GEORGIA ADULT HIV/AIDS CONFIDENTIAL CASE REPORT FORM (Patients \geq 13 years of age at time of diagnosis)

Mail completed form to: Georgia Department of Public Health, Epidemiology Section P.O. Box 2107 Atlanta, GA 30301 For additional information: Phone: 1-800-827-9769 or visit our website at http://health.state.ga.us/epi/hivaids

All health care providers diagnosing and/or providing care to a patient with HIV are obligated to report using Georgia HIV/AIDS Case Report. Case reports should be completed within seven (7) days after diagnosing or providing care to a patient with HIV/AIDS. Providers are required to submit reports on any patient new to his or her care, regardless if they have previously received care elsewhere.

Patients <13 should be reported on a Pediatric Case Report Form (https://dph.georgia.gov/hivaids-case-reporting)

Patient Identification (record all dates as mm/dd/yyyy) *Information NOT transmitted to CDC

*First Name		*Middle Nan	ne		*Last Name		Las	t Name Soundex
Alternate Name Type (ex: /	Alias, Married)		*First Name	9	*Middle Name	*Last	Nam	10
Address Type Residentia Foster home Homeless				*Current Addres	s, Street			Address Date
*Phone ()	City		County		State/Country		*ZIP	Code
*Medical Record Number				*Other ID Type		*Number		

Facility Providing Information (record all dates as mm/dd/yyyy)

Facility N	lame				*Phone	
*Street A	ddress					
City		County		State/Country	*ZIP Co	de
Facility Type	<u>Inpatient</u> : □ Hospital	<u>Outpatient</u> : □ □ Adult HIV cli	Private physician's office	<u>Screening. Diagnostic. Referral A</u> □ CTS □ STD clinic	gency:	<u>Other Facility</u> : □ Emergency room □ Laboratory □ Corrections □ Unknown
	☐ Other, specify			□ Other, specify		Other, specify
Date For	m Completed		*Person Completing F	Form	*Phone	

Patient Demographics (record all dates as mm/dd/yyyy)

Sex Assigned at Birth	Country of Birth
Male Female Unknown	US Other/US dependency (please specify)
Date of Birth///	Alias Date of Birth///
Vital Status 1-Alive 2-Dead Date o	Death/ / State of Death
Current Gender Identity Male Female Transger Additional gender identity (spe	der male-to-female (MTF)
Ethnicity Hispanic/Latino Not Hispanic/Latino	Inknown Expanded Ethnicity
	□ Asian □ Black/African American Expanded Race slander □ White □ Unknown

Residence at Diagnosis (add additional addresses in Comments) (record all dates as mm/dd/yyyy)

Address Type				
(check all that apply to address below	v) 🗆 Residence at HIV diagnosis	□ Residence at stage 3 (AIDS) diagnosis	Check if SAM	<u>1E</u> as current address
*Street Address				
City	County	State/Country		*ZIP Code

Facility of Diagnosis (add additional facilities in Comments)

Diagnosis Typ	e (check all that apply to	o facility below) □ HIV	□ Stage 3 (AIDS)	Check if <u>SAME</u> as facilit	ty providing info	rmation
Facility Name					*Phone()
*Street Addres	S					
City		County	State	e/Country	*ZIP Co	de
Facility Type	<u>Inpatient</u> : □ Hospital □ Other, specify	Outpatient: □ Private physic □ Adult HIV clinic □ Other, specify		ning, Diagnostic, Referral Agen S □ STD clinic er, specify	□ Laborat	<u>sility</u> : □ Emergency room tory □ Corrections □ Unknow specify
*Provider Nam	e	*Provider Pho	one ()		Specialty	

Patient History (respond to all questions) (record all dates as mm/dd/yyyy)

Pediatric Risk (please enter in Comments)

