Patient Identification	(record all da	ates as	s mm/dd/yyy	у)									
*First		Middle								Last Name			
Name Name **Fir		1	e *First			Name *Middle			*1 :	*Last			
Alternate Name Type   Diltil   Alias   Maldell					Name				Name				
<b>Address Type</b> □ Residential	□ Bad address	□ Corr	rectional facility	*Curre	nt Addres	ss, Street					Address Date		
	e □ Homeless Shelter □ Temp		ry □ Other								/ /		
Postal Shelter Temporary  *Phone  City			County			te/Country				*ZIP Code			
( )													
*Medical Record Number				*Other ID	Туре			Alia	*SSN as *SSN				
U.S. Department of Health and Human Services	(Patien	nts <u>&gt;</u> 13 y	t HIV Co	ime of diag	nosis) *Ir						Centers for Disease Control and Prevention (CDC)		
Health Department U					<u>')</u>					ved OM	B no. 0920-0573 Exp. 11/30/202		
Date Received at Health De			KY Testing/E Number (KY N				KY State Number						
Reporting Health Dept—Cit			·	<u> </u>		City/Cour	nty Nu	ımber					
Document Source			Surveillance	Method	□ Active	□ Passive	□ F	ollow u	p □ Rea	bstracti	ion 🗆 Unknown		
Did this report initiate a ne □ Yes □ No □ Unknown	_		1? Report Medium  □ 1-Field visit □ 2-Mailed □ 3-Faxed □ 4-Phone □ 5-Electronic transfer □ 6-CD/disk										
Facility Providing Info	ormation (re	cord al	l dates as n	nm/dd/vvv	v)								
Facility Name	, , ,				<u> </u>			,	*Phone(	)			
*Street Address													
City	County				Statol	Country		,	*ZIP Code				
City			State/Coun			Journal y	Li Code						
Type ☐ Hospital ☐ Adult HIV			IV clinic CTS ST				Diagnostic, Referral Agency:       Other Facility:       □ Emergency room         ITD clinic       □ Laboratory       □ Corrections       □ Unknown						
☐ Other, specify _		Other, spe				, specify					pecify		
Date Form Completed  ☐ Same as Date Received	/ /		*Person Co	ompleting I	-orm			ľ	*Phone (	)			
			Surv. Inves	stigator:									
Patient Demographic	s (record all	dates a	as mm/dd/yy	уу)									
Sex Assigned at Birth				ntry of Birt									
☐ Male ☐ Female ☐ Unk	nown			S  Unknov	vn □ Oth	er/US depen	ndency	/ (please	e specify) _				
Date of Birth/	/	Alias D	ate of Birth	/	_/								
Vital Status □ 1-Alive □			Date of Dea		/			State of					
•	<ul><li>□ Male □ Fem</li><li>□ Additional ger</li></ul>		•	nale-to-fema	ale (MTF)	□ Transge	nder f	emale-to	o-male (FT -	M) 🗆	Unknown		
Ethnicity													
Race													
Residence at Diagnos					nts) (rec	ord all dat	tes as	s mm/c	dd/vvvv)				
	•									□ Che	ck if <u>SAME</u> as current address		
Address Type □ Resident			<u>'</u>					` '					
*Street Address				,							1 7		
HIV: City		Coun	County			State/Country				*	*ZIP Code		
AIDS: 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.** 

STATE/LOCA	AL USE ONLY										
*Provider Nam	*Provider Name (Last, First, M.I.)										
