

ECDC OPERATIONAL SUPPORT

Handbook on tuberculosis laboratory diagnostic methods in the European Union

Updated 2026



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This report was sent for consultation to the members of ERLTB-Net (see Annex 3 for list of contributors).

The first version of this ECDC technical report, previously published as 'Mastering the basics of TB control: Development of a handbook on TB diagnostic methods' (Stockholm 2011), concerned the development of the handbook. This report was then revised and renamed 'Handbook on TB laboratory diagnostic methods in the European Union' in 2016, and subsequently in 2018 'Handbook on tuberculosis laboratory diagnostic methods in the European Union – Updated 2018'. As new scientific evidence became available, a further revision was undertaken in 2022, and four chapters were updated (chapters 6, 7, 8, and 10). A new version was published in 2025, in which two chapters were updated (chapters 1 and 2) and one new chapter was added (chapter 11). This version, published in March 2026, has a revised chapter 3.

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Abbreviations

AFB	Acid-fast bacilli
AMR	antimicrobial resistance
AST	antimicrobial susceptibility testing
BCG	Bacillus Calmette-Guérin
BD	Becton and Dickinson
BSL	Biosafety level
CFU	Colony-forming units
CLSI	Clinical and Laboratory Standards Institute
CSF	Cerebral spinal fluid
CRI	Colorimetric redox indicator
CXR	Chest X-ray
DNA	Deoxyribonucleic acid
DST	Drug susceptibility testing
ECOFF	Epidemiological cut-off value
EEA	European Economic Area
EMA	European Medicines Agency
ENP	European Neighbourhood Policy
EPTB	Extrapulmonary tuberculosis
ERLN-TB	European Reference Laboratory Network for Tuberculosis (2010–2014)
ERLTB-Net	European Reference Laboratory Network for Tuberculosis (from 2014)
ERLTB-Net2	European Reference Laboratory Network for Tuberculosis (from 2018)
EQA	External quality assessment
EU	European Union
EUCAST	European Committee on Antimicrobial Susceptibility Testing
GC	Growth control
HEPA filter	High-efficiency particulate air filter
HPF	High-power field
IATA	International Air Transport Association
IFN- γ	Interferon-gamma
INSTAND e.V.	Society for promoting quality assurance in medical laboratories
IGRA	Interferon-gamma release assay
IQA	Internal quality assessment
ISO	International Organization for Standardization
LED	Light-emitting diode
LJ medium	Löwenstein-Jensen medium
LTBI	Latent TB infection
MGIT	Mycobacteria growth indicator tube
MIC	Minimum inhibitory concentration
MIRU	Mycobacterial interspersed repetitive units
MODS	Microscopically observed drug susceptibility
MTBC	<i>Mycobacterium tuberculosis</i> complex
NAAT	Nucleic-acid amplification test
NGS	Next generation sequencing
NRA	Nitrate reductase assay
NRL	National reference laboratory
NTM	Non-tuberculous mycobacteria
PBMC	Peripheral blood mononuclear cells
PCR	Polymerase chain reaction
PPD	Purified protein derivative
PPE	Personal protective equipment
PPV	Positive predictive value
pWT	Phenotypically wild-type
QC	Quality control

RFLP	Restriction fragment length polymorphism
RNI	Reactive nitrogen intermediates
RRDR	Rifampicin resistance-determining region
SDS	Sodium dodecyl sulfate
SNP	Single nucleotide polymorphism
SOP	Standard operating procedures
SS+	Sputum smear positive
SS-	Sputum smear negative
TB	Tuberculosis
TLA	Thin Layer Agar
tNGS	Targeted next generation sequencing
TST	Tuberculin skin test
UKAS	United Kingdom Accreditation Service
UK NEQAS	UKAS-accredited proficiency testing provider No. 4715
VNTR	Variable number tandem repeat
WGS	Whole genome sequencing
WHO	World Health Organization
WRD	WHO-recommended rapid diagnostics
ZN	Ziehl-Neelsen (staining method)

Background and introduction

Tuberculosis (TB) is a major cause of morbidity and mortality in Europe. High-quality laboratory diagnosis of TB is the basis for both individual patient treatment and surveillance.

In 2007, a survey of existing mycobacterial laboratory services and quality control practices throughout the European Union (EU) confirmed the key role of national reference laboratories (NRL) for TB and their services. The main conclusion of the survey was that a network of reference laboratories for TB could contribute to improving the performance of mycobacterial laboratories in Europe.

Based on these results, the European Reference Laboratory Network for Tuberculosis (ERLN-TB) was launched in January 2010 with the aim of strengthening TB diagnostics in the EU. ERLN-TB was funded and coordinated by ECDC. One or two officially nominated reference laboratories from each EU Member State joined the network, along with those in European Economic Area (EEA) and EU candidate countries.

Following the success of the ERLN-TB network, in 2014, 2018, and 2022 ECDC commissioned a renewal of the network (called the European Reference Laboratory Network for Tuberculosis, ERLTB-Net, ERLTB-Net-2, and ERLTB-Net-3 respectively) involving the same centres but anticipating the participation of institutions from the EU enlargement countries in the longer term.

The three main goals of the new network are: to support the harmonisation of laboratory methods within the EU/EEA; to develop External Quality Assurance (EQA) schemes; and to provide training activities within the network to ensure EU-wide capacity-building for TB diagnostics. One of the main activities conducted by the network was the development of a handbook of key diagnostic methods for TB, which was first published in 2011. It underwent updating in 2014 and 2018, as new scientific evidence became available, and this document represents the latest version, updated in 2022.

The aim of this handbook is to provide network members and other laboratories involved in the diagnosis of TB with an agreed list of key diagnostic methods, ranging from microbiological diagnosis of active TB to the diagnosis of TB infection. This handbook offers a single source of reference by compiling all methods, with a strong focus on standard (reference) and evidence-based methods.

The handbook will also contribute to the improvement of disease surveillance data for Europe: data sent to ECDC's EpiPulse (The European surveillance portal for infectious diseases) and other surveillance systems should be robust and backed by quality laboratory diagnostics.

This edition of the handbook addresses the changing technological landscape that has emerged over the last decade, particularly with regard to molecular-based assays and genome sequencing. Much of this technology has led to a major shift in TB diagnostic activities with the development of multiple, large, and well-equipped diagnostic centres with similar capacity and skills to national reference laboratories (e.g. rapid molecular diagnostic tests, WGS, NGS). Conversely, other molecular diagnostic developments have moved us away from 'big laboratory' approaches and closer to 'point-of-care' devices. The two approaches are not mutually exclusive, and both bring advanced diagnostics closer to the patient. The handbook is designed to meet the needs of both centralised and decentralised service delivery models and recognises that the role of national reference laboratories will change significantly over the next few years.

How this handbook relates to other work available in this field

This handbook presents a compilation of methods currently applied in EU/EEA countries. It describes common work carried out and endorsed by European laboratory experts. It also features methods and procedures developed or refined by ERLTB-Net network partners.

What this document is/is not

This document is a handbook of agreed methods in the field of TB diagnostics for laboratories serving reference functions in Europe. It provides a comprehensive compilation of key methods for the diagnosis of TB. Relevant stakeholders are encouraged to use this compilation as a basis for the validation, development, updating and dissemination of information.

The current document does not contain any formal recommendations for implementation of specific methods in EU/EEA countries. Recommendations and protocols contained within the handbook are not mandatory for EU/EEA laboratories.

Intended use and users

This handbook, which provides both basic- and reference-level methods for the diagnosis of TB, is intended for use by laboratory experts. It will also be of interest to public health professionals in the field of global TB control, particularly those involved in European initiatives to foster progress towards the elimination of TB.

History of the handbook

The first annual meeting of the ERLN-TB was held in Stockholm in 2010. During this meeting, the network partners came to a consensus agreement about the relevant topics to be included in a handbook on (reference) laboratory methods for the diagnosis of TB. The approach was to include several standardised and reliable methods, rather than to focus on one single method. A dedicated writing committee was formed to compile the first draft of these methods, using a handbook format. Each chapter includes descriptions of standardised diagnostic methods and highlights key considerations regarding operational characteristics, biosafety, and quality assurance (QA).

The first edition was published in 2011. The handbook was extensively revised during 2014–2015 following the formation of the new ERLTB-Net network and a new version titled 'Handbook on TB laboratory diagnostic methods in the European Union' was published in 2016. The handbook was further revised in 2018, with changes to Chapters 6.7 and 10.4 as new scientific evidence become available. More recently, based on the discussion made during the Network Annual meeting in January 2021, it was agreed on the need of updating the handbook to capture the expanding landscape of molecular diagnostics recommended by the World Health Organization (WHO) and new developments in this field based on the use of next generation sequencing (NGS) technology as an accurate method for *Mycobacterium tuberculosis* (MTB) drug resistance prediction (Chapter 6), as well as the use of NGS methods for MTB typing and relatedness analysis (Chapter 8). In addition, following on the revised critical concentrations of several anti-TB medicine, and the publication of a reference method for testing of minimum inhibitory concentrations (MIC) by EUCAST, an updated version of Chapter 7 on phenotypic drug susceptibility testing was deemed necessary. Lastly, in light of this dynamic scenario, Chapter 10 was updated to provide examples of best practices on how to maximise the contribution of the laboratory in the diagnostic process. See page ii for further details.

This edition

This handbook represents the fifth edition of a publication on the most reliable TB diagnostic methods, endorsed by the members of the ERLTB-Net. This publication is a compilation of methods for the laboratory diagnosis of TB, designed for laboratory experts and public health professionals. It was compiled to contribute to the harmonisation of methods in the field of TB diagnosis in the EU/EEA, EU enlargement countries and European Neighbourhood Policy (ENP) partner countries, with the goal of ensuring comparability of TB diagnoses in Europe, and provision of the best care possible for TB patients, based on a quality-assured diagnosis. This publication can also support laboratories in establishing a safe working environment for staff by minimising the risk of exposure to *M. tuberculosis*. This edition of the ERLTB-Net handbook consists of 11 chapters, each with a list of relevant references. Below is a summary of each chapter.

1 Biosafety and biosecurity in tuberculosis diagnostic laboratories

M. tuberculosis can cause laboratory-acquired infections. To ensure effective infection control, it is crucial that a comprehensive and strict biosafety policy is developed and followed. Such a policy includes standardised rules and regulations for containment, personal protective equipment (PPE), standard operating procedures (SOP) for different laboratory tasks, and a transparent structure for regulating safe working conditions in diagnostic TB laboratories.

2 Laboratory accreditation and quality management

National TB programmes are supported by laboratories that provide reliable and quality-assured results. The chapter provides a comprehensive overview of existing International Organization for Standardization (ISO) standards relevant for the laboratory diagnosis of TB and describes quality management and laboratory accreditation procedures and considerations.

3 TB infection

Two types of tests are currently used for the diagnosis of TB infection: tuberculin skin tests (TSTs) and Interferon-gamma release assays (IGRAs). The chapter describes in detail the two most commonly used IGRAs for the detection of TB infection and provides support for interpreting and reporting test results.

4 Smear microscopy

Two types of staining are most commonly used for the detection of mycobacteria: carbol-fuchsin staining (Ziehl-Neelsen, Kinyoun) and fluorochrome staining (auramine, auramine-rhodamine). The chapter describes the preparation of the required reagents and the sputum smear samples, as well as the staining procedures and the system for reporting results.

5 Culture for *Mycobacterium tuberculosis* complex

The use of cultures improves the sensitivity and specificity of TB tests, particularly at the early stages of the disease, in cases of extrapulmonary tuberculosis (EPTB) and in the event of treatment failure. The chapter provides an overview of key principles for sampling and transporting clinical specimens and processing sputum and other specimens before inoculation to solid and liquid culture media, culture incubation and examination. The issue of contamination is addressed, along with the measures necessary to prevent laboratory-acquired TB infections.

6 Molecular assays for rapid TB and drug-resistant TB detection

Molecular assays can speed up mycobacterium identification and drug susceptibility testing, and thus lead to faster and more specific treatment. This chapter provides an overview of the most commonly used molecular diagnostics for the rapid detection of TB and drug-resistant TB from culture and clinical specimens. In addition, it includes a description of the current applications of next-generation sequencing (NGS) for drug susceptibility and resistance prediction starting from clinical specimens (i.e. targeted NGS) and culture isolates (i.e. whole genome sequencing).

7 Phenotypic-based antimicrobial susceptibility testing for *Mycobacterium tuberculosis* complex

The main objectives of phenotypic-based antimicrobial susceptibility testing are to improve individual treatment management of TB cases and drug-resistance surveillance at the level of a hospital, city, region, or country. The chapter describes the methods for phenotypic antimicrobial susceptibility testing (AST) for *Mycobacterium tuberculosis* and other members of the *M. tuberculosis* complex (MTBC), as well as the EUCAST reference method for minimum inhibitory concentration (MIC) determination.

8 Molecular typing of *Mycobacterium tuberculosis* complex isolates

Various DNA fingerprinting methods are currently available that serve different purposes and have variable characteristics for specific applications. This chapter briefly describes the three traditional DNA fingerprinting methods: spoligotyping, variable number tandem repeat [VNTR] typing, IS6110 restriction fragment length polymorphism [RFLP] typing, while providing a more extensive description of the newest typing methodologies relying on the Next generation sequencing (NGS) technology.

9 Use and validation of disinfectants for *Mycobacterium tuberculosis*

In Containment Level 3 laboratories handling *M. tuberculosis*, liquid and gaseous methods are currently in use for disinfection and decontamination. The efficacy of any new disinfectants is analysed using a set of standard validation methods.

10 Information for physicians: the laboratory diagnosis of TB

The process of collecting material for the diagnosis of mycobacteria requires great care, as each step can influence the diagnosis. This chapter aims to discuss how the information generated in the laboratory should be shared with clinicians and how to maximise the contribution of the laboratory in the diagnostic process.

11 The diagnosis of nontuberculous mycobacteria

This chapter provides an overview of the principles of nontuberculous mycobacteria (NTM) diagnostics. It includes an introduction to nontuberculous mycobacteria (NTM) epidemiology and disease principles (background, when to suspect NTM, clinical presentations), and a more detailed discussion of microbiological diagnostic methods, including microscopy, culture, and molecular techniques. The content is intended for laboratories diagnosing mycobacteria infections and is based on the authors' experience, as well as the literature list at the end of the chapter.

Disclaimer

Some protocols included in the handbook list specific commercial products and assays. Such instances do not constitute endorsement of relevant products by ECDC.

1 Biosafety and biosecurity in tuberculosis diagnostic laboratories¹

Ljiljana Žmak, Kadri Klaos, Silja Mentula, Ramona Groenheit (2025)

1.1 Introduction

Mycobacterium tuberculosis complex pathogens are mostly spread by inhalation of infectious aerosol particles, and laboratory-acquired infections with these pathogens present a serious occupational hazard. These organisms are Risk Group 3 agents, reflecting their potential to cause severe disease through airborne transmission and the need for enhanced containment measures to prevent exposure. The potential risk of infection highly depends on the method performed and the concentration of bacteria present during handling. When working with infectious material, every step carries some level of risk; however, staff performing aerosol-generating procedures on positive cultures (e.g. drug susceptibility testing) are at high risk of exposure.

The fourth edition of the World Health Organization (WHO) Biosafety Manual [1] provides important guidelines on the safe handling of biological materials in laboratory settings. The purpose of biosafety procedures is to reduce the likelihood of exposure to infectious agents and to prevent laboratory-acquired infections. It is important to adhere to biosafety protocols in laboratories performing TB diagnostics, and to provide initial and periodical staff training on biosafety.

Laboratory biosecurity is closely linked to biosafety and is a crucial part of providing overall safety when working with potentially dangerous pathogens, especially high-containment biological agents like *M. tuberculosis* complex. While biosafety focuses on the safe handling of biological materials in laboratory settings, it also includes mitigating the possibility of accidental release or exposure to harmful biological agents, as well as the risk of intentional misuse of pathogens or laboratory facilities for malicious purposes. This includes the risk of unauthorised access to or theft of laboratory information. Securing and protecting laboratory facilities and infrastructure is an important element of biosecurity that should be incorporated into laboratory standard practices [2].

This manual emphasises a risk-based approach to biosafety and biosecurity, which integrates laboratory practices, safety measures and equipment, and facility design.

1.2 Principles of risk assessment for laboratories diagnosing tuberculosis

The first step in safely handling *M. tuberculosis* complex bacteria in a laboratory is to conduct a systematic risk assessment. The hazards (biological agents, procedures and other factors that pose risks) need to be identified by gathering and analysing information. The risks must be evaluated in terms of likelihood and consequence, and control measures need to be selected and implemented to mitigate the risk to an acceptable level (see chapter 1.6).

Compared with earlier editions of the WHO Biosafety Manual, the fourth edition [1] introduces a significant shift by emphasising a risk assessment framework as the core of laboratory biosafety practices. This approach underlines the importance of tailoring safety measures to specific laboratory contexts. Unlike earlier editions, which relied heavily on fixed biosafety levels and standard operating procedures, the fourth edition advocates for a dynamic, context-specific risk assessment process.

As risk can vary significantly depending on factors such as local infrastructure, the nature of the work, and staff expertise, this new approach encourages laboratories to implement control measures proportionate to their identified risks. A risk assessment should therefore consider, for example, whether a laboratory predominantly works with rapid molecular diagnostics at a clinical laboratory or regularly confronts multi-drug resistant TB strains at a reference laboratory.

The shift to a risk-based approach is critical because:

- it allows laboratories to implement measures tailored to specific activities and contexts, avoiding over- or under-protection;
- resources can be prioritised and directed towards the most significant risks;
- laboratories can adapt to new challenges by continuously evaluating and adjusting control measures, enhancing overall safety.

¹ This chapter largely consists of a summary of principles and procedures previously published by the World Health Organization [1,2]. It focuses on aspects relevant to the infrastructure of European diagnostic laboratories for tuberculosis (TB). For a more comprehensive view, please refer to the publications listed at the end of this chapter.

Regularly reassessing risks and control measures is essential to ensure that control measures remain effective as conditions change. Laboratory environments, procedures, and staff can evolve, introducing new risks or altering existing ones. Regular reassessments help to identify gaps, maintain compliance with updated standards, and enhance safety.

1.3 Risk control measures

Effective risk control in laboratory settings relies on a combination of good microbiological practices, safe facility design, appropriate personal protective equipment (PPE), and rigorous transport protocols.

1.3.1 Good microbiological practice and procedures

Laboratories must adhere to standardised procedures that minimise the potential for exposure to *M. tuberculosis* complex bacteria. This includes clear protocols for sample collection, preparation, and analysis, ensuring that no steps increase the risk of generating infectious aerosols. It is also vital that laboratory staff are well-trained in these practices, regularly updated on the latest safety protocols, and aware of the necessary precautions to prevent cross-contamination and environmental contamination. Staff should understand the specific circumstances requiring heightened precautions, including the risk levels of different procedures and diagnostic steps (e.g. differences in handling primary samples vs. enriched cultures, susceptible vs. multidrug- or extensively drug-resistant strains, or how to handle risks while pipetting, vortexing or centrifuging). This code of practice plays a key role in maintaining a safe working environment.

1.3.2 Facility design, equipment and maintenance

Incorporating safety in planning a TB laboratory facility requires a careful, multidisciplinary approach that considers biosafety principles together with workflow functionality and diagnostic efficiency. The first step involves conducting a risk assessment to determine the required level of containment (e.g. Biosafety Level 2 or Biosafety Level 3), the anticipated volume of testing, and the types of diagnostics to be performed.

The TB laboratory should ideally be located in a dedicated part of the building with a clearly visible biohazard sign displayed at the laboratory entrance. Appropriate physical containment systems are a crucial step in providing staff safety. These include an anteroom with interlocked doors and separate zones for clean and dirty clothing, appropriate biosafety cabinets situated away from walking areas, and specialised ventilation systems.

The anteroom reduces the risk of airborne contaminants escaping the containment zone and provides a controlled space for donning (putting on) and doffing (taking off) PPE, all of which should be described in protocols for working in the particular laboratory. The anteroom should be maintained under positive pressure relative to the containment area to prevent backflow of potentially contaminated air. High containment areas for manipulating *M. tuberculosis* complex cultures should be equipped with specialised ventilation systems that can maintain negative pressure in the room, with 6–12 air changes per hour. There should be an alarm system to alert staff if there is a ventilation failure.

The selection and positioning of laboratory equipment should be compatible with biosafety containment practices. Namely, centrifuges should be equipped with sealed safety cups or rotors, while all aerosol-generating procedures, especially pipetting and mechanical homogenisation, should be performed within biosafety cabinets to minimise exposure risk to laboratory staff. Reliable utilities, such as electricity, water supply, backup power systems, and internet connectivity for data management, are also critical.

Laboratory walls and floors should be easy to clean, impermeable and resistant to disinfectants. Special attention should be paid to the selection of disinfectants with mycobactericidal activity and disinfection protocols – which are vital for infection control – must be strictly adhered to.

Wastewater from TB laboratories should be collected in a dedicated effluent decontamination system. Effective waste management procedures, including proper sterilisation and disposal of infectious materials, need to be in place. Proper maintenance of the aforementioned systems – including routine checks of equipment, ventilation and air filtration – is essential for their continued effectiveness.

1.3.3 Personal protective equipment for heightened control measures

PPE that can mitigate the risk of TB exposure includes laboratory coats, gloves, face shields, goggles, and – most importantly – fit-tested respiratory protection such as N95 respirators or higher-level masks. In high-risk settings, powered air-purifying respirators may be necessary.

Individual laboratory protocols should describe the PPE necessary for working in that particular setting. These requirements may vary slightly between laboratories, depending on institutional guidelines, the nature of the work

being conducted and the outcome of local risk assessments. However, the core principles of minimising contamination and ensuring personal safety remain essential.

1.3.4 Transport of biological materials

Materials should be packaged securely according to WHO guidelines [1] and United Nations transportation recommendations [3]. Packaging must include primary containers that prevent leaks, secondary containers that protect from breakage, and outer packaging that ensures safety during handling. The shipment must be properly labeled to indicate the presence of infectious agents, and clear instructions for emergency procedures must be included.

1.4 Emergency preparedness

In case of accidental release or exposure to biohazardous material, laboratory workers need to be trained in emergency preparedness and response plans and have periodic refresher trainings. Control measures should also be in place for other emergencies caused by, for example, chemicals, fire, power blackouts, floods or personal health issues. The health status of laboratory staff who work with heightened control measures should be adequately checked by occupational healthcare, and confidential records kept.

The laboratory should have a system for reporting incidents and emergencies, both within the institution and to relevant external bodies, including communication protocols for alerting appropriate authorities (e.g. health agencies). The laboratory should also have a system for securing the area to mitigate potential risks to public health.

Emergency procedures should cover how to handle accidents such as spills, aerosol releases and risk control measure failures, as well as any accidents involving TB samples or staff exposures to pathogens (e.g. inhalation of TB aerosol or skin contact with infected material). The procedures should include information about the availability of spill kits, suitable disinfectants, the appropriate PPE for handling accidents, and how to ensure proper decontamination of affected areas and equipment. In addition to describing how to contain the release, procedures should also cover recovery measures and any necessary post-incident analysis, including root-cause analysis, corrective action and process improvement.

First aid equipment and protocols for providing immediate care and medical evaluation in the event of accidental exposure should be readily accessible. In high-risk situations, such as after exposure to TB cultures, staff may be offered TB prophylaxis or preventive treatment after evaluation by a physician.

1.5 Risk assessment

The fourth edition of the WHO Biosafety Manual [1], including the associated monograph on risk assessment [4], provides a framework for risk assessments. The process includes five steps:

1. Gather information;
2. Evaluate the risks;
3. Develop a risk control strategy;
4. Select and implement risk control measures; and
5. Review risks and risk control measures (Figure 1).

Figure 1. Risk assessment framework, according to the World Health Organization’s Biosafety Manual (fourth edition [1]) and associated monograph [4]



Source: Reproduced from Laboratory biosafety manual, 4th edition, World Health Organization, Section 2: Risk assessment, page 6, Copyright (2022).

A comprehensive risk assessment requires input from laboratory staff who understand the processes and procedures within the scope of the work being assessed. The members of the risk assessment team should have demonstrated skill in working with *M. tuberculosis* complex and understand all the hazards associated with the protocols and procedures to be carried out in the laboratory. The team members must be familiar with the layout and condition of the laboratory facility, as well as the equipment to be used in the procedures.

The risk assessment requires consideration of many factors – for instance, routes of transmission, infectious dose, availability of prophylactic treatment, disease severity, contagiousness, high-risk laboratory procedures, competency of laboratory staff, susceptibility of individual staff and biosecurity.

Below is a short example of what a risk assessment for drug susceptibility testing (DST) of *M. tuberculosis* complex isolates could include. By systematically identifying hazards, evaluating risks, and implementing targeted control measures, laboratories can maintain a safe environment while conducting TB DST. The monograph on risk assessment [4] provides other examples.

Example risk assessment for drug susceptibility testing of *M. tuberculosis* complex isolates

1. Identify the hazards

Tuberculosis (TB) drug susceptibility testing involves handling *M. tuberculosis* complex pathogens, which are classified as Risk Group 3 agents due to their potential to cause serious disease through airborne transmission. Procedures such as specimen processing, culture, and manipulation can generate aerosols, increasing the risk of exposure to inhalation.

2. Evaluate the risks

Assess the likelihood and potential impact of exposure during each step of the DST process:

- **Specimen reception and processing:** Handling sputum samples may release infectious droplets.
- **Culture procedures:** Culturing *M. tuberculosis* complex can amplify the pathogen load, heightening exposure risk.
- **DST methods:** Manipulations during DST, such as setting up assays or interpreting results, can produce aerosols. Factors also influencing the risk include the volume of specimens, proficiency of staff, and existing safety protocols.

3. Implement control measures

Based on the risk assessment, apply the following controls:

- **Engineering controls:**
 - Use certified biosafety cabinets for all aerosol-generating procedures to contain airborne pathogens.
 - Ensure proper ventilation systems with HEPA filtration to maintain directional airflow and prevent contamination.
- **Administrative controls:**
 - Develop and enforce standard operating procedures that outline safe handling practices.
 - Provide comprehensive biosafety training for laboratory staff, emphasising the importance of adherence to protocols.
 - Implement a robust waste management system for the decontamination and disposal of infectious materials.
- **Personal protective equipment:**
 - Require laboratory staff to wear appropriate PPE when handling *M. tuberculosis* complex cultures.
- **Review and monitor:**
 - Continuously monitor the effectiveness of implemented control measures.
 - Conduct regular audits and inspections to ensure compliance with biosafety protocols.
 - Establish a system for reporting and investigating incidents or near-misses to identify areas for improvement.
 - Update risk assessments periodically or when changes occur in procedures, equipment, or regulations.

1.6 Laboratory biosecurity

Laboratory biosecurity is closely linked to biosafety. It is a crucial part of providing overall safety when working with potentially dangerous pathogens, especially high-containment biological agents like *M. tuberculosis* complex.

While *biosafety* is more focused on the safe handling of biological materials in laboratory settings to mitigate the possibility of an accidental release or exposure of harmful biological agents, *biosecurity* addresses the risk of intentional usage of pathogens or facilities for malicious purposes. This includes the deliberate misuse of biological agents or toxins, either for bioterrorism or bio crime, as well as the risk of unauthorised access to or theft of laboratory information which could lead to harmful consequences.

Strong biosecurity is therefore largely about securing the facility and preventing unauthorised use of laboratory facilities. Securing a BSL-3 laboratory involves ensuring that only authorised personnel can access sensitive areas, and includes implementing protocols for entry and exit, continuous monitoring, and maintaining detailed logs of all activities within the facility. National and/or international biosecurity regulations and guidelines should be implemented and strictly followed in all TB laboratories [2].

This section of the handbook focuses on the key elements needed for an efficient biosecurity management programme in diagnostic TB laboratories. The biosecurity challenge posed by scientific laboratories conducting research on *M. tuberculosis* complex strains that may fall within the dual-use research of concern area is beyond the scope of this section.

1.6.1 Institutional biosecurity policy

All laboratories that perform TB diagnostics, especially culturing and/or handling isolates, should take into consideration the potential of deliberate misuse of Risk Group 3 agents or unwanted access to laboratory information. Although all staff should be aware of the potential risks associated with handling the bacteria, one of the most prominent roles in achieving a safe environment is a nominated biosafety/biosecurity officer and/or committee. The biosafety/biosecurity officer is responsible for overseeing compliance with safety regulations, conducting risk assessments, and ensuring that all safety measures are in place. The person and/or committee also facilitates training programmes for staff to raise awareness about the risks associated with *M. tuberculosis* complex and how to manage them safely.

Institutions performing TB diagnostics should adopt a security-conscious culture and proactively minimise biosecurity risks. The institutional biosecurity policy includes all measures taken to mitigate the risk of malicious situations. It encompasses several key components for safeguarding laboratory operations, including staff security and access control, secure storage and handling, and data management. All these elements must be rigorously screened through a comprehensive biosecurity risk assessment.

1.6.2 Biosecurity risk assessment and control measures

The biosecurity risk assessment considers all the potential situations for microbial isolates and/or laboratory information and/or laboratory facilities to be misused, stolen, or exploited for malicious purposes. Regular audits and inspections are needed to ensure compliance with biosecurity standards and to identify any potential vulnerabilities that could be exploited. TB laboratories should evaluate access control, staff screening, as well as inventory control and data management to reduce the likelihood of diversions or theft.

1.6.3 Access control

One of the most critical components of laboratory biosecurity is ensuring that only authorised staff have access to sensitive materials. Zones with TB laboratories should have restricted access provided by access control systems.

The access can be controlled simply by key-locked doors or by electronic access control systems enabled by keycards or biometric scanners. Key-locked doors are a good option to provide basic access control in settings where resources are limited. However, electronic access control systems can also provide detailed information about access – such as who has entered and when. Intrusion detection systems (alarms, surveillance cameras) and security guards can also reduce the probability of unauthorised or criminal activity.

1.6.4 Staff security

Staff security includes doing background checks and screening of laboratory staff and visitors, as well as clearance procedures for staff working with high-containment biological agents. National or institutional regulations may call for such screening protocols for TB laboratories. If such procedures are not part of locally adopted regulations, whether or not staff security is needed will depend on the level of risk for that individual laboratory, as evaluated through a comprehensive risk assessment.

Regardless of variation in staff security protocols, it is important for all staff to have regular security training to ensure that they understand biosecurity risks and the protocols in place to mitigate them.

1.6.5 Inventory control and data management

Inventory control practices are essential to maintain the integrity of biological materials and prevent diversion or theft. All biological material should be clearly marked to avoid confusion or misuse and placed in locked and monitored secure areas, preferably in locked storage. An accurate tracking system of all biological materials should be in place that includes information such as quantities, storage locations, records of those who accessed the materials and movements. Regular inventory checks should be performed to ensure that all materials are accounted for and that no unauthorised transfers have occurred. Accurate record-keeping not only facilitates biosecurity but also ensures compliance with regulations and enhances transparency and accountability in laboratory operations.

The laboratory should also maintain secure systems for transporting biological materials within the institution, ensuring they are properly labeled, documented, and tracked.

As laboratory data are stored on computers/servers, an important part of ensuring laboratory biosecurity is evaluating vulnerabilities in the laboratory's data management system, including the security of computer networks, software systems, storage solutions, and data transfer methods. It is crucial to identify potential threats to laboratory data, including unauthorised access, cyberattacks, data corruption, or accidental loss. These threats can lead to breaches in security, loss of data integrity, violations of the General Data Protection Regulation and the dissemination of sensitive information, which could compromise public health and safety.

1.6.6 Emergency response protocols

Laboratories must have emergency response protocols in place for biosecurity breaches (unauthorised access, theft, or loss of biological materials, or cyberattack). These protocols should ensure swift containment of the incident, protect laboratory staff and the surrounding community, and guide the appropriate decontamination and reporting procedures.

In the event of a laboratory theft or break-in, immediate actions include alerting authorities, securing the scene, and evacuating personnel if necessary. Key personnel should be notified, and a lockdown of sensitive areas can be initiated. An inventory check should be conducted to account for all materials, and evidence should be preserved for forensic investigation. A biosecurity risk assessment should be performed to evaluate potential exposure and implement containment measures if needed.

Communication in the event of a biosecurity breach is crucial. Inform all staff and report to national and international health authorities as required. Cooperate with law enforcement, review and upgrade security protocols, and provide support to affected staff. Finally, document the incident thoroughly and follow-up with training to reinforce biosecurity awareness and emergency response procedures.

Having well-established emergency plans in place for biosecurity breaches reduces confusion, facilitates a coordinated response, and ensures compliance with regulatory requirements, ultimately safeguarding public health. Regular drills and training exercises should be conducted to ensure that all staff are familiar with emergency procedures and can act quickly in the event of a biosecurity incident.

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2 Laboratory Accreditation and Quality Management

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2.1 Introduction: the purpose of laboratory accreditation

National TB programmes depend on laboratories that deliver reliable and high-quality results, forming the backbone of effective tuberculosis control. Although quality assurance (QA) systems have long been integral to laboratory practice, the growing complexity of diagnostics and the need for international comparability have increasingly made formal accreditation essential. Accreditation is not simply a bureaucratic exercise, but a strategic investment that drives continual improvement, and ensures that laboratory results are reproducible and defensible.

A key accreditation outcome is the assurance of patient safety through consistent and reliable diagnostic results. Accredited laboratories operate under rigorously assessed standards, meaning that every test result can be trusted to guide clinical decisions and public health actions. This reliability is crucial for National Reference Laboratories (NRLs), whose data inform national policies, guide patient management, and underpin surveillance systems that shape global TB strategies. Accreditation also ensures quality and reliability, facilitating the comparability of results across laboratories and borders.

Moreover, accreditation increases confidence among all stakeholders, including clinicians, public health authorities, and patients. It promotes a culture of transparency and accountability, requiring laboratories to document procedures, monitor performance, and systematically address errors. This approach improves day-to-day operations, promotes continuous quality improvement, and enhances resource efficiency. In addition, accredited laboratories are better equipped to participate in international collaborations and research, where harmonised data are needed to monitor trends, detect outbreaks, and evaluate interventions, as their practices meet globally recognised quality standards.

Ultimately, accreditation is a strategic investment that contributes to the effectiveness, safety, and sustainability of TB diagnostic services within national and international health systems.

2.2 Accreditation standards: ISO 15189

The accreditation of tuberculosis and other medical laboratories benefits patients by ensuring that diagnostic results are trustworthy, impartial, consistent, and suitable for clinical decision-making. Laboratory accreditation is based on an International Organization for Standardization (ISO) standard (ISO 15189:2022), which outlines the specific quality and competence requirements for medical laboratories. This standard, customised for the clinical context and patient care, builds on ISO 17025, which pertains to testing and calibration laboratories and is generally not applied in medical laboratories.

To achieve and maintain accreditation, laboratories must comply with the rigorous requirements of ISO 15189:2022. These requirements include establishing a robust, risk-based quality management system, validating and verifying assays, and strictly controlling biosafety, biosecurity, and contamination risks. Laboratories must also demonstrate staff competence, ensure regular training, and maintain calibrated equipment and reliable laboratory information systems. Continuous quality monitoring, internal audits, and transparent handling of non-conformities are also necessary to ensure that any issues identified result in lasting improvements. Although the requirements are demanding, they help transform informal practices into reliable, standardised processes that protect patients and promote long-term operational excellence.

2.3 Key components of a TB laboratory quality management system

Component	Description
Organisational structure	Roles and responsibilities are clearly defined from top management to technical staff. The laboratory structure is well described. The laboratory director and quality manager are designated.
Document control	A document control system is implemented. Controlled documents include quality manual, Standard Operating Procedures (SOPs), records, and external documents (e.g. regulations, standards, manufacturer's instructions). SOPs are approved, version-controlled, accessible and readily available to authorised personnel, reviewed periodically and updated if necessary. Records are legible, traceable and securely stored. Retention periods are defined for documents and records following applicable legal or accreditation requirements. Obsolete document control is implemented to prevent unintended use.
Personnel competence	The laboratory has defined the required education, training, experience and skills for each role. Job descriptions are available, defining responsibilities and authorities. Laboratory staff have appropriate qualifications, training and experience. Initial and ongoing competency assessments are conducted. Training records are maintained and staff have access to SOPs. Ongoing training and continuous education are provided to maintain and improve competence.
Equipment management	The laboratory has assigned unique identifiers to all equipment and keeps an inventory registry. It ensures that equipment is properly installed, calibrated and maintained, defines preventive maintenance and calibration schedules, and keeps equipment records. Isothermal equipment critical to the analysis is monitored with calibrated probes to ensure accurate and reliable temperature control.
Inventory control	The laboratory has implemented an inventory control system. A list of approved suppliers is maintained. Inventory is monitored to prevent stock-outs and expiry. The quality of supplies is verified. Storage conditions meet requirements.
Process control	Laboratory procedures are defined and validated for each workflow step, covering pre-examination, examination, and post-examination processes. Test methods are validated or verified before use. Internal Quality Control (IQC) and External Quality Assessment (EQA) maintain quality, while Corrective and Preventive Actions (CAPA) address non-conformities. Quality indicators, such as turnaround times, error rates, and client satisfaction, are monitored to assess performance and identify trends for appropriate action. Laboratories systematically track key quality metrics, analyse trends and address issues.
Quality control	Internal (IQC) and external (EQA) quality controls are scheduled and conducted to ensure the reliability, accuracy and quality of examination results. IQC is routinely used to monitor analytical performance. EQA programmes ensure inter-laboratory comparability and support ongoing performance evaluation.
Data management	The laboratory ensures data integrity, confidentiality, availability and traceability throughout the laboratory workflow, including manual and electronic systems, such as Laboratory Information Systems (LIS), used to capture, process, store, transmit and retrieve patient and laboratory information. Proper data management and reporting systems must be established and maintained to ensure accurate and timely communication of test results.
Non-conformance management	The laboratory identifies and documents deviations from established procedures, requirements, or expected outcomes through internal or external audits, external assessments, quality control failures, incidents, complaints and staff observations. The root cause is investigated, the impact is evaluated, and corrective actions are taken. This includes monitoring implemented actions, evaluating the effectiveness of those actions, and taking preventive actions to eliminate the cause of a potential non-conformity or incident to prevent the occurrence/recurrence of similar issues.
Internal audit	The laboratory conducts systematic, independent, scheduled and documented self-assessments to evaluate the effectiveness of the Quality Management System (QMS), ensure compliance with standard requirements, and drive continual improvement. The selection of auditors should ensure objectivity and impartiality of the process (if possible, auditors shall be independent of the activity to be audited). Audit findings are documented and addressed.
Management review	The laboratory must conduct a structured, periodic evaluation of the entire QMS to ensure effectiveness, compliance, alignment with the laboratory's strategic direction, and continual improvement.

Component	Description
Continual improvement	The laboratory actively seeks to improve processes, services, and management systems to improve patient care and laboratory outcomes and to comply with evolving regulations. The laboratory should carry out regular reviews and updates of policies, procedures, and training programmes to ensure it stays up-to-date with advancements in microbiology and technology. Opportunities for improvement (improvement goals) are identified using quality indicators, audit results, non-conformities, and management reviews.
Biosafety and Biosecurity	The laboratory adheres to rigorous biosafety protocols at every stage of the process (transportation, packaging, safe and proper handling, disposal of infectious materials, waste management, etc.). Policies and procedures are implemented to protect laboratory operations. Areas handling biohazards require controls to prevent cross-contamination and unauthorised access (e.g. BSL-3). Personal protective equipment is available and properly used, and equipment for handling biological materials (e.g. biosafety cabinets) must be appropriately maintained and calibrated. Environmental conditions are monitored and controlled. Training on infection control and proper handling and storage of TB samples is provided. Risk assessments must include biological hazards and biosecurity risks, and include actions to mitigate them. The health of the staff is monitored. Established incident response and reporting mechanisms are in place [3].

2.4 Step-by-step accreditation guide

Stepwise implementation of ISO 15189 in TB laboratories

Step	Action	Tools/Resources
1	Conduct baseline assessment.	Global Laboratory Initiative (GLI) accreditation checklist [4].
2	Train staff on QMS principles.	SLMTA toolkit [5], World Health Organization (WHO) e-learning modules [6].
3	Develop QMS documentation.	ISO 15189 guidance, GLI templates [4].
4	Implement internal audits.	WHO audit checklist [6], mentor review.
5	Submit application to accrediting body.	National ISO-accredited agencies.
7	Undergo final accreditation audit.	Accrediting agency audit process.
8	Maintain accreditation status.	Regular audits, updates, training, internal audits.

2.5 Quality assessment

2.5.1 Internal quality assessment

Internal quality assessment is a systematic process involving the internal monitoring and evaluation of laboratory activities to ensure that procedures, personnel and equipment performance comply with predefined quality standards. It ensures the effectiveness, consistency, and reliability of laboratory processes, results, and overall performance. In addition to internal quality controls (IQC), internal quality assessment includes internal audits, method verification and validation (during implementation of new methods), monitoring internal quality indicators (to analyse trends, conduct root cause analysis, and implement corrective and preventive actions when the assessment reveals non-conformities or deviations), staff competency assessment, process reviews (including documentation), and continuous improvement evaluation.

Internal Quality Control (IQC) refers to the procedures and materials used in the laboratory to continuously monitor and evaluate the performance of examination systems in the detection of errors or variations in the analytical process before results are released, with the aim of ensuring the quality of examination results. Control materials should be well characterised, with known stable values, and must resemble patient samples. IQC materials must provide a clinically relevant challenge to the analytical method, with concentration levels close to the clinical decision limits. If IQC material is not available, alternative approaches may be used, for example, periodic reanalysis of previously tested, stored patient samples. The frequency of IQC must be defined based on the stability, complexity, and robustness of the analytical method and the potential risk of harm to the patient resulting from the reporting of an incorrect result. It is critical to record and analyse the resulting data, monitor internal quality control indicators to detect trends or shifts, and review the data against predefined acceptance criteria to prevent the release of results if these criteria are not met. If the requirements are not met, the results must be rejected, and the affected patient samples must be reanalysed after the error has been corrected. Patient results generated after the last acceptable IQC outcome must be evaluated to determine their validity. It is essential to establish IQC failure protocols and maintain records of all controls, deviations and actions taken.

2.5.2 External quality assessment

EQA is an essential element of quality assurance in accredited TB laboratories. It provides independent verification of testing accuracy and enables comparison with peer laboratories, supporting continuous improvement and diagnostic reliability. EQA is particularly important in TB diagnostics, where accurate detection and drug resistance testing are critical for patient management and public health.

According to ISO 15189, laboratories must participate in EQA for each major test area at defined intervals, typically at least twice per year. TB EQA programmes commonly include smear microscopy, culture, drug susceptibility testing (DST), and molecular methods. EQA participation helps identify technical issues, assess staff competency, and ensure consistent performance over time.

Laboratories must document their participation, analyse results, and implement corrective actions when needed. Performance trends should be reviewed during regular management meetings to guide quality improvement.

A list of internationally recognised EQA providers for TB diagnostics is available through the Global Laboratory Initiative (GLI) EQA dashboard [7]: <https://www.stoptb.org/who-we-are/stop-tb-working-groups/global-laboratory-initiative-gli/gli-eqa-dashboard>.

2.6 Implementation of laboratory methods

The laboratory should select and implement examination procedures which have been validated for their intended use to ensure clinical accuracy of the analysis and the quality of examination results. The specified requirements (performance specifications) for each examination procedure should relate to the intended use of that examination and must be selected based on the intended clinical purpose of the test. Authorised personnel must periodically evaluate the analytical methods provided by the laboratory to ensure that they are clinically appropriate for the examination request received. Therefore, validation and verification are critical to ensure that laboratory methods are suitable for specific purposes.

The table below provides a comparative analysis of both concepts, with the purpose of determining the appropriate application of each, based on the type of method selected by the laboratory (Table 1).

