

Patient's Name _____

**REPORT OF VERIFIED CASE
OF TUBERCULOSIS**

Street Address _____ (Last) _____ (First) _____ (M.I.) _____ (ZIP CODE)



**Centers for Disease
Control and Prevention**
National Center for HIV/AIDS,
Viral Hepatitis, STD, and
TB Prevention

FORM APPROVED OMB NO. 0920-0026 Exp. Date 05/31/2014

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1. Date Reported Month Day Year		3. Case Numbers Year Reported (YYYY) State Code Locally Assigned Identification Number State Case Number City/County Case Number Linking State Case Number Linking State Case Number	
2. Date Submitted Month Day Year		Reason: Reason:	

4. Reporting Address for Case Counting City Within City Limits (select one) ☐ Yes ☐ No County ZIP CODE		8. Date of Birth Month Day Year	
5. Count Status (select one) Countable TB Case ☐ Count as a TB case Noncountable TB Case ☐ Verified Case: Counted by another U.S. area (e.g., county, state) ☐ Verified Case: TB treatment initiated in another country Specify _____ ☐ Verified Case: Recurrent TB within 12 months after completion of therapy		9. Sex at Birth (select one) ☐ Male ☐ Female 10. Ethnicity (select one) ☐ Hispanic or Latino ☐ Not Hispanic or Latino	
6. Date Counted Month Day Year		11. Race (select one or more) ☐ American Indian or Alaska Native ☐ Asian: Specify _____ ☐ Black or African American ☐ Native Hawaiian or Other Pacific Islander: Specify _____ ☐ White	
7. Previous Diagnosis of TB Disease (select one) ☐ Yes ☐ No If YES, enter year of previous TB disease diagnosis:		12. Country of Birth "U.S.-born" (or born abroad to a parent who was a U.S. citizen) (select one) ☐ Yes ☐ No Country of birth: Specify _____	
14. Pediatric TB Patients (<15 years old) Country of Birth for Primary Guardian(s): Specify _____ Guardian 1 _____ Guardian 2 _____ Patient lived outside U.S. for >2 months? ☐ Yes ☐ No ☐ Unknown (select one) If YES, list countries, specify: _____		13. Month-Year Arrived in U.S. Month Year	
15. Status at TB Diagnosis (select one) ☐ Alive ☐ Dead If DEAD, enter date of death: If DEAD, was TB a cause of death? (select one) ☐ Yes ☐ No ☐ Unknown		16. Site of TB Disease (select all that apply) ☐ Pulmonary ☐ Pleural ☐ Lymphatic: Cervical ☐ Lymphatic: Intrathoracic ☐ Lymphatic: Axillary ☐ Lymphatic: Other ☐ Lymphatic: Unknown ☐ Laryngeal ☐ Bone and/or Joint ☐ Genitourinary ☐ Meningeal ☐ Peritoneal ☐ Other: Enter anatomic code(s) (see list): ☐ Site not stated } 1 2 3	

