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TB Diagnostic Methods



- IGRA (Interferon Gamma Release Assay)
- AFB Smear
- Nucleic Acid Amplification
- AFB Culture
- Clinical Presentation
- TST (Tuberculin Skin Test)
- X-ray

Modified from Denise Dunbar

IGRAs





- T-spot (heparin green top tube)
- QFT tubes that come with the kit or a heparin green top (if your lab will let you.....)



- Which lab do I send it to? How do I get it there?
- How quickly do I need to have it to the lab?
- How do I store the tube until transport?

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Milogen – Positive Control Low response may indicate including to generate IPPy Mil – Negative Control Adjust for bodgerand IPPy TB1 – Primarily detects CD4 Tcell response TB2 – Optimized for detection of CD4 and CD8 T cell responses - Essentially 2 tests in one blood draw - TB1 and TB2 should be close in value

Sputum specimens



- Who should I collect sputum from
 - Patients with respiratory symptoms
 - Patients with an abnormal CXR

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Bacteriologic and histologic Examinations



Especially when lung or larynx is site of disease:

- 3 sputum specimens for AFB smear and culture
- Collected 8-24 hours apart with at least 1 early morning specimen



Bacteriologic and histologic Examinations



- Sputum collection should be directly supervised
- For patients unable to cough up sputum, deep coughing may be induced



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Bacteriologic and histologic Examinations

Extrapulmonary Specimens



- Urine
- Cerebrospinal fluid *
- Pleural fluid *
- Pus
- Biopsy specimens

≭recovery poor



Laboratory Examination



AFB Smear

- First clue
- Presumptive diagnosis only
- Fluorochrome staining preferred method
- Results available in 24 hours
- Many patients have negative AFB smears

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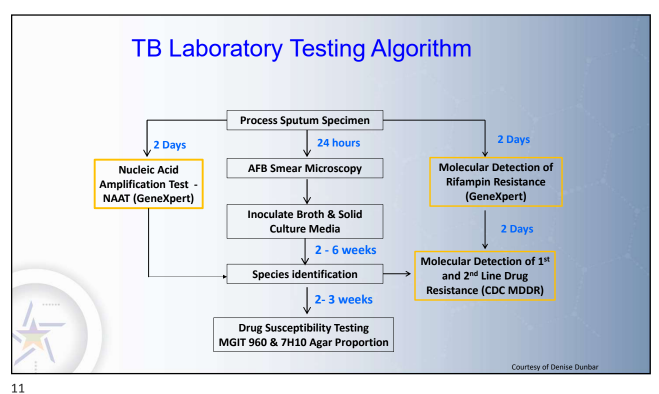
Specimen Quality



 Accurate laboratory results are directly proportional to the quality of the specimen

- Sputum
 - Recently discharged material from the bronchial tree, with minimal amounts of upper respiratory tract secretions
 - Well coached patient, collect at least 3 ml
 - Label tube, form, and indicate test:
 - Initial Dx: Smear, NAAT, & Culture
 - Follow-up: Smear and Culture
 - · Release from respiratory isolation?
 - Order Smear only
- Transport to lab cool and quickly

Courtesy of Denise Dunbar



	AFB Smear					
	САР	ATS	Interpretation	AFB/ml sputum	Infectiousness of patient	
	negative	negative	negative	<5,000	probably not infectious	
	1 or 2 per smear	1 or 2 per smear	weakly positive	~5,000	probably infectious	
	<1 per field	1+	moderately positive	~10,000	probably infectious	
one microscopic field	1.10 [:]	2+	moderately positive	~100,000	probably infectious	
	1-10 per field >10 per field	3+	strongly positive	~1,000,000	probably very infectious	
		4+	strongly positive	>1,000,000	probably very infectious	
					Courtesy of Denise Dunk	par

Nucleic Acid Amplification Tests (NAAT)



- Tiny amounts of DNA/RNA are amplified (copied) until there is enough for easy detection
- DNA/RNA is examined
 - Identification
 - **Detection of Drug Resistance**
- Test turnaround time measured in hours

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Nucleic Acid Amplification Test (NAAT)



- Detects M. tuberculosis complex nucleic acids; does not distinguish between live and dead bacilli
 - For initial diagnosis only
 - Not suitable for follow-up specimen or monitoring; cured patients may be NAAT + for years!
- Sensitivity compared to TB culture

 - >95% for AFB smear-positive
 Only 55-75% for AFB smear-negative
- · Does not replace culture for bacteriological Dx

Courtesy of Denise Dunbar

Laboratory Examination



Cultures

Used to confirm diagnosis

- Perform on ALL specimens regardless of AFB smear results
- Results available in 10 to 14 days (on liquid media, e.g. BACTEC)

TB may be diagnosed on the basis of signs and symptoms in the absence of a + culture

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AFB Culture



- More sensitive than smear
 - 5,000 to 10,000 AFB/ml for smear
 - ~10 viable AFB/ml for culture
- Positive for only ~85% of Pulmonary TB
 - Requires a quality specimen
 - May be invalid due to contamination
- Used to monitor patient response to treatment (like smear)
- · Required for drug susceptibilities & genotype
- Lengthy
 - 1-6 weeks by liquid media
 - 2-8 weeks by solid media

Courtesy of Denise Dunbar

Drug Susceptibility Testing (DST) of M. tuberculosis complex



Current Recommendations

- Initial isolate should be tested against first-line drugs (FLD)
 - Isoniazid, Rifampin, Ethambutol, Pyrazinamide
 - Repeat test if patient cult+ after 3 mo. Rx
- For isolates resistant to Rifampin or to any 2 FLDs: test second-line drug panel
 - Minimum: Fluoroquinolone, Ethionamide, & Injectable (Amikacin, Capreomycin, Kanamycin)

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Molecular Detection of Drug Resistance



- Examining DNA of specific genes for mutations known to be associated with phenotypic resistance
- Rapid analysis takes less than 1 day
- Can be done on culture isolates or directly on NAAT+ specimens

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CDC Molecular Detection of Drug Resistance (MDDR)



• Test Indications

- Known/suspect DR case or contact to DR case
- Previous TB Treatment
- Patient from area with high rate of DR TB
- Large public health impact
- Mixed or nonviable culture

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CDC Molecular Detection of Drug Resistance (MDDR)



- Provides 2-3 day DNA sequence analysis for drug resistance prediction
 - 7 classes of anti-TB drugs sequenced
- MDDR complements conventional DST
 - · Used alone, MDDR and conventional DST are imperfect
 - Used together, accuracy of drug resistance or susceptibility detection can be improved.
- Conventional DST results are still needed to confirm susceptibility to individual drugs.

Courtesy of Denise Dunbar

Summary



- Make friends with the laboratory that processes you specimens. Often
 if you can tell them what you are trying to do, they will help you get
 there
- Like most things we do, quality matters. That goes for the specimens that are sent to the laboratory
- Molecular tests are one of the biggest jumps forward in information informing patient decisions

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Questions?

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