Table 1. Comparison of method validation and verification in TB laboratories

	Verification	Validation
Definition	Process of confirming that a standard or validated method performs as expected for its intended use in the laboratory.	Process of confirming/proving that a new non-standardised or modified method is fit for purpose and performs as intended, based on predefined criteria.
Application	For standardised or validated methods already approved: <ul style="list-style-type: none"> Commercial CE-IVD* marked kits; Published peer-reviewed protocols; Standard methods (e.g. EUCAST, WHO); if used within the intended scope.	For non-standard, modified (commercial kits or procedures), and laboratory-developed methods (laboratory-developed tests, in-house methods), or standard methods outside the original intended scope (intended parameters: different sample types, use of third-party reagents, etc.).
Purpose	Confirm that the method performs adequately in the local context and in the specific laboratory's environment (using the specific laboratory's personnel, equipment and settings).	Generate evidence to prove the method works for the intended purpose. Demonstrate method is suitable for clinical use.
Goal	To ensure suitability and compliance with expected performance.	To prove effectiveness and reliability of a method before implementation.
Scope	Limited. Ensures that the method works in your laboratory under your conditions. Less extensive study (comparisons, performance checks).	Broader. Demonstrates method suitability, especially when no prior approval exists. More extensive study (assessment of the required validation parameters)
Parameters	Precision, accuracy, detection limits, sensitivity or specificity, any performance claims made by the manufacturer, etc.	Performance characteristics must be selected based on the intended clinical purpose of the test. Precision: intra-assay precision (within-run precision) evaluates repeatability; inter-assay precision (between-run precision) evaluates reproducibility; accuracy; analytical and diagnostic sensitivity and specificity; linearity; robustness; measurement uncertainty; interference and cross-reactivity (if relevant).
Documentation	Verification plan, results of performance characteristics, acceptance criteria and statement confirming that the method is fit for use in the laboratory.	Validation protocol (pre-defined), raw data and analysis, full study report, performance summary with acceptance criteria and outcomes, and final statement declaring that the method is validated, suitable for the intended use and fit for clinical purpose.

Examples	<ul style="list-style-type: none"> • GeneXpert MTB/RIF assay: confirm that the assay performs as expected with local staff, samples and equipment. Test known positive and negative controls; assess detection limit and reproducibility. • MGIT 960 System: confirm the system detects <i>M. tuberculosis</i> as per manufacturer's claims using known control strains. Evaluate time detection and contamination rate. 	<ul style="list-style-type: none"> • Validating a new lab-developed real time PCR for TB genotyping. Validate accuracy, specificity, analytical sensitivity, cross-reactivity and linearity. • Validating a second-line DST using a new or locally-adapted protocol: Validate Minimum Inhibitory Concentration (MIC) reproducibility, and compare with reference method or genotypic result.
ISO 15189:2022 Clause	7.3.2 Verification	7.3.3 Validation

**CE-IVD stands for Conformité Européenne-In Vitro Diagnostic, which indicates that an IVD medical device complies with the applicable European legislation (IVDR-Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices. Before May 2022, devices were CE-marked under IVD Directive 98/79/EC (IVDD)) and may be marketed in the European Economic Area (EEA). A CE-IVD mark is a symbol that manufacturers place on in vitro diagnostic devices to show that they conform to the IVDR, have passed the required conformity assessment and can be freely marketed in the EU/EEA. Laboratories in the EU must use the CE-IVD test unless they meet specific exemptions (e.g. validated in-house tests under Article 5(5)). CE-IVD ensures that the test has been independently reviewed, validated, and complies with the highest safety and quality standards. It is crucial to distinguish between commercially available CE-IVD-marked kits and those labelled for Research Use Only (RUO), as RUO are not approved for clinical use and cannot be used for patient diagnosis, unless they are fully validated by the laboratory and the manufacturer makes no claims about diagnostic accuracy or safety. RUO kits are only intended to be used for research and not for clinical diagnostics.*

2.7 Additional considerations

2.7.1 Human resources

ISO 15189:2022 requires laboratories to ensure that staff are qualified, trained, and competent for their assigned duties. Sustaining quality depends on staff retention, continuous training, and motivation. Competency assessments should therefore be conducted regularly and may include direct observation, proficiency testing, or review of results accuracy. For example, a technician performing smear microscopy may be assessed by blind rechecking of slides. Clear job descriptions, ongoing education, and recognition of staff contributions support a culture of quality. A stable and motivated workforce is essential for maintaining consistent and reliable diagnostic services.

2.7.2 Digital tools

Digital tools are essential for strengthening the quality management system (QMS) in TB laboratories. A QMS improves traceability, standardises documentation, and enhances data quality across core functions, such as equipment maintenance, inventory management, incident reporting, and staff training. This type of tool supports compliance with ISO 15189 by enabling real-time tracking of processes, automated reminders for quality tasks, and centralised storage of records. Digital solutions also facilitate internal audits, monitor corrective actions, and ensure the timely review of quality indicators. By integrating such systems, laboratories can improve efficiency, transparency, and consistency, supporting continuous improvement and long-term sustainability of diagnostic quality.

2.7.2 Accreditation of Whole Genome Sequencing (WGS)

WGS is increasingly used for both genotypic drug susceptibility testing (gDST) and molecular epidemiological surveillance of *Mycobacterium tuberculosis* (MTB). Laboratories seeking ISO 15189 accreditation for WGS-based methods must ensure that the entire workflow, from culture and DNA extraction to sequencing, bioinformatics, and reporting, is validated and assessed as part of the quality management system.

Accreditation typically covers two key analyses derived from WGS data:

- SNP-based cluster analysis for transmission surveillance
- Genotypic drug susceptibility testing (gDST).

In both cases, accreditation is based on the identification and interpretation of sequence variants: single nucleotide polymorphisms (SNPs) for cluster analysis and mutations catalogued for drug resistance prediction. Before accreditation, laboratories must perform thorough validation studies. For SNP-based cluster analysis, this includes retrospective analysis of isolates previously typed (e.g. MIRU-VNTR-defined clusters) and epidemiologically linked cases. For gDST, comparison with phenotypic DST (pDST) is required to assess concordance. A structured validation plan, including the scope, performance targets, and acceptance criteria, is strongly recommended to guide the process. The mutation catalogue used for gDST must be version-controlled and traceable, particularly when internal adaptations of public resources are applied. Similarly, all software tools used in the bioinformatics

pipeline should also be versioned and validated. Quality control parameters must be clearly defined, including thresholds that trigger resequencing. Participation in EQA schemes is essential. For example, MTB-specific WGS EQAs coordinated through international networks provide benchmarking opportunities and support accreditation. Routine trend monitoring should be implemented. This may involve sequencing a standard panel of control isolates (e.g. including H37Rv) at regular intervals (e.g. quarterly) to detect potential performance drift.

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3 The diagnosis of TB infection

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Summary

This chapter provides an overview of the diagnosis of Tuberculosis infection (TBI). It outlines the key concepts of infection with *Mycobacterium tuberculosis* and summarises current diagnostic methods. Particular emphasis is placed on the most widely used tests in Europe. The content is intended to serve as inspiration for laboratories involved in TBI diagnosis and is based on the authors' experiences as well as the literature listed at the end of the chapter. The chapter does not necessarily cover all aspects or methods related to the diagnosis of TBI.

3.1 Background and principles

TBI can occur following exposure to *Mycobacterium tuberculosis*, but most people who are infected do not develop TB disease. This condition has previously been referred to as latent TB infection (LTBI), but to better reflect the complexity and dynamic nature of TB pathogenesis, the broader term TBI is now preferred [1]. This terminology recognises TBI as part of a continuum that may progress through asymptomatic TB disease (also referred to as subclinical TB), before culminating in TB disease [2]. TBI is defined as a state of persistent immune response to *M. tuberculosis* antigens in the absence of symptoms, without radiological or microbiological evidence of TB disease.

It is estimated that around one quarter of the global population has been infected with *M. tuberculosis*, and while some people clear the infection the bacteria can persist for life in others. Primary infection leads to active disease in 5 to 10% of infected people, mostly within two years of infection [1]. The risk of progression to TB disease increases if the host's immune response is weakened, for example by HIV infection, malnutrition, use of corticosteroids or other immunosuppressive agents, or advanced age.

Identifying and treating TBI are key components of global TB control and form a major part of the WHO End TB Strategy [3]. Preventive therapy for TBI significantly reduces the risk of progression to TB disease, particularly when delivered through shorter, better-tolerated regimens that improve treatment completion rates.

Ideally, diagnostic tools for TBI should demonstrate:

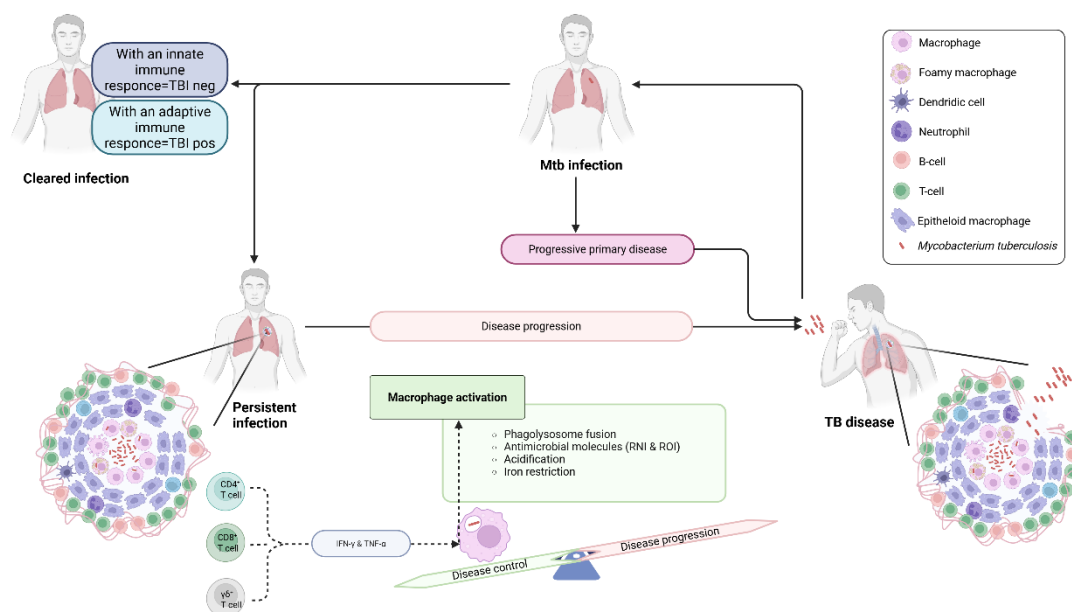
- high sensitivity in all at-risk populations;
- high specificity, regardless of BCG vaccination and exposure to environmental mycobacteria;
- reliability and stability over time;
- objective criteria for positive result, affordability, and easy administration; and
- ability to identify recently infected individuals at increased risk of progression to TB disease.

The diagnosis of TBI currently relies on two main groups of immunologic tests: skin tests and interferon- γ (IFN- γ) release assays (IGRAs). Newer IGRA platforms have expanded to include biomarkers such as IP-10 (IFN- γ inducible protein 10), and novel *M. tuberculosis* antigen-based skin tests (TBSTs) have emerged as alternatives to the traditional tuberculin skin test (TST) [4].

3.1.1 Immune response to *M. tuberculosis*

The immune response to *M. tuberculosis* is multifaceted. Immunological mechanisms involved in maintaining TBI are complex but important for clearing the infection and/or preventing progression to TB disease [5]. When the human host is infected by *M. tuberculosis*, there are three potential outcomes (Figure 2):

- cleared infection with spontaneous healing [6,7];
- persistent infection where mycobacteria are contained in a granuloma that consists of macrophages, which can differentiate into epithelioid macrophages or multinucleate giant cells, CD4+ and CD8+ T-cells, and B-cells. The T cells produce IFN- γ , which activates macrophages. CD8+ T-cells can lyse infected macrophages or kill intracellular bacteria with cytotoxic molecules such as perforin, granzymes, and granulysin [8,9]. Tumour necrosis factor (TNF) is produced by macrophages and T-cells. Dendritic cells are also present, and a mature granuloma is surrounded by fibroblasts. *M. tuberculosis* is present within the macrophages and extracellularly;
- development of TB disease due to disease progression following persistent infection or directly after infection in an immunocompromised host: On depletion of CD4 T-cells (e.g. during HIV infection), the granuloma does not function as well, production of IFN- γ may decrease, and macrophages are less activated. As a result, *M. tuberculosis* begins to multiply and TB disease develops. In the case of TNF neutralisation, the cells within the granuloma are no longer as tightly clustered, perhaps owing to chemokine or adhesion-molecule dysregulation. In addition, the macrophages are not as activated. These defects lead to a disorganised granuloma that is less able to control infection and greater immunopathology [10].

Figure 2. Potential outcomes of human host infection by *M. tuberculosis*

Following infection with *Mycobacterium tuberculosis*, some people are able to eliminate the bacterium through the innate or adaptive immune response [5]. However, many develop a persistent infection that may progress to TB disease, with approximately 5% doing so within the first two years [11]. The risk of TB disease is particularly elevated in the context of immune suppression, including impaired IFN- γ or TNF- α signalling, which are important macrophage activators [12]. Importantly, current tests for TBI are unable to differentiate people who have cleared the infection through the adaptive immune response from those with persistent infection or TB disease. reactive nitrogen intermediates (RNI), reactive oxygen intermediates (ROI) [12].

Image: ECDC. Created in BioRender. Lorentsson, H. (2026) <https://BioRender.com/xqli1y0>

3.2 The tuberculin skin test (TST)

First introduced in 1890, the TST is an intradermal injection of purified protein derivative (PPD). The PPD is a crude antigenic mixture, shared among *M. tuberculosis*, *M. bovis*, and other non-tuberculous mycobacteria (NTM) [13].

The test measures in vivo a delayed-type hypersensitivity reaction based on immunological recognition of mycobacterial antigens in exposed individuals. Mycobacterial antigens are injected below the epidermal layer, causing infiltration of antigen-specific lymphocytes and the elaboration of inflammatory cytokines. This inflammatory reaction results in the characteristic indurated area at the site of injection.

Before the introduction of IGRAs, the TST was the only tool for detecting TBI. Limitations of the test include:

- a high proportion of false positive and false negative results;
- difficulty in separating true infection from the effects of BCG vaccination and NTM infection;
- technical problems in intradermal administration;
- immune response boosting when repeated TST;
- potentially complicated and subjective interpretation; and
- a need for a second visit for measurement.

The TST is not described in this handbook as a laboratory procedure because it is an in vivo diagnostic method performed directly on the patient rather than in the mycobacteriology laboratory.

3.3 Interferon- γ release assays (IGRAs)

3.3.1 Introduction

WHO currently recommends the following IGRAs for the diagnosis of TB infection: the QuantiFERON-TB Gold Plus (QFT-Plus), by Qiagen GmbH, Hilden, Germany, the T-SPOT.TB (T-SPOT) by Revvity Inc., Waltham, the United States, the Wantai TB-IGRA (WANTAI), by Beijing Wantai Biological Pharmacy Enterprise Co. Ltd., Beijing, China, the STANDARD E TB-Feron ELISA (TBF), by SD Biosensor, Gyeonggi-do, Republic of South Korea, and the LIAISON QFT-Plus CLIA (LIAISON), by Diasorin, Saluggia, Italy [4,14,15]. The tests use an in-vitro ex-vivo approach for measuring cell-mediated immune responses to peptide antigens that simulate mycobacterial proteins. All platforms include antigens associated with ESAT-6, CFP-10, and TB7.7 (TB7.7 in TBF only). These antigens are absent from all BCG strains and most NTM.

Certain NTM, such as *M. gastri*, *M. kansasii*, *M. marinum*, *M. riyadhense*, *M. szulgai*, and *M. flavescens*, still contain homologous genes that may lead to cross-reactivity and false-positive results. The interpretation of IGRA results should therefore be made with caution in people with suspected NTM infection [16–18]. People infected with *M. tuberculosis* complex organisms, such as *M. tuberculosis*, *M. bovis*, *M. africanum*, *M. microti*, *M. canetti*, have mononuclear cells in their blood that recognise these mycobacterial antigens. This recognition process leads in vitro to the stimulation and secretion of IFN- γ from sensitised T-cells. The detection and subsequent quantification of IFN- γ , measured by enzyme-linked immunoassay (QFT-Plus/WANTAI/TBF), chemiluminescent immunoassay (LIAISON), or enzyme-linked immunospot (T-SPOT), forms the basis of these tests [14]. The tests are intended for use in conjunction with risk assessment, radiography, and other medical and diagnostic evaluations. Potential advantages of IGRAs over TST include: greater sensitivity, higher specificity (less influenced by BCG vaccination and NTM), less influence by technical problems in administration and interpretation, and the need for only one visit.

3.3.2 Current national guidelines and the clinical use of IGRAs

With the growing body of evidence, many national guidelines for the diagnosis of TBI now include IGRAs, although the TST continues to be used in many countries. TBI guidelines are predominantly available in high-income countries with established screening programmes. While considerable variation exists, four general approaches are commonly adopted:

- a two-step approach: TST performed first, followed by IGRA, either when the TST is negative (to increase sensitivity, particularly in immunocompromised individuals), or when the TST is positive (to increase specificity, particularly in BCG-vaccinated individuals);
- either TST or IGRA, but not both;
- IGRA and TST performed together (to increase sensitivity);
- IGRA only, replacing the TST.

Some guidelines do not recommend IGRA in children under five years old. Most current guidelines do not apply objective or transparent methods to grade the quality of evidence and strength of recommendations, and few disclose conflicts of interests. Existing national guidelines on TBI diagnosis in EU/EEA countries are listed in Table 2.

Table 2. List of national guidelines for the diagnosis of TB infection

Country	Guideline
Austria	Schmidgruber B. Guidelines for the diagnosis of latent tuberculosis in canton Vienna; Gesundheitsdienst (Department of Public Health) of Vienna; 2011; Austria.
Belgium	Wouter O. et al. Diagnose en behandeling van tuberculose; VGRT; 2010; Belgium. Link Callens S. et al. Belgian Guidelines on the Diagnosis and Management of Tuberculosis Infection; BELTA/FARES/VRGT; 2025; Belgium. Link
Bulgaria	Pharmacotheapeutic guide to pneumology and phtisiology; State Gazette 60/16 July 2024; 2024; Bulgaria. Link
Croatia	Programmatic guidelines for the control and prevention of tuberculosis; MoH; 2020; Croatia.
Cyprus	Prevention of transmission of Mycobacterium tuberculosis in healthcare settings; Ministry of Health Medical Services and Public Health Services; 2025; Cyprus. Link
Czechia	Recommendation of Czech Thoracic Society for QuantiFERON- TB Gold test; Research Institute for Tuberculosis and Respiratory Diseases; 2005; Czechia
Denmark	Andersen Å. et al. Tuberkulose Diagnostik og behandling; DSI; 2023; Denmark. Link Vejledning om Forebyggelse af Tuberkulose; Sundhedsstyrelsen; 2015; Denmark. Link Ravn P. et al. Tuberkuloseinfektion hos personer, som skal i immunsupprimerende behandling Rekommandationer for udredning og behandling; DSI/DLS; 2023; Denmark. Link Estmann A., Nielsen R. Retningslinje for infektionsscreening og profylakse ved inflammatorisk sygdom hos børn og unge; DPS; 2018; Denmark. Link
Estonia	Management of pulmonary and extrapulmonary tuberculosis; 2025; Estonia. Link
Finland	National Tuberculosis Program 2020: Recommendations for the prevention, treatment and monitoring of tuberculosis; 2020; Finland. Link
France	Infections tuberculeuses latentes Détection, prise en charge et surveillance; HCP; 2019; France. Link Dépistage de la tuberculose; Health insurance; 2025, France. Link La tuberculose; Ministry of health; 2025; France. Link
Germany	Richter E. et al. MIQ 05: Tuberkulose Mykobakteriose Qualitätsstandards in der mikrobiologisch-infektiologischen Diagnostik; 2019; Germany. Link
Hungary	The diagnosis of latent TB, care and TB prevention; NEAK; 2021; Hungary. Link
Ireland	Diagnosing and managing TB Infection; HSE Public Health: National Health Protection Office; 2025; Ireland. Link

Country	Guideline
Italy	Italian Ministry of Health; 2017; Italy. Link
Latvia	Procedure for the diagnosis of tuberculosis and tuberculosis infection; Latvian Association of Tuberculosis and Lung diseases; 2019, Latvia. Link
Malta	Pace-Asciak, A Prevention, Control and Management of Tuberculosis: A National Strategy for Malta; 2012; Malta.
The Netherlands	Commissie voor Praktische Tuberculosebestrijding- Screeningsbeleid; 2023; The Netherlands. Link
Portugal	DGS publica Referencial para o Diagnostico de Tuberculose em Crianças; DGS; 2024; Portugal. Link Manual de tuberculose e micobactérias não tuberculosas recomendações; DGS; 2020; Portugal. Link
Romania	Homorodean D., Moisiu A. National Guide for the TB Laboratory Network; 2022; Romania.
Slovakia	Slovakian Guidelines on Latent Tuberculosis Testing; Ministerstva Zdravonictva Slovenskej Republiky; 2020; Slovakia.
Slovenia	Svetina P. et al. Latent infection with Mycobacterium tuberculosis; 2015; Slovenia.
Sweden	Rekommendationer för preventiva insatser mot tuberkulos Hälsokontroll, smittspårning och vaccination; Folkhälsomyndigheten; 2022; Sweden. Link Björkman P. Tuberkulos – Vårdprogram från SILF; SILF; 2023; Sweden. Link
Switzerland	Handbuch Tuberkulose (Rohfassung); Kompetenzzentrum Tuberkulose 2011; Switzerland.

Information from guidelines and recommendations indicates that IGRAs are increasingly being adopted, particularly in low-incidence settings, as they offer higher specificity and logistical advantages [19]. The TST remains the preferred method in high-incidence and resource-limited settings. Importantly, screening for TBI, using TST and/or IGRA, should ideally be restricted to people with a sufficiently high risk of progressing to TB disease and who are likely to benefit from chemoprophylaxis if they test positive¹.

Overall, contact tracing practices in adults suggest a clear trend towards increased use of IGRAs, particularly in TB low-incidence countries, most commonly as a two-step testing strategy [20].

For the diagnosis of TB disease, ECDC does not suggest IGRAs as a replacement for standard diagnostic methods. In most situations, IGRAs provide limited or no additional value when combined with conventional diagnostic approaches. However, in specific clinical contexts, such as extrapulmonary TB, smear and/or culture negative TB, paediatric TB diagnosis, or in differential diagnosis of infection with NTM, IGRAs may offer supplementary information to support the diagnostic process and laboratory management [21].

3.3.3 Predictive value of IGRA for progression to TB disease

Existing evidence suggests that both TST and IGRA are acceptable but imperfect tests, and neither test can accurately differentiate between TB disease and TBI, distinguish recent from remote infection, or reliably predict progression from TBI to TB disease [19].

Overall, the available evidence indicates that the predictive value of IGRAs for progression to TB disease is very low and only marginally (and non-significantly) higher than that of the TST [22]. For high-risk populations (e.g. people living with HIV), current data are insufficient to suggest that IGRAs outperform the TST in predicting progression to TB disease [22].

In conclusion, both TST and IGRA have limited predictive value, and their main utility lies in identifying individuals who may benefit from preventive therapy. Ongoing and future studies may further clarify the role of IGRAs in TB clinical management and contribute to the identification of novel, highly predictive biomarkers.

3.3.4 IGRA performance and reproducibility

The performance characteristics (sensitivity and specificity) of the two IGRAs most commonly used in Europe (QFT-Plus and the T-SPOT) have been extensively reviewed, together with those of the TST and earlier versions of the QFT-Plus [19]. In the absence of a gold standard, surrogate markers are used to estimate test performance. The specificity of both assays (>95%) is not affected by BCG vaccination and is comparable to that of the TST in non-BCG-vaccinated individuals. In populations where BCG vaccination is routinely administered, the specificity of the TST is significantly lower (around 60%). Overall, the QFT-Plus and T-SPOT.TB assays show comparable diagnostic accuracy. WHO continues to endorse the use of either IGRA or TST, along with more recent Mtb antigen-based skin tests for the diagnosis of TBI.

IGRAs exhibit certain reproducibility issues. Variability may result from biological factors (such as immunomodulation and functionality of T-cells), manufacturing variability, or pre-analytical and analytical deviations. This has led to the reconsideration of revised cut-off values and the introduction of borderline zones (especially for the QFT-Plus), which may help improve both reproducibility and diagnostic performance of the assays [19,23].

3.3.5 Procedure 1: QuantiFERON- TB Gold PLUS

General principles

The QFT-Plus uses blood collection tubes containing antigens representing specific *M. tuberculosis* proteins or controls.¹ After blood collection (nil control, two antigen tubes, and a mitogen tube), the tubes are incubated at 37°C ± 1°C for 16 to 24 hours. Once incubation is complete, the tubes are centrifuged, plasma is harvested, and the amount of IFN-γ produced is measured by ELISA. Test results are reported in International Units (IU) relative to a standard curve generated using serial dilutions of a manufacturer supplied standard. Interference from heterophile antibodies is minimised by the addition of normal mouse serum to the green diluent and by using F(ab')₂ monoclonal antibody fragments as the IFN-γ capture antibody coated onto the microplate wells.

Baseline epidemiological data

Before performing the QFT-Plus, baseline epidemiological data should be recorded, including: name, full address, contact information, gender, occupation, place of birth, time since immigration (if applicable), travel history, history of BCG vaccination and TST, relevant clinical data (e.g. medication use, immunosuppression, weight loss, night sweats, fever, cough, abnormal chest X-ray (CXR), and previous TB treatment or chemoprophylaxis). These baseline data should be recorded on the patient data sheet accompanying the specimen (see Chapter 3, Annex 2).

Safety

All materials of human origin must be handled with care, as blood samples should be regarded as potentially infectious and may contain blood-borne pathogens such as HIV, hepatitis B, and hepatitis C. Handling, use, storage, and disposal of blood samples and assay components must comply with applicable national, regional, or local biohazard and safety regulations. Appropriate personal protective equipment – including eye protection, gloves, and standard laboratory protective clothing – should be worn. Correct laboratory procedures must be followed at all times to ensure safety and reliability.

Report interpretation

The predictive value of IGRA results depends on the prevalence of *M. tuberculosis* infection in the tested population. Each QFT-Plus result and its interpretation should be evaluated in conjunction with relevant epidemiological information, clinical history, physical findings, and diagnostic results. A negative test result **must be** repeated and confirmed at least two months after the exposure. The magnitude of the measured IFN-γ level cannot be used to determine the stage or degree of infection, the level of immune responsiveness, or the likelihood for progression to TB disease. QFT-Plus results are interpreted using the criteria in Table 3 below.

Table 3. QuantiFERON-TB Gold PLUS results interpretation

Nil (IU/ml)	TB1 minus Nil or TB2 minus Nil (IU/ml)	Mitogen minus Nil (IU/ml)*	QFT-Plus Result	Report/interpretation
≤8.0	≥0.35 and ≥ 25% of Nil	Any	Positive	<i>M. tuberculosis</i> infection likely
≤8.0	<0.35	≥0.5	Negative	<i>M. tuberculosis</i> infection NOT likely
≤8.0	≥0.35 and <25% of Nil	≥0.5	Negative	<i>M. tuberculosis</i> infection NOT likely
≤8.0	<0.35	<0.5	Indeterminate	Results are indeterminate for TB-antigen responsiveness
≤8.0	≥0.35 and <25% of Nil	<0.5	Indeterminate	Results are indeterminate for TB- antigen responsiveness
>8.0	Any	Any	Indeterminate	Results are indeterminate for TB- antigen responsiveness

When reporting results close to the clinical cut-off (0.35 IU/ml), it is important to acknowledge the inherent variability of the test within this borderline zone. Studies have shown higher rates of conversion and reversion near the threshold, raising concerns about reproducibility [23]. Previous investigations have revealed that laboratory practices across Europe vary. Some laboratories report results strictly as positive or negative per manufactures instructions, while others define an equivocal or borderline range, and may recommend repeat testing. Approaches include retesting the original specimen, requesting a new sample, or interpreting the result in context of clinical history and individual risk factors [24].

¹ The QuantiFERON-TB Gold package insert is available at: [QuantiFERON-TB Gold Plus ELISA Kit Package Insert - QIAGEN](#)

Limitations

- Diagnosis of TBI requires that TB disease is excluded through appropriate medical evaluation.
- A negative result must always be interpreted in the context of the individual's clinical and epidemiological information, particularly for individuals with impaired immune function.
- Several technical factors can lead to unreliable or indeterminate results, including:
 - incubation initiated more than 16 hours after blood collection;
 - storage of filled blood collection tubes outside the recommended temperature range ($22^{\circ}\text{C} \pm 5^{\circ}\text{C}$) prior to $37^{\circ}\text{C} \pm 1^{\circ}\text{C}$ incubation;
 - insufficient or incorrect mixing of blood collection tubes; and
 - incomplete washing of the ELISA plate.

If technical issues are suspected during sample collection or handling, the entire QFT-Plus procedure should be repeated. Note that responses to the mitogen positive control (and occasionally *M. tuberculosis* antigen) can be outside the range of the microplate reader. This does not affect the interpretation of test results.

Quality control

Internal quality assessment (IQA)

Test accuracy depends on the generation of an accurate standard curve. The results obtained from the standards must be reviewed before interpreting test sample results. Each laboratory should define the appropriate types of control materials and frequency of testing in accordance with local, regional, national, or other relevant accrediting or regulatory requirements (e.g. positivity rate, indeterminate rate).

External quality assessment

Comprehensive quality management, including both internal and external quality assessment (EQA), is a keystone of TB laboratory diagnosis and is essential for laboratory accreditation. The UK National External Quality Assessment Service (UK NEQAS, www.uknegas.org.uk) operates an EQA scheme for QFT-Plus tests, which is also available for customers outside the UK. Qiagen offers a QFT control panel (Cat No 0594-0805), comprising three sets of IFN- γ controls made up of recombinant human IFN- γ . Each complete set consists of three individual controls representing different IFN- γ concentration levels (levels 1, 2, and 3) within the linear range of QuantIFERON assays.

3.3.6 Procedure 2: T-SPOT

General principles

The T-SPOT assay, unlike the QFT-Plus, uses an enzyme-linked immunospot (ELISPOT) technique based on enumeration of activated specific T-cells responding to stimulation by specific antigens (ESAT-6 and CFP10) and resulting in IFN- γ secretion. Stimulation by ESAT-6 and CFP10 antigens occurs in separate microtitre plate wells.

During the procedure, peripheral blood mononuclear cells (PBMCs) are separated from a whole blood sample and counted to ensure a standardised number of cells is used in the assay. The PBMCs are incubated with the antigens to stimulate any sensitised T-cells present. Secreted IFN- γ is captured by specific antibodies coated on the membrane at the base of each well. A secondary antibody, conjugated to alkaline phosphatase and targeting a different epitope on the IFN- γ molecule, is then added and binds to the captured cytokine. Finally, a soluble substrate is added to each well. This substrate is cleaved by the enzyme, forming a spot of insoluble precipitate at the site of the reaction. Each spot represents the footprint of an individual cytokine-secreting T-cell. Counting the number of spots provides a measure of the abundance of *M. tuberculosis*-sensitive effector T-cells in the peripheral blood.

Baseline epidemiological data

In regard to the QFT-Plus assay, baseline epidemiological data are essential for the correct clinical interpretation of the test results. Data should include name, full address, contact details, gender, occupation, place of birth, time since of immigration (if applicable), travel history, history of BCG vaccination and TST, relevant clinical data (medication uptake, immunosuppression, weight loss, night sweats, fever, cough, abnormal CXR, and previous TB treatment or chemoprophylaxis). These data should be recorded on the patient data sheet accompanying the specimen (see Chapter 3, Annex 2).

Safety

All materials of human origin must be handled with care, as blood samples should be regarded as potentially infectious and may contain blood-borne pathogens such as HIV, hepatitis B, and hepatitis C. Handling, use, storage, and disposal of blood samples and assay components must comply with applicable national, regional, or local biohazard and safety regulations. Appropriate personal protective equipment, - including eye protection, gloves, and standard laboratory protective clothing - should be worn. Correct laboratory procedures must be followed at all times to ensure safety and reliability.

Important notes

- The T-SPOT assay involves cultivation of human PMBC. Therefore, strict aseptic technique must be applied throughout the procedure to avoid contamination of reagents, wells, cell cultures, and nourishing media;
- Blood samples should be processed within eight hours of collection. This time can be prolonged by using the T-cell *Xtend* reagent (previously available from Oxford Immunotec). If the T-cell *Xtend* reagent is used, the sample storage time before assay is increased to 32 hours. Only lithium-heparin tubes can be used in conjunction with T-cell *Xtend*.

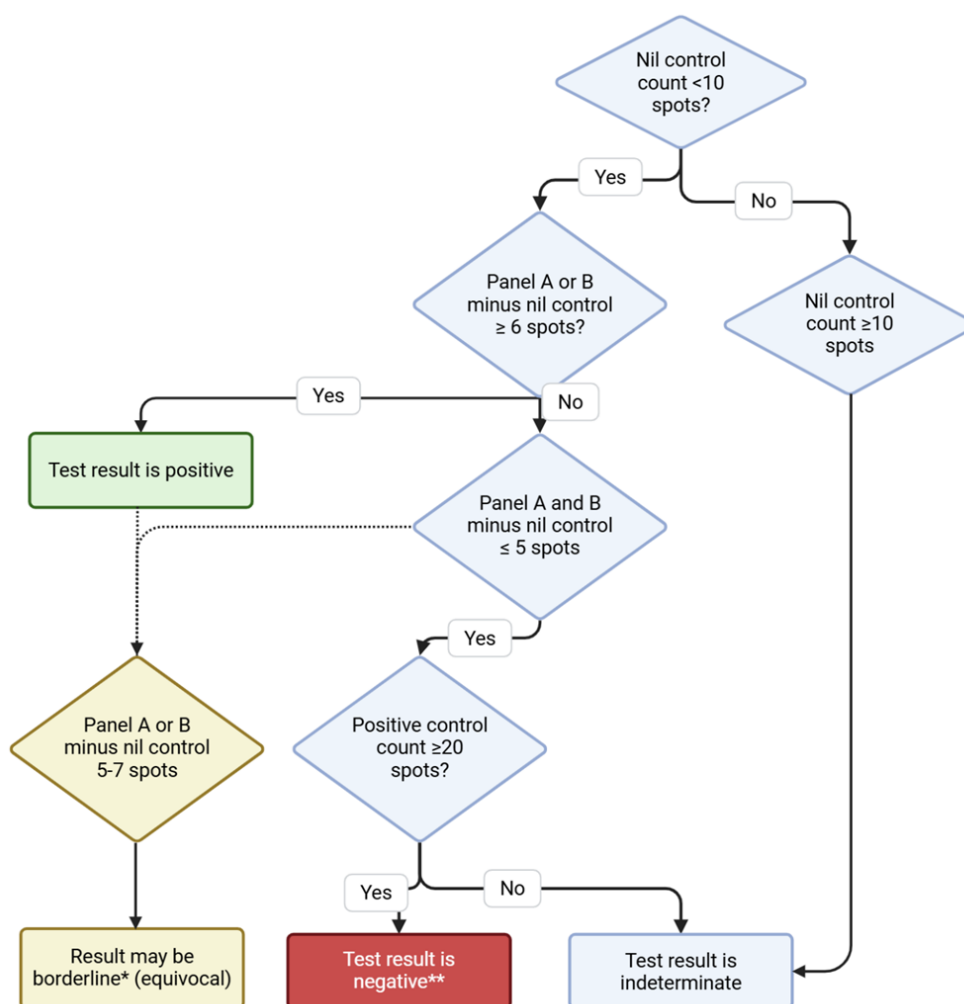
Reading and results interpretation

T-SPOT results are interpreted by subtracting the spot count in the nil control well from the spot count in each of the panels, according to the following algorithm (Figure 3):

- The test result is 'positive' if (Panel A minus nil control) and/or (Panel B minus nil control) ≥ 6 spots, AND a nil control count < 10 spots;
- The test result is 'negative' if both (Panel A minus nil control) and (Panel B minus nil control) ≤ 5 spots (including zero or negative values), AND a nil control count < 10 spots AND a positive control count ≥ 20 spots (or shows saturation);
- The test result is 'indeterminate' if either:
 - a nil control count ≥ 10 spots regardless of spot counts in Panel A and Panel B; or a positive control count < 20 spots and (Panel A minus nil control) and (Panel B minus nil control) ≤ 5 spots.
- In cases where the highest value of (Panel A minus nil control) and (Panel B minus nil control) is 5, 6, or 7 spots, the result may be considered as borderline (equivocal) due to potential biological and systematic variability. While borderline results are valid, they are less reliable than results further from the cut-off. Retesting using a new sample, is therefore recommended in such cases. If the repeat result is still borderline additional diagnostic tests and epidemiological information should be used to help determine TB infection status of the patient. A negative test result must be repeated and confirmed at least two months after the exposure.

Figure 3. T-SPOT interpretation algorithm

Note: This figure is a visualisation of the text above. Result interpretation should always be performed in accordance with the manufacturer's instructions.



* Borderline results are valid but less reliable. Retesting using a new sample is therefore recommended in such cases. If the repeat result is still borderline, additional diagnostic tests and epidemiological information should be used to help determine TB infection status of the patient.

** A negative test result must be repeated and confirmed at least two months after the exposure.

Created in BioRender. Lorentsson, H. J. N. (2026) <https://BioRender.com/0825twc>

Reporting

The manufacturer recommends using the following wording in the laboratory reports:

- A 'positive' result indicates that the sample contains effector T-cells reactive to *M. tuberculosis*.
- A 'negative' result indicates that the sample probably does not contain effector T-cells reactive to *M. tuberculosis*.

Quality control

Internal quality assessment (IQA)

Appropriate means of internal quality assurance and control should be determined, developed and implemented by each laboratory in accordance with local and governmental regulations. This can include:

- blind re-testing of specimens on a regular basis;
- keeping records on dates when kits are opened and finished, kit lot numbers; and
- fridge and freezer temperature sheets.

External quality assessment (EQA)

No formal EQA schemes for T-SPOT TB tests currently exist. These should be established and implemented as soon as possible.

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Annex 2. Patient data sheet

Sender’s contact details:

Name Laboratory/hospital
 Postal address
 Phone
 Fax
 E-mail

For laboratory use

Patient’s number
 Date received
 Time received
 Date of test

Please circle/mark/tick appropriate answer(s). Please do not leave any fields blank. If answer requires further details, please specify. Please remember that complete answers are essential for the correct interpretation of the test results.

Baseline epidemiological data

Patient’s first name
 Surname
 Date of birth
 Male ___ Female ___
 Postcode
 Occupation
 Was the patient born abroad?
 No ___
 Yes ___ Born in _____
 If yes, when did the patient come to _____(country)? _____(year)
 Has patient lived, or spent more than two months travelling in another country?
 Yes ___
 No ___
 Don’t know

History of BCG vaccination and TB skin tests

Has patient ever received a BCG vaccination? Yes ___ No ___ Don’t know ___
 If yes, please specify age: .
 BCG scar: Yes ___ No ___
 TB skin test done? Yes ___ No ___ Don’t know ___ Reading ___ mm

Clinical data

Is patient taking any of the following medications?
 ___ Oral steroids
 ___ Cytotoxic drugs
 ___ Other immunosuppressive drugs (please specify)
 ___ None of the above.
 Is the patient immunocompromised? Yes ___ No ___ Don’t know ___
 Is the patient HIV positive? Yes ___ No ___ Don’t know ___
 Does the patient have diabetes? Yes ___ No ___ Don’t know ___
 Does the patient have any of the following:
 ___ Fever
 ___ Night sweats
 ___ Loss of weight
 ___ Cough
 Is the patient’s CXR abnormal? Yes ___ No ___. If yes, please specify the location:

	R	L
Upper	_____	_____
Middle	_____	_____
Lower	_____	_____

Cavities? Yes ___ No ___ Don’t know ___
 Consolidation? Yes ___ No ___ Don’t know ___. Unilateral/bilateral?
 Other relevant clinical data:

4 Smear microscopy

Susana David, Vera Katalinić-Janković, Daniela Cirillo

Revised by Emanuele Borroni and Enrico Tortoli (2015)

4.1 Background and principles

Early laboratory diagnosis of TB still relies on the examination of stained smears. For universal application in resource-limited countries, microscopy of stained sputum smears is the best choice among diagnostic methods [7, 12]. This technique is based on the fact that the cell wall of the *Mycobacterium spp.* genus is rich in complex lipids that prevent access to common aniline dyes, but when stained with carbol-fuchsin or fluorochromes under special staining conditions, these are not easily decolourised, even with alcohol-acid solutions. Because of this characteristic, all members of *Mycobacterium spp.*, not only *M. tuberculosis*, are referred to as acid-fast bacilli (AFB).

At present, two types of acid fast stains are used to detect mycobacteria in clinical specimens:

- Carbol-fuchsin staining (Ziehl-Neelsen [ZN] method and its modification performed without heating the dye [Kinyoun cold staining]); and
- Fluorochrome (auramine or auramine-rhodamine) staining.

Kinyoun staining is a modification of the classic ZN staining which excludes the heating step during the staining procedure and uses a higher concentration of carbol-fuchsin. Mycobacteria appropriately stained by ZN and Kinyoun appear as red rods. Kinyoun staining is not as effective as ZN, therefore this procedure is not recommended [6].

Methods which apply a fluorochrome have been used to stain acid fast bacteria for many years. Using this method, mycobacteria are detected as bright fluorescent rods against a darker background. Fluorochrome staining has an increased sensitivity and less time is required to screen the slides when compared to Kinyoun or ZN staining because slides are screened at lower magnification [6].

Smear microscopy is simple, inexpensive and efficient in detecting those cases of pulmonary TB that are most infectious. Since its yield is highly dependent on its execution, the quality of smear microscopy is crucial in the fight against TB in resource-limited settings [6,7].

A major limitation of smear microscopy is its low sensitivity (25–75% compared to culture) and the high number of bacilli required for positivity (in the range of 5×10^3 – 10^4 bacilli per ml). Sensitivity and the positive predictive value (PPV) of smear microscopy are influenced by numerous factors [7,12,13] such as the prevalence and severity of the disease, the type and quality of the specimen, the number of mycobacteria in the sample and the quality of the smear preparation, staining and reading process. Smear microscopy does not allow for mycobacterial species identification, nor does it give an indication of the viability of mycobacteria in the sample. HIV co-infected TB patients may have disseminated paucibacillary disease with fewer AFB. Smear microscopy is often negative or may require more scrutiny in screening to identify lower numbers of AFB.

4.2 Procedure 1: Ziehl-Neelsen (ZN)

Each batch of prepared reagent should be recorded in a reagent preparation workbook which includes: the signature of the technician who prepared it, the date of preparation and the results of quality control testing [2,6].

4.2.1 Ziehl-Neelsen (ZN) reagent preparation

Good staining reagents, made with high-quality carbol-fuchsin dye are essential for detecting AFBs [6]. Contamination of reagents by environmental mycobacteria should be prevented by using freshly distilled water.

Standard reagents:

- Basic fuchsin powder
- Phenol crystals (the crystals should be almost colourless)
- Alcohol (denaturated 95% ethanol)
- Water (distilled or purified).

Decolourising solution:

- Concentrated sulphuric acid ($\geq 95\%$)
- Water (distilled or purified) or
- Hydrochloric acid (37%)
- Alcohol (denaturated 95% ethanol).

The counterstain solution:

- Methylene blue powder
- Water (distilled or purified).

A. Carbol-fuchsin (CF) reagent

The quality of basic fuchsin varies among different manufacturers with regard to its purity and solubility. The basic fuchsin content should represent 85–88% of the weight. If carbol-fuchsin purity is known, it should be used to calculate the final stain concentration of 0.3%. To calculate the required amount of basic fuchsin, divide the actual amount by the dye content. For instance, if the dye content is 75%, you must divide the amounts by 0.75. So $3\text{ g}/0.75 = 4$ grams will be weighed for the 0.3% stain. If powder with a dye content of $>85\%$ is used, there is no need to calculate the correction factor. If the dye purity is unknown or if the basic fuchsin dissolves poorly or precipitates are still visible after filtration, it may be wise to use the higher concentration (1%) when preparing the staining reagent.