Hospital/Facilit	ty										
Facility of D	iagnosis (add a	dditional facil	ities in Comments)								
	(check all that app		-	(AIDS)	□ Check if <u>SAME</u> a	s facility prov	iding info	rmation			
*Facility Name	,	, , , , , , , , , , , , , , , , , , , ,	,	/		*Pho		)			
*Street Address	 S						`	,			
City											
Facility Type	<i>Inpatient:</i> □ Hospit	al <i>Outpatient:</i> $\square$	ent: □ Private physician's office Screening, Diagnostic, Referral Agency: Other Facility: □ Emergency room								
' ''	□ Other, specify □ Adult HIV clinic □ CTS □ STD clinic □ La				□ Labora	ooratory   Corrections  Unknown					
								specify			
*Provider Name											
Patient Hist	ory (respond to	all questions)	(record all dates as i	mm/dd/y	ууу) 🗆	Pediatric	Risk (	please e	nter	in Com	ments
After 1977 and	before the earliest	known diagnos	is of HIV infection, this p	oatient had	d:						
Sex with male								Yes □	No	□ Unkno	wn
Sex with female								Yes □	No	□ Unkno	wn
Injected nonpres	scription drugs							Yes □	No	□ Unkno	wn
	g factor for hemophi	lia/coagulation dis	sorder	5				Yes □	No	□ Unkno	wn
Specify clotting f	factor: AL relations with a	ny of the followi	na:	Date	received /	/					
	AL contact with intra							Yes 🗆	No	□ Unkno	)W/D
	AL contact with bisex		arug user					Yes 🗆		Unkno	
			a/coagulation disorder with	h documen	nted HIV infection			Yes 🗆		Unkno	
HETEROSEXUAL contact with person with hemophilia/coagulation disorder with documented HIV infection  HETEROSEXUAL contact with transfusion recipient with documented HIV infection									No	Unkno	
HETEROSEXUAL contact with transfusion recipient with documented HIV infection  HETEROSEXUAL contact with transplant recipient with documented HIV infection								Yes 🗆		□ Unkno	
HETEROSEXUAL contact with transplant recipient with documented HIV infection, risk not specified								Yes 🗆		□ Unkno	
	Received transfusion of blood/blood components (other than clotting factor) (document reason in Comments)							wn			
			date received/								
	olant of tissue/organs							Yes □	No	□ Unkno	wn
Worked in a hea	Ithcare or clinical lab	poratory setting						Yes □	No	□ Unkno	wn
	exposure is being inv										
as primary mode of exposure, specify occupation and setting:  Other decumented risk (please include detail in Comments)								Voc	No	□ Halma	
Other documented risk (please include detail in Comments)							VVII				
			rtunistic Illnesses	<u> </u>							
			items below; enter document in HIV Testing History section.		HIV test data in Labo	oratory Data sec	ction, and	□ Yes	<b>3</b> 🗆 l	No 🗆 Ur	nknown
Clinical signs/sv	mptoms consistent	with acute retrovi	ral syndrome (e.g., fever, i		tigue, myalgia, pha	ryngitis, rash	,	□ Yes	_ N	lo □Un	known
Clinical signs/symptoms consistent with acute retroviral syndrome (e.g., fever, malaise/fatigue, myalgia, pharyngitis, rash, lymphadenopathy)? Date of sign/symptom onset/MUST INCLUDE DATE  Other evidence suggestive of acute HIV infection? If YES, please describe: Yes No Unknown							known				
Date of evidence Opportunistic II		MUST	INCLUDE DATE								
Diagnosis		x Date	Diagnosis		Dx Date	Diagnosis			Dx	Date	
Candidiasis, bronchi,	trachea, or lungs		Herpes simplex: chronic ulcers duration), bronchitis, pneumoni esophagitis			M. tuberculosis	, pulmonary	y <sup>1</sup>			
Candidiasis, esophaç	geal		Histoplasmosis, disseminated of	or		M. tuberculosis, extrapulmonary		ed or			
Carcinoma, invasive	extrapulmonary extrapulmonary extrapulmonary rcinoma, invasive cervical Isosporiasis, chronic intestinal (>1 mo. Mycobacterium, of oth duration) species, disseminated					, of other/u					
	occidioidomycosis, disseminated or Kaposi's sarcoma Pneumocystis pneum							,			
extrapulmonary Cryptococcosis, extra	apulmonary		Lymphoma, Burkitt's (or equiva	alent)		Pneumonia, red	current, in 1	2 mo. period	士		
Cryptosporidiosis, chronic intestinal (>1						ıltifocal					
Cytomegalovirus disease (other than in Lymphoma, primary in brain Salmonella septicemia							current	+			
liver, spleen, or nodes)  Cytomegalovirus retinitis (with loss of Mycobacterium avium complex or M. Toxoplasmosis of brain, onset at >1 mo.											
