMPLETED FORMS CAN BE SENT VIA THE U.S. POSTAL SERVICE OR YOU CAN USE THE FORM TO COLLECT THE REQUIRED DATA AND THEN CALL US. FAXING OR EMAILING OF COMPLETED FORMS IS NOT ALLOWED!

By Mail:

Iowa Department of Public Health 321 East 12th Street Des Moines, IA 50319 "03 Confidential"

By Phone: Al Jatta at 515-281-6918/515-322-8819 or Nabila Adamu at 515-721-8486

By Fax: 515-725-1278 Attention: Confidential 03

INSTRUCTIONS

Don't panic. The form can seem a little foreboding. We do not expect you to know everything on the form, but we do expect you to provide all the information that is known to you.

Additional information. The form is based on the standard CDC report form. It asks for standard CDC information. In addition to the standard information, please use the comment and local field sections of the form to provide any information you may have on the following:

- Is patient newly diagnosed? If yes:
 - Why was patient tested?
 - Has the patient been informed they are HIV positive?
 - How did the patient respond to the news that they were HIV positive?
 - How is the patient coping with the news?
- Patient's living situation (Who knows about patient's HIV status? Who doesn't?)
- Patient's marital or relationship status (including HIV status of partners)
- Maintaining patient confidentiality is supremely important. With this in mind, what is the best way (e.g., a specific phone telephone number, time or place) for a trained Disease Prevention Specialist to contact the patient to deliver partner services, i.e., education about HIV, linking to care and services, and assistance in notifying partners?
- Other places the patient has lived (including military deployments if any)
- Social Security Number

Questions? Please call 515-721-8486 or 515-322-8819.

Thank you! Thank you for complying with Iowa's HIV reporting statutes!

I. Patient Identification (record all dates as mm/dd/yyyy)

*First Name		*Middle Na	me			*Last Name		- P	Last Name Soundex			
Alternate Name Type (ex: Al	ias, Married)		*First Nam	e		*Middle Name	le Name *Last N			Name		
Address Type Residential Foster home	e 🗆 Homeles	s 🗆 Military		*Current	t Addres	s, Street				Address Date		
	Postal Shelter Temporary City County State/Country				4	*ZIP Code						
() *Medical Record Number				*Other ID Type *NumI			nber)er				
U.S. Department of Health and Human Services Adult HIV Confidential Case Report Form (Patients ≥13 years of age at time of diagnosis) *Information NOT transmitted to CDC Contexpondent CDC II. Health Department Use Only (record all dates as mm/dd/yyyy) Form approved OMB no. 0920-0573 Exp. 02/28/2026												
Date Received at Health Department				eHARS Document UID			State I	State Number				
// Reporting Health Dept—City	/County			City/County Number								
Document Source			Surveillar	Surveillance Method Active Passive Follow up Reabstraction Unknown								
Did this report initiate a new □ Yes □ No □ Unknown		tigation?	Report Mo		lailed	□ 3-Faxed □ 4	-Phone □ 5-E	lectron	ic tra	nsfer □ 6-CD/disk		
III. Facility Providing I	nformatio	n (record	all dates a	as mm/dd/y	уууу)							
Facility Name							*Phone					
*Street Address												
City	Count	у			State/C	ountry	*ZIP Co	de				
Type 🛛 Hospital 🗌 Adult HIV clini			Private physician's office Screening, Diagnostic, Referral Age ic CTS STD clinic Other, specify			ferral Agency:	gency: <u>Other Facility</u> : □ Emergency room □ Laboratory □ Corrections □ Unknown □ Other, specify					
Date Form Completed			*Person Co			<u></u>	*Phone		er, sp	ecny		
IV. Patient Demograpi	//	<u> </u>	s as mm/de	d/www)			()					
Sex Male Female					Birth 🗆	US 🗆 Other/U	S dependency (specify)			
Date of Birth//					Alias D	ate of Birth	//		_			
Vital Status 1-Alive 2	-Dead		ate of Death/// State			State of Death	of Death					
Sexual Orientation Straight or heterosexual Lesbian or gay Bisexual Additional sexual orientation (specify)												
Date Identified	Declined to	answer □	Unknown									
Date Identified // Ethnicity □ Hispanic/Latino □ Not Hispanic/Latino □ Unknown												
Race□ American Indian/Alaska Na(check all that apply)□ Native Hawaiian/Other Pac							e					
V. Residence at Diagnosis (add additional addresses in Comments) (record all dates as mm/dd/yyyy)												
Address Event Type (check all that apply to addres	s below)	Residence at	t HIV diagnos	sis 🗆 Resid	dence at s	stage 3 (AIDS) dia	gnosis 🗆 Cheo	ck if <u>SA</u>	<u>ME</u> a	as current address		
Address Type Residential Bad address Correctional facility Foster home Homeless Military Other Postal Shelter Temporary												
*Street Address												
City County				State/Country					*ZIP Code			
Public reporting burden of this collection of information is estimated to average 20 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to CDC, Project Clearance Officer, 1600 Clifton Road, MS D-74, Atlanta, GA 30333, ATTN: PRA (0920-0573). Do not send the completed form to this address.												