0.3% Carbol-fuchsin	
Basic fuchsin	3.0 g
95% ethanol	100 ml
Phenol crystals	50 g
Distilled water	900 ml

- Weigh 3.0 g of basic fuchsin powder and 50 g of phenol crystals separately.
- Add 100 ml of alcohol (denaturated ethanol) to a 1-l conical flask.
- Add 50 g of phenol and swirl the flask until it is dissolved.
- Add 3.0 g of basic fuchsin powder and continue to mix well until the fuchsin powder completely dissolves.

Check for remaining powder or crystals on the bottom. If there are any, continue swirling with slight heating. Only after the fuchsin is completely dissolved, add 850 ml of water and mix by continuing to swirl.

If precipitates are visible, the carbol-fuchsin staining reagent should be filtered. Filter the carbol-fuchsin again during the staining process, using a funnel with filter paper (or by placing a piece of filter paper directly on the slide). Other staining reagents do not need to be filtered. If any particles are detected in the carbol-fuchsin solution, the solution must be refiltered.

B. Decolourising solution

25% sulphuric acid	
Concentrated sulphuric acid	250 ml
Distilled water	750 ml

- Add 750 ml of distilled water to a 2-l conical flask.
- Measure 250 ml of concentrated sulphuric acid in a cylinder.
- Pour it slowly into the flask containing the water, directing the flow of acid gently along the inner side of the flask. **Always add the acid slowly to the water, not vice versa.**
- Mix well by swirling the flask.

3% HCl ethanol	
Concentrated hydrochloric acid	30 ml
95% ethanol	970 ml

- Add 970 ml of 95% ethanol to a 2-l conical flask.
- Measure 30 ml of concentrated hydrochloric acid into a cylinder.
- Pour it slowly into the flask containing alcohol, directing the flow of acid gently along the inner side of the flask with constant swirling. **Always add the acid slowly to the alcohol, not vice versa.**
- Mix well by swirling.

C. Counterstain

Methylene blue chloride	3.0 g
Distilled water	1 000 ml

- Weigh 3 g of methylene blue powder.
- Add the powder to 0.5 l of pure water in a conical flask.
- Swirl the contents of the flask to dissolve the dye.
- Add 0.5 l of water and mix again.

4.2.2 Storage of reagents

The flasks of freshly-prepared reagents should be covered until quality control procedures have been performed and the results have been evaluated [7]. Solutions should be stored in clean brown bottles and clearly labelled. The label should indicate the reagent name, concentration, and the preparation date. Reagents preserved in tightly closed bottles can be used for up to one year. Bottles should be kept out of direct sunlight. If clear bottles are used, stocks of reagents should be stored in a closed cabinet.

4.2.3 Quality control of freshly-prepared staining reagents

After preparing staining reagents, quality control checks should be performed on every batch [7,14]. Quality control is essential to ensure the effectiveness of staining reagents and the complete absence of AFB contamination.

Quality control results should be recorded in a logbook, with every batch clearly identified by the name of the reagent and the preparation date on the bottle labels. Perform quality control by using one or more freshly-prepared staining reagents, carrying out the usual staining procedure as described for positive controls. Test the performance of cabol-fuchsin by staining and examining two scanty or 1+ smears stained once, and two negative smears stained three times [2].

4.3 Procedure 2: Fluorochrome staining

4.3.1 Quality control of freshly-prepared staining reagents

A. Fluorochrome reagents

Auramine O (solution 1)	
Auramine	0.1 g
95% ethanol	10 ml

Phenol (solution 2)	
Phenol crystals	3.0 g
Distilled water	87 ml

Dissolve phenol crystals in water.

Mix solutions 1 and 2 and store in a tightly stoppered, dark-coloured bottle away from heat and light. Label bottles with the name of the reagent, the date of preparation and expiry date. Store at room temperature for three months. When left standing turbidity may develop but this does not affect the staining reaction.

B. Decolourising solution

0.5% Acid alcohol	
Concentrated hydrochloric acid	0.5 ml
70% ethanol	100 ml

Carefully add concentrated hydrochloric acid to the ethanol. Always add acid slowly to the alcohol, not vice versa. Store in a dark-coloured bottle. Label bottles with the name of the reagent, the date of preparation and expiry date. Store at room temperature for three months. For each volume of stain, two to three volumes of decolourising solution are needed.

C. Counterstains

Either potassium permanganate or acridine orange may be used as counterstains.

Potassium permanganate	
Potassium permanganate (K ₂ MnO ₄)	0.5 g
Distilled water	100 ml

Dissolve potassium permanganate using distilled water in a tightly stoppered, dark-coloured bottle. Label bottles with the name of the reagent, the date of preparation and expiry date. Store at room temperature for up to three months.

Acridine orange	
Anhydrous dibasic sodium phosphate (Na ₂ HPO ₄)	0.01 g
Distilled water	100 ml
Acridine orange	0.01 g

Dissolve sodium phosphate in distilled water. Add acridine orange and mix until dissolved. Store in a tightly stoppered dark-coloured bottle away from heat and light. Label bottles with the name of the reagent, the date of preparation and expiry date. Store at room temperature for three months.

4.3.1 Safety measures

Never add water to acid. To reduce exposure to toxic phenolic fumes, reagents and staining solution containing phenol should be prepared in a well-ventilated area or under a chemical hood. Always wear protective laboratory coats, gloves and safety glasses when handling a strong acid. In the event of an accident with acid, rinse the affected body part immediately with plenty of water.

4.4 Sample collection

Smear microscopy for *Mycobacterium spp.* detection can be used for a wide variety of biological samples. For the diagnosis of respiratory TB, sputum is the most commonly used sample. To ensure optimal recovery of TB bacilli from sputum, at least two specimens should be collected and processed for mycobacterial microscopy and culture [7,15].

Country guidelines will provide information on the number of recommended samples. Early morning specimens have the highest yield of AFB; however, it is now proven that good diagnostic specimens can be collected at any time. It is not recommended to perform smear microscopy from blood or very bloody samples due to the low sensitivity of the procedure. It is also not recommended to routinely perform smear microscopy from urine samples due to the frequent detection of saprophytic mycobacteria colonising the urogenital tract.

Samples should be collected in clean, wide-mouthed and leak-proof specimen containers [2,6]. Single use disposable plastic containers (50 ml capacity) are preferred in order to avoid transferring the specimens from one container to another. Alternatively, 50 ml disposable sterile conical tubes can be used.

Patients should receive clear written instructions on the proper collection of the sputum specimen for TB diagnosis. For patients on treatment, specimens should be collected at intervals specified in accordance with the country's guidelines [7]. Sputum collection should never be performed in the laboratory. It is a procedure generating infectious aerosols and should only be performed at a distance from other people, preferably in open spaces where possible, or in rooms with negative pressure and adequate air exchange [7,14].

A good specimen should be approximately 3–5 ml in volume [5]. Sputum specimens should appear thick and mucoid or clear but with purulent grains [7]. The colour varies from opaque white to green. Bloody specimens will appear reddish or brown. Note: clear saliva or nasal discharge is not suitable as a TB specimen [2,15].

Specimen handling

For optimum patient management, process the specimen as soon as possible (i.e. < 24 hours). For microscopic examination, the interval between collection and staining is not critical. Acceptable results can be obtained even if specimen delivery has been delayed.

Criteria of acceptability

Upon arrival in the laboratory, the quality of sputum samples should be assessed and reported in the referral form [7]. TB-positive sputa can vary in colour and aspect. If the sample is liquid and is clear and water-like, without particles or streaks of mucous material, process the sample but ensure that the poor quality of the sample is reported on the result form. When possible, encourage the patient/physician to submit a new specimen; however, even saliva can yield

positive results. All specimens should be processed except for broken or leaking containers, which should be discarded and another specimen requested.

Accept very small quantities if the patient has difficulty producing sputum. Blood-streaked sputum is suitable, but pure blood should not be examined [1,2].

4.5 Smear preparation

Although smear preparation for AFB detection [7] is a relatively safe procedure in terms of infected aerosol production, it is recommended that the slides be prepared in a class I or IIB biological safety cabinet [14,15] if available. If the smear is prepared after centrifugation of the sample (concentrated smear), the centrifuge holder must be opened within a biological safety cabinet.

- Smears should be prepared using new, clean, grease-free and unscratched slides. Using a pencil, record the laboratory register serial number and order number of the specimen on the frosted end of the slide. If plain unfrosted slides are used, label them using a diamond pencil.
- If smear is prepared directly from a fresh sample (without prior centrifugation) use an applicator stick or wire/disposable loop, select and pick up the yellowish purulent particles of sputum. For re-suspended pellets (after the centrifugation) a disposable loop is advisable.
- Prepare the smear in an oval shape in the centre of the slide. The smear size should be 2–3 cm in length and 1–2 cm wide, which will allow 100–150 fields to be counted in one length.
- For good spreading of the sputum, press the stick firmly perpendicular to the slide and move in small concentric circles or coil-like patterns.
- Place the used stick in a discard container.
- Use a separate stick for each specimen.
- Thorough spreading of the sample is very important, especially in the case of thick or purulent material; it should be neither too thick nor too thin. Prior to staining, hold the smear about 4–5 cm over a piece of printed paper. If letters cannot be read, it is too thick.
- For concentrated samples (after centrifugation at 3 000x g for 20 minutes, see sample preparation for TB culture) one or two drops of sediment should be smeared on the slide.
- Allow the smear to air-dry completely at room temperature within the biological safety cabinet.
- Do not dry smears in direct sunlight or over a flame.
- Pass the slide over a flame 2–3 times for about 2–3 seconds each time. Do not heat the slide for too long or keep it stationary over the flame or else the slide will be scorched.
- Alternatively, slides can be fixed for two hours on hot plates (65–75°C), within the biological safety cabinet. Table 4 lists the equipment needed for direct (unconcentrated) smear microscopy.

Table 4. Equipment required for smear preparation and staining

Equipment required for smear preparation and staining
Container to store specimen
Wire loop with an inner diameter of 3 mm to spread sputum on the slide
Microscope slide (grease-free and unscratched)
Marking pen to put the identification number on the microscopy slide
Forceps to hold smear slide
Bunsen burner to fix the smear slide and flame the smear during staining
Staining rack to hold the smear slide
Slide rack in which to place stained smear slide for air-drying

4.6 Staining procedures

4.6.1 Ziehl-Neelsen staining method

- Cover the entire surface of each heat-fixed slide with carbol-fuchsin.
- Using a Bunsen burner, gently heat the slides until vapour rises. Do not allow them to boil.
- Allow the stain to remain on the slide for ten minutes. Adequate time is required for the carbol-fuchsin to penetrate and stain the cell wall.
- Gently wash the stain from each slide with a stream of cold water until all the free stain has washed away.
- Cover each slide with acid alcohol; wait three minutes.
- Rinse slides again carefully with water and tilt each slide to remove excess water.
- Flood the slide with the methylene blue counterstain for one minute.
- Rinse slides again carefully with water, drain and air dry.

4.6.2 Fluorochrome staining method

- Prepare and heat fix smears.
- Place the numbered smears on a staining rack in batches (maximum 12).
- Flood the slides with auramine O stain and allow them to stain for 15 minutes.
- Be sure that the stain stays on the smear. Do not heat and do not use paper strips.
- Rinse the slide with water. Aim the flow of water at the edge of the slide and slowly peel the stain from the slide.
- Flood the slides with 0.5% acid alcohol and allow them to decolourise for three minutes.
- Ensure that the slides are flooded thoroughly with acid alcohol.
- Rinse off the 0.5% acid alcohol with water, drain the excess water from the slide.
- Flood each slide with potassium permanganate and allow it to quench for two minutes.
- Note: It is critical that the potassium permanganate remains on the slides for no longer than two minutes as over-quenching of fluorescence can occur.
- Wash off the potassium permanganate. Drain the excess water from the slide.
- Allow smears to air dry. Do not blot. Read as soon as possible after staining.

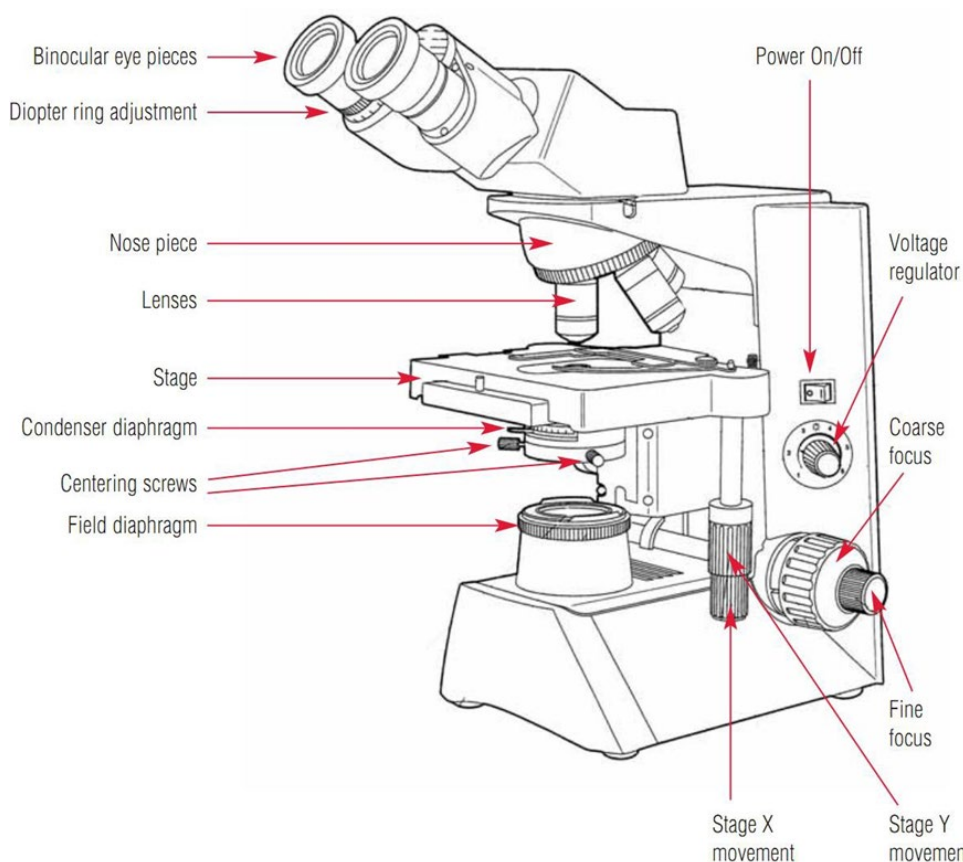
4.6.3 Automatic staining

Automated stainers that can process a large number of samples are commercially available. The machines require dedicated reagents and are able to perform both ZN and fluorochrome staining. Accurate and appropriate maintenance after each staining session is required to maintain consistent, high-quality staining.

4.7 Microscopy

Figure 4 shows the requirements of a microscope for smear examination.

Figure 4. Microscope components



Source: Lumb R, Bastian I. Laboratory diagnosis of tuberculosis by sputum microscopy. Adelaide: Institute of Medical and Veterinary Science; 2005. p. 38 [17].

4.7.1 Maintenance

Install the microscope on a rigid, flat, level surface, away from direct sunlight, dust, vibration (e.g. from centrifuges), water (sink, spray from a tap), chemical reagents or humidity.

The modern light microscope needs no particular daily maintenance, but considerable care is required in its use. For further information, please refer to the microscope manual for care and maintenance information.

4.7.2 Fluorescence microscopy

The identification of mycobacteria with the fluorescent dye auramine O is based on the affinity of the fluorochrome to the mycolic acids in the cell wall. Auramine O is excited by blue light and emits in the region of ~500 nm to ~650 nm.

Fluorescence microscopy has some significant advantages:

- High contrast fluorescence images allow for easier detection of AFB.
- The use of low- to medium-power lenses (typically 10x, 20x and 40x) permits a larger field of view than conventional microscopy, where typically a 100x lens is used.
- The fluorochrome staining method is simpler than the ZN method.

A binocular microscope equipped with a fluorescent light source and suitable filter set is used for auramine-stained smears. Fluorescent light is provided by a vapour lamp (such as mercury or xenon lamps). The mercury vapour lamp provides the strongest light, but it has a limited lifespan of about 100 to 200 hours, which must be monitored with a timer. Moreover, these lamps are very expensive and fragile.

4.7.3 Light-emitting diode (LED) microscopy

There is a compelling base of evidence promoting ultra-bright LED microscopy as a substitute for both conventional fluorescence microscopy and direct ZN microscopy [11]. LED-based microscopy facilitates identification of acid-fast bacilli in comparison with ZN, can be used with auramine staining, is cost-effective (lifespan of the lamp is over 10 000 hours), has low power requirements, and can be easily introduced in microscopy centres, including peripheral facilities. In addition, light intensity can easily be regulated.

Since LED-based microscopy has been acknowledged as a significant development in direct fluorescence microscopy, WHO has recommended that it replace conventional fluorescence microscopy and that it be phased in as an alternative to conventional ZN microscopy in both high- and low volume laboratories. During the implementation of LED microscopy, the following issues are of importance: training requirements, validation during the introductory phase, monitoring of trends in case detection and treatment outcomes. Adapted systems may need to be introduced for internal quality control as well as external quality assurance.

4.8 Recording and reporting

Recording and reporting of results [2,6,7] is summarised in Table 5.

Table 5. Reporting of microscopy smears

IUATLD/WHO scale (1000x field = HPF)	Microscopy system		
	Bright field (1 000x magnification: 1 length = 2 cm = 100 HPF)	Fluorescence (200–250x magnification: 1 length = 30 fields = 300 HPF)	Fluorescence (400x magnification: 1 length = 40 fields = 200 HPF)
Result			
Negative	Zero AFB/1 length	Zero AFB/1 length	Zero AFB/1 length
Scanty	1–9 AFB/1 length or 100 HPF	1–29 AFB/1 length	1–19 AFB/1 length
1+	AFB/1 length or 100 HPF	30–299 AFB/1 length	20–199 AFB/1 length
2+	FB/1 HPF in at least 50 fields	10–100 AFB/1 field on average	5–50 AFB/1 field on average
3+	3/1 HPF in at least 20 fields	>100 AFB/1 field on average	>50 AFB/1 field on average

4.8.1 ZN smear examination

Examine one length of the smear (2 cm) or 100 fields with light microscope, using 1 000x magnification. If less than 10 AFB are found in 100 fields, the number of AFB should be counted. For high positives, examination of only 20 to 30 fields is sufficient.

4.8.2 Auramine (fluorochrome) smear examination

Examine one length of an auramine slide with a fluorescent microscope, using 200–250x magnification, to cover 30 fields in one length, equivalent to 300 fields at 1 000x magnification. Alternatively, 400x magnification can be used, covering 40 fields at this magnification.

Negative report: Negative for acid-fast bacilli where no organisms observed in 100 fields. Positive report: Positive for acid-fast bacilli; provide AFB quantification.

The results should be recorded in the TB laboratory register, and recorded on the sample examination request form as well as forwarded to the person requesting the sample examination [10].

Figures 5–6 give examples of smear microscopy using auramine and ZN staining.

Figure 5. AFB smear microscopy of biological specimens

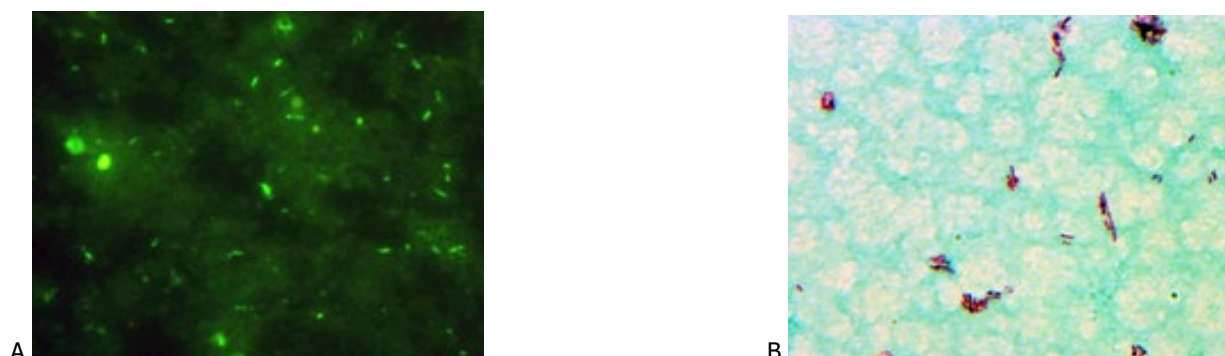
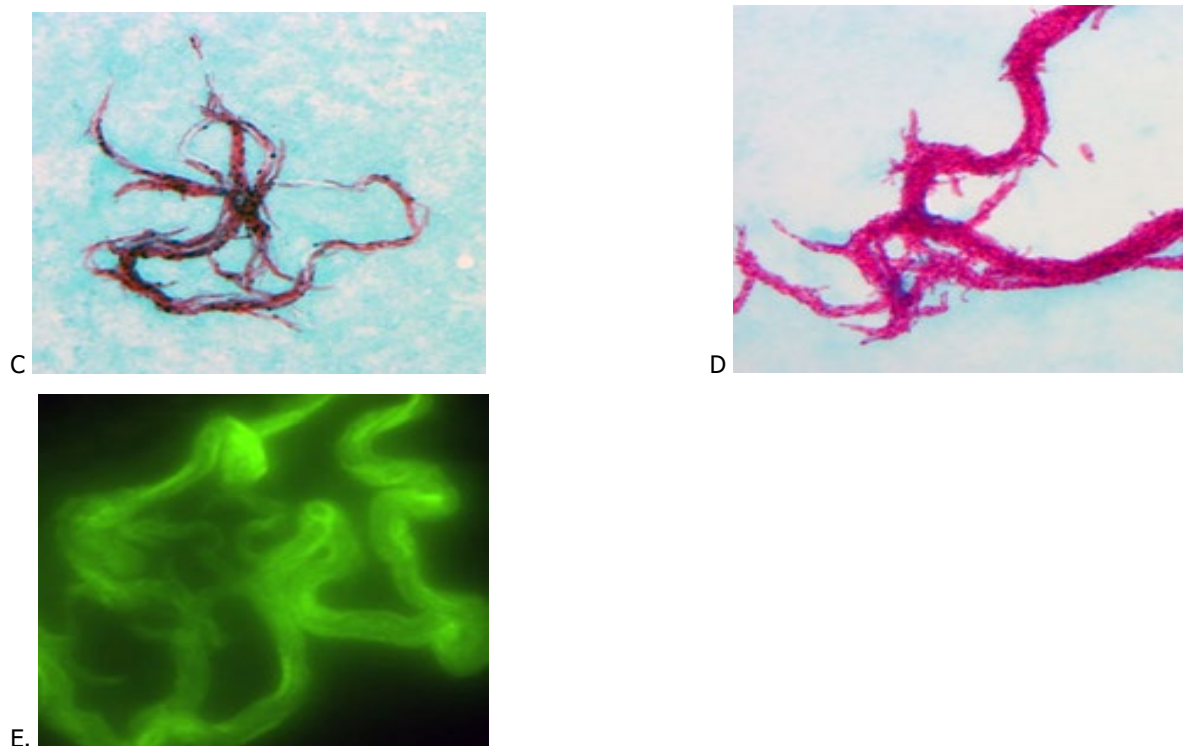


Figure 6. AFB smear microscopy of *M. tuberculosis* cultures



A and E: Auramine stain B–D: Ziehl-Neelsen stain

Images show cords. Pictures were kindly provided by Professor Zofia Zwolska, Head of the Microbiology Department, National Tuberculosis Reference Laboratory, National Tuberculosis and Lung Diseases Research Institute, Warsaw, Poland.

4.9 Quality control

4.9.1 Quality control parameters

Quality control in microscopy is a process for internally monitoring the performance of bench work in the laboratory. It consists of an effective and systematic process, ensuring that laboratory work is accurate, reliable and reproducible. This is done by assessing the quality of specimens; monitoring the performance of microscopy procedures, reagents and equipment; reviewing microscopy results and documenting the validity of microscopy methods.

A positive and a negative control slide should be included in each run of stains, verifying the correct performance of the procedure as well as the staining intensity of the acid-fast organisms [14,15].

Table 6 shows the most common causes of error in smear microscopy. Control slides should be assessed prior to reading the patient smears to confirm the correctness of staining. If quality control slides are acceptable, patient smears can be read and reported. If the control slide(s) are unacceptable, the procedures and reagent preparations should be reviewed. After identifying and correcting the problem, all patient slides should be repeated with a new set of controls. The results of the quality control of reagents should be reported in the reagent preparation workbook.

Table 6. Common causes of error in smear microscopy

Errors	Cause	Action to be taken
False negative	Smear too thick, detaching during staining	Improve homogenisation, reduce the material deposited.
	Smear too thin	Increase the amount or make smear in 1x2 cm area only.
	Poor staining	Check quality control of reagents, prepare new reagents, check dilution.
False positive	Cross-contamination	Avoid contact between slides during staining procedure, do not use staining jars. Clean objective lens after reading each slide. Check water/solutions for environmental contamination.
	Red precipitates	Prepare new solution. Filter before use.

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5 Culture tests for *Mycobacterium tuberculosis* complex

Susana David, Vera Katalinić-Janković, Lanfranco Fattorini and Daniela Cirillo

Revised by Emanuele Borroni, Enrico Tortoli, and Vladyslav Nikolayevskyy (2015)

5.1 Background and principles

Bacteriological cultures can provide a definitive diagnosis of TB. The primary advantage of culture tests over sputum microscopy is their higher sensitivity, allowing for the detection of very low numbers of bacilli (approximately 10 bacilli/ml of sputum compared with at least 5 000 bacilli/ml of sputum for microscopy). The use of cultures increases the potential of diagnosing TB at early stages of the disease. Culture tests are also used for the detection of treatment failures and for diagnosing extrapulmonary TB. The use of culture tests increases the number of TB cases found by 30–50%. Moreover, cultures are used for species identification and drug susceptibility testing (DST) [1,2].

As the EU has adopted the culture-based case definition, the main distinction in TB cases is between culture-positive and -negative, and not sputum smear status. The first section of the European Standards for Tuberculosis Care (ESTC) 2012 [1] dedicated to the Standards for Diagnosis (ESTC 1 through ESTC 6) specifies the minimum requirements for obtaining a valid TB diagnosis in different settings or with various suspect types, both microbiological and clinical [1,2]. If this is not feasible, culture tests should at least be performed for:

- diagnosis of cases with clinical and radiological signs of pulmonary TB where smears are repeatedly negative;
- diagnosis of extrapulmonary TB;
- diagnosis of childhood TB;
- diagnosis of TB among HIV-positive adults and children; and
- diagnosis and monitoring of MDR- and XDR-TB.

TB, although mainly a pulmonary disease, can affect any organ of the body. The isolation of the aetiological agent for effective microbiological diagnosis is dependent on:

- selection of the correct type of specimen;
- the quality of the sample; and
- adequate use of storage and transportation procedures.

Processing of inappropriate clinical specimens for mycobacterial cultures is a waste of both financial and human resources [6]. Clinical staff should be properly trained and accept only suitable specimens.

Because mycobacteria are usually slow growing and require a long incubation time, a variety of other microorganisms can overgrow the cultures of specimens obtained from non-sterile sites. Appropriate pre-treatment and processing of samples, as well as the use of selective culture media is critical for eliminating contaminants while not seriously affecting the viability of mycobacteria [10].

5.2 Biohazards and biosafety in the TB laboratory

Good microbiological techniques (GMT) – working methods designed to eliminate or minimise exposure to pathogens via, for example aerosols, splashes or accidental inoculation – are essential for minimising biohazards [13]. Nosocomial transmission of *M. tuberculosis* from specimens is a major concern for laboratory workers. All specimens suspected of containing *M. tuberculosis* should be handled with appropriate precaution at all times and opened only within an appropriate biosafety cabinet. Infectious aerosols are produced in the TB laboratory whenever a liquid suspension containing tubercle bacilli is handled. Biosafety measures in the laboratory are essential to protect workers against exposure to infectious aerosols. Please refer to Chapter 1 for more details on procedures and laboratory safety practices.

Because of their viscosity, sputa are a minimal source of infectious aerosols. By contrast, aerosols produced during processing (especially during centrifugation) of homogenised sputa and during culture handling, must be minimised and therefore processed and contained in a biological safety cabinet.

Classification of laboratory practices used for *M. tuberculosis* diagnosis should be based on a risk assessment (number and type of tests, prevalence of tuberculosis and of MDR-TB). Specimen processing for mycobacterial culture should be performed in a biological safety cabinet in at least a biosafety level 2 (BSL2) laboratory, whereas procedures involving manipulation of *M. tuberculosis* cultures (identification, sub-culturing, and DST) must be performed in a biological safety cabinet in laboratories complying with BSL3 standards. All aerosol-generating procedures should be performed in a class I or II biological safety cabinet [9-11].

The health of laboratory workers should be regularly monitored by the employer. They should be educated about the symptoms of TB and, if symptoms arise, they should be provided with readily accessible free medical care in accordance to the national regulations.

5.2.1 Minimum WHO recommendations for TB culture/drug susceptibility testing facilities

WHO recommends that all specimen processing procedures are carried out in a laboratory built and equipped for BSL2. The minimum requirements for a BSL2 TB laboratory are: restricted access to the laboratory, the presence of a fully functional and maintained biological safety cabinet and an autoclave or other means of decontamination available in the same building. More information on biosafety is given in Chapter 1 [9-14].

Identification, subculturing, and drug susceptibility testing should be performed in a BSL3 containment room with an anteroom and directional airflow from functionally clean to dirty areas, with at least six to 12 air exchanges per hour. The containment room may be the blind end of a corridor or formed by constructing a partition and door so that access to the laboratory is through an anteroom (e.g. double-door entry) or through the basic BSL2 laboratory. The autoclave should be in the vicinity of the laboratory so that the movement of contaminated materials is minimised. Biological safety cabinets must be ducted to the outside or vented through a thimble. Recirculation of air from biological safety cabinets into the laboratory room and recirculation to other areas within the building are not permitted. Please refer to Chapter 1 for further detail on laboratory safety levels and conditions [14].

The decision to use additional PPE should be based on risk assessments. Risk assessments should be reviewed routinely and revised when necessary [7].

Masks and respirators

One of the most common misconceptions is that a standard surgical mask can provide protection against *M. tuberculosis* infectious aerosols. Surgical masks made from poorly fitting porous material leave large gaps between the face and mask and therefore only help to prevent the spread of microorganisms from the wearer to others by capturing the large wet particles in the exhaled air.

Although biological safety cabinets and airflow in the laboratory are the main means of protection against exposure to contaminated aerosols generated during culture and drug susceptibility testing activities, the need for additional personal protection must be considered in certain settings, such as when MDR-TB and/or HIV are prevalent. Staff may be HIV-infected and highly susceptible to contaminated aerosols [14].

Protection from inhalation of infectious aerosols can be provided by respirators, which are devices with the capacity to filter particles of 0.3–0.4 µm diameter and fit closely to the face to prevent leakage around the edges. The N95 (FFP2) respirator is a lightweight, disposable nose and mouth respirator; it effectively filters out more than 95% of particles of diameter 0.3 µm and above. The FFP3 respirator removes more than 98% of such particles. Each user should be instructed in the proper use of the respirator and informed about its limitations. Respirators should be correctly fitted to the face to prevent leakage from around the face seal. This is done by placing the mask over the nose and mouth with the top elastic band over the crown of the head and the bottom elastic band over the back of the neck. The metal strip covering the nose should be firmly moulded over the bridge of the nose. Facial hair between the wearer's skin and the sealing surfaces of the respirator will prevent a good seal. Respirators should also be worn during emergency cleaning of spillages involving the release of viable organisms into the work area. Respirators should be stored in a convenient, clean, and sanitary location and discarded after eight hours of (cumulative) use and not be kept for more than one week [7].

Gloves

In accordance with international, universal procedures/guidelines, appropriate gloves should be worn for all procedures that involve the handling of body fluids. Gloves must be worn in case of hand injury/skin disease or when the risk of exposure to blood-borne pathogens is high; consequently, specimens resulting from invasive clinical investigation must be handled with gloves [12,14].

Gloves must be changed after every session that requires their use and after every interruption of the activity. Never wear gloves outside the laboratory. Every time hands are removed from the biological safety cabinet, gloves must be pulled off and discarded in a waste container in the safety cabinet [7,12,14].

Disposable latex, latex-free vinyl (clear) or nitrile gloves can be used and the correct size (small, medium, or large) should be available for all individuals. Hypoallergenic gloves should be provided in case of allergy to latex proteins and/or to the corn-starch used for powder. Re-using single-use gloves is not advised. Used gloves should be

discarded as contaminated material. Following the safe removal of gloves, wash hands immediately with water and liquid soap. Proper hand-washing with soap and adequate care in the handling of contaminated materials are critical elements of safe laboratory practice [14].

Gowns

Always wear a gown inside the laboratory (never outside) and change at least weekly. Long-sleeved back-opening gowns or overalls with narrow cuffs give better protection than laboratory coats and are preferred in microbiology laboratories. When worn, laboratory coats should be fully buttoned. An area of the laboratory must be designated for storage of used and new clothing. Laboratory gowns must be stored away from personnel clothing. Laundering services should be provided at/near the facility. Extra clothing should be available suitable for visitors, maintenance and emergency response personnel [7,12,14].

Always remove personal protective equipment in the following order:

- disposable gloves;
- gown/coat/suit/overalls; and
- respirator/mask.

5.3 Specimen collection, storage and transport

5.3.1 Sample collection

Proper specimen collection procedures and containers, adequate specimen volumes and appropriate transport conditions can all affect TB culture results. Correct labelling of specimens is critical. This includes patient and sample identification, sample type and date of collection.

As a general rule, it is preferable that specimens are collected before starting specific treatments. Specimens should always be collected with care to avoid contamination by host or environmental microorganisms and submitted in sterile, leak-proof, disposable, appropriately labelled, laboratory-approved containers without any fixatives.

If centrifugation is used for culture tests, the use of collection containers suitable for centrifugation should be considered. Decontamination and centrifugation in the collection container avoids having to transfer samples to another container [1-8].

Sputum samples

Most specimens received by the laboratory are sputum samples. Patients should be clearly instructed on how to collect the sputum specimen; written instructions must be provided.

A systematic review of 37 eligible studies [6] clearly showed that most TB cases (average 85.8%) were detected with the first sputum specimen. With the second sputum specimen, the average incremental yield was 11.9%; with the third specimen (when the first two were negative) the incremental yield was 3.1%. It is expected that laboratory analysis of two sputum smear samples will improve case-finding, reduce time to diagnosis, accelerate initiation of treatment and decrease the number of patients lost during the diagnostic process. Based on this evidence, WHO has recommended that two sputum samples in a single day be used to diagnose pulmonary TB in settings where a well-functioning EQA system is in place, the workload is high and human resources are limited.

A good sputum specimen should be approximately 3–5 ml of recently-discharged material from the bronchial tree. It is usually thick and mucoid. It may be fluid and contain pieces of purulent material. The colour may vary from opaque white to green. Bloody specimens will appear reddish or brown. Clear saliva or nasal discharge is not suitable as a TB specimen, although saliva should not automatically be rejected: induced and follow-up sputa resemble saliva. To avoid contamination or dilution of a good sample, specimens should not be pooled.

Other specimens

Body fluids (spinal, pleural, pericardial, synovial, ascitic, blood, pus, and bone marrow) should be aseptically collected in sterile containers using aspiration techniques or surgical procedures. Pleural effusion is a suboptimal specimen: tubercle bacilli are mainly in the pleural wall and not in the fluid. The minimum volume for pleural effusion is 20–50 ml. A pleural biopsy specimen is ideal.

For fluids that may clot, sterile potassium oxalate (0.01–0.02 ml of 10% neutral oxalate per ml fluid), heparin (0.2 mg/ml), or sodium citrate (two drops of 20% sodium citrate for every 10 ml of fluid) should be added as an anticoagulant to the culture.

Aseptically collected tissues should be placed in sterile containers without fixatives or preservatives and transported quickly to the laboratory. For prolonged transportation, dehydration should be prevented by adding sterile saline and maintaining a temperature of 4–15°C.

Urine is expected to be contaminated. To minimise excessive contamination of urine specimens, external genitalia should be washed before specimen collection. Once received in the laboratory, a urine sample must either be

processed immediately or centrifuged and the pellet refrigerated. As excretion of tubercle bacilli is intermittent, three consecutive early-morning midstream specimens must be collected.

Other respiratory specimens that can be submitted to the laboratory for mycobacterial culture are bronchial secretion (minimum volume 2–5 ml) and bronchial alveolar lavage samples (minimum volume 20–50 ml). Transbronchial and other biopsies taken under sterile conditions should be kept wet during transportation by adding 0.5–1 ml sterile 0.9% saline.

In children who produce little, if any sputum, aspiration of the early-morning gastric juice can be used for TB diagnosis. The gastric aspirate should be transported immediately to the laboratory and neutralised by adding 100 mg of sodium bicarbonate.

5.3.2 Storage of specimens

Specimens should be correctly collected and delivered as quickly as possible to the laboratory. Every effort must be made to organise and expedite specimen transportation and processing. Although TB bacilli can survive in sputum for one week in the absence of preservatives, the probability of successfully culturing the bacilli decreases with time and this is especially critical for paucibacillary specimens. If specimens cannot be transported to the laboratory within one hour, it is recommended to store them at 4°C. This does not apply to whole blood specimens, which are not to be refrigerated. On arrival at the laboratory, specimens should again be refrigerated until they can be processed. The delay between collection and inoculation should not exceed seven days.

5.3.3 Transportation of specimens

Packaging of infected specimens that are to be sent by surface or air mail must be carried out according to national biosafety and biosecurity guidelines or international rules. For international transfer of infectious substances, the International Air Transport Association (IATA) should be contacted [7].

Specimens and cultures should be packaged in a three-component packaging consisting of:

- a leak-proof primary receptacle(s);
- a leak-proof secondary packaging; and
- an outer packaging of adequate strength for its capacity, mass and intended use.

For the purposes of transport, infectious substances are defined as substances which are known or reasonably expected to contain pathogens [7,8]. Category A (UN2814) corresponds to an infectious substance which is transported in a form that, when exposure to it occurs, is capable of causing permanent disability, and/or life-threatening or fatal disease in otherwise healthy humans or animals. All other infectious substances as well as human biological specimens belong to Category B (UN3373).

Cultures of *M. tuberculosis* belong to Category A. However, for surface transport, when *M. tuberculosis* cultures are intended for diagnostic or clinical purposes, they may be classified as category B. For surface transport there is no maximum quantity per package.

For air transport:

- no primary receptacle should exceed 1 l (for liquids) or the outer packaging mass limit (for solids); and
- the volume shipped per package should not exceed 4 l or 4 kg.

These quantities exclude ice, dry ice or liquid nitrogen when used to keep specimens cold.

5.4 Homogenisation and decontamination of specimens

Most (but not all) specimens are considered contaminated. Pulmonary specimens including sputum, bronchial secretions, bronchoalveolar lavage, bronchial aspirates and brushings are usually contaminated by normal host microbiota. Extrapulmonary specimens may be divided into two main groups according to the extent of contamination:

- Aseptically collected specimens, usually free from other microorganisms (sterile);
- Specimens contaminated by normal flora or specimens not collected aseptically (not sterile).

Normally, contaminated extrapulmonary specimens are gastric lavage, laryngeal aspirates, urine, skin, autopsy materials, and uterine mucosa. Sterile specimens include pus from cold abscess, CSF, synovial or other cavity body fluids, as well as surgical biopsies.

Contaminated specimens must be subjected to rigorous decontamination procedures that liquefy the organic debris and eliminate the unwanted normal flora. Normal flora would rapidly overgrow the entire surface of the medium and consume it before the TB bacilli started to grow. Specimens must be homogenised to free the bacilli from the mucus, cells or tissue in which they may be embedded.

Digesting/decontaminating agents are to some extent toxic to tubercle bacilli and therefore to minimise the number of dead mycobacteria, the digestion/decontamination procedure must be followed precisely. A proportion of cultures will be contaminated by other organisms: a contamination rate of 3–5% is acceptable on solid media. Cultures in liquid media may show higher contamination rates (5–10%). Furthermore, if specimens (especially sputum) take several days to reach the laboratory, the contamination rate may be higher. At present, new commercial kits are available, containing transportation media supplemented with decontamination solutions for longer shipments between countries or cities. These may help to reduce the proportion of contaminated cultures due to long shipments and increase the proportion of positive cultures among those samples which are not inoculated at the point of collection. A contamination rate that approaches 0 indicates that the decontamination procedure was too harsh.

5.4.1 Digestion and decontamination of sputum samples

Digestion and decontamination using the sodium hydroxide (modified Petroff) method

Sodium hydroxide is toxic, both for contaminants and for tubercle bacilli; strict adherence to the indicated timings is therefore essential. This decontamination procedure can only be used for samples which will then be inoculated on solid media.

Reagents:

- Sodium hydroxide (NaOH) solution, 4%;
- Phosphate buffer 0.067 mol/l, pH 6.8.

Sodium hydroxide (NaOH) solution, 4%:

- Sodium hydroxide pellets (analytical grade): 4 g;
- Distilled water: 100 ml.

Dissolve NaOH in the distilled water. Aliquot in 2 ml quantities. Sterilise by autoclaving at 121°C for 20 minutes.

Phosphate buffer, 0.067 mol/l, pH 6.8:

- Stock solution A: disodium phosphate, 0.067 mol/l.

Dissolve 9.47 g of anhydrous Na₂HPO₄ in 1 l of distilled water.

- Stock solution B: monopotassium phosphate, 0.067 mol/l.

Dissolve 9.07 g of KH₂PO₄ in 1 l of distilled water.

Mix 50 ml of solution A and 50 ml of solution B. Use a pH meter to confirm that the correct pH for the buffer is reached. Adjust as necessary, using 10% phosphoric acid or 10% sodium hydroxide.

Aliquot in the volumes required for adding to a single centrifugation tube (e.g. 50 ml amounts if 50 ml centrifuge tubes are used), discarding the extra volume. Sterilise by autoclaving at 121°C for 20 minutes. Leftover volumes of buffer can then be pooled and sterilised again for further use.

Procedure:

- If sputum has not already been collected in centrifuge tubes, select sterile plastic screw-top centrifuge tubes (one for each specimen).
- With a permanent marker, write the number of the specimen on the wall of the tube (not on the cap).
- Write the number of each specimen and the inoculation date on two tubes of media.
- Mark the volume of sputum on the centrifuge tube (at least 2 ml, not more than 5 ml). Add an equal volume of 4% NaOH and tighten the screw-cap.
- Vortex to digest.
- Allow to stand for 15 minutes at room temperature.
- Fill the tube to within 2 cm of the top (e.g. to the 50 ml mark on the tube) with phosphate buffer.
- Centrifuge at 3 000 g for 15 minutes.
- Carefully pour off the supernatant through a funnel into a discard can containing 5% phenol or another mycobacterial disinfectant.
- Re-suspend the deposit in approximately 0.3 ml phosphate buffer.
- Inoculate the deposit on two slopes of egg-based medium labelled with the ID number. Use a pipette to inoculate each slope with 3–4 drops (approximately 0.1–0.15 ml).
- Smear one drop on a slide, marked with the ID number, for microscopic examination.

Digestion and decontamination using the N-acetyl-L-cysteine-sodium hydroxide (NALC-NaOH) method

The most widely used method for the digestion and decontamination of contaminated specimens is the N-acetyl-L-cysteine-sodium hydroxide (NALC-NaOH) method.

Decontamination using NALC-NaOH is based on the mucolytic properties of N-acetyl-L-cysteine (NALC) which enable the decontaminating agent, sodium hydroxide, to be used effectively at a low, final concentration.

Consequently, the NALC method results in more positive cultures than other methods, as it only kills about 30% of the tubercle bacilli in clinical specimens; a lower NaOH concentration means that contamination rates may be higher than for other decontamination methods. The time needed to process a single specimen is approximately 40 minutes, while 20 specimens would take approximately 60 minutes.

This method is suitable for cultures on both solid and liquid media. However, the disadvantages of the method are that NALC loses activity and must therefore be made fresh every day. Commercially prepared solutions are available, but expensive.

After exposure to the decontaminant and subsequent centrifugation, it is essential that the sediment is re-suspended in a 1:10 dilution of buffer (or water) to reduce the concentration of any toxic components that may inhibit the growth of TB bacilli.