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17. Sputum Smear (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown	Date Collected: Month Day Year 	
18. Sputum Culture (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown	Date Collected: Month Day Year 	Date Result Reported: Month Day Year
Reporting Laboratory Type (select one): ☐ Public Health Laboratory ☐ Commercial Laboratory ☐ Other		
19. Smear/Pathology/Cytology of Tissue and Other Body Fluids (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown	Date Collected: Month Day Year 	Enter anatomic code (see list):
Type of exam (select all that apply): ☐ Smear ☐ Pathology/Cytology		
20. Culture of Tissue and Other Body Fluids (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown	Date Collected: Month Day Year 	Enter anatomic code (see list):
Date Result Reported: Month Day Year 		
Reporting Laboratory Type (select one): ☐ Public Health Laboratory ☐ Commercial Laboratory ☐ Other		
21. Nucleic Acid Amplification Test Result (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown ☐ Indeterminate	Date Collected: Month Day Year 	Date Result Reported: Month Day Year
Enter specimen type: ☐ Sputum OR If not Sputum, enter anatomic code (see list):		
Reporting Laboratory Type (select one): ☐ Public Health Laboratory ☐ Commercial Laboratory ☐ Other		
Initial Chest Radiograph and Other Chest Imaging Study		
22A. Initial Chest Radiograph (select one) ☐ Normal ☐ Abnormal* (consistent with TB) ☐ Not Done ☐ Unknown * For ABNORMAL Initial Chest Radiograph: Evidence of a cavity (select one): ☐ Yes ☐ No ☐ Unknown Evidence of miliary TB (select one): ☐ Yes ☐ No ☐ Unknown		
22B. Initial Chest CT Scan or Other Chest Imaging Study (select one) ☐ Normal ☐ Abnormal* (consistent with TB) ☐ Not Done ☐ Unknown * For ABNORMAL Initial Chest CT Scan or Other Chest Imaging Study: Evidence of a cavity (select one): ☐ Yes ☐ No ☐ Unknown Evidence of miliary TB (select one): ☐ Yes ☐ No ☐ Unknown		
23. Tuberculin (Mantoux) Skin Test at Diagnosis (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown	Date Tuberculin Skin Test (TST) Placed: Month Day Year 	Millimeters (mm) of induration:
24. Interferon Gamma Release Assay for Mycobacterium tuberculosis at Diagnosis (select one) ☐ Positive ☐ Not Done ☐ Negative ☐ Unknown ☐ Indeterminate	Date Collected: Month Day Year 	
Test type: Specify _____		
25. Primary Reason Evaluated for TB Disease (select one) ☐ TB Symptoms ☐ Abnormal Chest Radiograph (consistent with TB) ☐ Contact Investigation ☐ Targeted Testing ☐ Health Care Worker ☐ Employment/Administrative Testing ☐ Immigration Medical Exam ☐ Incidental Lab Result ☐ Unknown		

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26. HIV Status at Time of Diagnosis (select one)

- ☐ Negative
 ☐ Indeterminate
 ☐ Not Offered
 ☐ Unknown
☐ Positive
 ☐ Refused
 ☐ Test Done, Results Unknown

If POSITIVE, enter:

State HIV/AIDS Patient Number:

City/County HIV/AIDS Patient Number:

27. Homeless Within Past Year (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

28. Resident of Correctional Facility at Time of Diagnosis (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

If YES, (select one):

- ☐ Federal Prison
 ☐ Local Jail
 ☐ Other Correctional Facility
☐ State Prison
 ☐ Juvenile Correction Facility
 ☐ Unknown

If YES, under custody of Immigration and Customs Enforcement? (select one)

- ☐ No
 ☐ Yes

29. Resident of Long-Term Care Facility at Time of Diagnosis (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

If YES, (select one):

- ☐ Nursing Home
 ☐ Residential Facility
 ☐ Alcohol or Drug Treatment Facility
 ☐ Unknown
☐ Hospital-Based Facility
 ☐ Mental Health Residential Facility
 ☐ Other Long-Term Care Facility

30. Primary Occupation Within the Past Year (select one)

- ☐ Health Care Worker
 ☐ Migrant/Seasonal Worker
 ☐ Retired
 ☐ Not Seeking Employment (e.g. student, homemaker, disabled person)
☐ Correctional Facility Employee
 ☐ Other Occupation
 ☐ Unemployed
 ☐ Unknown

31. Injecting Drug Use Within Past Year (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

32. Non-Injecting Drug Use Within Past Year (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

33. Excess Alcohol Use Within Past Year (select one)

- ☐ No
 ☐ Yes
 ☐ Unknown

34. Additional TB Risk Factors (select all that apply)

- ☐ Contact of MDR-TB Patient (2 years or less)
 ☐ Incomplete LTBI Therapy
 ☐ Diabetes Mellitus
 ☐ Other Specify _____
☐ Contact of Infectious TB Patient (2 years or less)
 ☐ TNF- α Antagonist Therapy
 ☐ End-Stage Renal Disease
 ☐ None
☐ Missed Contact (2 years or less)
 ☐ Post-organ Transplantation
 ☐ Immunosuppression (not HIV/AIDS)