vision) kansasii, disseminated or extrapulmonary of age											
HIV encephalopathy  1If a diagnosis date is	s entered for either tuber	rculosis diagnosis abo	vve, provide RVCT Case Numbe	er:		Wasting syndro	me due to l	HIV			

Laboratory Data (record additional tests and tests not specified	d below in Comments) (record all dates as mm/dd/yyyy)				
HIV Immunoassays (Nondifferentiating)					
TEST 1 - HIV-1 IA - HIV-1/2 IA - HIV-1/2 Ag/Ab - HIV-1 WB - HIV-1 I	FA □ HIV-2 IA □ HIV-2 WB				
Test brand name/Manufacturer	Lab name				
Facility name	Provider name				
	Collection Date// Doint-of-care rapid test				
TEST 2   HIV-1 IA   HIV-1/2 IA   HIV-1/2 Ag/Ab   HIV-1 WB   HIV-1 I					
Test brand name/Manufacturer	Lab name				
Facility name					
Result □ Positive □ Negative □ Indeterminate	Collection Date// Point-of-care rapid test				
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	Lab name				
Facility name					
Result <sup>1</sup> Overall interpretation:   HIV-1 positive  HIV-2 positive  HIV p					
☐ HIV-1 indeterminate ☐ HIV-2 indeterminate					
Analyte results: HIV-1 Ab: □ Positive □ Negative □ Indeterminate					
	<sup>1</sup> 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) ☐ Negative					
	ve 🗆 mvalid				
Collection Date/ Point-of-care rapid test					
☐ HIV-1/2 Ag/Ab and type-differentiating immunoassay (differentiates amon					
Test brand name/Manufacturer					
Facility name					
Result <sup>2</sup> Overall interpretation: □ Reactive □ Nonreactive □ Index value					
Analyte results: HIV-1 Ag: ☐ Reactive ☐ Nonreactive ☐ Not repor					
HIV-1 Ab: □ Reactive □ Nonreactive □ Reactive	undifferentiated Index value				
HIV-2 Ab: □ Reactive □ Nonreactive □ Reactive					
Collection Date//   Point-of-care rapid test 2	Complete the overall interpretation and the analyte results.				
HIV Detection Tests (Qualitative)					
TEST □ HIV-1 RNA/DNA NAAT (Qualitative) □ HIV-1 culture □ HIV-2 RNA/	DNA NAAT (Qualitative) □ HIV-2 culture				
Test brand name/Manufacturer	Lab name				
	Provider name				
Result □ Positive □ Negative □ Indeterminate	Collection Date / / / /				
HIV Detection Tests (Quantitative viral load) Note: Include earliest test a	t or after diagnosis.				
TEST 1 ☐ HIV-1 RNA/DNA NAAT (Quantitative viral load) ☐ HIV-2 RNA/DNA	NAAT (Quantitative viral load)				
Test brand name/Manufacturer	Lab name				
Facility name	Provider name				
	LogCollection Date///				
TEST 2 ☐ HIV-1 RNA/DNA NAAT (Quantitative viral load) ☐ HIV-2 RNA/DNA					
	Lab name				
	Provider name				
Result □ Detectable □ Undetectable Copies/mL					
Drug Resistance Tests (Genotypic)	LogCollection Date//				
TEST   HIV-1 Genotype (Unspecified)	Test brand name/Manufacturer				
Lab name					
Provider name Immunologic Tests (CD4 count and percentage)					
	_ CD4 percentage % Collection Date / /				
Test brand name/Manufacturer					
Facility name	Provider name				
First CD4 result <200 cells/µL or <14%: CD4 countcells/µL	CD4 percentage % Collection Date / / /				
Test brand name/Manufacturer	Lab name				
Facility name					
	Lab name				
Facility name	Provider name				
Did documented laboratory test results meet approved HIV diagnostic algorithm criteria?   Yes No Unknown					
If YES, provide specimen collection date of earliest positive test for this algorithm / / / / Complete the above only if none of the following were positive for HIV-1: Western blot, IFA, culture, viral load, qualitative NAAT (RNA or DNA), HIV-1/2 type-					
differentiating immunoassay (supplemental test), stand-alone p24 antigen, or nucleotide sequence.					