As a measure of precaution, an aliquot of the sediments should be kept for one week in the refrigerator and re-decontaminated if the inoculated cultures show signs of contamination. Optionally the sediment can be frozen (- 20°C) in a screw-cap sterile 1.5-2 ml vial that is properly labelled.

Reagents N-acetyl-L-cysteine-sodium hydroxide (NALC-NaOH) method:

Sodium hydroxide-citrate solution

Solution A: Sodium hydroxide 4%	
Sodium hydroxide pellets (analytical grade)	40 g
Distilled water	1 000 ml

Dissolve NaOH in the distilled water.

Solution B: Trisodium citrate 3H₂O 2.94%	
Trisodium citrate 3H ₂ O	29.4 g
Distilled water	1 000 ml

Dissolve trisodium citrate 3H₂O in the distilled water.

Mix solutions A and B, aliquot in 100 ml quantities, and sterilise by autoclaving at 121°C for 15 minutes. Store at 4°C in refrigerator.

N-acetyl-L-cysteine (NALC)

NALC-NaOH solution should be freshly prepared for daily use only.

Prepare by adding 0.5g NALC to 100 ml of the sodium hydroxide-citrate solution just before use: aliquot in 4 ml amounts.

Phosphate buffer, 0.067 mol/l, pH 6.8

See above for preparation.

Sputum processing

Sputum specimens are not to be pooled because of the risk of cross-contamination. Always digest/decontaminate the whole specimen – do not attempt to select portions of the specimen as is done for direct microscopy. Gently decant from the specimen container into the centrifuge tube. If the specimen is too viscous to pour, an equal volume of digestant/decontaminant can be added to the sputum in the specimen container before the mixture is poured carefully into an appropriate screw-top centrifuge tube.

Sputa are not to be processed in batches of more than 6–8 as the method is strictly time-dependent. Procedure:

- If sputum has not already been collected in centrifuge tubes, select sterile plastic screw-top centrifuge tubes (one for each specimen).
- With a permanent marker, write the number of the specimen on the wall of the tube (not the cap).
- Write the number of each specimen and the inoculation date on two tubes of media.
- Transfer the sputum (at least 2 ml, but no more than 5 ml) into a centrifuge tube. Add an equal volume of NALC-NaOH solution.
- Tighten the cap of the tube and shake or vortex. Mix for no more than 20 seconds.
- Keep at 20–25°C for 15 minutes to decontaminate.
- Fill the tube to within 2 cm of the top (e.g. the 50 ml mark on the tube) with 0.067 mol/l phosphate buffer (pH 6.8) or distilled water. Vortex mix.
- Centrifuge at 3 000g for 15 minutes.
- Carefully pour off the supernatant into a discard bottle containing the appropriate disinfectant.
- Re-suspend the deposit and inoculate onto two slopes of LJ medium (and one slope of LJ with pyruvate if needed) or into liquid medium. Using a pipette (not a loop), inoculate each slope with 3–4 drops (0.2–0.4 ml). Smear one drop on a slide (marked with the ID number) for microscopic examination.

5.4.2 Digestion and decontamination of specimens other than sputum

Laryngeal swabs

Smear examination is not done for laryngeal swabs. Swabs yield little material: as much of the material as possible must be collected and not wasted.

- Swabs must be cultured on the day they are received using sterile precautions.
- Use sterile forceps to transfer the swab to a sterile centrifuge tube.
- Add 2 ml of sterile distilled water.
- Decontaminate according to NaOH-NALC method (see above, Section 5.4.1).
- Before adding the phosphate buffer solution, remove the swab from the tube using sterile forceps.
- Fill the tube to within 2 cm of the top (e.g. the 50 ml mark on the tube) with phosphate buffer, 0.067 mol/l, pH 6.8 and mix the contents by inversion.
- Centrifuge at 3 000g for 15 minutes.
- Carefully pour off the supernatant into a discard bottle containing an appropriate disinfectant.
- Inoculate the deposit on two slopes of LJ medium (and one slope of LJ with pyruvate, if needed) or in liquid medium. Using a pipette (not a loop), inoculate each slope with 3–4 drops.

Gastric lavages

Gastric lavage specimens should be processed as soon as possible after collection; acidity can kill mycobacteria in the specimen so gastric lavage specimens must be processed within four hours. The gastric aspirate should be collected in a tube containing 100 mg of sodium bicarbonate for neutralisation and should be transported immediately to the laboratory. Proceed as for sputum.

If the specimen is watery, centrifuge at 3 000g for 15 minutes, pour off the supernatant, re-suspend the sediment in 5 ml of sterile distilled water and proceed as for sputum.

Mucopurulent materials

Handle as for sputum when the volume is 10 ml or less.

Handle as for mucoid gastric lavage when the volume is more than 10 ml.

Fluid materials

If the specimen has been collected aseptically, centrifuge and inoculate the sediment directly onto culture media, preferably liquid medium.

Materials that should not be decontaminated are:

- spinal, synovial or other cavitory body fluids;
- bone marrow;
- pus from cold abscesses;
- surgically resected specimens (excluding autopsy material); and
- material obtained from pleural, liver and lymph nodes as well as biopsies (if not fistulised).

To maximise the recovery rate, the entire CSF volume (or other small volume of aseptically collected fluid) should be cultured, preferably in liquid medium.

If the specimen was not aseptically collected:

- Handle as for sputum when the volume is 10 ml or less;
- Handle as for fluid gastric lavage when the volume exceeds 10 ml.

Tissue

If a biopsy needs to be processed for smear and culture, it is necessary to homogenise the biopsy in a sterile porcelain mortar or preferably in a small, non-reusable tissue grinder with 2–5 ml of sterile saline.

Mortars, pestles, and tissue grinders must be cleaned and sterilised thoroughly to prevent false-positive results or contamination due to organisms left over from previous specimens. Lymph nodes, biopsies and other surgically resected tissue should be cut into small pieces with a sterile scalpel or scissors. Homogenise the specimen in a sterile porcelain mortar or tissue grinder using 5 ml sterile saline and a small quantity of sterilised sand. Inoculate the suspension onto culture media.

5.5. Culture media: principles

As *M. tuberculosis* grows slowly, with a generation time of 18–24 hours (other bacteria reproduce within minutes), usual bacteriology techniques are not applicable to mycobacterial cultures. Moreover, growth requirements are such that *M. tuberculosis* will not grow in primary isolation on simple, chemically-defined media. The only media that allows for abundant growth are egg-enriched media containing glycerol and asparagine, and agar or liquid media supplemented with serum or bovine albumin. Many different media have been developed for *M. tuberculosis* growth and are generally classified into two main groups: solid media (egg- and agar-based) and liquid media.

Antibiotics can be added to culture media in order to prevent the growth of non-specific flora.

Both solid and liquid media are recommended for *M. tuberculosis* isolation from biological samples. An advantage of solid over liquid media is that colonies of mixed cultures and contaminants can be observed while liquid media promotes a faster growth of mycobacteria.

The choice of media depends primarily on the type of specimen. Non-selective media are recommended for use with samples from normally sterile sites (bone marrow, tissue biopsy samples, cerebrospinal fluid and other body fluids etc.), while selective media, that contain antimicrobial agents to prevent growth by contaminating bacteria and fungi, are recommended for use with contaminated (or potentially contaminated) specimens (sputum, abscess contents, bronchial washings, gastric lavage fluid, urine, etc.) [1,2].

The most commonly used non-selective media are:

- egg-based media: Löwenstein-Jensen (LJ) medium and Ogawa medium
- agar-based media: Middlebrook 7H10 and Middlebrook 7H11; and
- liquid media: Middlebrook 7H9 broth.

Other commonly used selective media available in some countries are:

- Egg-based media: Gruft modification of LJ (containing malachite green, penicillin and nalidixic acid as selective agents, and Mycobactosel LJ (containing malachite green, cycloheximide, lincomycin and nalidixic acid as selective agents);
- Agar-based media: selective 7H11 (Mitchison's medium), containing carbenicillin, amphotericin B, polymixin B and trimethoprim as selective agents; and
- Liquid media: in general they contain a modified Middlebrook 7H9 broth plus a mixture of antimicrobial agents.

Several automated systems have been commercially developed for rapid detection of mycobacteria in liquid medium:

- BACTEC µGIT 960 system (BD [Becton, Dickinson and Company] Diagnostic Systems);
- ESP Culture System II (Trek Diagnostic Systems);
- MB/BacT (bioMérieux).

5.6 Solid media

5.6.1 Egg-based media

LJ medium, which contains malachite green as an inhibitor of non-mycobacterial organism, is the most commonly used egg-based medium, especially for sputum culture. LJ is user-prepared or commercially prepared in slant tubes. LJ containing glycerol favours *M. tuberculosis* growth, while LJ without glycerol but containing sodium pyruvate enhances *M. bovis* growth. Both media should be used in geographical regions where patients may be infected with either organisms [3]. Ogawa medium is LJ without asparagine. Non-selective egg-based media can be stored in the refrigerator for several months provided that the tube caps are tightly closed to minimise evaporation.

A disadvantage of egg-based media is that when contamination does occur it may involve the entire slant surface, so the culture is generally lost. If specimens contain few bacilli it may take three to eight weeks before cultures become positive.

5.6.2 Agar-based media

These media are prepared in slant tubes or plates and are less likely than egg-based media to become contaminated. Middlebrook 7H10 and 7H11 media are usually prepared in the laboratory from commercially available agar-powdered bases, with the addition of Middlebrook oleic acid-albumin-dextrose-catalase (OADC) enrichment. Because of the transparency of 7H10 and 7H11 plates, *M. tuberculosis* micro colonies with typical cord formation can be detected and counted using a microscope as early as one week after incubation. Moreover, visibility of colonial morphology on agar plates is better than on egg-containing slants, aiding the identification of mycobacteria. Middlebrook 7H11 is preferable to 7H10 because it contains 0.1% casein hydrolysate, a substance favouring the recovery of isoniazid-resistant mycobacteria. Furthermore, 7H11 is also better for growing multi-drug-resistant (MDR) strains as these may not grow at all on 7H10 agar plates.

A disadvantage of Middlebrook media is that the surface dries more rapidly than egg-based media. It is important to know that daylight, heating, and storage at 4°C for more than four weeks may cause the release of formaldehyde in a sufficient concentration to inhibit the growth of mycobacteria.

5.6.3 Liquid media

Liquid media offer a considerable time advantage over solid media: 7–14 days in Middlebrook 7H9 liquid medium, compared with 18–28 days in Middlebrook 7H11 agar, or 21–42 days in LJ medium [5].

One of the most widely used automated systems for rapid detection of mycobacteria in liquid medium is the BACTEC µGIT 960 system [2]. The system's culture tubes consist of modified Middlebrook 7H9 broth, a growth supplement, and an antimicrobial agent mixture. A similar principle is used in the ESP Culture System II and the MB/BacT system. In the BACTEC 960 system and ESP Culture System II, *M. tuberculosis* growth is detected by the rate of oxygen consumption within the headspace of the cultures; in the MB/BacT system, a colorimetric sensor detects the production of CO₂ dissolved in the culture medium.

5.6.4 Quality control of media

Quality control of the media is needed to ensure that the strain isolated from a specimen is from the patient and not a contaminant present in the ingredients of the medium. The description below mainly applies to quality control of solid media, as it is on such media that colonies are visible to the eye and species identification is therefore possible.

Commercially prepared media do not need to be quality controlled for sterility, growth and selectivity, provided that documentation of the manufacturer's quality control procedures is obtained [12]. The information should include the preparation date, the lot number, the expiration date, the test organisms used, the date of testing and the result. In all other cases (user-prepared media and when documentation of sterility, growth and selectivity is not provided), the media must be checked for:

- medium conditions: colour, dehydration, contamination, bubbles;
- sterility: incubating from 1–3% of each batch at 35–37°C in 5 to 10% CO₂ for up to 21 days;
- performance: by testing growth of positive and negative control strains.

The organisms used as positive controls are *M. tuberculosis* H37Ra (ATCC 25177), *M. kansasii* ATCC 12478, *M. scrofulaceum* ATCC 19981, *M. intracellulare* ATCC 13950 and *M. fortuitum* ATCC 2841. *Escherichia coli* ATCC 25922 is used to demonstrate partial inhibition by non-selective media and complete inhibition by selective media.

Procedure [1,2]:

- Prepare a 0.5 McFarland suspension of the organisms in 7H9 broth.
- Inoculate media with 10µl of the control suspension using a pipette or a calibrated loop. To test selective properties of the media, inoculate them with 10µl of 1:10 suspension in sterile 0.85% NaCl. Incubate all media at 35–37°C in 5–10% CO₂ for up to 21 days.

Expected results are as follows:

Positive controls	Result
<i>M. tuberculosis</i> ATCC 25177	Growth on all media
<i>M. kansasii</i> ATCC 12478	Growth on all media
<i>M. scrofulaceum</i> ATCC 19981	Growth on all media
<i>M. fortuitum</i> ATCC 2841	Growth on all media
<i>M. intracellulare</i> ATCC 13950	Growth on all media (not included when testing selective media containing penicillin or carbenicillin)
Negative control	Result
<i>Escherichia coli</i> ATCC 25922	Partial inhibition in non-selective media, total inhibition in selective media

5.7 Culture tube inoculation

5.7.1 Solid media

In media that is purchased ready-to-use, condensed moisture is frequently observed on the culture slants and it is advisable to remove it before use. Each slant should be inoculated with 0.2–0.4 ml (2–4 drops or four loopfuls) of the centrifuged sediment. The use of sterile disposable Pasteur pipettes is highly recommended. The inoculum should be distributed over the entire surface of the slant.

At least two slopes of LJ medium per specimen should be inoculated with 0.2 ml of each sediment. In areas where *M. bovis* is isolated, an additional slope containing pyruvate is recommended. Using a ready-made commercially available egg-based media supplemented with antibiotic mixture may help to reduce contamination rates. Too little inoculum is a common cause of false-negative results. In the upper part of the slant the medium is thin and dehydrates readily; if mycobacteria are seeded only on this upper section, they might not grow, again leading to false-negative results.

5.7.2 Liquid media

Inoculation on liquid media should be performed under rigorous sterile conditions to avoid the risk of contamination. Liquid media is more susceptible to contamination than solid media and therefore needs to be supplemented with a mixture of specific antibiotics to kill the contaminants. These antibiotic mixtures are available from commercial companies selling culture media for automated culture systems.

Each properly labelled liquid culture tube should be inoculated with 0.5 ml of sediment and the sediment must be deposited under the surface of the medium, keeping the tube tilted at an angle of 45°. The tube is then returned to a vertical position, leaving the inoculum below the surface of the liquid.

5.8 Culture incubation

All cultures should be incubated at 35–37°C. Always check the temperature indicator before incubating the cultures. The cultures should be incubated until growth is observed, or discarded as negative after eight to 12 weeks (six weeks if liquid media is used).

Inoculated solid cultures should be incubated with caps loosened in a slanted position for at least one week to ensure an even distribution of the inoculum. Caps should then be tightened to prevent desiccation of the media and, if space is needed in the incubator, the tubes can be placed upright. Tops should be tightened to minimise evaporation which can result in the media drying out.

5.9 Culture examination

All cultures should be examined 48 hours after inoculation in order to:

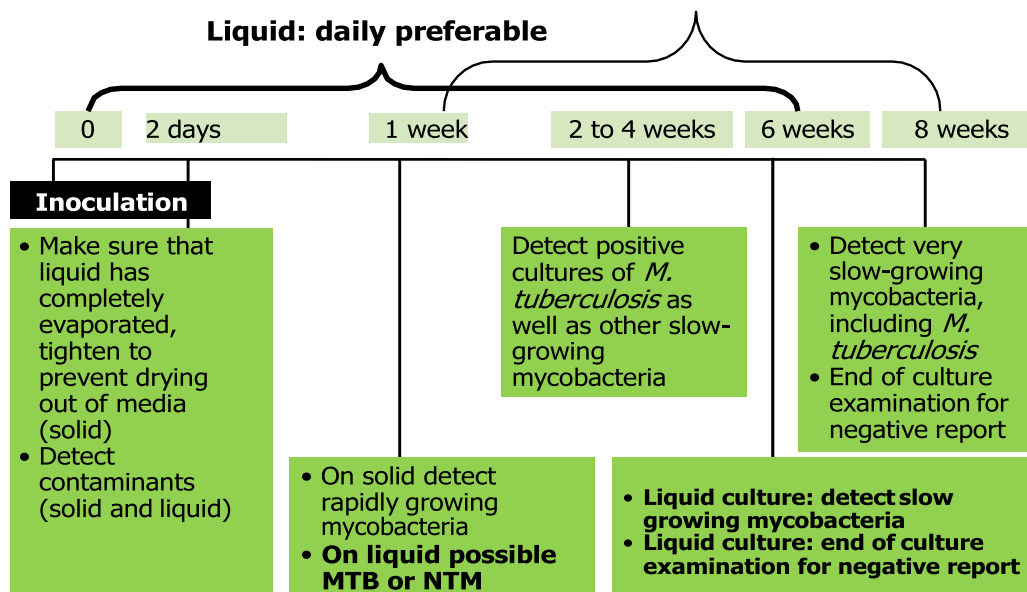
- check absorption of inoculated liquid;
- tighten caps to prevent drying out of media; and
- detect early contaminants.

Cultures should then be examined on a weekly basis or, if this is not feasible, at least three times during the eight-week incubation period (Figure 7).

- Seven-day check: To detect rapidly growing mycobacteria.
- Three-to-four-week check to detect positive cultures of *M. tuberculosis* as well as other slow-growing mycobacteria.
- End of culture check (after eight weeks) to detect very slow-growing mycobacteria, including *M. tuberculosis*, before discarding and reporting the culture as negative.

Figure 7. Minimal examination schedule for solid cultures

(solid: weekly preferable)



Source: Culture, DST and quality assurance package, WHO [14]

The different kinds of contaminants that should be considered are non-tuberculous mycobacteria, fungi, bacteria and yeasts.

After ZN staining, the culture should be handled according to the results:

- Presence of AFBs only in the deposit with no non-AFBs indicates pure growth of mycobacteria – the deposit should be processed for identification and drug susceptibility testing (inoculation of a non-selective agar plate, such as blood agar, can be used to check for purity).
- Presence of AFBs with non-AFBs in the deposit indicates contamination of the possible growth of mycobacteria – the deposit should be processed for decontamination and culture on solid media.
- No AFBs and only non-AFBs in the deposit indicate growth of contaminants – the deposit should be discarded.
- Any presence of contaminants should be recorded in the laboratory register and if the culture is discarded, it should be reported as a 'contaminated culture'.
- Evaluation of the contamination rate should be performed every three to six months for quality assurance purposes. A contamination rate of 3–5% is considered a good balance between the need to kill contaminating bacteria and the need to keep the majority of tubercular mycobacteria present in the sample. A contamination rate of 0–1% may indicate too strong a decontamination process. The contamination rate should refer to the number of contaminated tubes, not to the number of registered specimens.

Common contaminants are detailed below [4]:

Non-tuberculous mycobacteria:

- Fast- or slow-growers;
- Acid-fast bacilli;
- Not usually arranged in cords.

Fungi:

- Usually slow-growers;
- Non-acid-fast;
- Hyphae are thicker than mycobacteria.

Bacteria:

- Usually non-acid-fast except for some closely related genera (*Gordonia*, *Tsukamurella*, *Nocardia*, *Rhodococcus*, *Dietzia*) and *Legionella micdadei*.

Yeast:

- Usually non-acid-fast.

Oocystis:

- Usually non-acid-fast except for *Cryptosporidium*, *Isospora*, *Cyclospora*.

5.10 Chromatographic immunoassay for the qualitative detection of *Mycobacterium tuberculosis* complex from cultures

Assays used: Capilia TB-neo, TAUNS Laboratories Co, Shizuoka, Japan, BD μ GIT TBc Identification Test, BD (Becton, Dickinson and Company) Diagnostic Systems, Sparks, MD, USA, and SD BIOLINE TB Ag MPT64 Rapid test, Abbott, Chicago, IL, USA.

5.10.1 Introduction

Definite diagnosis of TB can be made by identifying *M. tuberculosis* complex organisms from a clinical sample after growth in solid or liquid media. Since *M. tuberculosis* complex strains (with the exception of some sub-strains of *M. bovis* BCG) but not non-tuberculous mycobacteria specifically and predominantly secrete the MPB64 protein (mycobacterial protein fraction from BCG of Rm 0.64), this can be used to discriminate between *M. tuberculosis* complex and non-tuberculous mycobacteria. Immunochromatographic assays based on the reaction of monoclonal antibodies against MPB64 have been developed and evaluated [15-17].

5.10.2 Materials

No special equipment is required for the test; it is sufficient to use the test provided by the manufacturer, a 100 μ l pipette, and a timer. The test consists of a sample placing area, a testing area containing the anti-MPB64 antibodies, and a control area where anti-species immunoglobulin antibodies are fixed.

5.10.3 Methods

The testing method is based on immunochromatographic principles, in which antibodies labelled with colloidal particles (such as colloidal gold) react with target antigens to form a migrating antigen-antibody complex, which is captured by a second fixed antibody. A colour reaction takes place where the labelled particles are fixed.

The tests can be used with positive liquid media tubes or visible colonies grown on solid media. In the case of liquid cultures, a 100 μ l volume is dropped on to the test device. For solid cultures, 1 μ l bacteria (=1 mm loop) or 1 AFB+ colony (at least 1 mm) are re-suspended in the respective buffer, then vortexed and a 100 μ l volume of these suspensions is used. The results should be read after 15 minutes but within 60 minutes of contact.

5.10.4 Results interpretation

For a specific MPB64 antigen-antibody reaction, a red-purple colour band becomes visible within 15 minutes. The culture is interpreted as positive for *M. tuberculosis* complex if the colour reaction takes place in the test and control area. The intensities of the bands may vary. The specimen is interpreted as negative if a colour reaction takes place only in the control area. The test is invalid if no band is visible in the control area or if the background colour inhibits the test interpretation. Figure 8 gives examples of chromatographic immunoassays.

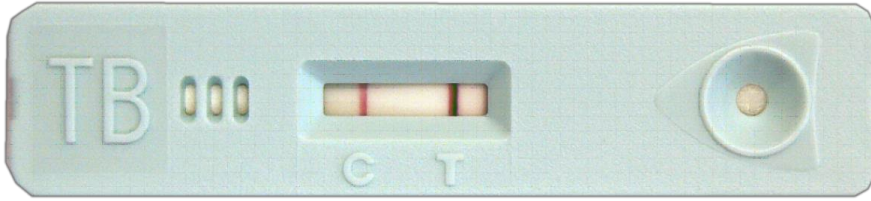
Although most *M. tuberculosis* complex strains may be correctly identified with the tests exhibiting a high sensitivity (92.4–99.2%) and specificity (100%), some test-negative strains have been isolated [16,17]. In some instances, the test misses the detection of *M. tuberculosis* (false negative result) as a result of mutations in the *mpb64* gene and consequent lack of secretion of the MPB64 protein in the culture media [17].

5.10.5 Biosafety

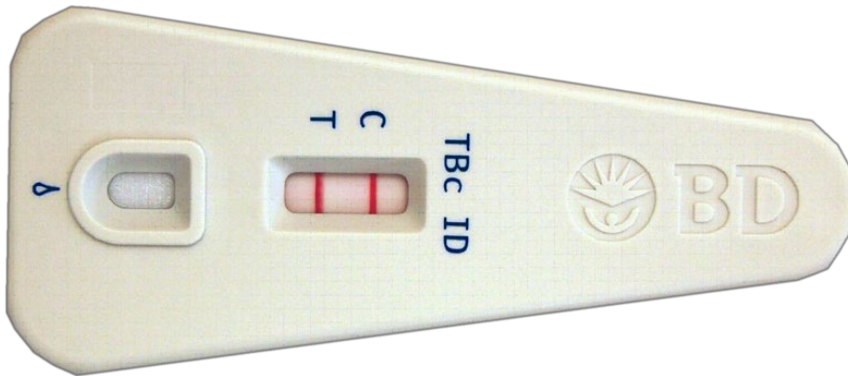
Appropriate biosafety precautions for handling mycobacteria must be used. The dropping procedure, development, and reading of the test should be carried out in an appropriate biological safety cabinet in a BSL 3 laboratory. As used test devices may contain viable mycobacteria, they should be discarded safely according to institutional guidelines for handling BSL-3 material (see Chapter 1).

Figure 8. Examples of chromatographic immunoassays for qualitative detection of *M. tuberculosis* complex

Capilia TB-neo



BD MGIT™ TBc Identification Test (TBc ID)



Developed with 100 µl of an AFB+ smear-positive liquid culture after 15 minutes incubation time.

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6 Molecular assays for TB and drug-resistant TB rapid detection

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Revised by Elisa Tagliani and Doris Hillemann (2022)

6.1 Background and principles

Over the past decade, molecular biology has gained a central role in the diagnosis of TB thanks to the development of new molecular tests, often referred to as nucleic acid amplification tests (NAATs). These assays rely on the amplification of targeted genes of *M. tuberculosis* complex enabling faster detection of TB and drug-resistant TB than conventional/culture-based methods. In addition, several NAATs are suitable to be performed at the more devolved levels of the health care system thus contributing to improving accessibility and quality of TB care.

This chapter provides an overview of the most commonly used molecular diagnostics for the rapid detection of TB and drug-resistant TB from culture and clinical specimens. In addition, it includes a description of the current applications of next-generation sequencing (NGS) for drug susceptibility and resistance prediction starting from clinical specimens (i.e. targeted NGS) and culture isolates (i.e. whole genome sequencing).

6.2 Line probe assays

Line probe assays (LPAs) are a family of tests based on reverse-hybridisation DNA•STRIP technology. The procedure involves three steps: (i) DNA extraction from the decontaminated pulmonary specimens or cultured material (solid or liquid media); (ii) multiplex amplification with biotinylated primers; and (iii) reverse hybridisation.

The assay membrane strips are coated with specific probes complementary to the amplified nucleic acids. After chemical denaturation, the single-stranded amplicons bind to the probes (hybridisation). Highly specific binding of complementary DNA strands is ensured by stringent conditions thus allowing the probes to reliably discriminate several sequence variations in the gene regions examined. The streptavidin-conjugated alkaline phosphatase binds to the amplicons' biotin via the streptavidin group. Finally, the alkaline phosphatase transforms an added substrate into a dye which becomes visible on the membrane strips as a colored precipitate.

6.2.1 Line probe assays for *Mycobacterium* genus species identification

Several test options are available for the detection and differentiation of mycobacteria directly from clinical specimens (i.e. GenoType cMdirect) and cultivated samples (e.g. GenoType Mycobacterium CM / AS, GenoType NTM-DR) (Table 7).

Table 7. Types and specifics of LPAs used for detection and differentiation of *Mycobacterium*

Differentiation	
GenoType MTBC	Detection of <i>M. tuberculosis</i> complex from cultures
GenoType cMdirect	Detection of <i>M. tuberculosis</i> and 20 clinically relevant NTM from patient specimens
GenoType Mycobacterium CM	Detection of <i>M. tuberculosis</i> complex and more than 20 clinically relevant NTM from cultures
GenoType Mycobacterium AS	Detection of 19 additional NTM from cultures
Differentiation and drug susceptibility testing	
GenoType NTM-DR	Detection of important NTM and their resistance to aminoglycosides and macrolides from cultures
GenoType MTBDR _{plus} V2	Detection of <i>M. tuberculosis</i> complex and resistance to rifampicin and isoniazid
GenoType MTBDR _s /V2	Detection of <i>M. tuberculosis</i> complex and resistance to fluoroquinolones and aminoglycosides/cyclic peptides
LPA Genoscholar PZA-TB II	Detection of resistance to pyrazinamide in isolates from patients with bacteriologically confirmed pulmonary TB
Leprosy	
GenoType LepaeDR	Detection of <i>M. leprae</i> and its resistance to rifampicin, ofloxacin and dapsone from patient specimens

The GenoType series all use the same kit content including the membrane strips coated with specific probes, and various solutions for denaturation, hybridisation, washing, and colorimetric reaction. In addition, the kit includes the amplification mixes AM-A (containing buffer, nucleotides, and Taq polymerase) and AM-B (containing salts, specific primers, and dye).

Sample decontamination (e.g. with sodium hydroxide and NALC) and concentration of the specimen by centrifugation is required before DNA isolation. This procedure must be conducted following adequate biosafety requirements. DNA extraction can be performed using a kit (i.e. GenoLyse) produced by the same manufacturer.

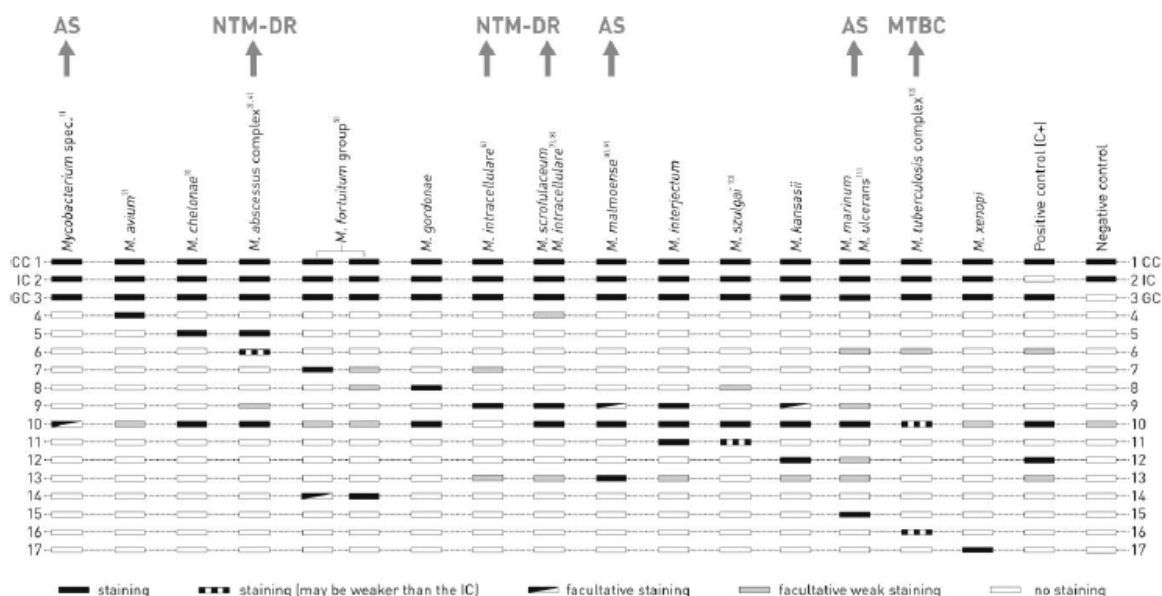
Importantly, the preparation of the amplification reaction must be conducted in a laboratory dedicated pre-amplification area ('clean area') to reduce the risk of cross-contamination or carry-over contamination.

The amplification reaction relies on a standard protocol where the number of cycles varies depending on whether the test is performed directly on the clinical specimen or on a cultivated one.

The hybridisation reaction of the amplified products can be done manually using a water bath and TwinCubator, or automatically using for example a GT-Blot 48 machine. For the interpretation of the test results refer to the assay specific Interpretation Chart available in each kit (an example is provided below, Figure 9). For more detailed information on the assay procedure, refer to the specific instruction for use available in the kit and on the manufacturer website.

Figure 9. Interpretation sheet for GenoType cMdirect assay

Interpretation Chart



Band No. 1 (CC): Conjugate Control
 Band No. 2 (IC): Internal Control
 Band No. 3 (GC): Genus Control

AS: **GenoType Mycobacterium AS** (from cultured material)
 MTBC: **GenoType MTBC** (from cultured material)
 NTM-DR: **GenoType NTM-DR** (from cultured material)

6.2.2 Line probe assays for identification of *M. tuberculosis* complex and drug resistance detection from pulmonary clinical specimens and cultivated samples

The two most widely used LPAs for *M. tuberculosis* complex and drug resistance detection in the ERLTB Network are the GenoType MTBDR_{plus} V2 and GenoType MTBDR_s/V2 assays (Bruker-Hain Lifescience, Nehren, Germany). In both cases, drug resistance is detected through the binding of amplicons to probes targeting the most common mutations to first- and second-line drugs or inferred by the lack of binding to wild-type probes.

They can be used for testing of smear-positive sputum specimens (i.e. GenoType MTBDR_{plus}) or of sputum specimens irrespective of the smear status (i.e. GenoType MTBDR_s/V2), as well as on culture isolates (indirect testing) [1].

Given the complexity of the assay, the necessity of multiple pieces of equipment and the laboratory infrastructure requirements, LPAs are usually implemented at middle- and upper-tier health facilities. The use of LPAs allows obtaining drug susceptibility testing (DST) results in 24-48 hours.

The GenoType MTBDR_{plus} V2 assay allows the simultaneous detection of *M. tuberculosis* complex and resistance to rifampicin and isoniazid by targeting mutations in the *inhA* promoter (from -15 to -8 nucleotides upstream) and *katG* (codon 315) regions, and in the rifampicin resistance-determining region (RRDR) of the *rpoB* gene (from codon 424 to 452, *M. tuberculosis* H37Rv nomenclature) for rifampicin resistance [2].

The GenoType MTBDR_s/V2 assay allows the simultaneous detection of *M. tuberculosis* complex and resistance to fluoroquinolones (e.g. levofloxacin and moxifloxacin) and aminoglycosides/cyclic peptides (e.g. amikacin, kanamycin and capreomycin). The assay includes the quinolone-resistance determining region (QRDR) of *gyrA* (from codon 85 to 96) and *gyrB* (from codon 536 to 541) genes for detection of resistance to fluoroquinolones and the *rrs* (nucleic acid positions 1401, 1402 and 1484) and the *eis* promoter regions (from -37 to -2 nucleotides upstream) for detection of resistance to second line injectable agents [3].

The overall sensitivity and specificity of LPAs for different drugs are reported in detail in the WHO consolidated guidelines on TB, Module 3 [1].

In 2021, WHO recommended the use of the LPA Genoscholar PZA-TB II (Nipro) for detection of resistance to pyrazinamide in isolates from patients with bacteriologically confirmed pulmonary TB [1]. The assay targets a 700-base pair fragment covering the entire *pncA* coding region and 18 nucleotides upstream. The assay comprises a total of 48 probes including three probes that allow the detection of silent mutations not associated with pyrazinamide resistance. Resistance is inferred by the lack of binding to the wild-type probes. Practical considerations for implementation of the Nipro Genoscholar PZA-TB II assay are available in the Information sheet (Annex 2.6) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

As for the LPAs used for *Mycobacteria* identification and drug resistance detection, the testing procedure for this class of LPAs involves three steps including DNA extraction, multiplex amplification using biotinylated primers and reverse hybridisation on nitrocellulose strips.

Appropriate biosafety precautions must be taken when handling mycobacteria. The transfer of bacteria into tubes and heat inactivation during DNA preparation should be carried out in an appropriate biosafety cabinet in a BSL3 laboratory. Upon microorganisms' inactivation, the process can be continued outside the BSL3.

The preparation of the amplification reaction and the mix amplification protocol are identical to the ones described in section 6.2.1. More detailed information on the assay's procedures is provided in the kit instructions for use [2, 3].

The configuration of the of GenoType MTBDR_{plus} V2 and GenoType MTBDR_s/V2 strips is illustrated in Figure 10.

Practical guidance on the interpretation of the GenoType MTBDR assays including information of the association of specific mutations with phenotypic drug resistance, instances in which specific resistance-conferring mutations are not identified and resistance can only be inferred, actions to be performed when certain mutations are detected, and the clinical implications of specific LPA mutations for the selection of appropriate TB treatment is provided in the Global Laboratory Initiative/WHO document 'Line probe assays for detection of drug-resistant tuberculosis: interpretation and reporting manual for laboratory staff and clinicians' [5].

Figure 10. Configuration of GenoType MTBDRplus V2 (a) and GenoType MTBDRsl V2 (b) strips

(a)	(b)
Line 1 ----- Conjugate Control	Line 1 ----- Conjugate Control
Line 2 ----- Amplification Control	Line 2 ----- Amplification Control
Line 3 ----- <i>M. tuberculosis</i> complex TUB	Line 3 ----- <i>M. tuberculosis</i> complex TUB
Line 4 ----- <i>rpoB</i> Locus Control <i>rpoB</i>	Line 4 ----- <i>gyrA</i> Locus Control <i>gyrA</i>
Line 5 ----- <i>rpoB</i> wild type probe 1 <i>rpoB</i> WT1	Line 5 ----- <i>gyrA</i> wild type probe 1 <i>gyrA</i> WT1
Line 6 ----- <i>rpoB</i> wild type probe 2 <i>rpoB</i> WT2	Line 6 ----- <i>gyrA</i> wild type probe 2 <i>gyrA</i> WT2
Line 7 ----- <i>rpoB</i> wild type probe 3 <i>rpoB</i> WT3	Line 7 ----- <i>gyrA</i> wild type probe 3 <i>gyrA</i> WT3
Line 8 ----- <i>rpoB</i> wild type probe 4 <i>rpoB</i> WT4	Line 8 ----- <i>gyrA</i> mutation probe 1 <i>gyrA</i> MUT1
Line 9 ----- <i>rpoB</i> wild type probe 5 <i>rpoB</i> WT5	Line 9 ----- <i>gyrA</i> mutation probe 2 <i>gyrA</i> MUT2
Line 10 ----- <i>rpoB</i> wild type probe 6 <i>rpoB</i> WT6	Line 10 ----- <i>gyrA</i> mutation probe 3A <i>gyrA</i> MUT3A
Line 11 ----- <i>rpoB</i> wild type probe 7 <i>rpoB</i> WT7	Line 11 ----- <i>gyrA</i> mutation probe 3B <i>gyrA</i> MUT3B
Line 12 ----- <i>rpoB</i> wild type probe 8 <i>rpoB</i> WT8	Line 12 ----- <i>gyrA</i> mutation probe 3C <i>gyrA</i> MUT3C
Line 13 ----- <i>rpoB</i> mutation probe 1 <i>rpoB</i> MUT1	Line 13 ----- <i>gyrA</i> mutation probe 3D <i>gyrA</i> MUT3D
Line 14 ----- <i>rpoB</i> mutation probe 2A <i>rpoB</i> MUT2A	Line 14 ----- <i>gyrB</i> Locus Control <i>gyrB</i>
Line 15 ----- <i>rpoB</i> mutation probe 2B <i>rpoB</i> MUT2B	Line 15 ----- <i>gyrB</i> wild type probe <i>gyrB</i> WT
Line 16 ----- <i>rpoB</i> mutation probe 3 <i>rpoB</i> MUT3	Line 16 ----- <i>gyrB</i> mutation probe 1 <i>gyrB</i> MUT1
Line 17 ----- <i>katG</i> Locus Control <i>katG</i>	Line 17 ----- <i>gyrB</i> mutation probe 2 <i>gyrB</i> MUT2
Line 18 ----- <i>katG</i> wild type probe <i>katG</i> WT	Line 18 ----- <i>rrs</i> Locus Control <i>rrs</i>
Line 19 ----- <i>katG</i> mutation probe 1 <i>katG</i> MUT1	Line 19 ----- <i>rrs</i> wild type probe 1 <i>rrs</i> WT1
Line 20 ----- <i>katG</i> mutation probe 2 <i>katG</i> MUT2	Line 20 ----- <i>rrs</i> wild type probe 2 <i>rrs</i> WT2
Line 21 ----- <i>inhA</i> Locus Control <i>inhA</i>	Line 21 ----- <i>rrs</i> mutation probe 1 <i>rrs</i> MUT1
Line 22 ----- <i>inhA</i> wild type probe 1 <i>inhA</i> WT1	Line 22 ----- <i>rrs</i> mutation probe 2 <i>rrs</i> MUT2
Line 23 ----- <i>inhA</i> wild type probe 2 <i>inhA</i> WT2	Line 23 ----- <i>eis</i> Locus Control <i>eis</i>
Line 24 ----- <i>inhA</i> mutation probe 1 <i>inhA</i> MUT1	Line 24 ----- <i>eis</i> wild type probe 1 <i>eis</i> WT1
Line 25 ----- <i>inhA</i> mutation probe 2 <i>inhA</i> MUT2	Line 25 ----- <i>eis</i> wild type probe 2 <i>eis</i> WT2
Line 26 ----- <i>inhA</i> mutation probe 3A <i>inhA</i> MUT3A	Line 26 ----- <i>eis</i> wild type probe 3 <i>eis</i> WT3
Line 27 ----- <i>inhA</i> mutation probe 3B <i>inhA</i> MUT3B	Line 27 ----- <i>eis</i> mutation probe 1 <i>eis</i> MUT1
Colored marker	Colored marker

6.3 Real time PCR-based assays for the identification of *M. tuberculosis* complex and drug resistance detection

The pipeline for new real time PCR based assays for diagnosis of TB with or without drug resistance detection has quickly progressed over the past years. WHO has recommended and supported the development and application of new rapid and accurate diagnostic methods focused on the simultaneous detection of *M. tuberculosis* complex and presence of drug resistance as the best strategy to advance TB laboratory diagnosis. Several WHO recommended rapid diagnostics (WRDs) are now providing options for different settings, but considerations on testing capacity, diagnostic yield, and feasibility of performing the test in the laboratory setting must be taken into account to ensure the diagnostic molecular method selected is appropriate for the specific clinical setting [6].

As an increasing number of WRDs serve similar purposes, WHO has grouped them into classes defined by the type of technology (e.g. automated or reverse hybridisation, nucleic acid amplification tests [NAATs]), the complexity of the test for implementation (e.g. low, moderate, or high – considering the requirements of infrastructure, equipment and technical skills of laboratory staff) and the target conditions (e.g. diagnosis of TB, and detection of resistance to first-line or second-line drugs) [4].

In this section we provide a description of the real-time PCR based assays most commonly used within the ERLTB Network including considerations on their key operational and implementation aspects. Reference and links to policy guidance documents, implementation manuals and training packages is also provided.

This section is not intended to provide the detailed description of the test procedure for which we refer to the specific instructions for use and operating manuals released by the assays' manufacturers.

6.3.1 Single-use sample-processing cartridge system with integrated multicolour real-time PCR capacity for detection of *M. tuberculosis* complex and drug resistance detection

Xpert MTB/RIF and Xpert MTB/RIF Ultra

Xpert MTB/RIF and Xpert MTB/RIF Ultra (Ultra) (Cepheid, Sunnyvale, USA) are cartridge-based real time PCR assays for the simultaneous detection of *M. tuberculosis* complex and rifampicin resistance from clinical specimens. These are fully automated assays as the cartridge contains all reagents required for bacterial lysis, nucleic acid extraction, amplification, and amplicon detection. The only manual step is the addition of a bactericidal lysis buffer to the specimen, which largely eliminates concerns about biosafety during the test procedure. Overall, the assays take approximately 2 hours from sample processing to automatic interpretation and release of the results.

The Xpert MTB/RIF and Xpert Ultra run on the same GeneXpert platform (i.e. six-colour optic instrument), as well as on the newly released 10-colour optic GeneXpert instrument [7].

The Xpert Ultra assay launched in 2017 as a next-generation assay, has an increased sensitivity for *M. tuberculosis* complex detection compared to Xpert MTB/RIF as it targets two multicopy genes (IS6110 and IS1081) instead of one (*rpoB*) and it has a larger DNA reaction chamber, as well as improved fluidics and enzymes [8]. Xpert Ultra uses the same semi-quantitative categories for *M. tuberculosis* complex detection as the Xpert MTB/RIF (i.e. High, Medium, Low and Very Low), with the additional category 'trace' to identify the paucibacillary samples positive to IS6110/IS1081 targets but negative to *rpoB*. In this case no result for rifampicin susceptibility or resistance is provided.

Simultaneous detection of rifampicin resistance is achieved by targeting the RRDR of the *rpoB* gene. In the Xpert MTB/RIF assay, resistance is detected in case of lack or delayed binding of one or more of the five probes to the RRDR, while in the Xpert Ultra four sloppy molecular beacons have been designed to detect mutations by measurable shifts in the probes melting temperatures peaks. This results in an increased capacity of Xpert Ultra to detect mutations at specific codons (e.g. 452), to correctly identify mutations in mixed and paucibacillary samples, and to differentiate synonymous mutations from those associated with resistance [8].