35. Immigration Status at First Entry to the U.S. (select one)

- ☐ Not Applicable
 ☐ Immigrant Visa
 ☐ Tourist Visa
 ☐ Asylee or Parolee
 • "U.S.-born" (or born abroad to a parent who was a U.S. citizen)
 ☐ Student Visa
 ☐ Family/Fiancé Visa
 ☐ Other Immigration Status
 • Born in 1 of the U.S. Territories, U.S. Island Areas, or U.S. Outlying Areas
 ☐ Employment Visa
 ☐ Refugee
 ☐ Unknown

36. Date Therapy Started

Month Day Year

37. Initial Drug Regimen (select one option for each drug)

	No	Yes	Unk		No	Yes	Unk		No	Yes	Unk
Isoniazid	☐	☐	☐	Ethionamide	☐	☐	☐	Moxifloxacin	☐	☐	☐
Rifampin	☐	☐	☐	Amikacin	☐	☐	☐	Cycloserine	☐	☐	☐
Pyrazinamide	☐	☐	☐	Kanamycin	☐	☐	☐	Para-Amino Salicylic Acid	☐	☐	☐
Ethambutol	☐	☐	☐	Capreomycin	☐	☐	☐	Other	☐	☐	☐
Streptomycin	☐	☐	☐	Ciprofloxacin	☐	☐	☐	Specify _____			
Rifabutin	☐	☐	☐	Levofloxacin	☐	☐	☐	Other	☐	☐	☐
Rifapentine	☐	☐	☐	Ofloxacin	☐	☐	☐	Specify _____			

Comments:

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Initial Drug Susceptibility Report

(Follow Up Report – 1)

Year Counted 	State Case Number
	City/County Case Number

Submit this report for all culture-positive cases.

38. Genotyping Accession Number

Isolate submitted for genotyping (*select one*): ☐ No ☐ Yes

If YES, genotyping accession number for episode:

39. Initial Drug Susceptibility Testing

Was drug susceptibility testing done? (*select one*) ☐ No ☐ Yes ☐ Unknown

If NO or UNKNOWN, do not complete the rest of Follow Up Report – 1

If YES, enter date FIRST specimen collected on which initial drug susceptibility testing was done:

Enter specimen type: ☐ Sputum

OR

If not Sputum, enter anatomic code (see list):

Month	Day	Year

40. Initial Drug Susceptibility Results (*select one option for each drug*)

	Resistant	Susceptible	Not Done	Unknown
Isoniazid	☐	☐	☐	☐
Rifampin	☐	☐	☐	☐
Pyrazinamide	☐	☐	☐	☐
Ethambutol	☐	☐	☐	☐
Streptomycin	☐	☐	☐	☐
Rifabutin	☐	☐	☐	☐
Rifapentine	☐	☐	☐	☐
Ethionamide	☐	☐	☐	☐
Amikacin	☐	☐	☐	☐
Kanamycin	☐	☐	☐	☐

	Resistant	Susceptible	Not Done	Unknown
Capreomycin	☐	☐	☐	☐
Ciprofloxacin	☐	☐	☐	☐
Levofloxacin	☐	☐	☐	☐
Ofloxacin	☐	☐	☐	☐
Moxifloxacin	☐	☐	☐	☐
Other Quinolones	☐	☐	☐	☐
Cycloserine	☐	☐	☐	☐
Para-Amino Salicylic Acid	☐	☐	☐	☐
Other	☐	☐	☐	☐
Specify _____				
Other	☐	☐	☐	☐
Specify _____				

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Case Completion Report

(Follow Up Report – 2)

Year Counted 	State Case Number
City/County Case Number	

Submit this report for all cases in which the patient was alive at diagnosis.