VI. Facility	of Diagnosis (add	additional	facilities in Comment	s)						
Diagnosis Typ	e (check all that apply to	o facility belo	w)	(AIDS) □ Check if <u>SAME</u>			formatio	n		
Facility Name *Phone ()										
*Street Addres	;S									
City	ity County State/Country *ZIP				*ZIP C	Code				
Facility Type	Inpatient: □ Hospital □ Other, specify	□ Adult HIV	□ Private physician's office / clinic ecify	CTS STD clinic		Labo	<i>er Facility</i> : □ Emergency room aboratory □ Corrections □ Unknown ther, specify			
*Provider Name *Provider Phone () Specialty				ialty						
VII. Patient	History (respond f	to all ques	tions) (record all dates	s as mm/dd/yyyy)	🗆 Pe	diatri	c Risk	(ente	er in C	comments)
			sis of HIV infection, this p							
Sex with male							□ Yes	🗆 No	U	Inknown
Sex with female							□ Yes	□ No		Inknown
Injected nonpre	scription drugs	1					□ Yes	🗆 No		Inknown
Received clottir	ng factor for hemophilia/o	coagulation c	Jisorder				□ Yes	🗆 No		Inknown
Specify clotting				Date received /	/					
	IAL relations with any									
HETEROSEXUAL contact with person who injected drugs						nknown				
HETEROSEXUAL contact with bisexual male							□ Yes	🗆 No	U	Inknown
HETEROSEXUAL contact with person with hemophilia/coagulation disorder with documented HIV infection							□ Yes	□ No	U	Inknown
HETEROSEXU	HETEROSEXUAL contact with transfusion recipient with documented HIV infection						nknown			
HETEROSEXUAL contact with transplant recipient with documented HIV infection							□ Yes	□ No	🗆 U	Inknown
HETEROSEXUAL contact with person with documented HIV infection, risk not specified							□ Yes	□ No	🗆 U	Inknown
Received transfusion of blood/blood components (other than clotting factor) (document reason in Comments)						□ Yes	🗆 No	🗆 U	Inknown	
First date received/ / Last date received//										
Received transplant of tissue/organs or artificial insemination						□ Yes	🗆 No	🗆 U	Inknown	
Worked in a healthcare or clinical laboratory setting						□ Yes	🗆 No	🗆 U	Inknown	
If occupational exposure is being investigated or considered as primary mode of exposure, specify occupation and setting:										
Other documen	ted risk (include detail in	1 Comments)					□ Yes	□ No	🗆 U	Inknown
VIII. Clinica	I: Acute HIV Infe	ction and	Opportunistic Illne	sses (record all dates	as mm/dd/y	ууу)				
and enter patient of	or provider report of previou	us negative HIV	V test result in HIV Testing Histo	ted negative HIV test result data ory section malaise/fatigue, myalgia, ph					□ No □ No	Unknowr Unknowr
lymphadenopat Other evidence	thy)? Date of sign/sym suggestive of acute HIV	nptom onset V infection?				ı, 				
Date of evidence Opportunistic I	ce//									
Diagnosio		Dy Data	Diagnosia	Dx Data	Diagnosia					v Doto

opportunistic infesses							
Diagnosis	Dx Date	Diagnosis	Dx Date	Diagnosis	Dx Date		
Candidiasis, bronchi, trachea, or lungs		Herpes simplex: chronic ulcers (>1 mo. duration), bronchitis, pneumonitis, or esophagitis		M. tuberculosis, pulmonary ¹			
Candidiasis, esophageal		Histoplasmosis, disseminated or extrapulmonary		M. tuberculosis, disseminated or extrapulmonary ¹			
Carcinoma, invasive cervical		Isosporiasis, chronic intestinal (>1 mo. duration)		Mycobacterium, of other/unidentified species, disseminated or extrapulmonary			
Coccidioidomycosis, disseminated or extrapulmonary		Kaposi's sarcoma		Pneumocystis pneumonia			
Cryptococcosis, extrapulmonary		Lymphoma, Burkitt's (or equivalent)		Pneumonia, recurrent, in 12 mo. period			
Cryptosporidiosis, chronic intestinal (>1 mo. duration)		Lymphoma, immunoblastic (or equivalent)		Progressive multifocal leukoencephalopathy			
Cytomegalovirus disease (other than in liver, spleen, or nodes)		Lymphoma, primary in brain		Salmonella septicemia, recurrent			
Cytomegalovirus retinitis (with loss of vision)		Mycobacterium avium complex or M. kansasii, disseminated or extrapulmonary		Toxoplasmosis of brain, onset at >1 mo. of age			
HIV encephalopathy				Wasting syndrome due to HIV			