Pooled sensitivities and specificities of Xpert Ultra and Xpert MTB/RIF for TB and drug resistance detection in pulmonary and extrapulmonary specimens in adult and children have recently been published [9-12]. Briefly, for pulmonary TB, Xpert Ultra pooled sensitivity and specificity against culture were 90.9% (86.2 to 94.7) and 95.6% (93.0 to 97.4), versus Xpert MTB/RIF pooled sensitivity and specificity of 84.7% (78.6 to 89.9) and 98.4% (97.0 to 99.3). For rifampicin resistance detection, the pooled sensitivity and specificity were 94.9% (88.9 to 97.9) and 99.1% (97.7 to 99.8) for Xpert Ultra versus 95.3% (90.0 to 98.1) and 98.8% (97.2 to 99.6) for Xpert MTB/RIF. A higher number of rifampicin resistance indeterminate results was detected with Xpert Ultra, pooled proportion 7.6% (2.4 to 21.0) compared to Xpert MTB/RIF pooled proportion 0.8% (0.2 to 2.4), with an estimated difference in the pooled proportions of indeterminate results for Xpert Ultra versus Xpert MTB/RIF of 6.7% (1.4 to 20.1)[1].

In 2020, WHO updated the recommendations on the use of Xpert MTB/RIF and Xpert Ultra as initial tests in adults and children with signs and symptoms of pulmonary and extrapulmonary TB [1]. Updated guidance on the interpretation of Xpert Ultra 'trace' result in HIV-negative people was also provided, recommending that an initial 'trace' call should be considered a true-positive result in those without a prior TB episode or a recent history of TB treatment. In these patients, it is no longer recommended repeating the Xpert Ultra test in case of a 'trace' positive result. In contrast, interpretation of 'trace' results in patients with previous TB may require further clinical assessment and additional tools to exclude a false-positive result. In the case of an indeterminate rifampicin result due to the low bacterial load in the specimen (i.e. MTB detected 'very low' or 'trace'), additional investigations should be performed to confirm or exclude resistance to the drug.

Xpert MTB/XDR

The Xpert MTB/XDR test (Cepheid, Sunnyvale, USA) detects *M. tuberculosis* complex DNA and genomic mutations associated with resistance to isoniazid, fluoroquinolones, ethionamide and second-line injectable drugs (amikacin, kanamycin and capreomycin) in a single cartridge, in approximately 90 minutes [13]. The sample processing procedure and cartridge handling are the same as for Xpert MTB/RIF and Xpert Ultra, however, the Xpert MTB/XDR runs on a 10-colour optics GeneXpert instrument instead of the six-colour optics traditionally used for Xpert MTB/RIF and Xpert Ultra testing. Xpert MTB/XDR targets the genes, codon regions and nucleotide sequences detailed in Table 8.

Table 8. Genes, codon regions and nucleotide sequences targeted by the Xpert MTB/XDR test

Drug	Gene target	Codon regions	Nucleotide
Isoniazid	<i>inhA</i> promoter	Not applicable	–1 to –32 intergenic region
	<i>katG</i>	311–319	939–957
	<i>fabG1</i>	199–210	597–630
	<i>oxyR-ahpC</i> intergenic region	Not applicable	–5 to –50 intergenic region (or –47 to –92) ^a
Ethionamide	<i>inhA</i> promoter	Not applicable	–1 to –32 intergenic region
Fluoroquinolones	<i>gyrA</i>	87–95	261–285
	<i>gyrB</i>	531–544 (or 493–505) ^a	1596–1632
Amikacin, kanamycin, capreomycin	<i>rrs</i>	Not applicable	1396–1417
Amikacin, kanamycin	<i>eis</i> promoter	Not applicable	–6 to –42 intergenic region

^a Codon numbering system according to Camus et al. (2002), as reported in Cepheid, Clinical evaluation of the Xpert MTB/XDR assay, Report R244C2 Xpert MTB/XDR Rev 1.0.

Xpert MTB/XDR assay is intended for use as a reflex test (a follow-up test automatically initiated by a clinical/microbiological laboratory when an initial test result meets pre-determined criteria; e.g. positive or outside normal parameters) in TB specimens, both unprocessed sputum and concentrated sputum sediments, determined to be positive for *M. tuberculosis* complex. The limit of detection for *M. tuberculosis* complex by Xpert MTB/XDR (136 cfu/mL in unprocessed sputum) is similar to that of Xpert MTB/RIF (112.6 cfu/mL), but higher than that of Xpert Ultra (15.6 cfu/mL). In the assay package insert is stated that specimens with a 'trace' result by Xpert Ultra are expected to be below the limit of detection of Xpert MTB/XDR assay and are thus not recommended for testing with this assay [14].

Mutations associated with resistance to isoniazid, fluoroquinolones, ethionamide, and second line injectables are detected by measurable shifts in the melting temperature peaks of the assay probes (i.e. sloppy molecular beacons). A detailed description of the assay design, development, and analytical performance has been published [15], while the diagnostic accuracy data for resistance detection to isoniazid, fluoroquinolones, ethionamide, and amikacin in people with TB detected by Xpert MTB/XDR derived from two multicentre studies have recently been reviewed [16].

Briefly, irrespective of rifampicin resistance, isoniazid resistance pooled sensitivity and specificity were 94.2% (87.5 to 97.4) and 98.5% (92.6 to 99.7) against phenotypic based DST, while fluoroquinolone resistance summary sensitivity and specificity were 93.2% (88.1 to 96.2) and 98.0% (90.8 to 99.6).

In people with known rifampicin resistance, ethionamide resistance summary sensitivity and specificity were 98.0% (74.2 to 99.9) and 99.7% (83.5 to 100.0) against genotypic based DST, while amikacin resistance summary sensitivity and specificity were 86.1% (75.0 to 92.7) and 98.9% (93.0 to 99.8) against phenotypic DST [16].

WHO recommendations on the use of Xpert MTB/XDR in people with bacteriologically confirmed pulmonary TB, with or without rifampicin resistance, are listed in the WHO consolidated guidelines, module 3: diagnosis, while additional considerations on the Xpert MTB/XDR operational features and implementation aspects are available in the information sheet (Annex 2.5) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

6.3.2 Chip-based real time micro-PCR for detection of *M. tuberculosis* complex and rifampicin resistance detection

Truenat MTB, MTB Plus and MTB-RIF Dx assays

The Truenat MTB, MTB Plus, and MTB-RIF Dx assays (Molbio Diagnostics, Goa, India) use chip-based real-time micro-PCR for the semiquantitative detection of *M. tuberculosis* complex (MTB and MTB Plus assays) and rifampicin resistance (MTB-RIF Dx assay) directly from sputum specimens (regardless of the smear status) and can report results in under an hour.

The Truenat testing system uses two portable, battery-operated devices, the Trueprep AUTO v2 Universal Cartridge based Sample Prep Device for the automated extraction and purification of DNA, and the Truelab Real Time micro-PCR Analyzer for performing real-time PCR. The system uses room temperature stable reagents (Trueprep™ AUTO Sample Pre-treatment and Prep kits) and Truenat micro PCR chips. The Truelab Analyzer is available with 1 (Uno)

chip port as well as with 2 (Duo) or 4 (Quattro) chip ports, which allow for independent testing of multiple samples at once. All reagents and consumables required for the test procedures are provided by the manufacturer, with the exception of personal protective equipment (same level of protection as required for microscopy or Xpert MTB/RIF), a timer, and hypochlorite-based disinfectant [17]. The system is designed to be operated in peripheral laboratories with minimal infrastructure requirements and can function at up to 40°C ambient temperature and up to 80% relative humidity [1].

Detection of *M. tuberculosis* complex by Truenat MTB is achieved by targeting a single copy gene (*nrdB*). The Truenat MTB Plus assay has a higher sensitivity than Truenat MTB as it targets multiple genes (*nrdZ* and IS6110). This results in a limit of detection of 30 cfu/ml for Truenat MTB Plus compared to 100 cfu/ml for Truenat MTB [17]. If a positive result is obtained with the MTB or MTB Plus assay, an aliquot of extracted DNA is run on the Truenat MTB-RIF Dx assay to detect mutations associated with RIF resistance. Resistance mutations are detected by a probe melt analysis of the real-time PCR products, which takes approximately an additional hour.

Detailed standard operating procedures for sputum sample preparation, DNA extraction and PCR amplification are described in the Stop TB Partnership Practical guide to Truenat tests implementation [17]. Briefly, DNA is extracted from pre-treated sputum specimens using the kit-specific cartridge and the Trueprep AUTO v2 Device. This process takes approximately 20 minutes. Automated PCR amplification and fluorescent probe-based detection of the extracted DNA on the Truelab Analyzer takes 35 minutes. Results of the Truenat MTB Plus are semiquantitative (i.e. MTB detected as high, medium, low, or very low).

WHO recommends using the Truenat MTB and MTB Plus assays in adults and children with signs and symptoms of pulmonary TB as an initial diagnostic test for TB rather than smear microscopy or culture. The Truenat MTB-RIF Dx is instead recommended for use in people with signs and symptoms of pulmonary TB and a Truenat MTB or MTB Plus positive result as an initial test for rifampicin resistance detection rather than culture and phenotypic DST [1]. Evidence reviewed by WHO on the use of the Truenat TB tests for the detection of *M. tuberculosis* complex and rifampicin resistance was generated through a multicenter prospective clinical evaluation study performed in the intended setting of use (i.e. microscopy centres). Compared to culture, the key performance characteristics of these tests among people with signs and symptoms of pulmonary TB are summarised in Table 9.

Table 9. Diagnostic accuracy of Truenat MTB, MTB Plus and MTB-RIF Dx tests relative to culture, in microscopy centre settings (derived by FIND evaluation study)^[18]

Test	Sensitivity (all patients)	Sensitivity (SS+ patients)	Sensitivity (SS- patients)	Specificity (all patients)
Truenat MTB	0.73	0.91	0.37	0.98
Truenat MTB Plus	0.80	0.96	0.46	0.97
Truenat MTB-RIF Dx	0.84	0.88	0.67	0.95

SS+ = sputum smear positive; SS- = sputum smear negative

Interestingly, findings from this multicentre accuracy study also showed that the rate of rifampicin resistance indeterminate results varied depending on the specimen bacillary load, suggesting that the sensitivity of Truenat MTB Plus to detect MTB is likely higher than that of the Truenat MTB-RIF Dx chip to detect rifampicin resistance [18].

6.3.3 Centralised assays for *M. tuberculosis* complex detection and detection of resistance to rifampicin and isoniazid

A new class of technologies that allow for testing of different conditions using disease-specific assays on the same platform has recently come to market. Several manufacturers have developed moderate complexity automated nucleic acid amplification tests (NAATs) for detection of *M. tuberculosis* complex and resistance to rifampicin and isoniazid on these high throughput instruments. These assays are faster and less complex to perform than phenotypic culture-based DST and LPA as they are largely automated. However, they may require an initial manual specimen treatment step before the test material is transferred into the sample processing tube. Furthermore, adequate biosafety measures should be in place, as well as test-specific equipment, well trained and qualified laboratory staff to set up the tests and carry out the necessary equipment maintenance [1]. For reasons of cost and economy of scale, these assays are generally provided as a centralised service.

In 2019, WHO evaluated the accuracy of four different centralised assays for detection of *M. tuberculosis* complex detection and resistance to rifampicin and isoniazid [19, 20], and in 2020, issued recommendations for their use on respiratory samples as initial tests, rather than culture and phenotypic DST, for detection of *M. tuberculosis* complex and resistance to both rifampicin and isoniazid in people with sign and symptoms of pulmonary TB [1].

The four technologies and assays currently included in the moderate complexity automated NAATs class by WHO are: i) Abbott RealTime MTB and MTB RIF/INH assays, ii) Roche cobas MTB and MTB-RIF/INH assays, iii) Hain FluoroType MTBDR assay, and iv) BD MAX MDR-TB assay.

Based on the systematic review commissioned by WHO, the overall pooled sensitivity for *M. tuberculosis* complex detection compared to culture was 93.0% (90.9–94.7%) and specificity 97.7% (95.6–98.8%). Overall pooled sensitivity for detection of rifampicin resistance compared to phenotypic DST was 96.7% (93.1–98.4%) and specificity was 98.9% (97.5–99.5%), while overall pooled sensitivity for detection of isoniazid resistance compared to phenotypic DST was 86.4% (82.8–89.3%) and specificity was 99.2% (98.1–99.7%) [1, 21].

Abbott RealTime MTB and MTB RIF/INH assays

Abbott Molecular (Chicago, IL, USA) has one NAAT for detection of *M. tuberculosis* complex (RealTime MTB test) and one for the simultaneous detection of rifampicin and isoniazid resistance (RealTime MTB RIF/INH), which can be performed in a standalone mode (starting from the specimen) or in reflex mode (as follow up of the RealTime MTB test). The RealTime MTB test targets the multicopy gene IS6110 and the *pab* gene, while in the RealTime MTB RIF/INH, eight probes covering the RRDR of the *rpoB* gene detect mutations associated with resistance to rifampicin, and four probes detect mutations in the *katG* gene and *inhA* promoter region for isoniazid resistance. The reported limit of detection for *M. tuberculosis* complex is 17 cfu/ml, and 60 cfu/ml for drug resistance detection [1].

The assays are run on Abbot m2000 RealTime system, including two instruments, the m2000sp and m2000rt. Automated DNA extraction and PCR plate preparation starting from inactivated raw or processed sputum specimens is performed on the m2000sp. Alternatively, DNA can be manually extracted using the Abbott mSample Preparation System DNA kit, wherein cells are lysed and DNA is captured by magnetic microparticles. Then, the PCR plate is manually sealed and transferred to the Abbott m2000rt for real time PCR. The Abbot m2000 RealTime system allows for the high-throughput detection of *M. tuberculosis* complex (96 samples including two assay controls), with positive specimens reflexed to the MTB RIF/INH assay (24 samples including two assay controls) for full MDR-TB diagnosis within 10.5 hours.

Detailed information on the assays' procedure and analytical performances is available in the assays' instruction for use and instrument manuals provided by the manufacturer, and publicly available [22]. The accuracy and clinical performance of the Abbott-RT and Abbott-RIF/INH for the detection of TB and DR-TB have been evaluated in systematic reviews and prospective clinical studies [23-25]. Additional considerations on the operational capacity and implementation aspects of the test are available in the information sheet (Annex 2.1) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

Becton Dickinson MAX MDR-TB assay

Becton Dickinson (BD, Sparks, MD, USA) has a multiplexed real-time PCR assay (BD MAX MDR-TB) for the simultaneous detection of *M. tuberculosis* complex and resistance to both rifampicin and isoniazid that can be run on the BD MAX System (Figure 11). The test targets the multicopy genomic elements IS6110 and IS1081, as well as a single copy genomic target (*dev*) for *M. tuberculosis* complex detection, the RRDR of the *rpoB* gene (426 – 452) for rifampicin resistance detection and the *inhA* promoter region and the 315 codon of the *katG* gene for isoniazid resistance. The reported limit of detection is 0.5 cfu/ml for *M. tuberculosis* complex detection, and 6 cfu/ml for drug resistance detection [1].

The test is performed on the BD MAX platform, in which the DNA is automatically extracted starting from a pretreated NALC/NaOH decontaminated sputum (recommended option) or raw sputum, followed by real-time PCR. Bacterial cell lysis is done chemically and by heat, and the released DNA is then captured by magnetic affinity beads. Up to 24 samples can be tested per run on the computer-controlled benchtop system, with the automatic released of results within 4 hours. The assay includes master mixes, reagent strips, extraction tubes, sample tubes, transfer pipettes, and septum caps, while the sample pretreatment reagent (BD MAX STR) and PCR cartridges are provided separately.

Detailed information on the assay procedure and BD MAX System operation is provided in the assay instruction for use and User's Manual. The accuracy and clinical performance of the BD MAX MDR-TB assay have been evaluated in several prospective clinical studies [26-29]. Additional considerations on the operational capacity and implementation aspects of the test are available in the Information sheet (Annex 2.2) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

Figure 11. BD MAX system at the National Research Center, Borstel, Germany

Roche cobas MTB and cobas MTB-RIF/INH assays

Roche Molecular Systems, Inc. (RMS, Roche, Basel, Switzerland) has two NAATs, the cobas MTB and the cobas MTB-RIF/INH tests, to detect *M. tuberculosis* complex and resistance to rifampicin and isoniazid, respectively. The MTB assay uses real-time PCR for *M. tuberculosis* complex detection by targeting the 16S rRNA and five *esx* genes (*esxJ*, *esxK*, *esxM*, *esxP*, *esxW*) starting from inactivated human respiratory samples, including raw and NALC-NaOH-treated sputum and bronchoalveolar lavage fluid. It can generate results for 96 tests (including assay controls) in a 3.5 hour run. The MTB-RIF/INH assay targets the RRDR of *rpoB* gene for rifampicin resistance and the *inhA* promoter region and the *katG* gene for detection of resistance to isoniazid. This assay can be used as a reflex test of cobas MTB-positive samples providing the rifampicin and isoniazid resistance profile for up to 96 samples (including assay controls) in an additional 3.5 hours. The limit of detection reported by the company for this test is 7.6 cfu/ml for decontaminated sediments and 8.8 cfu/ml for unprocessed sputum [1].

The tests are run on the cobas 6800/8800 Systems, which automatically extract DNA for qualitative real-time PCR. Prior to running the cobas MTB or the cobas MTB-RIF/INH assays on the cobas 6800/8800 Systems, samples must be inactivated using the manufacturer specific solution followed by sonication and centrifugation for which additional instrumentation is needed. Therefore, bacterial cell lysis is done chemically (lysis reagent), enzymatically (proteinase) and physically (sonication). Subsequently, the released bacterial DNA is captured by magnetic glass particles and undergoes amplification. Fluorescent-labelled probes allow for the specific detection of the MTB and RIF/INH targets.

Detailed information on the assay procedure and cobas 6800/8800 Systems operation is provided in the assay instruction for use and User's Manuals. The accuracy and clinical performance of the cobas MTB and cobas MTB-RIF/INH tests have been evaluated in several prospective clinical studies [30-32]. Additional considerations on the operational capacity and implementation aspects of these tests are available in the Information sheet (Annex 2.3) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

Bruker-Hain Lifesciences FluoroType MTB and FluoroType MTBDR assays

Bruker-Hain Diagnostics has two real-time nucleic acid amplification tests, the FluoroType MTB and the FluoroType MTBDR to detect *M. tuberculosis* complex and resistance to rifampicin and isoniazid, respectively. The assays use the LATE-PCR amplification and lights-on/lights-off chemistry and target the IS6110 DNA insertion element for *M. tuberculosis* complex detection (FluoroType MTB), and the *rpoB* gene, the *inhA* promoter and *katG* gene for detection of *M. tuberculosis* complex and associated resistance to rifampicin and isoniazid, respectively (FluoroType MTBDR) [1].

The assays run on the FluoroCycler System. Two platforms with different throughput capacity are currently available: the FluoroCycler 12 for the amplification and detection of up to 12 samples and the FluoroCycler XT, which allows the testing of up to 96 samples (including assays controls) and provides results within 4 hours (Figure 12). The limit of detection reported by the company for *M. tuberculosis* complex detection by FluoroType MTB (version 2) is 2.6 cfu/ml and 9 cfu/ml by FluoroType MTBDR, while for rifampicin and isoniazid resistance detection is 14 cfu/ml [1].

Bacterial nucleic acids extraction from decontaminated sputum specimens (FluoroType MTB), or decontaminated sputum specimens and culture isolates (FluoroType MTBDR), can be performed manually using the specific kit

provided by the manufacturer (FluorLyse) or be fully automated using the GenoXtract instrument. This allows for the processing of up to 96 samples as well as the preparation of the PCR plate, which is then transferred on the FluoroCycler XT instrument. DNA extraction by GenoXtract relies on the capturing of intact cells to magnetic beads, from where the cells are washed and then lysed. Both assays use a high-resolution melt curve analysis to detect and automatically report fluorescent detection associated with the probes specific for the *M. tuberculosis* complex and drug resistance targets. The assay differentiates between high-level and low-level isoniazid resistance, and the FluoroSoftware automatically reports the specific mutations identified for each gene target, including mutations that are rare or are associated with unknown resistance profiles.

Detailed information on the procedure and FluoroCycler System operation is provided in the assay instruction for use and User's Manuals available upon request to the manufacturer. The accuracy and clinical performance of the FluoroType MTB and FluoroType MDRTB tests assays have been evaluated in prospective clinical studies [33, 34]. Additional considerations on the operational capacity and implementation aspects of these tests are available in the Information sheet (Annex 2.4) of the WHO operational handbook on tuberculosis. Module 3: diagnosis [4].

Figure 12. FluoroCycler XT at the Ospedale San Raffaele TB Supranational Reference laboratory, Milan, Italy



6.4 Next generation sequencing based approaches for drug-resistant tuberculosis detection

Next-generation sequencing (NGS) has emerged as a powerful tool to improve TB management and control through the rapid, accurate, and comprehensive detection of clinically relevant mutations. This is crucial for clinicians to make prompt decisions regarding the best treatment options, in particular for patients with multi- and extensively drug-resistant TB. Furthermore, NGS-based DST overcomes many of the limitations of the currently available molecular assays which target only a limited number of genomic regions for the testing of a restricted number of drugs, often do not provide information on the specific mutation detected and have limited ability to detect subpopulations of resistant organisms including heteroresistance.

Rapid, reliable and increasingly affordable NGS technologies can now guide all components of TB control, from international surveillance of prevalence and drug-resistant TB to determination of the species or subspecies of *M. tuberculosis* complex isolates and their resistance profile based on the identification of single nucleotide polymorphisms (SNPs), as well as the identification of transmission clusters and outbreaks investigation in different settings (see chapter 8 on molecular typing).

Several comprehensive reviews have been recently published describing the current landscape of NGS applications and recent developments of NGS as a tool for the diagnosis and clinical management of TB [35, 36].

In 2018, WHO published a technical guide that summarises the characteristics of available NGS technologies and provides guidelines for NGS technology selection, procurement and implementation by TB reference laboratories in low- and middle-income countries for the diagnosis of drug-resistant TB in clinical samples [37]. In 2021 WHO published a catalogue of mutations to serve as a global standard for interpreting molecular information for drug resistance prediction [38], and an updated version is expected in early 2023.

In this section, we focus on the two most used NGS workflows for *M. tuberculosis* complex drug resistance prediction: whole genome sequencing and targeted-NGS.

6.4.1 Whole Genome Sequencing

Whole genome sequencing (WGS) approaches use NGS platforms to reconstruct the complete DNA sequence of the bacterial genome allowing the identification of SNPs and insertions/deletions (indels) within regions associated with resistance to anti-TB drugs. The generated data not only allow for the prediction of drug-resistant or -sensitive phenotype, that can be used to inform treatment decisions, but also contribute to our understanding of novel resistance mechanisms for both current and newer drugs as well as to the identification of compensatory mutations.

Currently, WGS is generally performed starting from bacterial strains grown in culture due to the need for a relatively high quantity of good quality high molecular weight DNA. Regardless of the platform and/or chemistry employed for sequencing, WGS workflow is subdivided into four main steps: 1) high molecular weight DNA extraction and quantification; 2) library preparation (including DNA fragmentation, adapter linkage, and PCR amplification); 3) automated single- or paired-end NGS leading to the generation of millions of reads; 4) data analysis. The first three steps are often referred to as the 'wet' laboratory portion of the process while the bioinformatics step as the 'dry' laboratory portion. Detailed information on each of these steps can be found in the WHO technical guide on NGS for *M. tuberculosis* complex [37].

Multiple sequence data analysis solutions exist that differ widely in scope, pipelines and output formats, with little standardisation among them [35]. Generally, sequencing analysis for *M. tuberculosis* complex involves input data validation and quality control, followed by mapping to a reference genome (e.g. *M. tuberculosis* H37Rv) and detection of genomic variants including SNPs and indels. This could be done using a variety of free online tools (e.g. TB profiler, PhyReSE, Mykrobe predictor TB, etc.) [39], or alternatively using commercial software or in-house developed pipelines, some of which are available in public repositories such as GitHub. For a more detailed description of in-house pipelines refer to available publications [40,41,35,42].

In general, as clinical decisions such as the design of the most appropriate treatment regimen rely on the bioinformatic analysis of the sequencing data, it is crucial to use robust and validated pipelines for drug resistance prediction. Resistance profiling is achieved by comparing *M. tuberculosis* complex mutations detected to published curated up-to date databases such as the WHO catalog of mutations in *M. tuberculosis* complex [38]. This is the largest catalogue of TB mutations that has been developed to date, with a total of 17 000 mutations, the inclusion of more than 38 000 *M. tuberculosis* isolates, with phenotypic DST results and WGS data contributed from 41 countries [43]. Efforts to expand the current catalogue with gene regions and mutations implicated in resistance to newer drugs are ongoing.

Several publications on examples of integration of WGS-based systems into routine TB diagnostic algorithms in selected high-income settings in Europe and North America, including the benefits and clinical impact of WGS on treatment decisions are now available [44-47].

6.4.2 Targeted Next Generation Sequencing

An alternative approach for obtaining a comprehensive resistance profile is the design of rapid, targeted genomic panels that include only genes known to be involved in drug-resistance. This approach allows to bypass the need for *M. tuberculosis* complex culture as it can be performed directly on primary specimens, which is invaluable to reference laboratories because it reduces the time to TB and DR-TB diagnosis and the costs associated to culture [37].

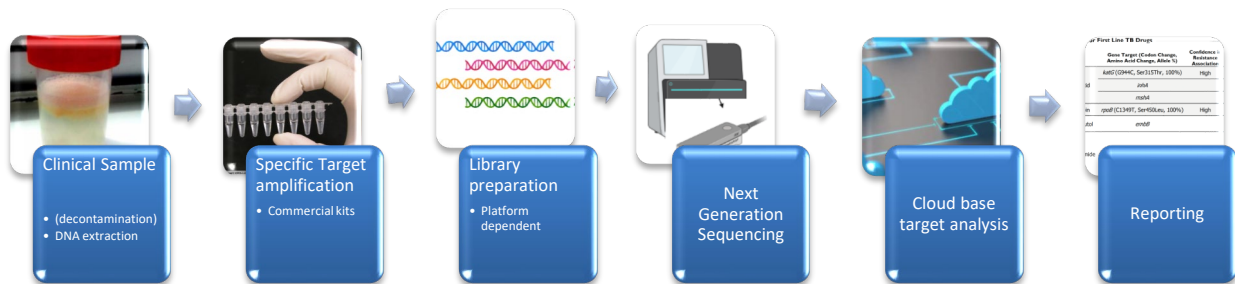
Targeted NGS requires the pre-existing knowledge of the targets (i.e. known resistance markers) but it has the advantage of being customisable and scalable to include additional targets of interest. Furthermore, it allows to interrogate specific regions in the bacterial genome with a higher depth of coverage (i.e. deep/ultra-deep sequencing) thus offering high confidence for mutation detection, also enabling the detection of mixed populations and heteroresistance, within a sample. An additional advantage is that compared to WGS, targeted NGS is much less data intensive and requires significantly less data storage [36].

A few end-to-end solutions for targeted NGS-based DR-TB diagnosis running on both Illumina and Oxford Nanopore platforms are currently being evaluated in a Unitaid funded project (Seq&Treat) coordinated by FIND

(<https://www.genomeweb.com/Reprint-GW22009FIND>). Importantly, evidence from this project will inform WHO policy on the use of targeted NGS for DR-TB diagnosis.

One of these technologies is the Deeplex Myc-TB assay (Genoscreen, Lille, France) (CE-IVD), a 24-amplicon mix that has capacity to screen for mutations in 18 genes known to be associated with *M. tuberculosis* complex resistance to 15 anti-TB drugs including the new and repurposed drugs bedaquiline, clofazimine and linezolid, allowing the simultaneous detection of mycobacterial species identification and genotyping. Samples can be pooled and analysed in a single bench top NGS platform run (e.g. up to 48 samples including controls on an Illumina MiSeq). The targeted NGS workflow includes five steps: 1) DNA extraction from a decontaminated and heat inactivated specimen; 2) multiplex amplification including clean-up of amplicon mixtures and quantification; 3) library preparation including tagmentation, amplification, clean-up and quantification; 4) sequencing; 5) data analysis [37]. The kit is linked to a secured cloud-based application, for fast and easy analysis and interpretation of the sequencing data. More detailed information on the procedure is available in the kit User Manual.

Figure 13. Deeplex-MycTB Workflow



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7 Phenotypic susceptibility testing to anti-tuberculous agents for *Mycobacterium tuberculosis* complex

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7.1 Background and principles

This chapter describes phenotypic antimicrobial susceptibility testing (AST) for *Mycobacterium tuberculosis* and other members of the *M. tuberculosis* complex (MTBC) [1, 2].

Major objectives for AST in MTBC include: (i) ensuring effective individual treatment and management of a TB case; (ii) anti-tuberculosis drug resistance surveillance at the level of a hospital, city, region, or country; (iii) identification of the need for isolation of patients; and (iv) determination of the scope of institutional and community outbreak investigations required [1, 2]. The implementation of the WHO End TB Strategy requires provision of susceptibility data for all patients with signs and symptoms of TB [3].

Because of the development of antimicrobial resistance (AMR) in TB worldwide, rapid and reliable AST to antituberculous agents is required. Specimens for culture and AST should be obtained at or before the start of treatment and testing should be performed rapidly for at least rifampicin and isoniazid. Baseline susceptibility testing for fluoroquinolones would be highly recommended for patients coming from countries with a high prevalence of resistance to fluoroquinolones in new TB patients [4]. AST may be repeated if the patient is still culture positive after two months of treatment [5].

The AMR rates for antituberculous agents are higher in re-treatment cases (acquired resistance or secondary resistance) than in new cases (primary resistance), the difference varying according to the country or epidemiological situation [6]. AMR observed during treatment, designated as 'acquired' or 'secondary resistance', results from the selection and multiplication of resistant mutant subpopulations pre-existing in the tubercle bacillus isolate before therapy. AMR observed before treatment, designated as 'primary resistance', is the consequence of exposure to a drug-resistant source of infection [7].

Identification of the mycobacterial culture or MTBC acid-fast bacilli is necessary before AST is undertaken to avoid false resistant results due to the presence of non-tuberculous mycobacteria (NTM) which may have similar growth patterns but harbour intrinsic resistant characteristics. The knowledge of the exact MTBC species and in some cases the lineage is important when interpreting the AST results, as some strains show intrinsic resistance to a drug (e.g. *M. bovis* and *M. canettii* being resistant to pyrazinamide [8, 9]), or increased MIC values for a specific drug (e.g. Lineage 1 strains having increased MIC values of pretomanid [10]).

Laboratories should only use reliable methods for MTBC AST using the WHO-recommended drug critical concentrations [11-13]. The laboratory should have considerable experience in the chosen method, and it should be carried out by skilled technicians. It is also extremely important to periodically perform internal and external quality controls (QC) for AST. According to ISO 15189 (standards 5.3.2), all new reagents (lot or shipment) should be verified for performance before being used, therefore, all batches of media (LJ and MGIT), as well as the other reagents used for AST should be tested. If the lot/batch QC fails, all results obtained within that batch, as well as the new batch of a reagent should be thoroughly reviewed, and the testing might be repeated. All laboratories performing AST have to participate in an external QC programme [1, 14].

AST is usually performed on MTBC cultures (indirect testing) but it could also be performed on specimens containing acid-fast bacilli known to belong to MTBC (direct testing) [15] as long as special conditions are fulfilled as described in section 7.3.3. Results are obtained faster for direct testing; however, there is a lower rate of success due to possible contamination or wrong estimation of the inoculum.

Given the high risk of generating infectious aerosols from specimens and the high concentration of infectious particles, all procedures described in this chapter should be performed in a high risk (TB-containment BSL3) laboratory [16].

7.2 General method – AST by culture

7.2.1 Definitions

Critical concentration: the lowest concentration of an anti-TB agent that will inhibit the in vitro growth of 99% of phenotypically wild type isolates of *M. tuberculosis* complex.

Critical proportion: the proportion of resistant organisms within a particular cultured isolate that is used to determine resistance to a particular drug. Any culture that shows less than 1% growth on a medium containing a critical concentration of the agent being tested when compared with the growth on a control without the agent is considered to be susceptible; a culture that has 1% or more growth on the medium containing the critical concentration of the agent is considered to be resistant.

Minimum inhibitory concentration (MIC): the lowest concentration of an antimicrobial agent that prevents visual growth of the isolate in a solid medium or broth dilution susceptibility test, under specific conditions of testing (see EUCAST reference protocol below).

Clinical breakpoint: the concentration/s of an antimicrobial agent equal or above the critical concentration that separates isolates that will likely respond to treatment (S categorisation) from those which will likely not respond to treatment (R categorisation). This concentration is determined by correlation with available clinical outcome data, MIC distributions, genetic markers, and pharmacokinetic/pharmacodynamic data including drug dose. An increase dosing can be used in some defined cases (I categorisation). The clinical breakpoint is used to guide individual clinical decisions in patient treatment.

Epidemiological cut-off value (ECOFF): the upper end of the Gaussian-shaped MIC distribution of a species when only phenotypically wild-type (pWT) isolates are tested. The ECOFF typically encompasses 99% of pWT isolates [17].

Growth control: Number of colonies obtained upon inoculation of dilutions of the mycobacterial suspension in a culture medium that does not contain any anti-TB agent.

Resistance is defined as a decrease in sensitivity of sufficient degree to be reasonably certain that the strain concerned is different from drug susceptible wild-type reference strains. In this case, desired clinical effect of the drug is unlikely.

Susceptibility is defined by a level of sensitivity not significantly different from wild-type strains that have never come into contact with the drug. In this case, expected clinical responsiveness by the use of the drug is likely [18, 19].

7.2.2 Introduction

The methods introduced in this chapter aim not only to detect resistance but also to assess susceptibility [19, 1, 2]. Applying these methods should lead to high sensitivity for detecting resistance (i.e. a low rate of false susceptibility results) and high specificity (i.e. a low rate of false-resistance results). Additionally, these methods should also offer high sensitivity for assessing susceptibility (i.e. a low rate of false-resistance results) and a high specificity (i.e. a low rate of false susceptible results). Performance studies on the various methods are referenced in each method section.

Phenotypic DST methods rely on the use of critical concentrations of anti-TB agents. Definitions and usefulness of critical concentrations as well as other cut-off values (clinical breakpoints, MICs and epidemiological cut-off values) in phenotypic DST have considerably evolved over the last decade and are considered in detail elsewhere [20, 21, 2].

In 2021, WHO published a technical report with the revised critical concentrations for culture-based phenotypic AST to first-line anti-TB drugs isoniazid and rifampicin [13]. Critical concentrations for rifampicin have been lowered while those for isoniazid have been maintained at the present level. This update helps addressing the discordance observed between phenotypic and molecular methods to detect rifampicin resistance and improves the accuracy of AST [13]. The updated concentrations are reported in Table 10. All concentrations are in mg/L and apply to the proportion method with 1% as the critical proportion. Slightly different critical concentrations are provided by the Clinical and Laboratory Standards Institute (CLSI) [22].

Table 10. Critical concentrations for isoniazid and the rifamycins [13]

Medicine	LJ	7H10	7H11	MGIT
Isoniazid	0.2	0.2	0.2	0.1
Rifampicin	40.0	0.5	1.0	0.5
Rifabutin	-	-	-	-
Rifapentine	-	-	-	-

LJ = Löwenstein–Jensen medium; 7H10 = Middlebrook 7H10 growth medium; 7H11 = Middlebrook 7H11 growth medium; MGIT = Mycobacteria Growth Indicator Tube

In 2021, WHO recommended the use of a four-month regimen for drug-susceptible TB composed of rifapentine, isoniazid, pyrazinamide, and moxifloxacin [23]. Given the lack of available minimum inhibitory concentration (MIC) data for rifapentine, according to WHO, complete cross-resistance with rifampicin should be assumed until sufficient data to the contrary become available (i.e. genotypic AST and phenotypic AST results for rifampicin should be used as the surrogate for rifapentine) [13].

Drugs for the treatment of rifampicin-resistant (RR)/multidrug-resistant (MDR) TB are grouped by WHO based on their relative benefits and harms into [24]:

- **Group A:** fluoroquinolones (levofloxacin and moxifloxacin), bedaquiline and linezolid (considered highly effective and strongly recommended for inclusion in all regimens unless contraindicated);
- **Group B:** clofazimine and cycloserine or terizidone (conditionally recommended as agents of second choice);
- **Group C:** all other medicines that can be used when a regimen cannot be composed with Group A and B agents. The medicines in Group C (ethambutol, delamanid, pyrazinamide, imipenem-cilastatin/ meropenem, amikacin, ethionamide/prothionamide, *P*-aminosalicylic acid) are ranked by the relative balance of benefit to harm usually expected of each.

In May 2022, WHO released a rapid communication stating that the six-months all-oral regimen of bedaquiline, pretomanid, linezolid, and moxifloxacin (BPaLM) may be used programmatically for patients (aged ≥ 15 years) with rifampicin-resistant TB not yet exposed to bedaquiline, pretomanid (not categorised), and linezolid. This regimen may be used without moxifloxacin (BPaL) in the case of documented resistance to fluoroquinolones (in patients with pre-XDR-TB) [25]. Given this new recommendation, building capacity to perform AST to bedaquiline and linezolid has become an urgent priority.

For many of the recommended drugs for use in the longer RR/MDR-TB treatment regimen, there are no validated AST protocols described yet, and acquired resistance under treatment is not well characterised.

The critical concentrations and clinical breakpoints for medicines recommended for the treatment of rifampicin-resistant and multidrug-resistant TB according to WHO are reported in Table 11. All concentrations are in mg/L and apply to the proportion method with 1% as the critical proportion.

Table 11. Critical concentrations and clinical breakpoints for medicines recommended for the treatment of RR/MDR-TB [24]

Groups and steps	Medicine	LJ	7H10	7H11	MGIT
Group A: Include all three medicines	Levofloxacin	2.0	1.0	-	1.0
	Moxifloxacin (CC)	1.0	0.5	0.5	0.25
	Moxifloxacin (CB)	-	2.0	-	1.0
	Bedaquiline	-	-	0.25	1.0
	Linezolid	-	1.0	1.0	1.0
Group B: Add one or both medicines	Clofazimine	-	-	-	1.0
	Cycloserine / terizidone	-	-	-	-
Group C: Add to complete the regimen and when medicines from Groups A and B cannot be used	Ethambutol	2.0	5.0	7.5	5.0
	Delamanid	-	-	0.016	0.06
	Pyrazinamide	-	-	-	100.0
	Imipenem–cilastatin	-	-	-	-
	Meropenem	-	-	-	-
	Amikacin	30.0	2.0	-	1.0
	Ethionamide	40.0	5.0	10.0	5.0
	Prothionamide	40.0	-	-	2.5
<i>P</i> -aminosalicylic acid	-	-	-	-	

LJ = Löwenstein–Jensen medium; 7H10 = Middlebrook 7H10 growth medium; 7H11 = Middlebrook 7H11 growth medium; MGIT = *Mycobacteria Growth Indicator Tube*

For pretomanid, a provisional breakpoint of 1 mg/L using the MGIT System has been proposed by the European Medicines Agency (EMA) [26], while EUCAST has proposed 2 mg/L as a provisional screen value to be used for AST in MGIT (EUCAST, September 2022) until sufficient MIC data using the EUCAST reference method is available (<http://www.eucast.org>). WHO has not yet endorsed a critical concentration for this drug.

7.2.2 Materials

For direct testing: smear-positive specimens after classical decontamination (see Section 5.6) should be used [15].

For indirect testing: clinical isolates of MTBC, as a positive liquid culture or colonies on a solid media, after checking for purity should be used [1, 27].

7.2.3 Methods

In this chapter we will focus on the AST methods primarily used in the ERLTB-Network, namely the proportion method on both solid media (i.e. Löwenstein-Jensen) [28] and liquid Middlebrook (7H9) media using the BD BACTEC MGIT automated mycobacterial detection system [29-32]. In addition, we will describe the EUCAST reference method for minimum inhibitory concentration (MIC) determination used to define epidemiological cut-off values (ECOFFs) and clinical breakpoints (CBs) [33, 34].

Other solid culture methods (i.e. the resistance ratio method and the absolute concentration method) are still used, although rarely, in some laboratories as they are relatively inexpensive. However, these methods have been standardised for testing of first-line drugs only (rifampicin, isoniazid, ethambutol and pyrazinamide) [35].

Several additional phenotypic assays, such as the nitrate reductase assay (NRA), microplate microdilution using resazurin (colorimetric redox indicator assay, CRI) or commercial plate (Sensititre MYCOTB MIC plate), microscopic observed direct susceptibility testing (MODS) and the Thin Layer Agar methods (TLA) are less reproducible than the proportion methods. WHO has endorsed the use of the NRA and MODS assays for direct AST, and the NRA, MODS and CRI assays for indirect AST, in reference laboratories under clearly defined operational conditions following strict laboratory protocols [36]. These methods have been described and recently reviewed elsewhere [37-43].

7.2.4 Report/interpretation

Reporting of susceptibility results is easily done for each tested drug, if the method was performed according to the standard protocol. Unexpected resistant results should be checked by molecular confirmation, as described in Chapter 6 or by repeated testing. This may be done as soon as a resistance pattern is observed. If a mutation known to confer resistance is observed, this confirms resistance. Conversely, if no mutation in the genes known to confer resistance is detected, this implies that the strain needs to be retested with a reference method. If the strain still appears resistant but no mutation is found, colonies growing in the presence of the particular antibiotic should be retested [8].

Emergency and priority reporting should be carried out for cases of smear-positive pulmonary TB and for all cases detected as RR/MDR-TB. These results should be reported according to the institutional protocol, preferably to the attending clinical team. Routine reporting should contain the start date of testing and the date of reporting.

For antibiotics for which several concentrations can be tested, the level of resistance can be determined and reported [44]. Usually, a report of low resistance does not imply that the drug will not be given; conversely, a report of a high-level of resistance implies that the drug is of no clinical use. However, more clinical studies should be done correlating patient outcome with levels of resistance.

7.3 Proportion method on Löwenstein-Jensen medium

7.3.1 Introduction

The proportion method on Löwenstein-Jensen (LJ) medium was one of the first methods developed for susceptibility testing of MTBC and it is still considered a reference method. Subsequently, it was adapted to be used with other media such as the Middlebrook agar (7H10 or 7H11) as described in the WHO technical manual for drug susceptibility testing [11]. It should be emphasised that anti-TB drug critical concentrations may be different between the media (see Tables 10 and 11) [13, 45].

The proportion method calculates the proportion of resistant bacilli present in the clinical isolate [35]. In this method, the growth (i.e. the number of colonies) on a control LJ tube that does not contain an anti-TB agent (i.e. growth control) is compared with the growth present on LJ tubes containing the critical concentration of the anti-TB drug being tested. In general, at least two dilutions of a culture suspension are tested on both the growth control and the drug containing tube. The ratio of the number of colonies on the medium containing the anti-TB agent to the number of colonies on the medium without the anti-TB agent is calculated, and the proportion is expressed as a percentage. Below a critical proportion of 1%, the strain is classified as susceptible; above that proportion, it is classified as resistant (see section 7.2.1). The critical proportion was assessed by a study of a cohort of patients in the 1960s who failed TB treatment [18] and is particularly useful for patients in whom heteroresistance is suspected. The proportion method on LJ media is well established for first-line (i.e. rifampicin, isoniazid, ethambutol) and some second-line drugs (i.e. levofloxacin, moxifloxacin, amikacin, ethionamide/prothionamide).

7.3.2 Materials

For direct testing, smear-positive specimens are decontaminated as per the usual procedure (see Section 5.4.1.) and the sediment is used either pure or diluted.

For indirect testing, a pure, well-characterised culture of *M. tuberculosis* complex (MTBC) bacteria in the active growth phase is necessary.

LJ tubes (preferably with screw caps) with the incorporated anti-TB agent to be tested, and LJ tubes without any anti-TB agent. Pure formulation (no patient tablets) of the anti-TB drugs being tested must be always used. Other materials include sterile pipettes and McFarland turbidity standard no. 1.