41. Sputum Culture Conversion Documented (select one) ☐ No ☐ Yes ☐ Unknown If YES, enter date specimen collected for FIRST consistently negative sputum culture: Month Day Year If NO, enter reason for not documenting sputum culture conversion (select one): ☐ No Follow-up Sputum Despite Induction ☐ Patient Refused ☐ Patient Lost to Follow-Up ☐ No Follow-up Sputum and No Induction ☐ Other Specify _____ ☐ Died ☐ Unknown	
42. Moved Did the patient move during TB therapy? (select one) ☐ No ☐ Yes If YES, moved to where (select all that apply): ☐ In state, out of jurisdiction (enter city/county) Specify _____ Specify _____ ☐ Out of state (enter state) Specify _____ Specify _____ ☐ Out of the U.S. (enter country) Specify _____ Specify _____ If moved out of the U.S., transnational referral? (select one) ☐ No ☐ Yes	
43. Date Therapy Stopped Month Day Year	44. Reason Therapy Stopped or Never Started (select one) ☐ Completed Therapy ☐ Not TB If DIED, indicate cause of death (select one): ☐ Lost ☐ Died ☐ Related to TB disease ☐ Unrelated to TB disease ☐ Uncooperative or Refused ☐ Other ☐ Related to TB therapy ☐ Unknown ☐ Adverse Treatment Event ☐ Unknown
45. Reason Therapy Extended >12 months (select all that apply) ☐ Rifampin Resistance ☐ Non-adherence ☐ Clinically Indicated – other reasons ☐ Adverse Drug Reaction ☐ Failure ☐ Other Specify _____	
46. Type of Outpatient Health Care Provider (select all that apply) ☐ Local/State Health Department (HD) ☐ IHS, Tribal HD, or Tribal Corporation ☐ Inpatient Care Only ☐ Unknown ☐ Private Outpatient ☐ Institutional/Correctional ☐ Other	

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State Case No. _____

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Case Completion Report - Continued

(Follow Up Report - 2)

47. Directly Observed Therapy (DOT) (select one)

- ☐ No, Totally Self-Administered
☐ Yes, Totally Directly Observed
☐ Yes, Both Directly Observed and Self-Administered
☐ Unknown

Number of weeks of directly observed therapy (DOT)

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48. Final Drug Susceptibility Testing

Was follow-up drug susceptibility testing done? (select one) ☐ No ☐ Yes ☐ Unknown

If NO or UNKNOWN, do not complete the rest of Follow Up Report -2

If YES, enter date FINAL specimen collected on which drug susceptibility testing was done:

Enter specimen type: ☐ Sputum

OR

If not Sputum, enter anatomic code (see list):

--	--

Month	Day	Year

49. Final Drug Susceptibility Results (select one option for each drug)

	Resistant	Susceptible	Not Done	Unknown
Isoniazid	☐	☐	☐	☐
Rifampin	☐	☐	☐	☐
Pyrazinamide	☐	☐	☐	☐
Ethambutol	☐	☐	☐	☐
Streptomycin	☐	☐	☐	☐
Rifabutin	☐	☐	☐	☐
Rifapentine	☐	☐	☐	☐
Ethionamide	☐	☐	☐	☐
Amikacin	☐	☐	☐	☐
Kanamycin	☐	☐	☐	☐

	Resistant	Susceptible	Not Done	Unknown
Capreomycin	☐	☐	☐	☐
Ciprofloxacin	☐	☐	☐	☐
Levofloxacin	☐	☐	☐	☐
Ofloxacin	☐	☐	☐	☐
Moxifloxacin	☐	☐	☐	☐
Other Quinolones	☐	☐	☐	☐
Cycloserine	☐	☐	☐	☐
Para-Amino Salicylic Acid	☐	☐	☐	☐
Other	☐	☐	☐	☐
Specify _____				
Other	☐	☐	☐	☐
Specify _____				

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