After 1977 and before the earliest known diagnosis of HIV infection, this patient had:			
Sex with male	□ Yes	□ No	Unknown
Sex with female	□ Yes	🗆 No	Unknown
Injected nonprescription drugs	□ Yes	🗆 No	Unknown
Received clotting factor for hemophilia/coagulation disorder Specify clotting factor: Date received//	□ Yes	□ No	Unknown
HETEROSEXUAL relations with any of the following:			
HETEROSEXUAL contact with intravenous/injection drug user	□ Yes	□ No	Unknown
HETEROSEXUAL contact with bisexual male	□ Yes	□ No	Unknown
HETEROSEXUAL contact with person with hemophilia/coagulation disorder with documented HIV infection	□ Yes	□ No	Unknown
HETEROSEXUAL contact with transfusion recipient with documented HIV infection	□ Yes	🗆 No	Unknown
HETEROSEXUAL contact with transplant recipient with documented HIV infection	□ Yes	□ No	Unknown
HETEROSEXUAL contact with person with documented HIV infection, risk not specified	□ Yes	□ No	Unknown
Received transfusion of blood/blood components (other than clotting factor) (document reason in Comments)	□ Yes	🗆 No	Unknown
First date received// Last date received//			
Received transplant of tissue/organs or artificial insemination	□ Yes	🗆 No	Unknown
Worked in a healthcare or clinical laboratory setting	□ Yes	□ No	Unknown
If occupational exposure is being investigated or considered as primary mode of exposure, specify occupation and setting:			
Other documented risk (please include detail in Comments)	□ Yes	□ No	Unknown

Clinical: Acute HIV Infection and Opportunistic Illnesses (record all dates as mm/dd/yyyy)

Suspect acute HIV infection? If enter patient or provider report of prev		two items below; enter documented negatives in HIV Testing History section.	re HIV test data i	n Laboratory Data section, and	□ Yes	□ No	Unknown
lymphadenopathy)? Date of sig	n/symptom onset		atigue, myalgia	a, pharyngitis, rash,	Yes	□ No	Unknown
Other evidence suggestive of act Date of evidence//	ute HIV infection?	If YES, please describe:			Yes	□ No	Unknown
Opportunistic Illnesses							
Diagnosis	Dx Date	Diagnosis	Dx Date	Diagnosis		Dx Da	ite
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 extrapulmonary ¹	or		
Carcinoma, invasive cervical		Isosporiasis, chronic intestinal (>1 mo. duration)		Mycobacterium, of other/unid species, disseminated or extra			
Coccidioidomycosis, disseminated or extrapulmonary		Kaposi's sarcoma		Pneumocystis pneumonia			
Cryptococcosis, extrapulmonary		Lymphoma, Burkitt's (or equivalent)		Pneumonia, recurrent, in 12 r	no. 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, recur	rent		
Cytomegalovirus retinitis (with loss of vision)		Mycobacterium avium complex or M. kansasii, disseminated or extrapulmonary		Toxoplasmosis of brain, onset of age	at >1 mo.		
HIV encephalopathy				Wasting syndrome due to HIV	1		
¹ If a diagnosis date is entered for either tu	uberculosis diagnosis	above, provide RVCT Case Number:					

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

TEST 1 🗆 HIV-1 IA 🗆 HIV-1/2 IA 🗆 HIV-1/2 Ag/Ab 🗆 HIV-1 WB 🗆 HIV-1 IF	FA 🗆 HIV-2 IA 🗆 HIV-2 WB
Test brand name/Manufacturer	Lab name
Facility name	
Result Positive Negative Indeterminate	Collection Date/ / Deint-of-care rapid test
TEST 2 🗆 HIV-1 IA 🗆 HIV-1/2 IA 🗆 HIV-1/2 Ag/Ab 🗆 HIV-1 WB 💷 HIV-1 IF	
Test brand name/Manufacturer	l ah name
Facility name	Provider name
Facility name Result Positive Negative Indeterminate	Collection Date / Deint of care rapid text
HIV Immunoassays (Differentiating)	
□ HIV-1/2 type-differentiating immunoassay	Role of test in diagnostic algorithm
(differentiates between HIV-1 Ab and HIV-2 Ab)	□ Screening/initial test □ Confirmatory/supplemental test
Test brand name/Manufacturer	
Facility name	Provider name
Result¹ Overall interpretation : □ HIV-1 positive □ HIV-2 positive □ HIV positive □ HIV-2 positive □ HIV po	
□ HIV-1 indeterminate □ HIV-2 indeterminate	
	Collection Date// Point-of-care rapid test
	Always complete the overall interpretation. Complete the analyte results when available.
□ HIV-1/2 Ag/Ab differentiating immunoassay (differentiates between HIV Ag	
Test brand name/Manufacturer	
Facility name	
Result □ Ag positive □ Ab positive □ Both (Ag and Ab positive) □ Negativ	
Collection Date// Dent-of-care rapid test	
HIV-1/2 Ag/Ab and type-differentiating immunoassay (differentiates among	
Test brand name/Manufacturer	
Facility name	
Result ² Overall interpretation: □ Reactive □ Nonreactive □ Index value	
Analyte results: HIV-1 Ag: Reactive Nonreactive Not report	able due to high Ab level Index value
HIV-1 Ab: Reactive Nonreactive Reactive	Indifferentiated Index value
HIV-2 Ab: Reactive Nonreactive Reactive	Indifferentiated Index value
Collection Date / / / Deint-of-care rapid test ²	Complete the overall interpretation and the analyte results.
Collection Date// Dent-of-care rapid test ² (Complete the overall interpretation and the analyte results.
HIV Detection Tests (Qualitative) TEST	DNA NAAT (Qualitative)
HIV Detection Tests (Qualitative) TEST	DNA NAAT (Qualitative)
HIV Detection Tests (Qualitative) TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/E Test brand name/Manufacturer Facility name	DNA NAAT (Qualitative)
HIV Detection Tests (Qualitative) TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/D Test brand name/Manufacturer Facility name Result Positive Negative Indeterminate	DNA NAAT (Qualitative) HIV-2 culture Lab name
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HIV Detection Tests (Qualitative) TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/D Test brand name/Manufacturer	DNA NAAT (Qualitative) HIV-2 culture Lab name Provider name Collection Date// or after diagnosis. NAAT (Quantitative viral load) Lab name Provider name Log Collection Date/_/ NAAT (Quantitative viral load) Lab name Provider name Collection Date/_/ Test brand name/Manufacturer Facility name Collection Date/_/ Collection Date/_/
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HIV Detection Tests (Qualitative) TEST HIV-1 RNA/DNA NAAT (Qualitative) HIV-1 culture HIV-2 RNA/D Facility name	DNA NAAT (Qualitative) HIV-2 culture Lab name