7.3.3 Methods

Media preparation

A detailed description of the procedure used for the preparation of LJ media for AST is available in the WHO technical manual for drug susceptibility testing of medicines used in the treatment of TB [11]. For the preparation of the medium containing the agent to be tested, the agents are incorporated into the liquid mixture according to their specific critical concentrations before the media is dispensed into tubes and inspissated. LJ medium with and without incorporated drugs can be stored at 4–8 °C for one month.

Drugs

The drugs to be tested should be stored according to the manufacturer's instructions. Substances, solvents and diluents are listed in Table 12 along with critical concentrations for testing on LJ media.

Table 12. Solvents and diluents

Drugs	Substance	Critical concentration in LJ (mg/l)	Solvents	Diluents
Isoniazid	Isoniazid	0.2	DW	DW
Rifampicin	Rifampicin	40	DMSO	DW
Ethambutol	Ethambutol dihydrochloride	2	DW	DW
Amikacin	Amikacin sulfate	40	DW	DW
Levofloxacin	L-ofloxacin	2	0.1N NaOH	DW
Moxifloxacin	Moxifloxacin	1	0.1N NaOH	DW

DW = Sterile distilled water; DMSO = dimethyl-sulfoxide; NaOH = Sodium-hydroxide.

Bacterial suspension and inoculation

Indirect drug susceptibility testing

This is carried out on a primary isolate or a subculture on LJ medium. A representative portion of the isolate is obtained by sampling as many colonies as possible within one or two weeks after the appearance of growth.

The colonies are transferred to a glass tube or an Erlen flask without residual culture medium. Homogenisation of the suspension can be done using glass beads (3.0 mm in diameter) or with a glass rod with a molten rounded tip by rubbing the bacteria onto the glass wall. The suspension is then made by adding 0.9% sodium chloride or distilled water. After thorough mixing and homogenisation of the suspension, the tubes should rest for ten minutes, after which the supernatant is pipetted into another tube.

The turbidity of the suspension should be visually adjusted to 1 mg wet bacterial mass/ml (about one full loop with an inner diameter of 3 mm), or to the reference suspension McFarland standard 1.0 (9.9 ml sulphuric acid [1% volume concentration] with 0.1 ml barium chloride solution [1% mass concentration]). The latter is about 10-fold less concentrated than the former.

Serial dilutions of 10^{-1} to 10^{-5} of the standard suspension are prepared by diluting sequentially 1.0 ml of the standard suspension in tubes containing 9 ml of sterile distilled 0.9% sodium chloride.

Usually, two dilutions of the inoculum are inoculated onto the control LJ tubes (without drug), the second inoculum being a 1:100-fold dilution of the first inoculum: 10^{-1} and 10^{-3} of the 1 mg/ml suspension, or 10^{-2} and 10^{-4} of the McFarland 1.0.

The drug containing LJ tubes are then inoculated with the two dilutions. The volume of the inoculum in both the control and drug containing tubes is 0.1 ml.

Direct testing using primary specimens

Decontaminated smear-positive specimens are inoculated directly onto LJ slopes using two dilutions, as for the indirect testing, the second inoculum being a 1:100 dilution of the first inoculum. Dilutions are made according to the number of acid-fast bacilli per microscopic field (x1 000 magnification):

- Undiluted (inoculum 1) and 10^{-2} (inoculum 2) if there is less than 1 AFB per field;
- 10^{-1} and 10^{-3} if there are 1 to 10 AFB per field;
- 10^{-2} and 10^{-4} if there are more than 10 AFB per field.

The drug containing LJ media tubes are inoculated with both dilutions if possible, or at least with dilution 1 for first-line drugs (1% critical proportion) and dilution 2 for the second-line drugs amikacin and moxifloxacin. The volume of the inoculum is 0.1 ml per tube.

Incubation

After inoculation, the tubes are incubated at $36\pm 1^\circ\text{C}$ in a slanted position, with the screw caps slightly loosened to allow for the evaporation of the inoculum. This does however depend on the type of screw caps; those that can be closed immediately after inoculation may be preferred. After 2 to 4 days, screw caps are tightened, and the tubes are further incubated.

7.3.4 Report interpretation

The following three steps are used for reading the results:

- Counting of the colonies grown on the growth control tubes. For dilution 1 (e.g. 10⁻²) a confluent growth is often observed since circa 10⁴ colony-forming units (CFU) have been inoculated; for dilution 2 (e.g. 10⁻⁴) about 20–100 colonies are counted. This number may differ from strain to strain since some are dysgonic (i.e. grow with difficulty on the culture media);
- Counting of the colonies grown on the LJ tubes containing the anti-TB agent being tested. Usually the first dilution (i.e. 10⁻² dilution) is used for counting.
- Calculation of the proportion of resistant bacilli by comparing the counts on growth control dilution 2 (i.e. 10⁻⁴ dilution) with the counts on the tubes containing the anti-TB agent (usually dilution 1). If the proportion is equal or above 1 %, the strain is reported as resistant; if the proportion is below 1%, the strain is reported as susceptible.

The reading of results is carried out after 28 days (early reading) and 42 days (final reading) after inoculation. If after four weeks of incubation the proportion of resistant colonies is higher than the critical proportion, the strain can be reported as resistant. Also, if the reading on day 28 shows that there are no colonies on the drug containing media and the colonies on the control tubes are mature, the strain can be reported as susceptible. Except for these two instances, all other results should be reported after the reading on day 42.

If the number of colonies in the control tubes inoculated with inoculum 1 is below to 100, the test should be repeated with a less diluted inoculum.

7.4 Drug susceptibility testing in liquid media (Bactec MGIT 960)

7.4.1 Introduction

The first liquid-based culture media were introduced commercially in the 1990s, and several evaluations have demonstrated good correlation with the solid media proportion method and significant time savings. One of the earlier disadvantages of this system was the use of a radioactive labelled substrate. Because of the strict regulations of handling and waste disposal of radioactive material along with the biosafety aspects of using syringes for bacterial inoculation, it became necessary to develop a non-radiometric broth-based technique.

Two liquid culture systems for AST are currently commercially available: the BACTEC MGIT 960 system (Becton Dickinson, Sparks, MD, USA) and the VersaTREK™ automated microbial detection system (Thermo Fisher Scientific, Waltham, MA, USA). The BACTEC MGIT 960 system uses 7H9 liquid media which contains an oxygen-quenched fluorochrome embedded in silicone at the bottom of the culture tube. During bacterial growth, the free oxygen in the medium is utilised and replaced with carbon dioxide. With the depletion of free oxygen, the fluorochrome is no longer inhibited, resulting in fluorescence and identification of bacterial growth (measured as growth units, GU), which can be detected manually or automatically [30, 32, 46-47].

7.4.2 Materials

For the inoculum preparation, a pure, well-characterised culture of *M. tuberculosis* bacteria in the active growth phase is required. Other material required include BACTEC MGIT culture tubes, growth supplements and drugs. Drugs are supplied either in lyophilised form or as pure powders by vendors and must be reconstituted following the manufacturer instructions making sure that the final test concentrations are the same as those recommended by WHO.

7.4.3 Methods

Media preparation

A total of 0.1 ml of a reconstituted drug solution and 0.8 ml supplement (e.g. OADC, consisting of oleic acid, albumin, dextrose and catalase) (available commercially) is added to each of the 7 ml of 7H9 media-containing tubes.

Drugs

Ready to use AST kits for first line drugs and pyrazinamide are commercially available (i.e. BACTEC MGIT 960 SIRE and IRE kits and BACTEC MGIT 960 PZA kit). Recently, Becton Dickinson has made lyophilised preparations available for second-line drugs like amikacin and moxifloxacin for use in the MGIT 960 as well as in other media (Table 13). Other second line drugs (e.g. Linezolid and Clofazimine) are supplied by specific manufacturers (e.g. Sigma- Aldrich, Caymen Chemical). New drugs such as Bedaquiline and Delamanid can only be procured through

official suppliers (i.e. NIH AIDS Reagent Program³, and BEI resources⁴, respectively). Drugs must be properly reconstituted to provide critical concentrations recommended by WHO (Tables 10 and 11). Detailed information on how to prepare the drugs stock and working solutions are available in the WHO technical manual on drug susceptibility testing [11].

Table 13. Second line drugs solvents, diluents and concentrations for AST in MGIT

Drugs	Stock solution concentration (mg/L)	Solvent	Diluent	Working solution concentration (mg/L)	Final concentration in MGIT (mg/L)
Levofloxacin	10 000	0.1N NaOH ¹	DW	84.0	1.0
Moxifloxacin (CC)	10 000	0.1N NaOH ¹	DW	21.0	0.25
Moxifloxacin (CB)	10 000	0.1N NaOH ¹	DW	84.0	1.0
Ethionamide	10 000	DMSO	DMSO	420.0	5.0
Amikacin	10 000	DW	DW	84.0	1.0
Linezolid	1 000	DW	DW	84.0	1.0
Bedaquiline	1 000	DMSO	DMSO	84.0	1.0
Delamanid	10 000	DMSO	DMSO	5.04	0.06
Clofazimine	10 000	DMSO	DMSO	84.0 ²	1.0

DW = Sterile distilled water; DMSO = dimethyl-sulfoxide; NaOH = Sodium-hydroxide.

¹ NaOH is added drop by drop until complete dissolution of the compound, thereafter DW is added up to the total volume.

² Clofazimine working solution must be prepared new before use starting from the stock solution (10 000 mg/L) as this drug is unstable at low concentration.

Concentrated solution in the appropriate diluent can be stored at -20°C for up to six months. Stability of the concentrated solution varies for the different drugs. Working solutions are usually unstable and should be prepared fresh before use.

Inoculation

The inoculum for indirect drug susceptibility testing in Bactec MGIT can be prepared from liquid or solid media, according to the manufacturer's guidelines [48]. Testing using inoculum from liquid media includes the following steps:

- A positive MGIT culture tube is used as the inoculum. The tube is inverted one to two times and then left undisturbed for about five to ten minutes to let big clumps settle to the bottom;
- The inoculum (supernatant) of the positive MGIT tube is used undiluted within 1-3 days after positivity has been flagged by the Bactec MGIT instrument for culture positivity, or diluted 1:5 if incubated 4-5 days after. A volume of 0.5 ml of the suspension has to be added aseptically into every drug-containing tube;
- For each set of AST, the control tube will be 1:100 dilution of the original inoculum (1% control): the inoculum is diluted 1:100 by adding 0.1 ml of the suspension to 10 ml of sterile saline. The tube has to be well mixed before adding 0.5 ml into the growth-control tube. For pyrazinamide testing performed in MGIT culture media with reduced pH that is provided by the manufacturer Becton Dickinson a 1:10 diluted growth control is used.

Incubation

After inoculation, the tubes are incubated at 37 °C in the Bactec MGIT instrument where fluorescence is detected automatically. In manual operation, tubes are incubated at 37 °C and are read under UV light every day.

7.4.4 Report interpretation

The BACTEC MGIT 960 instrument continually monitors all tubes for increased fluorescence. Fluorescence in the drug-containing tubes is compared to the fluorescence in the Growth Control (GC) tube to determine the susceptibility results. When the growth unit (GU) of the GC (i.e. 1% inoculum and 10% for PZA) reaches a predetermined threshold (i.e. 400 GU) within 4-13 days (SIRE) or 4-21 days (PZA), the GU values of the drug containing vials are evaluated.

Results are qualitative: if the GU of drug containing tube is less than 100, the strain is reported susceptible. Conversely, if the GU of the drug containing tube is more than 100, the strain is reported as resistant. If the GC is

³ NIH AIDS Reagent Program: www.hivreagentprogram.org/Catalog/HRPantimicrobialCompoundsandOtherChemicals/ARP-12702.aspx

⁴ BEI resources: www.beiresources.org/Catalog/BEIantimicrobialCompoundsandOtherChemicals/NR-51636.aspx

still negative after 14 days, the test should be repeated paying attention to the inoculum preparation procedure and considering the strain growth capacity (dysgony).

The MGIT 960 System coupled with the TB eXiST module (eXtended individual Susceptibility Testing) running on a separate computer equipped with the EpiCenter software can be used to perform AST for a panel of second line drugs for which critical concentrations have been established for this method [49]. The module also allows extending the protocol length beyond 13 days in order to accommodate slow-growing resistant MTBC isolates.

All positive tubes should be checked for contamination, by preparing a Gram-stained smear and/or adding one drop onto a blood plate, and for mycobacterial growth by preparing a Ziehl-Neelsen stained smear.

7.5 Minimum Inhibitory Concentration (MIC) determination following the EUCAST reference method

Although several methods have been described for the determination of minimal inhibitory concentrations (MIC) of antituberculous agents against MTBC isolates, no standardised methods were available before 2019 to enable measuring the epidemiologic cut off values (ECOFFs) and clinical breakpoints according to EUCAST strategies. In 2016, the EUCAST subcommittee for anti-mycobacterial drug susceptibility testing (AMST) was launched with a primary goal of defining a reference method for MIC determination of the MTBC. This reference method has been made publicly available on the EUCAST website in July 2019 (https://www.eucast.org/mycobacteria/methods_in_mycobacteria) with a protocol published in 2020 [34] along with preliminary results [50]. The main criteria of the protocol agreement were to produce MIC results with good reproducibility.

Briefly, the protocol is a broth microdilution method in Middlebrook 7H9-10% OADC medium. The final inoculum is a 105 CFU/mL suspension, obtained from a 10⁻² dilution of a 0.5 McFarland suspension prepared after vortexing bacterial colonies with glass beads before making a suspension in sterile water. The culture is maintained in a U-shaped 96-well polystyrene microtitre sterile plate with a lid incubated at 36 ± 1 °C (an example of plate configuration is shown in Figure 14). Reading is done using an inverted mirror. The MIC, expressed in mg/L, is the lowest concentration that inhibits visual growth, as soon as the 1:100 diluted control (i.e. 103 CFU/mL suspension) shows visual growth.

EUCAST-AMST used *Mycobacterium tuberculosis* H37Rv ATCC 27294 as a reference strain and its targeted MIC values were within the range 0.03-0.12 for isoniazid, 0.12-0.5 for levofloxacin and 0.25-1 mg/L for amikacin [50].

All drug solutions should be prepared according with the Good Manufacturing Practice (GMP) and powders must be obtained directly from the drug manufacturer or from reliable commercial sources together with appropriate documentation for quality assurance. Generally, drugs should be dissolved as described in ISO-20776-1 guidelines or if not listed, per recommendation of the manufacturer.

Since many antituberculous drugs are not soluble in water, the solvent should be used with caution with regard to its own potential inhibitory effect against MTBC. Consequently, for solvents other than water, such as DMSO, there should be growth controls containing the same proportion of solvent (e.g. in case of DMSO) as the drug containing medium, and the concentration of the solvent should be the same for all concentrations. For example, bedaquiline, delamanid, pretomanid, clofazimine are dissolved in DMSO and the final concentration in each well could be 1% or 0.5% DMSO, according to the manufacturer's recommendation. Due to limited solubility of these drugs, serial dilution of the drug should be made first with 100% DMSO before addition (1/100) to each well.

Figure 14. Scheme of the microtitre plate for the EUCAST AMST broth microdilution reference protocol

	1	2	3	4	5	6	7	8	9	10	11	12
A	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20
B	negative control	GC 100%	AA1 (10-2) C8	AA1 (10-2) C7	AA1 (10-2) C6	AA1 (10-2) C5	AA1 (10-2) C4	AA1 (10-2) C3	AA1 (10-2) C2	AA1 (10-2) C1	GC 1%	200ul dH20
C	negative control	GC 100%	AA1 (10-2) C8	AA1 (10-2) C7	AA1 (10-2) C6	AA1 (10-2) C5	AA1 (10-2) C4	AA1 (10-2) C3	AA1 (10-2) C2	AA1 (10-2) C1	GC 1%	200ul dH20
D	negative control	GC 100%	AA2 (10-2) C8	AA2 (10-2) C7	AA2 (10-2) C6	AA2 (10-2) C5	AA2 (10-2) C4	AA2 (10-2) C3	AA2 (10-2) C2	AA2 (10-2) C1	GC 1%	200ul dH20
E	negative control	GC 1%	AA2 (10-2) C8	AA2 (10-2) C7	AA2 (10-2) C6	AA2 (10-2) C5	AA2 (10-2) C4	AA2 (10-2) C3	AA2 (10-2) C2	AA2 (10-2) C1	GC 100%	200ul dH20
F	negative control	GC 1%	AA3 (10-2) C8	AA3 (10-2) C7	AA3 (10-2) C6	AA3 (10-2) C5	AA3 (10-2) C4	AA3 (10-2) C3	AA3 (10-2) C2	AA3 (10-2) C1	GC 100%	200ul dH20
G	negative control	GC 1%	AA3 (10-2) C8	AA3 (10-2) C7	AA3 (10-2) C6	AA3 (10-2) C5	AA3 (10-2) C4	AA3 (10-2) C3	AA3 (10-2) C2	AA3 (10-2) C1	GC 100%	200ul dH20
H	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20	200ul dH20

AA1 – AA3 Antituberculous agent 1-3; GC, growth control; GC 100% corresponds to the same inoculum as in the drug containing wells; GC 1% corresponds to the hundredfold diluted inoculum; negative control is 200 µL of 7H9-OADC; dH20, sterile distilled water.

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8 Molecular typing of *Mycobacterium tuberculosis* complex isolates

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8.1 Background and principles

In the last two decades, multiple molecular typing methods for *Mycobacterium tuberculosis* complex (MTBC) isolates have emerged, with different levels of reproducibility, discriminative power and demands on technical expertise.

Previously, phage typing, anti-TB drugs susceptibility profiling and other biochemical features allowed the differentiation of only a very limited number of strains, while nowadays, DNA-based typing techniques can potentially discriminate even individual transmission events. At present, DNA fingerprinting supports routine contact tracing in many countries as well as studies on person-to-person transmission, early disease outbreak identification, high transmission risk groups, laboratory cross-contamination [1, 2] and the distinction between reinfection and reactivation [3, 4]. In particular DNA fingerprinting of *M. tuberculosis* has greatly improved the understanding of TB transmission, which is especially important for drug-resistant strains. Moreover, the recognition of genotype families has facilitated studies on the population structure of MTBC and its transmission dynamics.

The various DNA fingerprinting methods available serve different purposes and have variable characteristics that enable their use in specific applications. Legacy typing approaches interrogate defined and highly variable targets in the genome, and include methods such as spoligotyping, MIRU-VNTR (Mycobacterial Interspersed Repetitive Units – Variable Number of Tandem Repeats) typing and IS6110 RFLP (Restriction Fragment Length Polymorphism). The advent of whole genome sequencing (WGS) based on Next Generation Sequencing (NGS) technology allows genome-based typing thus offering the highest possible discrimination and resolution power for molecular epidemiology of MTBC. Conversely, a targeted NGS approach interrogates small portions of the genome, but it can nevertheless yield phylogenetic classification, spoligotype data, and resistance profile prediction from primary patient samples. This chapter aims to describe the characteristics of these methods and their most important applications.

Although the application of DNA fingerprinting has improved our knowledge of the natural history of TB infections and the disease dynamics, there are still open questions. All DNA fingerprinting methods operate according to different molecular clocks⁵ and the stability of DNA profiles has been studied extensively, but is not fully understood [5]. Ideally, from the perspective of molecular typing, every transmission results in a slight change in the DNA fingerprint, while the strains remain recognisable, which makes it possible to distinguish primary sources in a chain of transmission from secondary and tertiary ones, enabling much more detailed analysis of transmission in a given area. Unfortunately, mutations in the genome of MTBC occur according to a stochastic process and therefore DNA fingerprinting will never be a perfect tool in studies on transmission. However, the application of e.g. WGS of *M. tuberculosis* isolates for the visualisation of transmission chains has demonstrated the discriminatory power of this approach. In addition, genome data have successfully been used to infer evolutionary dynamics especially towards resistance, and it may even help to unravel epidemiological linkages across time and space.

Below is an outline of the three traditional DNA fingerprinting methods: spoligotyping, MIRU-VNTR typing and RFLP typing, followed by a description of WGS and targeted NGS.

8.2 IS6110 RFLP typing

The restriction fragment length polymorphism (RFLP) method for typing bacterial strains was the first DNA-strain typing method that proved to be suitable for studying transmission [6]. It is based on the fact that the number of IS6110 mobile insertion sequences of 1.35kb present in the genome of strains differs from 0 to approximately 30. These genomic insertion sites themselves are also highly variable in MTBC strains, resulting in highly variable banding patterns.

The generation of RFLP patterns is technically demanding and time consuming, and requires a high amount (i.e. 2 µg) of purified genomic DNA as starting material. The process involves restriction enzyme cleavage of the DNA; fragment

⁵ A measure of evolutionary change over time at the molecular level that is based on the theory that specific DNA sequences spontaneously mutate at constant rates. This measure is chiefly used for estimating how long ago two related organisms diverged from a common ancestor.

separation by electrophoresis; the transfer of the fragments to a DNA membrane and hybridization with a DNA labelled probe complementary to the *IS6110* transposon sequence, and final visualisation of the results on a light-sensitive film. Each individual step of the process is crucial for the final result, which also gives an idea of the difficulties experienced with regard to inter-laboratory comparability.

In addition, the analysis of *IS6110*RFLP patterns with the specialised software like BioNumerics is complex, requiring experienced users. However, the difficulties associated with RFLP typing have not altered the fact that RFLP typing has revolutionised our understanding of TB transmission. Moreover, for strains of particular genotype families such as the Beijing clade, the level of discrimination of RFLP typing is still superior to that of the more recently introduced 24-loci MIRU-VNTR typing [7-9].

8.3 Spoligotyping

Spoligotyping is based on polymorphisms in the direct repeat (DR) locus of the mycobacterial chromosome [10, 11]. The well-conserved 36-bp direct repeats are interspersed with unique spacer sequences, varying from 35 to 41 bp in size. The order of the spacers has been found to be well conserved [12]. Currently, 94 different spacer sequences have been identified, 43 of which are used in the first-generation spoligotyping for *M. tuberculosis* complex strains [13]. After amplification, the denatured PCR products are applied in the reversed line on a membrane with covalently bound multiple synthetic spacer oligonucleotides deduced from DR region sequences of the two control strains (*M. tuberculosis* H37Rv and *M. bovis* BCG P3).

Clinical MTBC isolates can be differentiated by the presence or absence of one or more spacers. Almost all strains reveal a few of these spacers. The obtained patterns – 43 spacers present or not – are usually characteristic of a particular genotype family [14]. Spoligotyping is therefore a simple, cheap, rapid and reproducible [15] tool to study the phylogeny of MTBC strains or to associate phenotypic features of isolates with the genotype family the bacteria represent [11]. The level of discrimination of spoligotyping is generally low and it is necessary to be cautious when using this method to examine the TB transmission at the strain level [16, 8]. However, it is possible to use spoligotyping as a screening method in typing to rule out transmission when different spoligotyping profiles are obtained. On the other hand, when spoligotype patterns are identical, then no conclusion can be drawn regarding the epidemiological relationship between the respective patients. The SpolDB4 database is one of the largest publicly available databases on MTBC and contains spoligotype patterns from approximately 40 000 clinical isolates representing 122 countries.

8.4 MIRU-VNTR typing

The mycobacterial interspersed repetitive unit variable number tandem repeats (MIRU-VNTRs) assessment of bacterial strains has proven to be a suitable method for the detection of genetic polymorphisms within bacterial species [17]. Differences in the number of tandem repeats in the 24 stretches of the genome of MTBC strains are the basis for this internationally recognised typing method. In multiplex PCRs, up to 24 loci are amplified using primers specific for the flanking regions of each repeat locus, after which the size of the amplified stretches of tandem repeats are used to determine the number of tandem repeats present. The number of tandem repeats detected at the different loci results in a numerical code that serves as a DNA fingerprint of the respective TB bacteria and allows users to easily exchange the MIRU-VNTR typing results and perform inter-laboratory comparisons.

The determination of the PCR products in MIRU-VNTR typing can be done manually by performing all single-locus PCRs, and then interpreting the product length using electrophoreses. However, nowadays, MIRU-VNTR typing kits containing all the reagents necessary to perform MIRU typing on 24 VNTR loci, in accordance with the international standard are also commercially available (e.g. GenoScreen, Lille; France) and automated and higher throughput methods are widely available, which allow to determine the size of the PCR products by sequencing using dye labelled primers, thus saving time and increasing accuracy.

The 24 loci MIRU-VNTR typing allows to distinguish unrelated strains and provides the clonal stability to reliably identify isolates from the same transmission chain [18].

Open databases where users can add MIRU-VNTR patterns for the online analysis of the clonal identification of MTBC isolates are widely available (e.g. MIRU-VNTR*plus* database) [19]. Cluster analysis can also be done locally using the BioNumerics software (Applied Maths, Ghent, Belgium). National molecular epidemiology databases with embedded cluster analysis tools exist in several EU countries. Cluster analysis tools are also available within international databases and epidemiology reporting tools, e.g. TESSY.

Detailed protocols on multilocus MIRU-VNTR typing are available online at <https://www.miru-vntrplus.org/MIRU/files/MIRU-VNTRtypingmanualv6.pdf>

8.5 Whole genome sequencing

Although classic genotyping methods target highly variable genetic elements, they only interrogate a small fraction of the genome. Therefore, they cannot capture microevolution potentially occurring in other genomic regions. In contrast, WGS based on NGS gives access to nearly complete genome sequences. The extended use of such approach for research and epidemiological control has become possible through the rapid improvements and increasing affordability of NGS technologies. So-called 'benchtop' NGS systems generating sequencing data in the range of 10-20 Gb can be integrated into a routine workflow, with a throughput adapted to a routine microbiological laboratory [20, 21]. In addition, commercially available easy-to-use kits for DNA library preparation and sample multiplexing require only small amounts of genomic DNA inputs, which shortens the time needed to get enough material for DNA extraction from culture of slow-growing mycobacteria.

The overall workflow for WGS starts from purified genomic DNA extracted from bacterial culture. This is converted to a set ('library') of fragments of a defined size with specific adapter sequences at both ends of the fragment. Libraries are loaded onto NGS instruments resulting in millions of short sequence reads per sample. Using a dedicated bioinformatics pipeline, these data can be used to detect genomic mutations compared to a reference genome (usually that of *M. tuberculosis* H37Rv). For genotyping and similarity/relatedness analysis, the respective set of single nucleotide polymorphism (SNP) variants is often used. An interesting alternative from WGS data is the core genome multi locus sequence typing (cgMLST) approach, which offers inherent standardization and easy incorporation of new datasets into a comparison enabling ongoing real-time surveillance [22]. Implementation of a suitable analysis pipeline should follow accepted standards and include evaluation against reference datasets of established analysis solutions [23, 24].

Interestingly, due to the nature of NGS technology and its limitations, it is possible to extract spoligotype patterns from whole genome sequencing data, whereas MIRU-VNTR and IS6110 fingerprinting types cannot be reliably inferred from NGS data.

Several studies demonstrated that whole genome-based approaches provide more resolution than classical genotyping (e.g. 24-loci MIRU-VNTR typing and IS6110 fingerprinting) [25-28, 21]. In particular, whole genome sequence analysis allows for a more precise differentiation of isolates belonging to a specific recent transmission chain from other, closely but not directly related isolates, which is especially useful in resolving complex outbreak situations [27, 28, 21]. In such situations, the phylogenetic trees based on the obtained genome-wide SNP or cgMLST data of the isolates correlate much better with the available epidemiological data, and the spatio-temporal distribution patterns of the corresponding TB cases, than trees based on classic typing data [28, 21, 29-31]. In addition to a better time-dependent signal, the higher resolution offered by whole genome-based analysis also provides other valuable information. For example, the presence of a particularly contagious case (i.e. a super-spreader), leading to multiple secondary cases, can be inferred by the observation of star-like topologies in the phylogenetic tree, where clonal variants only differing by a few SNPs or alleles and corresponding to secondary cases branch directly from a central node, representing the same single source case.

Furthermore, the identification of vacant nodes in the tree topology is suggestive of undiagnosed cases in the population. The unidirectional accumulation of SNPs allows a more clear-cut association of new cases with previous cases in a longitudinal outbreak, which may allow for more targeted contact tracing investigations [28, 21]. This can be of use for the precise determination of the source cases for patient isolates of particular importance, such as MDR isolates.

To identify and delineate recent transmission chains based on whole genome sequences, a key parameter to calibrate is the level of genome-wide variation that occurs in *M. tuberculosis* strains within and between infected individuals over time. In different studies, the observed levels of divergence between such longitudinal isolates collected from chronically-infected patients or from epidemiologically-linked cases rarely exceeded three to five SNPs, thus defining a cut-off range for predicting recent transmission [26, 28, 21]. The calculated mean rate of change in DNA sequence was approximately 0.5 SNPs per genome per year, providing a quantitative estimate of the short-term evolution rate of *M. tuberculosis* in the human host population [28, 21]. Interestingly, this rate matches the mutation rate estimated for *M. tuberculosis* in a macaque infection model [32]. Given sufficient data, a dedicated analysis pipeline can unravel outbreak dynamics and estimate the temporal and spatial distribution and evolutionary history of a phylogenetic clade [33-37].

Compared to classic genotyping, another major advantage of whole genome sequencing is that it simultaneously provides direct and valuable information for predicting drug resistance, as well as highest resolution genotyping [38, 20]. This combination of diagnostic and epidemiological information in a single assay is a great benefit, especially with *M. tuberculosis* for which early detection of drug resistance is important. Analysis based on whole genome sequencing goes far beyond conventional molecular tests, focusing on known selected mutations in hotspot regions of genes involved in resistance to first and second-line anti-tuberculous drugs, thereby possibly missing novel resistance-associated mutations [39]. In principle, WGS captures most, if not all the gene sequences determining the so-called *M. tuberculosis* resistome, which allows interrogation of all known mutations associated with drug resistance [20]. Moreover, whole genome sequencing at high coverage may detect the emergence and

co-existence of different drug resistance- conferring mutations before selection and fixation of a final mutant, in possible combination with compensatory mutations [40, 41]. Such detection is of clinical relevance as the co-existence of wild-type and mutant subpopulations resulting in hetero-resistance may confound the current phenotypic and molecular drug resistance tests [40, 41] as well as conclusions on transmission or secondary acquisition of drug resistance. In general, the inference of a resistance profile from whole genome data relies on well curated databases capturing resistance-associated mutations and ideally also variants not linked to resistance. Interpretation of the molecular variants leading to resistance to bedaquiline and delamanid is more complex and requires the evaluation of a large number of resistant strains to reach the same confidence we have for rifampicin. Recent initiatives led by the ReSeqTB and CRyPTIC consortium together with WHO led to the implementation of curated and up-to-date resources for resistance-associated genetic markers [42-45].

8.6 Targeted NGS

In addition to whole genome characterisation, next generation sequencing can also be used together with a specific enrichment or amplification protocol to enable complex primary patient samples as input and vastly increased sensitivity. The option to work directly with patient samples reduces turn-around times and still reaching similar levels of sensitivity compared with classical genotyping and molecular resistance detection methods [46, 47]. Of course, the restriction to a defined set of genomic regions will not allow the same level of resolution power as WGS. Still, general classification such as mycobacterial species identification, lineage determination, and spoligotyping are possible given respective genomic regions are targeted (e.g. Deeplex MycTB, Genoscreen). This approach enables at least to rule out transmission chains when different spoligotypes and phylogenetic SNPs are detected among strains and may be suggestive of certain relatedness when same phylogenetic SNPs / spoligotypes (and DR patterns) are identified among samples epidemiologically linked or for clonal strains circulating in restricted settings. While commercially available targeted NGS solutions can be employed directly on DNA extracted from patient samples, the sensitivity threshold is currently still not well defined, but microscopy positive samples can commonly be analysed successfully.

8.7 General considerations on genotyping methods

The five DNA genotyping methods here described can be used for different applications. Spoligotyping is generally used in studies to reveal the genotype family of the respective bacteria; whereas this technique is less suitable for strain typing. Both RFLP and 24-loci-MIRU-VNTR typing have a higher level of discrimination and reproducibility and can be used for strain typing. The turn-around time of MIRU-VNTR typing is significantly shorter than that of RFLP typing and it is also technically far less demanding. In addition, MIRU-VNTR typing can be used in contact tracing and source-case finding and can reliably rule out transmission.

WGS has become available with the introduction of NGS, and the respective technology is still quickly evolving. Notably, WGS offers the highest possible resolution for outbreak analysis and tracing transmission chains. The compilation of extensive data about resistance-associated genomic variants also offers the possibility to infer a comprehensive resistance profile from both whole genome and targeted NGS data.

Although NGS-based molecular tracing of MTBC has a vast potential, its generalised use is still hampered by some remaining challenges. Despite rapid cost decreases, NGS analyses still remain too costly for many TB laboratories, especially since the implementation of NGS analysis carries high initial investment costs. For most NGS approaches, costs are also directly affected by the batch size and multiplexing, as performing incomplete sequencing runs significantly increases the cost per sample. Another important factor relates to the integration of NGS into existing laboratory workflows. It has been shown that when including resistance analysis, WGS can actually be cost-competitive with standard approaches and an interesting option also for low-middle income settings [48, 49].

In addition to the technical training and specialised skills required for performing the 'wet' component of the NGS workflow, the need for expert guidance on sequencing data analysis and interpretation should also be considered. Moreover, the 'dry' component of the NGS workflow requires a dedicated IT infrastructure for data management and data storage. In contrast to classic genotyping, technical and analytical modalities (e.g. the precise delimitation of the genome sequence that is taken into account in the analysis; the minimal sequence coverage; etc.) have not yet been standardised [24]. Consequently, datasets generated by different laboratories are not yet directly comparable, and universal databases are not yet available (e.g., for multicentric longitudinal epidemiological studies and surveillance). While WHO recently published a comprehensive catalogue of resistance-associated mutations together with the CRyPTIC consortium [45] the underlying bioinformatics functionality for the detection of resistance-conferring variants is not yet standardised [24].

As an important point, the input material required for WGS analysis of TB samples can only reliably be gained from cultures. Even modern workflows require at least 1 ng of pure DNA, and the process is relatively sensitive to contaminations as there is no specific amplification of genomic targets. This is partially overcome by targeted NGS, which includes specific amplification of targeted genomic regions. However, targeted NGS will not allow for high resolution genotyping.

Contact tracing complemented by genotyping is considered to be important for understanding person-to-person transmission. It is less clear, however, to which extent genotyping itself is cost-effective and if it has any added value beyond contact tracing from an immediate public health point of view [50].

The performance of DNA fingerprinting has also been used for predicting the size of future clusters following the detection of the first two cases of a new cluster. Time between the cases, age, and location are variables that become known shortly after the diagnosis of a new TB cluster. By combining the molecular data and the patient's registration data, new cluster episodes can be predicted using the risk factors. This information can contribute to early warning systems for the national health services [51].

DNA fingerprinting can detect possible cross-contamination occurred in the laboratory. If a laboratory detects two isolates with identical DNA profiles in only one week, this usually indicates a sampling or laboratory mishap [1]. In this case, the clinician should be asked to review the clinical picture of the patient, and the microbiologist should check the positivity rate of the culture and whether the cultures with identical DNA fingerprints have been contaminated at a particular stage. Regular checks of positive cultures are recommended to detect this common problem; in the Netherlands, about 3% of all positive cultures are cross-contaminations [52].

The widespread application of DNA fingerprinting has provided substantial insights into TB transmission, especially when conventional epidemiological investigation and molecular typing are combined [53]. The strong association of TB transmission with gender and lower age of the source case in a low-prevalence setting has been shown by molecular fingerprinting techniques [54]. Moreover, studies on transmission within and outside households in South Africa [29,31] has yielded important insights into the origin of TB infections in a high-prevalence setting [55, 56]. There is also an increased risk that previously treated and cured TB patients will develop TB again when reinfected [57].

Recent work demonstrated the high-resolution power of WGS especially for MDR-TB outbreaks, and enabled unravelling the evolution of phylogenetic lineages towards transmissibility and resistance acquisition [58, 33, 59, 36].

8.8 Materials

Purified DNA from MTBC bacteria is generally the best material for molecular typing. For methods based on DNA amplification, such as spoligo- and MIRU-VNTR typing, only a small amount is needed as starting material. Even purified DNA from a sufficient number of bacteria in clinical material will result in a typing pattern. However, briefly incubating these bacteria in a liquid culture medium will generally yield more reliable and reproducible results, which is advisable because of the costs and the time to perform the typing techniques. For RFLP typing, based on the specific restrictions of the isolated unamplified DNA, a fully grown culture is needed as 2 µg of highly purified genomic DNA are required. In a diagnostic setting, it will take several weeks to achieve this amount of growth after TB detection.

For WGS, state-of-the-art library preparation kits require as little as 1 ng of DNA as input if using for example the Illumina sequencing platforms, but it can be significantly higher if using e.g. the MinION sequencer from Oxford Nanopore Technologies. Genomic DNA suitable for WGS can be reliably extracted from early positive liquid (MGIT) cultures [48]. Still, subculture of a positive primary specimen is a sensible step to minimise DNA contaminants from the bacterial fauna and human material. Targeted NGS can be directly employed on DNA extracted from patient samples.

Membranes, reagents and positive control DNA for spoligotyping can be purchased. The performance of spoligotyping is better, especially in laboratories with molecular experience, and the results are more comparable among different laboratories [60].

For 24-loci MIRU-VNTR typing, a kit containing all reagents for eight multiplex amplification reactions can be obtained commercially. In addition, the PCR product fragments are analysed on an automated DNA sequencer. In the absence of an expensive analyser, the 24-loci MIRU-VNTR can be used with single or multiplex amplification reactions and detection of the product, either automated or manually (see standard operating procedure on MIRU-VNTRplus website at <http://www.miru-vntrplus.org>). The quality of international MIRU-VNTR typing performance and the inter-laboratory reproducibility has significantly improved over the last years [61].

Both WGS and targeted NGS represent complex multi-step protocols requiring specific oligomers, enzymes and reagents to convert the initial DNA sample into an NGS library, and of course the sequencing instrument itself as well as a variety of auxiliary equipment. A variety of commercial solutions exist that enable the preparation of a NGS library from bacterial DNA samples. Likewise, there are various NGS instruments available, most notably also several benchtop sequencers with smaller throughput and relatively higher sequencing costs but also smaller footprint in physical dimensions and investment costs [38].

8.9 Results/interpretation

The analysis of *IS6110* RFLP patterns by BioNumerics software is technically demanding [13]. The most important aspect is the inclusion of internal (a mixture of two molecular markers) and external standards (DNA of a control

strain with a suitable range of bands) for normalisation and accurate reading of the band sizes. RFLP typing consists of a lengthy, multi-step laboratory procedure that is prone to error. Poor quality in RFLP typing can be connected to laboratory technique, but very often is caused by the incorrect interpretation of results with the pattern analysis software.

Spoligotype patterns are codes of 43 possible digits and can be sorted with standard software such as Excel. However, most institutions use the BioNumerics software (Applied Maths, Kortrijk, Belgium), which is able to compare the results of any typing method. There is an international database of spoligotype patterns [13, 62] which holds tens of thousands of spoligotype patterns that can be used to compare the locally obtained typing results with patterns that have been found elsewhere and are already labelled with a genotype family designation.

The result of MIRU-VNTR typing is a numerical code of usually 12, 15, or 24 numbers which can be analysed relatively easily, as described above in the section on spoligotyping. In epidemiological investigations local comparisons are best done using the BioNumerics software for which special plugins are available. There are several international databases with MIRU-VNTR typing results in which locally obtained results can be compared with international collections (such as <http://www.miru-vntrplus.org> or http://www.pasteur-guadeloupe.fr:8081/SITVIT_ONLINE).

Due to the nature of the raw data produced by NGS instruments, any analysis requires a bioinformatics solution. This encompasses both dedicated software tools as well as suitable IT hardware for calculation and data storage. For genotyping based on whole genome data, a cgMLST approach is an interesting option with both open source and commercial software (e.g. Ridom SeqSphere+ Software, Ridom GmbH, Münster, Germany) solutions available for data analysis.

8.10 Quality control

Quality control of DNA fingerprinting is of the utmost importance. On all occasions, first-line controls should include strains with a known DNA fingerprint. In the case of spoligotyping, DNA of H37Rv and P3 should be included in each test to assess the performance of each of the 43 spacer oligonucleotide probes present on the blot. In MIRU-VNTR typing, a strain with a known typing profile should be included. In RFLP typing, a particular strain with a wide range of bands should be used in each test to check the normalisation [60].

A second-line control procedure is also advised and should include the blinded exchange of a set of DNA samples with another laboratory twice a year to test the reproducibility. As a third-line of control, a blinded set of DNA samples supplied by an international organisation to multiple institutes should be analysed to test proficiency in the given procedure [60].

For NGS, ideally, all steps of the workflow, from DNA extraction to sequencing, data analysis and reporting, should be standardised and well documented, and an external quality assessment programme should be in place to ensure the generated data meet international standards [24]. Since 2017, an EQA scheme for MTB WGS has been implemented within the ERLTB-Network to allow comparison of performance and results among different laboratories, to provide objective evidence of testing quality, and to identify areas for improvement and training needs [63].

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9 Use and validation of disinfectants for *Mycobacterium tuberculosis*

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Revised by Thomas Pottage and Allan Bennett (2015)

9.1 Liquid disinfection

There are two standard methods for the analysis of disinfectants against mycobacterial organisms including *Mycobacterium tuberculosis* (TB); they are the quantitative suspension and quantitative carrier test. These methods have been documented as industry standards: suspension testing EN 14348:2005 and carrier testing

EN 14563:2008. The principle of the suspension test is to determine the efficacy of a chemical disinfectant against mycobacterial organisms within a suspension at a given temperature and over a set period of time. The carrier test is used to determine the efficacy of a disinfectant against a dried solution of *Mycobacterium* on a glass slide over a set period and at a set temperature. The strains of *Mycobacterium* specified in both standards are *Mycobacterium avium* (ATCC 15769) and *Mycobacterium terrae* (ATCC 15755), with *M. terrae* specified as the surrogate for *M. tuberculosis*. These micro-organisms are specified due to their lack of pathogenicity and there seems to be no reason why the standards could not be adapted for use with TB strains, simply by replacing the surrogate organism with the desired TB strain. However, replacing the strains would mean that the test would not follow the standard method and could not be designated as such. The standard methods, using TB surrogates, are summarised in the following sections. For full details the standards can be obtained from the relevant national standards body.

This could not be stated as tested to the standard but to a method based closely on the standard, and would give more accurate TB disinfection results since the disinfectant would be tested on the TB strain that would be used in vitro.

9.1.1 Suspension test: EN 14348:2005

A test suspension of the mycobacterial agent(s) should be prepared to give a suspension concentration from

1.5×10^9 CFU/ml to 5.0×10^9 CFU/ml. This suspension should be used on the day it is prepared. To count the stock suspension, dilutions of 10^{-7} and 10^{-8} should be prepared and 1 ml of the dilutions should be distributed as 0.5 ml onto 2 MCO (Middlebrook and Cohn 7H10 medium with 10% OADC enrichment) agar plates. This process should be performed in duplicate. The plates should then be incubated at $37 \pm 1^\circ\text{C}$ for 21 days.

The disinfectant test solutions should be prepared at 1.25 times the required test concentration. This is because during testing, with the addition of the validation suspension (1 ml) and interfering substance (1 ml) to 8 ml of the disinfectant, the concentration will be diluted to 100% of the test concentration. A further two concentrations of the product should also be prepared for testing, with at least one of these concentrations being weaker than the active range concentration. The test product suspensions should be made up in hard water to the required concentration.

Test products which are supplied ready-prepared for application (e.g. pre-soaked wipes) can be used but it should be noted in the test report that the concentration during testing will be equal to 80% and that this is the highest concentration achievable. Further dilutions of these ready-to-use products should be made using distilled water instead of hard water.

Product test solutions should be prepared for each test and used within two hours of preparation.

The test should be performed under two different conditions relating to the loading of organic material. This clean solution is prepared by adding 0.3 g of bovine albumin fraction V to 100 ml of the diluent (Tryptone sodium chloride solution, 1 g Tryptone, 8.5 g sodium chloride in 1 000 ml water). The clean conditions will require the addition of 1 ml of a 3 g/l solution of Bovine albumin fraction V solution to 9 ml of the test solution and test product. This will give a final solution concentration of 0.3 g/l in the 10 ml test solution.