¹If a diagnosis date is entered for either tuberculosis diagnosis above, provide RVCT Case Number:

Result Desitive Desiti

IX. Laboratory Data (record additional tests and tests not specified below in Comments) (record all dates as mm/dd/yyyy) HIV Immunoassays TEST 🗆 HIV-1 IA 🗆 HIV-1/2 IA 🗆 HIV-1/2 Ag/Ab 🔅 HIV-2 IA Test Brand Name/Manufacturer _ Lab Name Facility Name Provider Name Collection Date ____/___/

Testing Option (if applicable)
Point-of-care test by provider
Self-test, result directly observed by a provider² Lab test, self-collected sample

CDC 50.42A Rev. 01/2023 (Page 2 of 4)

-ADULT HIV CONFIDENTIAL CASE REPORT-

IX. Laboratory Data (record additional tests and tests not specified below in Comments) (record all dates as mm/dd/yyyy) (cont)

TEST I HIV-1/2 Ag/Ab differentiating immunoassay (differentiates between HI	
Test Brand Name/Manufacturer	Lab Name
Facility Name Result Overall: Reactive	Collection Date / /
Analyte results: HIV-1 Ag: Reactive Nonreactive HIV-1/2 Ab	
Testing Option (if applicable) Point-of-care test by provider Self-test, res	ult directly observed by a provider ² Lab test, self-collected sample
TEST D HIV-1/2 Ag/Ab and type-differentiating immunoassay (differentiates an	nong HIV-1 Ag, HIV-1 Ab, and HIV-2 Ab)
Test Brand Name/Manufacturer	Lab Name
Facility Name	Provider Name
Result ³ Overall interpretation: Reactive Nonreactive Index Value Analyte results: HIV-1 Ag: Reactive Nonreactive Not reportation	CONECTION DATE///
HIV-1 Ab: □ Reactive □ Nonreactive □ Reactive □	ndifferentiated Index Value
HIV-2 Ab: 🗆 Reactive 🛛 Nonreactive 🗆 Reactive u	ndifferentiated Index Value
Testing Option (if applicable) Point-of-care test by provider Self-test, res	ult directly observed by a provider ² Lab test, self-collected sample
TEST I HIV-1/2 type-differentiating immunoassay (supplemental) (differentiate	,
Test Brand Name/Manufacturer	Lab Name
Facility Name	vith HIV-2 cross-reactivity
	/-1 indeterminate □ HIV-2 indeterminate □ HIV-1 positive □ HIV-2 positive
Analyte results: HIV-1 Ab: Positive Negative Indeterminate	Collection Date / /
HIV-2 Ab: Positive Negative Indeterminate	
Testing Option (if applicable) Point-of-care test by provider Self-test, res	ult directly observed by a provider ² Lab test, self-collected sample
TEST I HIV-1 WB I HIV-1 IFA I HIV-2 WB	l ah Namo
Test Brand Name/Manufacturer	Provider Name
Facility Name	Collection Date//
Testing Option (if applicable) Point-of-care test by provider Self-test, res	ult directly observed by a provider ² Lab test, self-collected sample
HIV Detection Tests	
TEST I HIV-1/2 RNA NAAT (Qualitative)	Lab Name
Test Brand Name/Manufacturer	_ Provider Name//
Result □ HIV-1 □ HIV-2 □ Both (HIV-1 and HIV-2) □ HIV, not differentia	
Testing Option (if applicable)	
TEST D HIV-1 RNA NAAT (Qualitative and Quantitative)	
Test Brand Name/Manufacturer	Lab Name
Facility Name	Provider Name
DESUL GUALIAUVE I DESCUVE I NOOLESCUVE	
Analyte results: HIV-1 Orantitative: □ Detectable above limit □ Detectable	conection Date / / /
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable abov	ectable within limits 🛛 Detectable below limit
Analyte results: HIV-1 Quantitative: Detectable Detectable Detectable Testing Option (if applicable) Detectable Point-of-care test by provider Self-test, rest	actable within limits □ Detectable below limit Copies/mL Log ult directly observed by a provider ² □ Lab test, self-collected sample
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT HIV-1 LIV-2 RNA/DNA	ctable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 HIV-2 RNA/ Test Brand Name/Manufacturer	ctable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 HIV-2 RNA/ Test Brand Name/Manufacturer	Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 HIV-2 RNA/ Test Brand Name/Manufacturer	cctable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 HIV-2 RNA/ Test Brand Name/Manufacturer	cctable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 Culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 Culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	ccable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name Collection Date //
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name Collection Date //
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name Collection Date //
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name Collection Date /
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name Provider Name Collection Date /
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cobies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit Detectable above limit Testing Option (if applicable) Point-of-care test by provider Self-test, res TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cobies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cobies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cobies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name
Analyte results: HIV-1 Quantitative: Detectable above limit HIV-1 culture HIV-2 RNA/ Test Brand Name/Manufacturer	cotable within limits Detectable below limit Copies/mL Log ult directly observed by a provider ² Lab test, self-collected sample DNA NAAT (Qualitative) HIV-2 culture Lab Name