Documentation of Tests					
Did documented laboratory test results meet approved HIV diagnostic algorithm criteria? Ves Unknown					
If YES, provide specimen collection date of earliest positive test fo					
Complete the above only if none of the following was positive: HIV-1 Will If HIV laboratory tests were not documented, is HIV diagnosis documented.					
in the laboratory tests were not documented, is the diagnosis doct		te of diagnosis////			
Date of last documented negative HIV test (before HIV diagnosis dat	e) / /				
Specify type of test:					
Treatment/Services Referrals (record all dates as mm/	dd/yyyy)				
Has this patient been informed of his/her HIV infection? This pa	tient's partners will be notified about	their HIV exposure and counseled by			
Yes No Unknown I-Hea	alth dept 🛛 2-Physician/Provider 🗖 🤅	3-Patient 🗆 9-Unknown			
Evidence of receipt of HIV medical care other than laboratory test					
□ 1-Yes, documented □ 2-Yes, client self-report, only Date of me For Female Patient	edical visit or prescription//				
This patient is receiving or has been referred for gynecological or	Is this patient currently pregnant?	Has this patient delivered live-born infants?			
obstetrical services Yes No Unknown	□ Yes □ No □ Unknown	□ Yes □ No □ Unknown			
For Children of Patient (record most recent birth in these boxes; re	cord additional or multiple births in Comr	ments)			
*Child's Name		Child's Date of Birth			
Obildie Leet Neme Coundary	Okildia Otata Numekan	<u> </u>			
Child's Last Name Soundex Facility Name of Birth	Child's State Number	*Phone			
(if child was born at home, enter "home birth")					
Facility Type Inpatient: Outpatient:	Other Facil	<i>lity</i> : □ Emergency room			
□ Hospital □ Other, speci	fy Correction	ons 🗆 Unknown			
□ Other, specify	□ Other, sp	1			
*Street Address		*ZIP Code			
City County	y	State/Country			
Antiretroviral Use History (record all dates as mm/dd/yy	(עע)				
Main source of antiretroviral (ARV) use information (select one)		Date patient reported information			
Patient interview Medical record review Provider repo	ort NHM&E Other	//			
Ever taken any ARVs? Yes No Unknown	ort 🗆 NHM&E 🗆 Other	1			
Ever taken any ARVs? Yes No Unknown If yes, reason for ARV use (select all that apply)					
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///			
Ever taken any ARVs? Yes No Unknown If yes, reason for ARV use (select all that apply) HIV Tx ARV medications PrEP ARV medications	Date began / / Date began / / /	Date of last use / / Date of last use / /			
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