The dirty conditions are achieved by combining a solution of bovine albumin fraction V and high concentration sheep erythrocytes. The solution is produced by dissolving 3 g of bovine albumin fraction V in 97 ml of diluent to give a solution of 30 g/l. This is diluted 1 in 10 during the test to give a final solution of 3 g/l. At least 8 ml of fresh defibrinated sheep blood should be centrifuged at 800 g for 10 minutes. The supernatant should be discarded and the erythrocytes re-suspended in the diluent. This step should be repeated at least three times until the supernatant has no colour. The erythrocytes can then be re-suspended and 3 ml added to 97 ml of the bovine albumin solution. The solution can be kept for seven days at a temperature ranging from 2°C to 8°C .

The suspension test is performed by first pipetting 1 ml of the clean or dirty interfering solution into a sterile tube followed by 1 ml of the test suspension. This tube should then be placed in a water bath at the set temperature of

20°C for two minutes. After the two minutes, 8 ml of the test product should be added to the tube and the time started. The tube should be mixed at the beginning of the 60-minute test period and just before the end of the period.

After the 60-minute test period, a 1 ml sample of the test solution should be removed from the tube and pipetted into a fresh sterile tube containing 1 ml of water and 8 ml of validated neutraliser. Pre-validation tests should have been conducted to ensure that the neutraliser is capable of neutralising the test product and does not inhibit the growth of the test organism(s). The tube should be placed in a water bath at 20°C for five minutes.

After the neutralisation step, a 1 ml sample of the neutralised test suspension should be pipetted and distributed evenly between two MCO plates, in duplicate. A further 500 µl of the neutralised test suspension should then be taken and added to a tube containing 4.5 ml of the neutraliser. This will give a 10⁻¹ solution of the test suspension which should be diluted further to 10⁻³ using the neutraliser. A 1 ml sample should be taken from each of the dilution tubes and divided between two MCO plates. This should be done in duplicate, to give four plates with 500 µl of diluted test suspension on them. The plates should be incubated for 21 days at 37±1°C.

The procedures for the other concentrations of the test product should be performed at the same time.

9.1.2 Carrier Test: EN 14563:2008

The carrier test should be performed using either *M. avium* (ATCC 15769) and *M. terrae* (ATCC 15755) or only *M. terrae*.

A test suspension of the mycobacterial agent(s) should be prepared to give a suspension concentration from 1.5 x 10⁹ CFU/ml to 5.0 x 10⁹ CFU/ml. This suspension should be used on the day it is prepared. To counting the stock suspension, dilutions of 10⁻⁷ and 10⁻⁸ should be prepared and 1 ml of the dilutions should be distributed as 0.5 ml onto 2 MCO (Middlebrook and Cohn 7H10 medium with 10% OADC enrichment) agar plates. This process should be performed in duplicate. The plates should then be incubated at 37±1°C for 21 days.

The disinfectant test solutions should be prepared using hard water. Three preparations shall be prepared, one in the active concentration range and at least one below the active concentration range.

Test products which are supplied ready-prepared for application (e.g. pre-soaked wipes) can be used directly for the testing. Any further dilutions of these products should use water in the place of hard water.

The product test solutions should be prepared for each test and used within two hours of preparation.

The disinfectant should be tested under two different conditions, with clean and dirty solutions. The procedure for producing these solutions is mentioned previously in the suspension test method (Suspension Test: EN 14348:2005).

The carrier for the test should be a glass carrier which has been frosted on one side (dimensions 15 x 60 x 1 mm). The carrier should be cleaned with 70% ethanol, and then a 10 mm square marked on the frosted side. Finally it should be sterilised in a dry heat oven.

To inoculate the carrier 9 ml of the test suspension should be added to 1 ml of the interfering substance in a clean tube. The tube should be mixed and 50 µl should be pipetted into the inoculation square on the carrier and distributed evenly around the square using the pipette tip. The carrier should then be placed in an incubator at 36±1°C for a maximum of 60 minutes or until it is visibly dry. The drying time should be recorded in the report sheet.

The carrier test is performed by pipetting 10 ml of the test product solution into a screw cap tube (wide enough to accommodate the carrier slide). The tube should then be placed in a water bath at 20°C. The carriers should be placed into the tubes immediately after the drying process has finished, making sure the inoculation square is covered by the test product solution. The timer should be started immediately on immersion and the tube left for the 60-minute contact time.

After the 60-minute exposure period, the carrier should be transferred to a screw cap tube filled with 10 ml of neutraliser and 1 ml of glass beads (diameter from 0.25 mm to 0.5 mm). Place in a water bath at 20°C then mix for 15 seconds. The tube should then be left for a further four minutes 45 seconds, giving a total neutralisation time of five minutes. Pre-validation tests should have been conducted to ensure the neutraliser is capable of neutralising the test product and does not inhibit the growth of the test organism(s). The tube should be placed in a water bath at 20°C for five minutes.

After the neutralisation period 1 ml of the neutralised solution containing the re-suspended test organisms from the carrier should be removed and divided equally between two MCO plates. This process should be performed in duplicate. Remove a further 500 µl and transfer to a tube containing 4.5 ml of neutraliser. This will give a 10⁻¹ solution of the test suspension which should be further diluted to 10⁻³ using the neutraliser.

Take a 1 ml sample from each of the dilution tubes and divide this between two MCO plates. This should be performed in duplicate to give four plates with 500 µl of diluted test suspension on them. The plates should be incubated for 21 days at 37±1°C.

The procedures for the other concentrations of the test product should be performed at the same time.

9.1.3 Choice of method to be used

The standard to be used depends on the application for which the disinfectant product is being used. For example, a surface disinfectant should preferably be tested using the carrier tests as that most closely matches this application. However, carrier tests are often thought to be more time consuming, even though there are less complications with the neutralisation step. The recommendation for using glass slides in the carrier test may raise concerns about potential for accidental abrasions in the laboratory and other materials may be considered. The specified 60-minute contact time will not always reflect the use of a product and it may be more realistic to reduce this to a shorter period, especially for the carrier test.

9.2 Gaseous disinfection

Gaseous decontamination has been used in microbiological laboratory facilities to decontaminate biological safety cabinets, equipment and the laboratory itself between experiments and before servicing [1-3]. It is often recommended for dealing with emergency situations such as uncontrolled releases of liquid culture in laboratories. Gaseous decontamination uses a vaporised chemical to contact, and in many cases condense onto, the exposed surfaces within the enclosure. Traditionally, gaseous lab decontamination has been undertaken using formaldehyde [4], where paraformaldehyde crystals are heated to sublimation or liquid formalin solution is boiled, both releasing formaldehyde vapour into the enclosure. This vapour then condenses onto the exposed surfaces and shows its disinfection properties by alkylating protein molecules when it binds to the primary amide and amino groups [2,5].

Although it is effective as a fumigant against *M. tuberculosis*, even in sputum [6] and against *M. bovis* [7], formaldehyde is a potential carcinogen [4], requires a long aeration period (for removal of the vapour from the enclosure, unless it can be neutralised, or external ventilation is available) and the paraformaldehyde residues left on the surfaces can be labour intensive to remove [5,8]. These drawbacks to formaldehyde have led to investigations into the use of alternative gaseous decontamination technologies for fumigation.

Of the candidates for a replacement fumigant in place of formaldehyde the best studied is perhaps hydrogen peroxide. The gaseous hydrogen peroxide systems were originally marketed for use in pharmaceutical clean room facilities, but their use has broadened to include microbiological laboratories [4], animal facilities [9], spacecraft assembly facilities [10] and in the hospital environment [11]. Hydrogen peroxide decontamination as a process has numerous advantages over formaldehyde fumigation. It leaves no residues, has better operator safety and is less damaging to the environment. Hydrogen peroxide works as an oxidising agent which produces hydroxyl radicals and superoxide anions [4] which can attack the cell's DNA and lipids, but being highly reactive the hydroxyl radicals will also react with other inorganic matter and materials [2].

Hydrogen peroxide is marketed for fumigations in three ways: Vaporised Hydrogen Peroxide (VHP), Hydrogen Peroxide Vapour (HPV) and aerosolised Hydrogen Peroxide (aHP). Hydrogen peroxide plasma sterilisation, which introduces vaporised hydrogen peroxide in a small vacuum chamber where the contaminated equipment is placed, has been shown to be effective for the decontamination of contaminated instruments, such as bronchoscopes [12].

9.2.1 Vaporised hydrogen peroxide and hydrogen peroxide vapour

Gaseous hydrogen peroxide is produced by heating and vaporising liquid hydrogen peroxide. Liquid hydrogen peroxide solutions have already been shown to be an effective decontaminant against TB [13,14]. Two of the more established companies have different approaches for the use of the vaporised hydrogen peroxide: Steris' generators use VHP and Bioquell's HPV.

The major differentiating factor between the two systems is the presence of microcondensation on the surfaces of the enclosure being fumigated. Steris' VHP technology dehumidifies the air within the enclosure prior to injection of VHP. This decreases the dew point in the enclosure and allows VHP to be injected without forming condensation on the surfaces, meaning although VHP is injected the system is designated 'dry'. The HPV technology designed by Bioquell operates in a similar fashion to formaldehyde fumigation, where HPV is injected into an enclosure above the dew point to allow for the formation of microcondensation on the surfaces. The microcondensate is a microscopic layer of hydrogen peroxide approximately 2–6 µm in thickness. Table 14 below describes more differences between the two technologies.

Table 14. Differences between the two major gaseous hydrogen peroxide technologies, Steris VHP and Bioquell HPV

Parameter	Steris	Bioquell
Description	Vaporised hydrogen peroxide	Hydrogen peroxide vapour
Condensation formed?	No	Yes
Requires dehumidification of enclosure?	Yes	No
Period of hydrogen peroxide injection	Continuous through exposure phase	One injection prior to decontamination phase
Volume of hydrogen peroxide needed	Large	Small
Generator located internally or externally in relation to enclosure	External	Internal or external (dependent on generator chosen)
Remote activation and monitoring	Yes (laptop needed)	Yes (control unit supplied extra)

Both VHP and HPV technologies have been shown to be efficacious against a wide range of bacteria, viruses and prions [2,9,11,15-19], often in company sponsored trials. Gaseous hydrogen peroxide has also been shown to be effective at killing TB within a biological safety cabinet and high-level containment laboratories [4,20]. The concentration of TB used by Hall on each indicator was approximately 103 cells, this is low in comparison with the 106 *G. stearothermophilus* spores on each of the other biological indicators. It was argued that the lower number of cells was more likely to equate to what would be remaining in a laboratory spillage after initial cleaning procedures were followed [20]. Kahnert's investigation found that even if a higher concentration of TB cells were dried onto biological indicators (ranging from 8.0x10⁴ to 2.3x10⁶ CFU), no organisms were recoverable after VHP exposure [4].

9.2.2 Dry mist hydrogen peroxide

Another way of aerosolising hydrogen peroxide within an enclosure for fumigation is to use an aerosolised hydrogen peroxide generator. An example of the generator, Glosair 400, is produced by Advanced Sterilisation Products. The aHP fumigation technique has been demonstrated to be effective against methicillin-resistant *Staphylococcus aureus*, *Acinetobacter baumannii*, *Clostridium difficile* and TB [21-24]. A comparison between aHP and 0.5% sodium hypochlorite solution was made by Barbut, which found the aHP more effective in killing *C. difficile* spores [22]. The aHP technology has also been successfully demonstrated against TB that had been dried onto stainless steel carriers (varying concentrations from 5x10⁵ to 5x10⁶ CFU/ml) and placed around a biosafety level 3 laboratory [25]. In Grare's study 5% hydrogen peroxide was used with a 60-minute exposure period to kill TB, whereas a study performed by Andersen under similar conditions demonstrated that the TB was not killed after aHP exposure [26]. It was hypothesised that the difference in results might be due to the preparation of the TB indicators. In Andersen's study, TB was dried from a saline solution whereas in Grare's experiments TB was dried from a distilled water solution which might have weakened the cell membrane, making it more susceptible to aHP decontamination [25,26].

9.2.3 Plasma hydrogen peroxide

A technology which can be employed to decontaminate smaller instruments rather than entire rooms or laboratories is hydrogen peroxide plasma sterilisation. The decontamination process uses a small vacuum chamber which is filled with vaporised hydrogen peroxide. After the vapour has diffused in the chamber, electromagnetic radiation is introduced to break the hydrogen peroxide molecules apart, inducing a plasma state and producing hydroxyl reactive species. This technology has been shown to be effective for sterilising bronchoscopes that had been contaminated with TB and initially decontaminated in a washer/disinfector, compared to bronchoscopes that had only been cleaned using a standard washer/disinfector cycle [12].

9.2.4 Use of gaseous disinfection in accident scenarios

Spillage of pathogenic microorganisms in a laboratory outside primary containment equipment should be an extremely rare event and should be prevented by employing the proper procedures and practices. The use of glass should be discouraged and samples should be contained when removed from primary containment using transport containers or bagging solutions. However, there may be occasions where these precautions are impractical [25].

The reaction to a spill of TB in a laboratory will depend on many factors, including the sample type (diagnostic specimen/positive, MDR TB, titre if known, staff exposed, location, etc.) and so any recommendation is based on a local risk assessment. However, the following course of events will occur:

- An aerosol will be generated which will be gradually removed by deposition or dilution.

- There will be uncontained liquid on the laboratory floor, localised in a pool where the container lands, and in the form of splashes and deposited aerosols which will be widely dispersed on the laboratory floor and potentially on other surfaces.
- Laboratory staff exposed to the spillage should leave the location, remove contaminated garments and wash any exposed areas of skin immediately. The laboratory should not be entered until any aerosol has been removed. The time allowed should be based on information about the sample volume, titre and room ventilation rate.
- There are three ways that the spill can be decontaminated.
- Immediate gaseous disinfection, preferably using a remote system (formaldehyde or hydrogen peroxide). The advantage of this system is that no operator is exposed by re-entering the area. However, the disadvantage is that the gaseous disinfectant may not penetrate into the highest concentration material of the spill. This area should be subjected to a final surface disinfection stage by an experienced laboratory worker wearing appropriate PPE before the laboratory is re-opened.
- After a suitable period of time an experienced laboratory worker wearing appropriate PPE enters the laboratory and decontaminates the spill site before setting off the gaseous decontamination. The laboratory can then be entered directly following the gaseous disinfection. The disadvantage of this approach is the exposure of the worker entering the laboratory.
- Surface disinfection alone could be undertaken by a worker in PPE including RPE. The disadvantages of this approach are that not all contaminated surfaces may be dealt with and the worker will be exposed.

Once again, the approach taken should be based upon a risk assessment carried out by experienced members of laboratory staff with biosafety expertise. It is preferable to have a risk assessment framework in place for such an eventuality.

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10 Information for physicians: laboratory diagnosis of TB

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10.1 Introduction

The TB laboratory provides crucial information for the diagnosis and management of people affected by TB disease.

Microbiologic data, based on culture or genomic tests, provide evidence to confirm clinical, radiological, epidemiological and pathology suspicions.

According to WHO guidelines for TB diagnosis, bacteriological confirmation should be attempted in all cases of pulmonary TB using WHO recommended diagnostics (WRDs) [1]. The same diagnostic tests could be used to provide bacteriological confirmation in cases of extrapulmonary TB (EPTB) when an appropriate specimen can be collected.

Therefore, optimal communication and understanding among the different clinical and laboratory specialists is of the utmost importance. The communication should include specification of the appropriateness of the different tests in the diagnostic algorithms, their negative and positive predictive values and how to translate the results into effective clinical management.

The whole process of collecting clinical specimens for the microbiological diagnosis is critical and each step may influence the result. Therefore, it is vital that collection and transport of diagnostic material should be optimal. The clinical material for laboratory analysis should always be accompanied by sufficient and correct patient information. In the case of presumptive pulmonary TB, the patient should be provided with the appropriate sputum pots and detailed instructions on how to produce the most suitable sputum for TB diagnostics. Only appropriate approaches for obtaining and transporting clinical specimens will result in the disease being successfully confirmed, which is also critical for surveillance and epidemiological purposes. The same rigorous process for collection and transport applies to EPTB specimens, often collected by invasive and not repeatable techniques. The material should be kept under well-defined temperature conditions and transported to the laboratory within defined time limits.

In the past 10 years several commercial molecular methods for detection of *M. tuberculosis* and identification of the drug-resistant variants have become the standard for the diagnosis of TB [1, 2]. Tests can be applied to a large variety of diagnostic specimens and can provide results in hours. With the implementation of the Next generation sequencing (NGS) technology coupled to target amplification, it is possible to identify markers of resistance to a large number of antimicrobials used in the therapeutic regimens directly from clinical specimens. Few commercial tests are available (see Chapter 6), and their applicability depends on the bacterial load present in the clinical specimen. Information on drug resistance to the main TB drugs becomes available in a few days from sample collection. The main limitations of those tests are linked to the bacillary load in the sample, to the lack of capacity to differentiate live from dead bacilli and to the suboptimal targets' inclusion in the tests.

Whole genome sequencing (WGS) from *M. tuberculosis* strains grown in culture has become the standard for epidemiological studies (see Chapter 8). Turn-around time is longer compared to targeted approaches, as it is linked to the availability of a positive culture. Data analysis is not fully automated, but many microbiology laboratories are now considering having a bioinformatic expert in the staff. Analysis of the sequencing data should be performed by standardised pipelines and a user-friendly report should be provided to the clinical team to optimise TB management (Figure 15). The report should include clear explanations on the determinants of drug resistance to the different drugs and how to manage the cases of discrepancies with phenotypic drug susceptibility testing (DST).

M. tuberculosis is a BSL3 pathogen, and each country has specific regulations on the safe transportation of these bacteria to referral laboratories. Communication on when and how a culture should be sent by a peripheral laboratory can help to reduce problems in transportation of this pathogen. Positive *M. tuberculosis* cultures can usually, but not always, reliably confirm a TB diagnosis. When a positive TB culture is reported from the laboratory from a sample collected from a person with very low suspicion of TB, it is very important to contact the laboratory to exclude clerical errors or laboratory cross-contamination. Cross-contamination rates (i.e. false-positives) of up to 3–5% of positive cultures have been reported. If misdiagnosis is suspected, communication between the laboratory and the physician is imperative to avoid unnecessary treatment. Often the presumptive TB diagnosis can be re-considered by the physician in the case of false-positive cultures. Noting and discussing cross-contamination and other problems in the diagnosis of TB helps to improve the quality of this procedure.

This chapter aims to discuss how the information generated in the laboratory should be shared with clinicians and how to maximise the contribution of the laboratory in the diagnostic process.

All the information included in this chapter is based on the tests in use today. The field of diagnostics for TB is rapidly evolving after many years of stagnation. We expect that in few years different samples and tools for diagnosis of pulmonary TB will become available.

10.2 General considerations regarding the diagnosis of TB

General considerations are based on WHO recommendations and guidelines [1, 2] and publications listed at the end of the current chapter.

Diagnosis of TB and drug-resistant TB (DR-TB) should only be performed in accredited laboratories with appropriate workload to maintain proficiency for all the tests offered. WHO recommends a molecular test (among the WRDs) to be performed on a respiratory specimen as a first test for the diagnosis of pulmonary TB in adults able to produce sputum [2]. On the same specimen liquid culture using automated machines should be performed. Clinical material (such as sputum) for the culture-based diagnosis of TB and the initial drug susceptibility testing should be collected before the start of treatment.

For children and people not able to produce sputum, gastric aspirate or molecular detection on stools can be attempted [3, 4].

For EPTB, highly sensitive molecular tests should be attempted [1]. Liquid culture in MGIT remains the most sensitive technique.

The microbiologist should inform their clinical counterpart that the diagnostic accuracy of a test may vary with the specimen, so for EPTB it is important, in the presence of an initial negative molecular test, to rely on the clinical judgement for treatment initiation, while waiting for culture results. Liquid cultures are reported negative if no growth is observed at 6 weeks. TB meningitis requires special attention because of the low volume of cerebrospinal fluid (CSF) often submitted to the laboratory, therefore, rapid molecular tests such as Xpert Ultra should be prioritised in combination with liquid culture, while microscopy is usually negative also when cytospin is used.

Mycobacterial blood culture is not performed in all laboratories. It is important to inform the clinical counterpart that the yield may depend on the proper sample collection and transport. Specific blood culture bottles or isolator tubes should be provided in advance. For laboratory diagnosis multiple blood cultures are needed.

Whenever possible, a rapid molecular test to identify resistance to rifampicin and isoniazid should be offered. If, for cost reasons, it is not possible to test all presumptive TB cases, then the following groups should be prioritised: relapses, retreatments, contacts of MDR TB patients or isoniazid-resistant cases, and cases from high prevalence DR-TB countries. If mutations conferring resistance to rifampicin and/or isoniazid are detected, a molecular test to evaluate resistance to at least fluoroquinolones (FQs) should be performed before starting the therapy. On smear positive samples, targeted NGS on rifampicin-resistant cases can provide a rapid and more comprehensive drug resistance profile. Isoniazid-resistant cases should also be tested for fluoroquinolones resistance before starting the WHO-recommended treatment containing levofloxacin. Wherever the short 4 months regimen for drug susceptible TB is implemented [5, 6], the laboratory should provide information on molecular susceptibility to rifampicin (rifampicin remains reference for rifapentine), fluoroquinolones and isoniazid.

When culture is available, phenotypic tests can be performed. If the strain is a known rifampicin- or isoniazid-resistant strain, then a fluoroquinolone phenotypic test should be set up together with first line drugs. Whenever possible, for rifampicin-resistant cases second line drugs should be tested and priority to the drugs used for treatment should be given.

The following are additional tips to maximise the diagnostic yield:

- Molecular tests could be used for diagnosis of TB and drug resistance also after few days of treatment but are not suitable for treatment follow-up (unless development of resistance is suspected and, in this case, the molecular test could identify the appearance of drug resistance determinants not identified at baseline).
- After collection of the sputum in a suitable container, the quality of the sputum should be directly checked in terms of quality and quantity. A considerable part of the sputum should be true sputum and not saliva.
- Clinical material should be collected aseptically in sterile containers of the right size and shape to avoid contamination with non-tuberculous mycobacteria and other microorganisms. It should be noted that tap water contains multiple mycobacteria of different species and therefore should not be used in this procedure.
- All patient material should be collected, stored and transported according to the national guidance or standards, and in sufficient quantity. There is still a knowledge gap on the *ex vivo* survival of mycobacteria in clinical material. In one study on Ziehl–Neelsen (ZN)-positive sputa kept at 4°C, 60% of the mycobacteria in the sputum appeared to be viable after four weeks, while at room temperature only 38% survived [7]. It is therefore advised to send pulmonary clinical material directly to the laboratory. If this is not possible, these should only be kept in the refrigerator for the minimum number of days. CSF should be processed immediately.

- Short transportation times, from specimen collection to arrival to the laboratory, can benefit diagnosis. Some extra-pulmonary fluids may even benefit from inoculation at the bedside into blood liquid culture to increase the probability of a positive *M. tuberculosis* diagnosis. However, if inoculated at the bedside, an additional sample should be collected for microscopy and great care should be made to avoid contamination by saprophytic flora.
- The yield of positive results will increase for some specimens if a higher amount of the material is provided (in case of CSF) or multiple samples are examined. In general, and especially for fluids with likely low concentration of bacteria, such as CSF, ascitic and pleural fluid, the largest possible volume should be collected and sent to the laboratory.
- All clinical material for TB diagnosis sent to a laboratory should be accompanied by a laboratory form including all the relevant information. Dedicated forms should be used, providing information such as the name of the patient, date of birth, gender, patient file number, probability of resistance to anti-tuberculosis drugs (i.e. previous history of TB, previous or current anti-tuberculosis treatment and country of birth), HIV status where possible, required diagnostic tests, date of sample collection, whether the sample was taken before or during treatment, and detailed information on the submitting physician, including telephone number.
- It is the responsibility of the sender to pack clinical samples appropriately. In most countries packaging is provided by the laboratories and generally consists of several layers of leak-proof material. The packaging should indicate that the material should only be opened inside an appropriate laboratory. *M. tuberculosis* cultures are a BSL3 microorganism, so special regulations apply. In principle, the biosafety regulations of directive 2000/54/EC [8] can be applied, but national authorities have also released individual regulations for the transportation of BSL3 microorganisms (see Chapter 1). It is recommended that the sender notifies the laboratory when the culture has been sent. Receiving laboratories should acknowledge receipt so that missing parcels can be traced.
- If *M. tuberculosis* culture is routinely performed at regional or peripheral laboratories, while additional laboratory procedures such as identification, drug susceptibility testing and genotyping are executed in larger laboratories and/or at National Reference laboratories (NRLs), it is of the utmost importance that positive cultures are sent to the centralised facility without delay.
- If the conditions of transport have compromised sterility and viability of *M. tuberculosis*, molecular diagnostics could still be applied. If samples are shipped for molecular tests only, inactivation before shipment could be considered. In this case samples can be shipped as 'non-infectious' reducing the shipping costs.
- Culture isolates can also be either inactivated or DNA can be extracted before shipment to reduce the cost of transport. In this case, the receiving laboratory should provide the preferred protocol for inactivation or DNA extraction.

10.3 Specific considerations regarding diagnosis of TB

10.3.1 Pulmonary material

Sputum (expectorated)

If there is a suspicion of pulmonary TB, >3 ml of early morning sputum should be collected in an appropriate sputum pot with a wide opening and a secure lid. This should be done on at least two consecutive days. As alternative, a second sample could be collected on spot if this is possible. The sputum should be freshly expectorated from the lung (rather than saliva) and the patient should be instructed on how to produce this material. The pooled collection of sputum over 24 hours should not be performed as the extended time of collection increases the chance of contamination by non-tuberculous mycobacteria and other bacterial microorganisms.

Sputum (induced)

If the patient is unable to produce sputum, sputum can be induced by supplying an aerosol of hypertonic saline solution (5%-10% NaCl) generated by a nebulizer. Such specimens may appear thin and watery and should be labeled 'induced sputum' so they will not be discarded by the laboratory as inadequate specimens. This procedure should be administered by trained personnel using appropriate respiratory protection in an isolation booth or in an area with appropriate environmental controls.

Bronchial lavage

The bronchial lavage (BAL) involves the instillation of sterile saline solution into a subsegment of the lung through the bronchoscope to wash the airways, followed by the suction and collection of the fluid sample. Although a routine procedure, it should be emphasised that carrying out a bronchoscopy in a patient suspected of TB can represent a risk to the person performing the procedure, and requires thorough disinfection of the bronchoscope (i.e. the procedure must be performed using appropriate safety standards). BAL should be centrifuged, and the pellet decontaminated in the laboratory before culture.

Gastric lavage

Investigation of gastric fluid is recommended in the diagnosis of pulmonary TB when examination of sputum or bronchial lavage fluid is not possible, for example in young children. Fasting gastric fluid should be collected after the administration of 20–30 ml of physiological saline in 5–10 ml of sodium carbonate (Na₂CO₃). The material should be transported to the laboratory within four hours or neutralised immediately at the site of collection. The investigation of gastric fluid can also be useful in the case of immunocompromised patients who are unable to provide sputum.

Nasopharyngeal aspirate

Nasopharyngeal aspiration (NPA) is another common method of obtaining clinical samples from children. Although it is less invasive than gastric lavage, NPA is still considered as an aerosol generating procedure⁶, and therefore, biosafety and infection prevention and control practices should be followed by the trained personnel. The procedure is performed by placing a flexible plastic catheter, connected to a suction pump, through the nostril into the posterior nasopharynx and by applying a gentle suction. The sample is collected into a sputum trap, and at least 2–5 mL of secretions should be collected.

Stool

Children with TB swallow sputum containing TB bacilli originating from the lungs, which then pass through the digestive tract, where they can be detected in stool samples. Stool is, therefore, regarded as a respiratory specimen for the diagnosis of TB.

Stool is a newly WHO recommended specimen for the diagnosis of pulmonary TB in children using Xpert MTB/RIF or Ultra [4]. It can be used as an alternative specimen, especially in situations when it is challenging to obtain adequate respiratory specimens for the diagnosis of pulmonary TB, such as in younger children. Testing stool may be more acceptable and feasible in certain settings, as it is less invasive than gastric or nasopharyngeal aspiration.

Testing a stool sample by Xpert MTB/RIF or Ultra requires a pre-processing step. Resources on the processing methods are widely available [9].

10.3.2 Extrapulmonary material

TB disease can occur in almost any anatomical site; thus, a variety of clinical specimens other than sputum (e.g. urine, cerebrospinal fluid, pleural fluid, pus, or biopsy specimens) may be submitted for examination when extrapulmonary TB disease is suspected. Procedures for the expeditious and recommended handling of the specimen must be in place or assured before the specialist performs an invasive procedure to obtain the specimen. Especially important is rapid transportation to the laboratory according to the laboratory's instructions. It is important to note that the portion of the specimen placed in formalin for histologic examination cannot be used for culture.

10.3.3 Pleural material

If pleural fluid is obtained, the chance of a positive culture can be improved by having increased volumes, which can then be concentrated. The largest volume possible should be taken and sent. However, recent studies have shown that the transport medium, place of inoculation and type of inoculation medium influence the yield of mycobacteria. Bedside inoculation in combination with a liquid *Mycobacterium spp.* medium appeared to be the best choice but at high risk of contamination. Alternatively, containers with heparin should be used to avoid clotting and trapping of mycobacteria [10]. Pleural fluid aspiration and pleural biopsy may increase diagnostic yield. A pleural biopsy is the recommended material for pleural TB.

10.3.4 Lymphadenitis material

For the diagnosis of tuberculous lymphadenitis, lymph node biopsy (ideal) and fine-needle aspiration are the first-choice diagnostic methods in both low-incidence and endemic countries [11, 12]. In the case of a negative fine needle aspiration, an excision biopsy should be considered, which often results in a higher chance of positive microscopy for mycobacteria. For TB detection, the use of molecular WRDs is recommended. A fine-needle aspirate should be taken using a 19- or 21-gauge needle; the sample needs to be transported directly to the microbiological/pathological laboratory to prevent evaporation. It is important to inform both the pathologist and the microbiologist beforehand that the delicate clinical material is to be dealt with immediately.

Under no circumstances should the material for microbiological examination be placed in formalin (used for the histopathological specimen) as it will kill any TB bacteria present in the sample.

⁶ Jackson T, Deibert D, Wyatt G, et al. Classification of aerosol-generating procedures: a rapid systematic review. *BMJ Open Res* 2020;7:e000730. doi:10.1136/bmjresp-2020-000730

10.3.5 Peritoneal fluid

If tuberculous peritonitis is suspected, at least 5–10 ml ascitic fluid can be collected and sent to the laboratory or inoculated at the bedside in a liquid mycobacteria culture medium. However, in a review involving more than a thousand patients, a peritoneal biopsy appeared more sensitive [13, 14]. In general, sending the largest possible volume for centrifugation at the laboratory is ideal.

10.3.6 Urine

An early morning specimen should be collected, and the entire sample sent to the laboratory. The specimen should be refrigerated until transport. Multiple specimens over several days may be required to obtain a positive specimen. Due to contamination and deterioration, 24-hour urine specimens are not acceptable. Microscopy examination of urine sediment is not recommended due to the high risk for false positivity because of the presence of saprophyte mycobacteria of the genitourinary tract.

10.3.7 Blood

Blood cultures for mycobacteria should be performed using specific blood culture bottles provided by the laboratory. If the laboratory does not have the automated incubator, specimens should be collected in isolator tubes. The tubes should be stored at room temperature and transported to the laboratory on the same day of collection. Multiple cultures should be attempted to increase the diagnostic yield.

10.4 Information flow from microbiologist to physician and instructions regarding TB diagnosis

Physicians should provide basic patient information and specimen(s) to the laboratory that needs this information to guide its work, interpret the results and participate in TB surveys at local, regional, national and international levels. When laboratory results are reported to the physician, a basic level of information should always be provided to avoid mistakes and confusion, including date of specimen reception at the laboratory, test date, patient identifiers, results, and indications how to translate results into clinical management. Physicians should seek an unambiguous clarification of the results if needed. In the case of microscopic examination, an indication of the semi-quantitative result according to the WHO scale should be provided. If a peripheral or regional laboratory has already performed diagnostic tests, it is very helpful if these results are sent together with the culture to a centralised laboratory to avoid duplication or to allow confirmation.

For cultures, the quantitative results in terms of time to positivity (in days) for liquid medium are helpful (particularly where the samples have been sent to indicate treatment progress) and possibly the number of colonies on solid medium could be stated with an interpretation of the likely clinical relevance of this. A low number of colonies on a primary culture, unusual timings for positive cultures (e.g. a higher than usual positivity rate associated with inoculation on or around the same day, or a very short time to positivity in a smear-negative extra-pulmonary sample) may raise serious concerns of cross-contamination, and the laboratory needs to be aware of this. If suspected, positive cultures found on consecutive days can be subjected to molecular typing using WGS (see Chapter 8), and if the same sequencing profile is obtained, cross-contamination is highly likely.

In addition, the results of nucleic-acid amplification tests (NAATs) should be provided to the physician using appropriate wording and context and semi-quantitative results should always be reported. Overall, results validity depends on the use of inhibition/amplification controls, negative and positive controls, and the positive and negative predictive value. For example, when a NAAT is positive with a low semi-quantitative value, while smear microscopy is negative, it should be explained that the positive predictive value of this result is highly dependent on the prevalence of TB in the respective patient category. Sensitivity of new platforms has highly increased with the next generation of molecular tests now available in the market (see Chapter 6). Increasing the sensitivity for the paucibacillary smear negative samples has come at the cost of decreasing the specificity. Samples that are reported as Xpert Ultra 'trace' should be interpreted based on the clinical and epidemiological situation. People at high risk for serious TB consequences for underlying diseases, age, immunological status or known contacts of infectious TB cases should be started on treatment while culture results are pending. If no information on the molecular drug resistance pattern is available, then the treatment should be initiated based on the epidemiological information.

There has been substantial progress in the development of rapid diagnostic tests for both TB and drug resistance (see Chapter 6). These tests reduce delays in diagnosing and initiating TB treatment [15], and can be used to direct the therapy and the choice of anti-tuberculosis drugs. WHO and other international consortia are providing information on how to interpret different mutations in drug resistance associated genes [16]. The capacity to interpret genomic variants with high confidence is high for drugs such as rifampicin, isoniazid, pyrazinamide, fluoroquinolones, injectable agents and linezolid. However, it is always possible to encounter discrepancies between phenotypic and genotypic test results and it is understandable if this generates confusion. The role of the clinical

microbiologist is to guide the correct results interpretation and to clearly explain why these discrepancies occur. The main reasons for genotypic/phenotypic discrepancies are: i) technical difficulties and errors in the performance of DST; ii) the presence of unknown mechanism conferring drug resistance; iii) mutation confers low level resistance and critical concentration is set too high; iv) mutation is outside the region targeted by the molecular assay; v) the presence of heteroresistance not detected by molecular methods; vi) molecular assay detect silent mutations; and vii) errors due to probe interaction/binding in LPA or other molecular assays.

Next generation sequencing is now becoming a diagnostic tool at the EU-NRL level. It can be performed as WGS on positive culture or as target NGS from diagnostic samples (containing at least 5 000 to 10 000 genomes per mL of sample) [17]. The difference among the two NGS-based strategies is shown in Table 15.

Table 15. Key differences between WGS and targeted NGS strategies

Whole genome sequencing	Targeted Next generation sequencing
Culture-based	Test for selected targets on direct clinical specimens
Detection of the total genomic changes (loss of function mutations role)	High coverage per sample, increased sensitivity for mutation detection, mixed infections
High volume of data	Detection of heteroresistance at low frequencies
Assembly and analysis challenging	High number of samples per run, lower cost per sample compared to standard Sanger sequencing
Expensive for higher coverage data	
No standardised kit available	
EQA available	

Both allow to evaluate a large number of determinants and are highly valuable in supporting the management of multidrug-resistant (MDR), pre-extensively resistant (pre-XDR), or extensively resistant (XDR) TB cases.

In general, the presence of mutations or large deletions or insertions in genes involved in the pathways of drug activation should be considered with suspicion particularly when mutations develop during the therapy, are associated to smear and culture positivity, or with lack of radiological improvement. In this case, minimum inhibitory concentration (MIC) testing on broth microdilution and phenotypic DST using the recommended drug critical concentrations should be requested and promptly reported.

Finally, it is important to establish a clear communication with the clinical counterpart: clinicians should understand advantages and limitations of all the tests that the laboratory offers and be guided in the interpretation of the results.

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11. The diagnosis of nontuberculous mycobacteria

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Summary

This chapter provides an overview of the principles of nontuberculous mycobacteria (NTM) diagnostics. It includes an introduction to nontuberculous mycobacteria (NTM) epidemiology and disease principles (background, when to suspect NTM, clinical presentations), and a more detailed discussion of microbiological diagnostic methods, including microscopy, culture, and molecular techniques. The content is intended for laboratories diagnosing mycobacteria infections and is based on the authors' experience, as well as the literature list at the end of the chapter.

11.1 Background and principles

The genus *Mycobacterium* is a cohesive taxonomic group distinguished by numerous phenotypic and genotypic characteristics that set it apart from other genera [1]. Nontuberculous mycobacteria (NTM) refers to all mycobacterial species other than those in the *M. tuberculosis* complex, and the *M. leprae* complex (*M. leprae* and *M. lepromatosis*). To date, over 200 NTM species have been identified, which can be classified into distinct groups based on phenotypic and genotypic characteristics.

Although new genera have been proposed for some mycobacteria (e.g. *Mycolicibacterium*, *Mycolicibacter*, *Mycolicibacillus*, and *Mycobacteroides*) [2], these are largely considered theoretical taxonomy constructs. In practice, genomic and clinical microbiology still uses the genus *Mycobacterium* for all NTM [3].

The currently accepted phylogeny groups closely related species into complexes or clusters, including the *M. avium*, *M. terrae*, *M. smegmatis*, *M. fortuitum*, *M. abscessus*, *M. celatum*, *M. kansasii*, and *M. simiae* complexes/groups [1].

NTM species can also be categorised by their growth rate:

- Rapidly growing mycobacteria (RGM) form colonies on solid medium within seven days.
- Slowly growing mycobacteria (SGM) require more than seven days to form mature colonies on solid media. According to the Runyon classification (4), SGM can be further divided based on their pigmentation:
 - Runyon I; Photochromogens: appear yellow-orange when exposed to light (e.g. *M. kansasii*, *M. marinum*, *M. simiae*);
 - Runyon II; Scotochromogens: are always yellow-orange, even if grown in the dark (e.g. *M. scrofulaceum*, *M. gordonae*, *M. szulgai*, *M. xenopi*);
 - Runyon III; Nonchromogens: do not produce pigment and appear pale-white (e.g. *M. avium* complex [MAC], *M. ulcerans*, *M. malmoense*, *M. terrae* complex).

The RGM (e.g. *M. fortuitum*, *M. abscessus*, *M. smegmatis*, and *M. chelonae*) were classified as Runyon IV and most of them are non-chromogenic [4].

Although molecular diagnostics have replaced many conventional laboratory tests, the growth rate remains an important criterion for preliminary classification and has implications for phenotypic drug susceptibility testing.

NTM are ubiquitous environmental organisms, commonly found in water or soil – e.g. NTM have been isolated in:

- tap water and shower heads
- swimming pools and hot tubs
- other humid environments [4,5].

Their ability to resist disinfection, adhere to surfaces, and form biofilms enables them to thrive in natural and urban water systems, air, soil and sediments. This widespread distribution, combined with population growth, has led to increased contact between humans, animals, and the environment [6]. Although nearly all NTM diseases result from environmental exposure, rare cases of person-to-person transmission have been documented (e.g. among individuals with cystic fibrosis) prompting enhanced infection control measures in specialised care settings [7].

NTM are increasingly being recognised as opportunistic pathogens of humans. Most often, NTM affect the lungs (NTM-pulmonary disease, NTM-PD) and may cause progressive disease in susceptible hosts, particularly individuals with functional or structural lung disease, such as:

- chronic obstructive pulmonary disease
- bronchiectasis
- emphysema
- cystic fibrosis
- idiopathic pulmonary fibrosis
- primary ciliary dyskinesia
- α 1-antitrypsin deficiency
- sequelae resulting from previous tuberculosis (TB) infection.

Extrapulmonary NTM disease (NTM-EP) is less common and primarily seen as cervical lymphadenitis in children, skin and soft tissue infections following inoculation, and disseminated disease in individuals with inherited or acquired immune deficiencies, which also increases the risk of NTM-PD [5].

Since many NTM are not 'true' pathogens, such as *M. tuberculosis* or *M. leprae*, their isolation from a human specimen does not automatically imply that there is an NTM infection or disease. Therefore, diagnosis is often difficult. In addition to NTM isolation, it requires clinical symptoms, paraclinical findings (most often radiological), and the exclusion of other diseases, particularly TB [11]. In addition, colonisation can be permanent in patients with bronchial defects and transient isolation can be seen at non-sterile body sites due to environmental exposure or contamination during sampling or handling procedures. Management is further challenged by NTM intrinsic multi-drug antimicrobial resistance and the need for combination therapy (often three or four drugs) over many months to years, because few antimicrobials are effective and almost none of them are bactericidal. Consequently, treatment is associated with significant side effects and high relapse rates. Furthermore, antimicrobial susceptibility testing (AST) is often not standardised, and poorly predictive of positive outcome (8)(9)(10). AST necessitates specialised laboratory services and the involvement of clinical experts with specific experience in NTM management [11,12].

NTM manipulate the body's immune response through various mechanisms, targeting not only the innate immune response, but also effectively evading the adaptive immune response [6]. This manipulation allows NTM to resist host immune elimination, achieve long-term survival, and even utilise pathways such as autophagy and cell death to facilitate its dissemination within the host.

Recently, NTM species with high genomic similarity to MTB have been described, such as *M. decipiens*, *M. shinjukuense*, *M. lacus*, and *M. riadhense*. These species may represent 'missing links' between low-virulence mycobacterial opportunists and the highly-virulent obligate pathogen MTB. They share many attributes with MTB in terms of pathogenicity, clinical manifestations, and host adaptation, but they test negative in PCR assays targeting MTB complex species [13].

Despite these diagnostic and clinical challenges, potentially leading to NTM disease being under-recognised, NTM disease is increasing globally. This increase is attributed to longer life expectancy; a greater burden of chronic diseases; the intensification of immunosuppressive therapies; increased availability and improvement in mycobacterial diagnostics; greater awareness of NTM disease, and climate-associated changes. The decline in TB in high income countries, along with the abandonment of the BCG vaccination, may also lead to an absence of adaptive immunity [14].

11.2 When to suspect an NTM infection

The clinical presentation of NTM-PD closely resembles that of other respiratory conditions, including lung cancer, TB, and pre-existing lung disease. Therefore NTM-PD will often only be considered after the TB, cancer and other chronic inflammatory diseases have been ruled out.

Symptoms of NTM-PD may include:

- persistent cough
- sputum production
- haemoptysis
- breathlessness
- fever
- night sweats
- unintentional weight loss
- significant fatigue.