If HIV laboratory tests were not documented, is HIV diagnosis documented by a physician?   Yes  No  Unknown					
and the state of t					
	If YES, provide date of diagnosis				
Date of last decumented negative HIV test (hefers HIV diagnosis date)	If YES, provide date of diagnosis / / /				
Date of last documented negative HIV test (before HIV diagnosis date) Specify type of test:	If YES, provide date of diagnosis/ //				

Treatment/Services Referrals (record all dates a	as mm/dd/yyyy)					
	This patient's partners will be notified abo  ☐ 1-Health dept ☐ 2-Physician/Provice					
Evidence of receipt of HIV medical care other than laborate		· · · · · · · · · · · · · · · · · · ·				
	e of medical visit or prescription/					
	nrolled at (Clinic):   HRSA Sponsored	□ Other □ None □ Unknown				
ID Facility Name:						
Antiretroviral Use History (record all dates as m						
Main source of antiretroviral (ARV) use information (select or □ Patient interview □ Medical record review □ Prov	one) vider report    NHM&E   Other	Date patient reported information				
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	Date began / /	Date of last use///				
□ PEP ARV medications	Date began / /	Date of last use///				
□ PMTCT ARV medications	Date began / /	Date of last use///				
□ HBV Tx ARV medications	Date began / /					
□ Other (specify reason)						
ARV medications		Date of last use / / /				
For Female Patient						
This patient is receiving or has been referred for gynecolog obstetrical services ☐ Yes ☐ No ☐ Unknown	gical or Is this patient currently pregnant	Has this patient delivered live-born infants?  □ Yes □ No □ Unknown				
For Children of Patient (record most recent birth in these b	poxes; record additional or multiple births in C	omments)				
*Child's Name		Child's Date of Birth				
Child's Last Name Soundex	Child's State Number					
Facility Name of Birth		*Phone				
(if child was born at home, enter "home birth")		( )				
		Facility:   Emergency room				
☐ Hospital ☐ Oth ☐ Other, specify		ections  Unknown r, specify				
*Street Address	3.00	*ZIP Code				
City	County	State/Country				
HIV Testing History (record all dates as mm/dd/y)	ann)	•				
Main source of testing history information (select one)	133)	Date patient reported information				
□ Patient interview □ Medical record review □ Provider re	port □ NHM&E □ Other	/ /				
Ever had previous positive HIV test?   Yes   No   Un	IIV test / /					
Ever had a negative HIV test?   Yes   No  Unknown  Date of last negative HIV test (if date is from a lab test with test type, enter in Lab Data section)//						
Number of negative HIV tests within the 24 months before		a III Lab Data Section)				
Comments						
*Local/Optional Fields						
*DATE REFERRED FOR PARTNER SERVICES (PS):/	//_ Already in NEDSS □	Yes   No NEDSS ID #:				
SOUNDEX:						
This report to CDC is authorized by law (Sections 304 and 306 of the Public Health under state and local statutes. Your cooperation is necessary for the understanding whom a record is maintained is collected with a guarantee that it will be held in con disclosed or released without the consent of the individual in accordance with Secti	g and control of HIV. Information in CDC's National HIV Surv ifidence, will be used only for the purposes stated in the assu	eillance System that would permit identification of any individual on				