X. Treatment/Services Referrals (record all dates as mm/dd/yyyy)							
Has this patient been informed of his/her HIV infection? This patient's partners will be notified about their HIV exposure and counseled by							
□ Yes □ No □ Unknown □ 1-Health dept □ 2-Physician/Provider □ 3-Patient □ 9-Unknown							
Evidence of receipt of HIV medical care other than laboratory test result (select one; record additional evidence in Comments)							
□ 1-Yes, documented □ 2-Yes, client self-report, only Date of medical visit or prescription///							
For Female Patient							
This patient is receiving or has been referred for gynecological or obstetrical services Is this patient currently pregnant? Has this patient delivered live-born infants? obstetrical services Yes No Unknown Image: Service servi							
For Children of Patient (record most recent birth in these boxes; record additional or multiple births in Comments)							
*Child's Name Child's Name							
Child's Last Name Soundex Child's State Number							
Facility Name of Birth		*Phone					
(if child was born at home, enter "home birth")		()					
Facility Type Inpatient: Outpatient:	Other Facili	it <u>y</u> : □ Emergency room					
□ Hospital □ Other, specify □ Corrections □ Unknown							
Other, specify	□ Other, sp	ecify					
*Street Address		*ZIP Code					
City County	,	State/Country					
XI. Antiretroviral Use History (record all dates as mm/de	d/vvvv)						
Main source of antiretroviral (ARV) use information (select one)		Date patient reported information					
□ Patient interview □ Medical record review □ Provider repo	rt □ NHM&E □ Other	/ /					
Ever taken any ARVs? Yes No Unknown							
If yes, reason for ARV use (select all that apply)							
HIV Tx ARV medications	_ Date began / / /	Date of last use/ /					
PrEP ARV medications							
PEP ARV medications							
PMTCT ARV medications							
HBV Tx ARV medications							
Other (specify reason)							
ARV medications Date began// Date of last use/ /							
XII. HIV Testing History (record all dates as mm/dd/yyyy	y)						
Main source of testing history information (select one) Date patient reported information							
Patient interview Medical record review Provider report NHM&E Other _//							
Ever had previous positive HIV test result? Yes No Unknown Date of first positive HIV test result//							
Was the first positive test result from a self-test performed by the patient? Yes No Unknown							
Ever had a negative HIV test result? Yes No Unknown Date of last negative HIV test result (if date is from							
a lab test with test type, enter in Lab Data section)//							
Was the last negative test result from a self-test performed by the patient? Yes No Unknown							
Number of negative HIV test results within the 24 months before the first positive test result □ Unknown							
How many of these negative test results were from self-tests performed by the patient?							
XIII. Comments							

XIV. *Local/Optional Fields

This report to CDC is authorized by law (Sections 304 and 306 of the Public Health Service Act, 42 USC 242b and 242k). Response in this case is voluntary for federal government purposes but may be mandatory under state and local statutes. Your cooperation is necessary for the understanding and control of HIV. Information in CDC's National HIV Surveillance System that would permit identification of any individual on whom a record is maintained is collected with a guarantee that it will be held in confidence, will be used only for the purposes stated in the assurance, and will not otherwise be disclosed or released without the consent of the individual in accordance with Section 308(d) of the Public Health Service Act (42 USC 242m).