It is crucial to look for clues, such as a person reporting an increased frequency of lower respiratory tract infections that does not improve despite antibiotics. In the context of NTM-PD, these symptoms are particularly relevant for individuals with functional or structural lung disease. Furthermore, NTM-PD is especially relevant in individuals with weakened immune systems, such as:

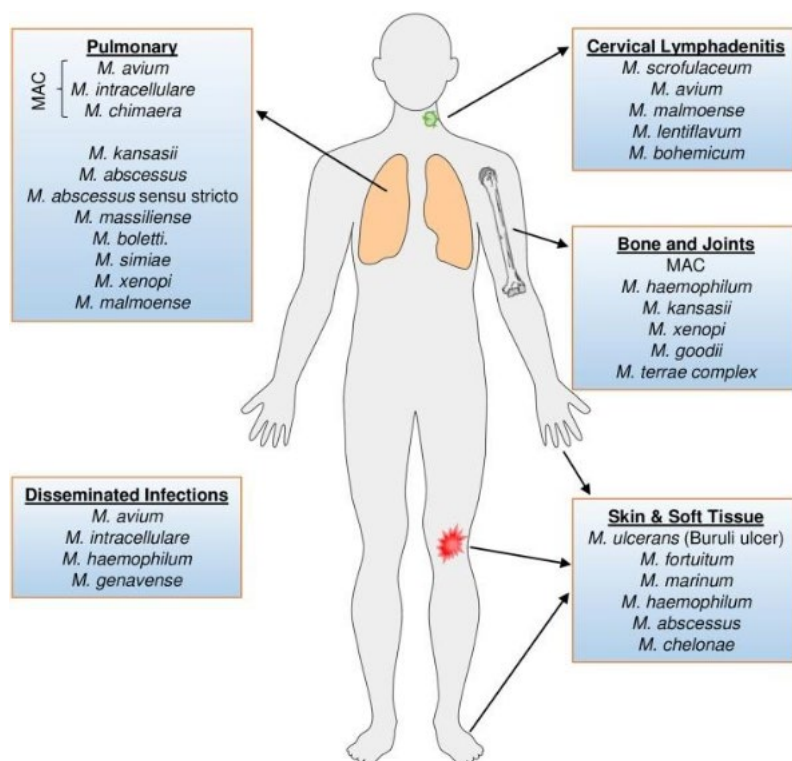
- children
- the elderly
- organ transplant recipients
- individuals with acquired immune deficiency syndrome (AIDS)
- those receiving immunosuppressive treatment.

In addition to patients with pre-existing lung diseases or immunocompromised individuals, postmenopausal women with low BMI and connective tissue-related abnormalities are also at higher risk of NTM disease, especially for nodular pulmonary disease (Lady Windermere syndrome) [15]. Other potential risk factors include a range of geographical and environmental variables, such as high evapotranspiration, a greater proportion of surface water, increased precipitation, lower elevation, warmer temperatures, and lower concentrations of aluminium and manganese in the soil [16].

11.3 Specific clinical presentations and NTM species

While NTM species can affect various body sites (Figure 16), certain clinical presentations are often associated with specific species. For example, pulmonary disease is most frequently caused by *Mycobacterium avium* complex, *M. kansasii*, and *M. abscessus*, whereas skin and soft tissue infections are more often caused by *M. chelonae* and *M. marinum* [17]. The *M. ulcerans* complex is involved in skin and soft tissue infections characterised by extensive, necrotic, painless ulcers, but only in certain geographical regions, such as West Africa and Australia [18].

Figure 16. Body sites affected by NTM species



Source: Baldwin SL, Larsen SE, Ordway D, Cassell G, Coler RN. The complexities and challenges of preventing and treating nontuberculous mycobacterial diseases [19]. Adapted.

Skin, soft tissue, and bone NTM disease

The NTM species most associated with localized skin and subcutaneous infections include:

- *M. fortuitum*
- *M. abscessus*
- *M. chelonae*
- *M. marinum*
- *M. ulcerans*
- *M. haemophilum*.

However, nearly all NTM species have been reported as causes of cutaneous disease.

Skin infections typically occur at the sites of:

- puncture wounds, open traumatic injuries, or fractures;
- long-term intravenous or peritoneal catheters;
- post-injection abscesses;
- tattooing;
- liposuction;
- surgical wound infections, such as those after mammoplasty;
- cardiac bypass surgery;
- corneal infections following laser procedures [20].

M. marinum infection, also referred to as 'fish tank granuloma' or 'swimming pool granuloma', typically occurs following exposure to aquatic environments [21]. Common sources of infection include:

- cleaning or maintaining of fish tanks and aquariums;
- swimming in pools, lakes, or oceans, particularly where water is untreated or contaminated;
- handling of marine life, such as fish, shellfish, or crustaceans.

Diagnostic principles and sampling for NTM infection

Given the large number of identified NTM species (>200), the wide spectrum of NTM virulence, and the variable host susceptibility described above, a single set of diagnostic criteria is neither useful nor accurate for all NTM species across all clinical scenarios.

The NTM-PD disease diagnosis is based on a combination of clinical, radiographical, and microbiological criteria [11] [20], including:

- pulmonary or systemic symptoms; AND
- radiological findings; AND
- appropriate exclusion of other diagnoses;

combined with:

- positive culture results for NTM from at least two separate expectorated sputum samples; OR
- positive culture results for NTM from at least one bronchial wash or lavage; OR
- transbronchial or another lung biopsy, with:
 - mycobacterial histologic features (granulomatous inflammation or Acid-Fast Bacillus (AFB)) AND positive culture for NTM; OR
 - biopsy showing mycobacterial histologic features (granulomatous inflammation or AFB), AND one or more sputum or bronchial washings that are culture-positive for NTM.

The diagnosis of NTM-EP is less well-defined compared to NTM-PD. It typically relies on a combination of:

- risk factors;
- history of exposure;
- clinical presentation;
- microbiological evidence;
- histopathological findings.

The specific diagnostic approach depends on the site of infection (e.g. skin, lymph nodes, bones, or disseminated disease).

For both NTM-PD and NTM-EP, we recommend that the diagnosis is made by clinicians with experience in NTM management, based on microbiological data from quality-assured laboratories with specific NTM competencies. These laboratories should participate in external quality assurance (EQA) programmes.

11.4 Specific sampling instructions

For the diagnosis of NTM-PD, the types of samples are similar to those used for the diagnosis of TB: sputum, bronchial aspirate and bronchoalveolar lavage. As NTM may be present in the upper respiratory tract and oral cavity, having several samples positive for the same NTM species is helpful in discriminating contamination from bronchial colonisation. Repeating the sampling after several weeks further supports the diagnosis of true colonisation or infection.

In fibrocavitary forms of respiratory NTM infections, repeated sputum sampling often leads to NTM isolation and diagnosis. In contrast, nodular bronchiectatic forms are frequently associated with limited spontaneous sputum production, requiring assistance from pulmonary physiotherapy or bronchoscopy with bronchoalveolar lavage (BAL) or bronchial aspirate, followed by examination of post-bronchoscopy sputum.

For the diagnosis of NTM-EP infections, the approach depends on the site of the infection. When the site is non-sterile, the collection and analysis of multiple samples is recommended. In skin and soft tissue NTM infections, aspiration of abscesses or tissue biopsy is necessary; superficial sampling is not considered adequate. For lymph node involvement, needle aspiration is usually sufficient, particularly in children.

In patients with immunodeficiency (see above) and in those suspected of disseminated infection, blood cultures should be obtained. It is important to note that this requires specific blood-culture bottles (e.g. F-lytic bottles from Becton-Dickinson) incubated for at least 42 days, depending on the suspected NTM species.

11.5 Laboratory-based methods used in the diagnosis of NTM infection

11.5.1 Interferon-gamma-release-assay (IGRAs)

Interferon-Gamma Release Assays (IGRAs) are in vitro blood tests primarily developed and validated for the detection of (latent) MTB infection. These assays measure the release of interferon-gamma (IFN- γ) by T-cells in response to MTB-specific antigens, such as ESAT-6 and CFP-10, which are encoded in the RD1 genomic region.

It is important to note that certain NTM species, such as *M. gastri*, *M. kansasii*, *M. marinum*, *M. riyadhense*, *M. szulgai*, and *M. flavescens*, carry homologue genes that may lead to cross-reactivity and cause false-positive IGRA results [22,23,24]. This underscores the need for cautious interpretation of IGRA results in patients with, or suspected of having NTM infections.

Recently, IGRA tests specifically targeting NTM have been developed using glycopeptidolipid (GPL) antigens [25]. These assays are now commercially available. However, despite their promise, additional validation is necessary before broader clinical implementation will be possible.

11.5.2 Smear microscopy

Smear microscopy using Ziehl-Neelsen or auramine-O staining does not differentiate between MTB and NTM. For Standard Operating Procedures (SOPs), please refer to Chapter 4 of this handbook: 'Smear Microscopy'.

Some NTM species exhibit distinctive morphological characteristics: *M. xenopi* tend to appear thinner and longer, *M. kansasii* is broader with visible striations, and *M. avium* is typically short. While these features were once helpful for preliminary differentiation, they are not sufficiently specific for reliable identification. Today, more accurate diagnostic methods, such as molecular identification techniques, or matrix-assisted laser desorption ionisation time-of-flight mass spectrometry (MALDI-TOF), are required to distinguish MTB from NTM species.

11.5.3 Decontamination of specimens

Procedures for the decontamination of specimens for NTM diagnosis are often similar to those used for TB. For respiratory specimens, or non-sterile sites, decontamination is processed following the N-acetyl-L-cysteine-sodium hydroxide (NALC-NaOH) method. For further details and SOPs, please refer to Chapter 5 of this handbook 'Digestion and decontamination of sputum samples'.

In the event of heavily contaminated specimens, such as sputum from patients with cystic fibrosis, the use of additional decontaminating agents can be beneficial in inhibiting the growth of Gram-negative bacilli. These agents may include 1% hydrochloric acid, 5% oxalic acid, or chlorhexidine [26].

Cetylpyridinium chloride (CPC) is recommended for decontaminating water samples in some settings to facilitate the cultivation of NTM, such as *M. chimaera* from heater-cooler units [27,28]. The rationale for using CPC is its selective antimicrobial activity: it effectively reduces non-mycobacterial contaminants while preserving the viability of NTM, thereby improving the recovery of NTM isolates [29,30].

11.5.4 Culture tests for NTM

Cultures for the detection of NTM include specific solid media, such as Middlebrook 7H10 or 7H11 agars with OADC enrichment, or egg-based media, such as Löwenstein-Jensen, Ogawa, Coletsos, and Stonebrink. Liquid media (e.g. enriched Middlebrook media, Youmans) are also used and allow faster culture positivity. Various automated reading systems for cultures in liquid media are available on the market (e.g. MGIT⁷ 960 from Becton-Dickinson, BacT/ALERT from bioMérieux, and VersaTREK™ Myco from Thermo Fisher).

Culture in the presence of antimicrobials (home-made solid media, MGITs with PANTA, or selective medium RGM) can also prevent the growth of bacteria and fungi other than mycobacteria. Cultures in solid media allow observation of colonies and categorisation of species according to pigmentation (see details on Runyon

⁷ MGIT – Mycobacteria Growth Indicator Tube

classification above). They can also help detect mixed infections and allow quantification of each NTM species based on the number of colonies. Liquid and solid cultures are therefore complementary and should be performed in all suspected NTM infection cases. It is also useful to report whether colonies are 'rough' (R) or 'smooth' (S), as rough colonies have been associated with increased virulence in pulmonary *M. abscessus* infections [31].

Some species, such as *M. genavense*, grow mostly in liquid medium or on 7H11 supplemented with mycobactin J after very prolonged incubation [32] while others, such as *M. marinum* and *M. chelonae*, grow better on solid media at 30°C. Some NTM are very difficult to grow, and the culture media should contain adjuvants or vitamins, such as *M. haemophilum* requiring haemin (Table 1 below).

The incubation temperatures for NTMs range between 28 and 37°C. Incubation at 37°C supports the growth of many NTM species. Some NTM, such as *M. chelonae*, *M. marinum*, *M. haemophilum*, and *M. ulcerans*, grow better or even exclusively at 30°C for primary cultures. Other species, such as *M. xenopi*, prefer the temperature range 40–42°C, although they can also grow at 37°C, but more slowly. Solid cultures are typically incubated for about 8–12 weeks, although some NTM, such as *M. avium subsp. paratuberculosis* and *M. ulcerans*, may require even longer incubation times (Table 1).

Table 1. Growth characteristics of difficult-to-grow NTM

Species	Special growth characteristics	References
<i>M. genavense</i>	Requires mycobactin J and extended incubation (8–12 weeks).	[32] [33]
<i>M. haemophilum</i>	Requires lower temperature (e.g. 30°C) and iron supplements (e.g. hemin or ferric ammonium citrate).	[34]
<i>M. ulcerans</i>	Requires lower temperature (e.g. 30°C), extended incubation (8–12 weeks), benefits from glycerol or pyruvate.	[35]
<i>M. marinum</i>	Requires lower temperature (e.g. 30°C).	[21]
<i>M. avium subsp. paratuberculosis</i>	Requires mycobactin J and extended incubation (12–20 weeks).	[36]
<i>M. conspicuum</i>	Requires lower temperature (e.g. 30°C).	[7][37]
<i>M. tilburgii</i>	Cannot be cultured on standard media.	[38]
<i>M. lepraemurium</i> (feline leprosy)	Cannot be cultured on standard media; Can be grown in rodents (e.g. mice) or mineral salt minimal medium supplemented with simple sources of C and N; Ogawa egg-yolk medium at 34°C in approximately 90 days of incubation.	[39][40]
CLG (canine leprosy)	Cannot be cultured on standard media.	[41][42]

For detailed step-by-step protocols for culturing mycobacteria, please refer to Chapter 1 of the book 'Mycobacteria protocols' [43].

11.6 Molecular assays for NTM

11.6.1 Direct commercial molecular assays performed on primary clinical samples

NTM is rarely detected in clinical specimens for several reasons. Firstly, overly sensitive detection methods risk identifying colonisation or contamination – particularly in paucibacillary samples – rather than true infection. Since clinically relevant NTM disease is often multibacillary, such high sensitivity may be unnecessary and even misleading in primary specimen testing. Secondly, few commercial tests allow differentiation between *M. tuberculosis* complex and NTM directly from primary specimens and often these tests have not been extensively validated and/or have shown inconsistent results. Details of the tests are set out below.

- **Anplex MTB/NTMe real-time detection** (Seegene, CE-IVD): allows differentiation of NTM (based on 16S rRNA) from MTBC (based on *IS6110* and *mpb64*) in a single well using primary specimens (e.g. sputum, BAL). This test does not allow identification of the exact NTM species.
- **NeoPlex™ TB/NTM Multiplex Real-time PCR Assay** (Genematrix, CE-IVD): Detects *M. tuberculosis*, *M. abscessus*, *M. massiliense*, *M. avium*, *M. intracellulare*, and *M. kansasii* directly from sputum. Results are available within two hours from automated nucleic acid extraction to result analysis.
- **GenoType™ CMdirect VER1.0** (Hain LifeScience Bruker; CE-IVD): allows identification of TB and clinically relevant NTM species, *M. avium*, *M. chelonae*, *M. abscessus*, *M. fortuitum*, *M. gordonae*, *M. intracellulare*, *M. scrofulaceum*/*M. intracellulare*, *M. malmoeense*, *M. interjectum*, *M. szulgai*, *M. kansasii*, *M. ulcerans* complex, and *M. xenopi*, directly from sputum.

An overview of these tests is also provided in Supplementary Table 2.

11.6.2 PCR and other line-probe assays

While whole genome sequencing provides the highest resolution for identifying NTM species, other molecular methods continue to be widely used in mycobacterial reference laboratories. These include restriction enzyme analysis (e.g. PRA-hsp65), line probe assays (LPA), multi-plex and real-time PCRs and Sanger sequencing of single or multiple marker genes such as 16S rRNA, *rpoB*, ITS, and hsp65. Supplementary Table 1 provides a list of primers used for PCR-based assays [44–61]. However, it should be noted that species or subspecies identification based on single gene targets can be inaccurate, particularly with *rpoB*, due to recombination events among NTM strains [62].

In recent years, line-probe assays (LPAs) have become increasingly popular in diagnostic laboratories for identification and drug resistance prediction of both TB and NTM. LPAs detect species and sub-species-specific PCR-amplified DNA sequences by hybridising them to oligonucleotide probes immobilised on a strip. Visualisation is achieved through an enzyme-substrate reaction that produces coloured bands (Figure 17).

Figure 17. Visualisation of LPAs, example of GenoType NTM-DR DNA strip assay

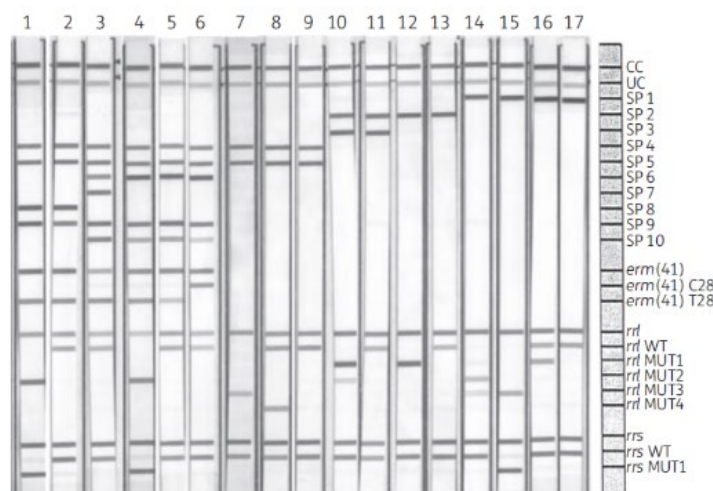


Figure 2. Representative GenoType NTM-DR DNA strip results obtained with NTM isolates showing representative profiles. Lanes 1 and 2, *M. abscessus* subsp. *massiliense* with *rrl* and *rrs* mutations in lane 1 and WT genes in lane 2; lane 3, *M. abscessus* subsp. *bolletii* WT for *rrl* and *rrs*; lane 4, *M. abscessus* subsp. *abscessus* *erm(41)* t28 genotype and *rrl* and *rrs* mutations (heterogeneous for *rrs* with a faint WT band); lane 5, *M. abscessus* subsp. *abscessus* *erm(41)* t28 genotype and WT for *rrl* and *rrs*; lane 6, *M. abscessus* subsp. *abscessus* *erm(41)* c28 genotype and WT for *rrl* and *rrs*; lane 7, *M. chelonae* with an *rrl* mutation and WT *rrs*; lane 8, *M. chelonae* heterogeneous for *rrl* (WT and *rrl* mutation) and WT *rrs*; lane 9, *M. chelonae* WT for *rrl* and *rrs*; lane 10, *M. chimaera* with a double mutation in *rrl* and WT for *rrs*; lane 11, *M. chimaera* WT for *rrl* and *rrs*; lane 12, *M. intracellulare* with an *rrl* mutation and WT for *rrs*; lane 13, *M. intracellulare* WT for *rrl* and *rrs*; lane 14, *M. avium* with a double *rrl* mutation and WT for *rrs*; lane 15, *M. avium* with one *rrl* mutation and one in *rrs* (heterogeneous profile with a faint WT band); lane 16, *M. avium* heterogeneous for *rrl* (WT and mutation) and WT for *rrs*; lane 17, *M. avium* WT for *rrl* and *rrs*. CC, conjugate control; UC, universal control.

Source: Macheras E, Roux AL, Ripoll F, Sivadon-Tardy V, Gutierrez C, Gaillard JL, et al. Inaccuracy of single-target sequencing for discriminating species of the *Mycobacterium abscessus* group. *J Clin Microbiol.* 2009 Aug;47(8):2596–600 [62]

The following commercial LPAs and multiplex real-time PCR assays are currently available for use on positive cultures:

- **GenoType NTM-DR** (Hain LifeScience; CE-IVD) [63]: differentiates between the three *M. abscessus* subspecies, *M. chelonae* and the three MAC species: *M. avium*, *M. intracellulare* and *M. chimaera*. In addition, it also differentiates between genes and mutations related to macrolide and aminoglycoside resistance, providing results within five hours of culture. Rare cases of *M. abscessus* subsp. *abscessus* or *bolletii* with truncated *erm(41)* genes might be misclassified as subsp. *massiliense* with this assay.
- **Genotype CM VER2.0** (Hain LifeScience; CE-IVD): common mycobacteria – differentiates between *M. tuberculosis* and 13 frequently isolated and/or clinically relevant NTM species or complexes, including *M. abscessus*, *M. avium*, *M. intracellulare*, *M. marinum/ulcerans*, *M. kansasii*, *M. xenopi*, *M. chelonae*, *M. gordonae*, *M. fortuitum*, *M. scrofulaceum*, *M. interjectum*, *M. szulgai*, and *M. malmoense*.
- **GenoType AS VER1.0** (Hain LifeScience; CE-IVD): identifies 17 less common NTM species: *M. simiae*, *M. mucogenicum*, *M. goodii*, *M. celatum*, *M. smegmatis*, *M. genavense*, *M. lentiflavum*, *M. heckeshornense*, *M. szulgai*, *M. intermedium*, *M. phlei*, *M. haemophilum*, *M. kansasii*, *M. ulcerans*, *M. gastri*, *M. asiaticum*, and *M. shimoidei*.
- **INNO-LiPA MYCOBACTERIA V2** (Fujirebio; CE-ICD): detects the *Mycobacterium* genus and differentiates between 16 mycobacterial species and two mycobacterial complexes: *M. tuberculosis* complex, *M. kansasii*, *M. xenopi*, *M. gordonae*, *M. genavense*, *M. simiae*, *M. marinum*, *M. ulcerans*, *M. celatum*, MAIS, *M. avium*, *M. intracellulare*, *M. scrofulaceum*, *M. malmoense*, *M. haemophilum*, *M. chelonae* complex, *M. fortuitum* complex, and *M. smegmatis*. This assay targets the 16S-23S rRNA gene spacer region [64].

- **Genoscholar™ NTM+MDRTB** (Nipro; World Health Organization (WHO) approved): distinguishes between *Mycobacterium* species (*M. tuberculosis* complex, *M. avium*, *M. intracellulare*, and *M. kansasii*) while also testing resistance to rifampicin and isoniazid.
- **FluoroType Mycobacteria Ver 1.0** with FluoroSoftware® XT-IVD software (Hain Lifescience - Bruker): differentiates between 32 clinically relevant NTM and MTB complexes in a single well, with results 2.5 hours after positive culture using multiplex qPCRs.

11.6.3 Targeted Next-generation sequencing

Targeted next-generation sequencing (tNGS) is a genomic technique that focuses on sequencing specific regions of interest rather than the entire genome, as with Whole Genome Sequencing (WGS). In 2023, WHO recommended the use of tNGS to detect known mutations conferring resistance to anti-TB medicines. The Deeplex MycTB tNGS assay (Genoscreen) is a target sequencing kit for detecting genotypic resistant MTB. However, it also detects NTM (more than 150) species based on sequence variation in the *hsp65* gene [65]. A recent study [66] showed that WGS and Deeplex can yield different results, underlining the need for further validation.

11.6.4 Challenges in using molecular assays

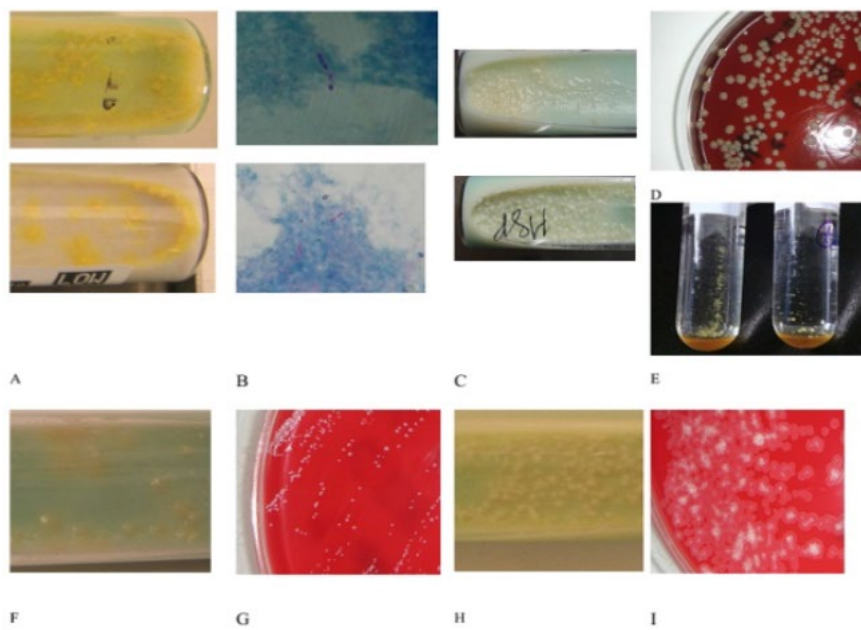
Certain NTM cannot be reliably distinguished by the molecular assays mentioned above. For a complete list of such cases, users should refer to the full documentation provided by the manufacturer. An overview of common difficulties is also provided in Supplementary Table 2, though it may not be exhaustive.

It is also important to note that molecular assays approved for use on cultured isolates may yield results when applied directly to primary specimens, especially those with a high bacillary load. However, performance in such cases can be variable, and results may not always be concordant with species identification obtained by WGS or MALDI-TOF spectrometry performed on cultured isolates.

11.6.5 Phenotypic tests and MALDI-TOF mass spectrometry

The appearance and pigmentation of colonies on a solid medium, along with the rate of growth, can provide initial guidance for the identification of NTM species, but does not allow precise species determination (Figure 18).

Figure 18. Macroscopic aspects of nontuberculous mycobacteria growing in liquid and on solid media



A: *M. kansasii* colonies on Loewenstein Jensen media; B: *M. kansasii* Ziehl-Neelsen staining; C: *M. chelonae* on Loewenstein Jensen media; D: *M. fortuitum* on blood soy agar; E: *M. xenopi*: grain appearance on liquid media (MGIT); F and G: *M. abscessus* smooth colonies on Loewenstein Jensen media (F) and on blood soy agar (G); H and I: *M. abscessus* rough colonies on Loewenstein Jensen media (H) and on blood soy agar (I).

Conventional methods based on biochemical and enzymatic phenotypic characteristics are no longer used. Identification is now primarily established using either molecular methods or MALDI-TOF mass spectrometry [67–69]. These two approaches are complementary and are often combined to achieve accurate identification at the species, or even subspecies level. Identification is useful for assessing the pathogenicity of the strain, helping to differentiate between contamination, colonisation and infection, and guiding appropriate epidemiological and therapeutic management [9][70].

MALDI-TOF spectrometry began to be applied to mycobacteria in the early 2010s and is mainly used for NTM identification in this context [41]. MTB complex isolates are easily identified by MPT64 antigen immunochromatography or specific PCR and are usually handled at BSL3 laboratories. The principle with MALDI-TOF is to compare protein profiles, mainly ribosomal proteins, extracted from an NTM colony. The colony is smeared onto a MALDI target plate, followed by the application of a matrix that facilitates protein extraction, either directly or via prior treatment with formic acid [67]. In the spectrometer, a laser disperses the proteins which are analysed in terms of their size and properties.

For practical purposes, two factors are important in NTM identification using MALDI-TOF: (i) the spectrometer must include an extensive database of NTM reference profiles to enable reliable comparison with the unknown isolate; (ii) it is often necessary to test three to five individual colonies to achieve a reliable identification score (above 2.0 using Bruker system) [69]. It should be noted that some NTM cannot yet be identified with current MALDI-TOF systems. Moreover, subspecies-level differentiation, such as within the *M. abscessus complex*, is rarely possible using MALDI-TOF alone.

Further analysis of lipid components, which requires a dedicated module, can enhance identification resolution [71].

11.7 NTM susceptibility testing

NTM susceptibility testing is recommended only for cases that meet the diagnostic criteria as outlined in the 2020 ATS/ERS/ESCMID/IDSA clinical practice guidelines for NTM-PD [11] and in [20], which includes NTM-EP. Therefore, antibiotic susceptibility testing (AST) is not advised on a routine basis, as NTM findings do not necessarily reflect NTM disease. Furthermore, for many NTM, AST is difficult, and results may therefore lead to misinterpretations, either false susceptibility (risking treatment failure) or false resistance (resulting in omission of potentially effective antibiotics). Consequently, initiating drug susceptibility testing and interpreting AST results requires specialised laboratory services and the involvement of clinical experts with specific experience in NTM management [72] [8]. AST should ideally be performed only when (i) the antibiotic has demonstrated sufficient activity against the species in question, and (ii) there is suspicion of acquired resistance, such as in relapse cases following prior antibiotic treatment.

AST methods and interpretation criteria used for MTB are not applicable to NTM, as growth conditions and drug activity profiles differ (e.g. pyrazinamide is inactive against NTM).

The objective of AST is primarily to distinguish clinical isolates with a wild-type pattern from those that have acquired resistance. Correlation between NTM AST results and clinical efficacy has so far only been established for selected species-drug combinations, as listed in Table 2.

Table 2. Antimicrobial susceptibility testing (AST) for non-tuberculous mycobacterial species

NTM species	Antimicrobial	Phenotypic AST needed**	Resistance gene to be studied*
<i>M. avium complex</i> (e.g. <i>M. avium</i> , <i>M. intracellulare</i> , <i>M. intracellulare subsp. chimaera</i> , and others)	Clarithromycin/azithromycin Amikacin	Useful in relapse cases Clarithromycin answers for azithromycin.	<i>rrl</i> <i>rrs</i>
<i>M. kansasii</i> , <i>M. szulgai</i>	Rifampicin/rifabutin	Useful in relapse cases Rifabutin answers for rifampicin	<i>rpoB</i>
<i>M. chelonae</i>	Clarithromycin/azithromycin Tobramycin	Useful in relapse cases Clarithromycin answers for azithromycin.	<i>rrl</i> <i>rrs</i>
<i>M. fortuitum complex</i> (e.g. <i>M. fortuitum</i> , <i>M. senegalense</i> , <i>M. peregrinum</i> , <i>M. mucogenicum</i> , and others)	Clarithromycin/azithromycin Levofloxacin, moxifloxacin	Useful in all cases, before starting treatment due to different wild-type profiles with regard to the species. Useful in relapse cases.	<i>erm (39)***</i> <i>gyrA</i>
<i>M. abscessus (subspecies massiliense, abscessus and bollettii)</i> .	Clarithromycin/azithromycin Amikacin	Useful for all cases <u>before</u> starting treatment due to different wild-type profiles among the species/subspecies. Clarithromycin answers for azithromycin. Useful in relapse cases.	<i>erm (41)</i> <i>rrl</i> <i>rrs</i>

* Gene involved in intrinsic or acquired resistance: intrinsic resistance is observed in wild-type strains, acquired resistance is observed mostly in strain isolates under or after treatment.

** In broth microdilution following CLSI recommendations (CLSI M24/M62) or manufacturers recommendations (e.g. Sensititre plates from ThermoFisher).

*** Although *erm (39)* is a gene involved in inducible macrolide resistance in the *M. fortuitum complex*, we do not have much evidence for its prevalence in the complex since it has not been extensively studied so far.

Although previous recommendations suggested testing a broad panel of antimicrobials, such as those listed in the RGM and SGM panels in Table 3, acquired resistance to some of these antimicrobials has not yet been described. Furthermore, there are no validated genotypic or phenotypic criteria for differentiating susceptible from resistant strains. For these antimicrobials, it is recommended to follow the clinical guidelines [11] [9].

Time to results depends on sufficient *in vitro* growth, comparing antibiotic-containing media with drug-free controls. An important exception is the detection of inducible clarithromycin resistance in rapidly growing NTM, due to the functional *erm* gene encoding erythromycin methylase. In such cases, incubation must be extended to 14 days, as false susceptibility is often observed when reading results after only 3–5 days, even if the growth control is already positive [73].

The standard protocol for NTM AST is broth microdilution (BMD), with guidance provided by the Clinical and Laboratory Standards Institute (CLSI) (<https://clsi.org/standards/products/microbiology/documents>) and by EUCAST (www.eucast.org/mycobacteria).

The preferred method for measuring the minimum inhibitory concentration (MIC) in liquid medium is to use microplates, as most of the clinical breakpoints have not been validated. The goal is to classify isolates as having a 'wild-type susceptibility pattern' for the NTM species tested. CLSI-supported clinically relevant breakpoints currently only exist for macrolides (clarithromycin and azithromycin) and amikacin, and only for MAC species and *M. abscessus*. The other breakpoints (i.e. those mentioned in 'CLSI M24S Ed.2E — Performance Standards for Susceptibility Testing of Mycobacteria, *Nocardia* spp., and Other Aerobic Actinomycetes') should be avoided, as they fall within the middle of the wild-type population and do not meet the EUCAST definition of an epidemiological cut-off value (ECOFF) [12]. The ECOFFs and tentative ECOFF (i.e. when MIC distributions are truncated) have been published for macrolides and amikacin against *M. avium*, *M. intracellulare*, and *M. abscessus*, based on multicentre data using these commercial CE-IVD microtiter plates [12].

Two CE-IVD commercial systems are available: RAPMYCO and SLOMYCO from ThermoFisher. Other microdilution plates are available for research use only (Table 3) [12].

Table 3. Commercial systems for phenotypic drug susceptibility testing of nontuberculous mycobacteria

Name of the microplate	SLOMYCO1	SLOMYCO2	RAPMYCO1	RAPMYCO2	FRATMYC1	FRATMYC2
<i>Status</i>	CE-IVD	RUO	CE-IVD	RUO	RUO	RUO
Amikacin	1–64	1–256	1–64	1–256	0.5–512	
Amoxicillin/Clavulanic Acid			2/1–64/32			
Bedaquiline						0.004–4
Cefepime			1–32			
Cefoxitin			4–128	1–128	0.5–256	
Ceftriaxone			4–64			
Ciprofloxacin	0.12–16	0.12–8	0.12–4	0.12–4		0.06–64
Clarithromycin	0.06–8	0.06–64	0.06–16	0.06–16	0.25–128	
Clofazimine		0.015–4		0.03–4		0.008–4
Doxycycline	0.12–16	0.12–8	0.12–16	0.12–8		
Ethambutol	0.5–16					
Ethionamide	0.3–20					
Imipenem/Relebactam						0.12/4–128/4
Imipenem			2–64	0.008–32	0.5–128	
Isoniazid	0.25–8					
Linezolid	1–64	1–32	1–32	1–32	0.25–128	
Meropenem/Vaborbactam						0.25/8–64/8
Minocycline		0.06–8	1–8		0.12–64	
Moxifloxacin	0.12–8	0.015–4	0.25–8	0.015–4	0.06–64	
Omadacycline						0.06–32
Rifabutin	0.25–8	0.12–4				0.03–32
Rifampicin	0.12–8	0.004–4			0.25–8	
Streptomycin	0.5–64	0.5–32				
Tedizolid						0.03–32
Tigecycline			0.015–4	0.03–2	0.03–8	
Tobramycin			1–16	0.12–16	1–32	
Trimethoprim/sulfamethoxazole	0.12/2.38–8/152	0.25/4.75–4/76	0.25/4.75–8/152	0.25/4.75–4/76	0.25/4.75–32/304	

Values indicate the ranges of antibiotic concentrations ($\mu\text{g/mL}$) included in the plates (two-fold dilutions).

* RUO: for research use only.

Molecular detection of resistance-associated gene mutations should be used in conjunction with phenotypic AST.

As mentioned previously, it is critical that personnel are trained and experienced in order to ensure accuracy and reproducibility in NTM AST. Laboratories that do not often encounter NTM are advised to refer samples to specialised reference centres with high AST through-put and proper quality control systems.

One commercial line probe assay, GenoType NTM-DR (Bruker), is available and can be used to detect macrolide and aminoglycoside resistance in *M. abscessus*, *M. chelonae*, and MAC [63]. For drugs used more recently in the

treatment of NTM infections, such as bedaquiline, clofazimine, tigecycline, and carbapenems, there is, as yet, no consensus on genotypic or phenotypic AST [10].

It is important to note that in vitro AST does not reflect antimicrobial activity against intracellular NTM, a key survival niche in the host. Furthermore, NTM are known to form biofilms in vivo, whereas standard AST only evaluates activity against planktonic bacilli and not biofilm-embedded forms.

11.8 Whole genome sequencing and bioinformatic analysis

The same DNA extraction and WGS protocols developed for *M. tuberculosis* can generally be applied to NTM. However, unlike *M. tuberculosis*, many NTM species carry plasmids, undergo genetic recombination, and may acquire genes from other bacteria through horizontal gene transfer. These features complicate genome assembly and can interfere with accurate phylogenetic analysis.

In recent years, there has been a notable increase in the development of WGS-based bioinformatic tools specifically tailored for NTM (Table 4), reflecting growing research interest and diagnostic demand. However, many of these tools remain under development, with limited benchmarking across diverse datasets and a lack of widespread adoption or external validation by independent laboratories. In addition, many free web-based platforms only store results temporarily, which can limit reproducibility and downstream analysis. When working with NTM, it is important to be aware that mixed infections, either involving different NTM species or distinct strains of the same species, are relatively common. Therefore, it is recommended that morphologically distinct colonies should be subcultured and sequencing data carefully reviewed for signs of mixed populations. Indicators may include genome assemblies with an unusually high number of contigs or larger-than-expected genome sizes.

Although Kraken2 is a powerful taxonomic classification tool and particularly useful for detecting contamination with non-NTM bacteria, its results should be interpreted with caution in the context of potential mixed infections. In some cases, Kraken2 may suggest the presence of multiple (sub)species that are not actually present in the sample, likely due to misclassification or artefacts in the reference database. This issue is particularly common when analysing novel species or subspecies within the *M. abscessus* complex.

Table 4. Overview of bioinformatic tools currently available for the analysis of whole genome sequencing data from nontuberculous mycobacteria

Tool Name	Interface	Key functionalities for NTM	Input formats	Specificity	Ref
NTM-Profiler	Command-line	(Sub)species identification, drug resistance prediction	FASTQ, BAM/CRAM, FASTA, VCF	NTM-specific	GitHub - jodyphelan/NTM-Profiler: Profiling NTM WGS data
MyCodentifier*	Command-line	Species identification	FASTQ	TB and NTM	GitHub - JordyCoolen/MyCodentifier: Nextflow pipeline to identify TB and NTM species from NGS data [74]
NTMseq	Command-line	Quality control, contamination detection, (sub)species identification, resistance and plasmid prediction, assembly, MLST, phylogenetics	FASTQ, FASTA	NTM-specific	https://github.com/ngs-fzb/NTMtools [75]
SAM-TB*	Web-based	(Sub)species identification	FASTQ	TB and NTM	[76]
GenoMycAnalyzer*	Web-based	Quality control, species identification	FASTQ	TB and NTM	[77]
Pathogenwatch	Web-based	Species identification, MLST, phylogenetics	FASTA, FASTQ	Microbes	[78]
TYGS	Web-based	Species identification, phylogenetics	FASTA	Microbes	[79]
SeqSphere+	Desktop-based	(cg)MLST, SNP analysis, phylogenetics, cluster analysis	FASTQ, FASTA	Microbes	[80]
pubMLST	Web-based	MLST, phylogenetics, cluster analysis	FASTA	Microbes	[81]

* All tools are available at no cost, with the exception of SeqSphere+, which is a commercial product.

**Not validated by authors.

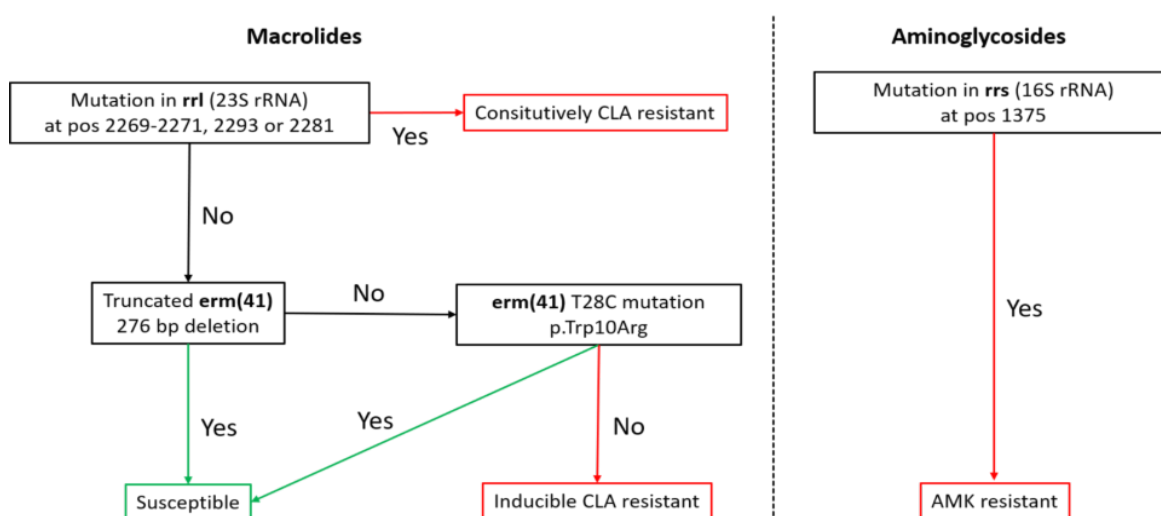
*** Ref: reference.

Tools for typing and transmission analysis often rely on the core genome multi locus sequence typing (cgMLST) approach, which offers inherent standardisation and easy incorporation of new datasets into a comparison enabling ongoing real-time surveillance. However, a validated and published cgMLST scheme in SeqSphere+, pubMLST and Pathogenwatch with thresholds for cluster analysis is currently only available for *M. abscessus* [82]. In addition to cgMLST, NTM isolates can also be compared using the single nucleotide polymorphism (SNP) approach, similar to that used for MTB. Therefore tools such as MTBseq [83] and Snippy can be repurposed for SNP-based analysis of NTM, provided that the reference genome is replaced with that of an NTM species.

NTM exhibit a wide array of intrinsic resistance mechanisms that significantly complicate antimicrobial therapy. These mechanisms include biofilm formation, low permeability of their cell wall, efflux pumps, and specific enzymatic modifications that inactivate antibiotics or alter their targets [84] [85]. In addition, NTM can acquire additional resistance, especially during antibiotic therapy. Unlike MTB, there are currently no established WHO-endorsed mutation catalogues available for NTM. However, initiatives to develop such catalogues are gaining momentum.

At present, validated WGS-based resistance prediction is only available for *M. abscessus*. This is based on a defined interpretation scheme (Figure 19) that incorporates a limited number of well-characterised mutations linked to resistance against macrolides and aminoglycosides, which are key antibiotics in *M. abscessus* treatment. Detection of these mutations in short-read WGS datasets can be done with bioinformatic tools, such as NTM-profiler.

Figure 19. Decision algorithm for resistance prediction of macrolides and aminoglycosides in *M. abscessus*



Adapted from [86].

In the NTM-DR genotype assay, the following positions are analysed: position 28 in the *erm(41)* gene, positions 1406 to 1409 in the *rrs* gene (based on *E. coli* numbering; equivalent to positions 1374 to 1376 in *Mycobacterium abscessus* ATCC19977), and positions 2058 to 2059 in the *rrl* gene (*E. coli* numbering; corresponding to positions 2270 to 2271 in *M. abscessus* ATCC19977). A truncated *erm(41)* is identified when the assay indicates the presence of subsp. *massiliense*.

Given that WGS analysis of NTM is still in its early stages, it may be advisable to use multiple tools in parallel to cross-validate results.

11.9 Conclusions

Diagnostics of NTM can be performed in laboratories across the European Union, as most diagnostic methods rely on those used for TB. These methods are typically available, as a minimum, in national reference laboratories.

Human samples submitted for TB testing are frequently positive for NTM instead. In such cases, when TB has been ruled out, laboratory strategies for handling positive NTM cultures can be inspired by the example provided in Supplementary Figure 1.

When NTM infection is suspected, this chapter offers guidance and examples of methods to support the diagnosis. Although expertise in NTM diagnosis is shared across the EU by microbiologists, molecular biologists, researchers, epidemiologists, and clinicians, further research is still needed in several areas.

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Supplementary material

Supplementary Table 1. Example of primers used for multiplex PCR, Sanger sequencing, real-time PCR and restriction fragment analysis

Target species	Target Region	Fw name	Fw sequence 5'-3'	Rv name	Rv sequence 5'-3'	Length (bp)*	Sequence primer	Reference (PMID)
Mycobacteria	ITS	MYCF	GATTGGGACGAAGTCGTAACAAG	MYCR	CTCGTTGACAGCTCCCGAG	438	MYCF/MYCR	20697162
Mycobacteria	ITS	MYITSF = MYCF	GATTGGGACGAAGTCGTAACAAG	MYITSR	AGCTCCCACGTCCTTCATCGGCT	400-500	MITSFS:TGGATAGTGGTTGCGAGCAT and MITSRS: GATGCTCGCAACCACTATCCA	18231831
Mycobacteria	ITS	ITS-F	TGGATCCGACGAAGTCGTAACAAG G	ITS-R	TGGATCTGCCAAGGCATCCACCAT	350-450	ITS-F/ITS-R	15814999
Mycobacteria	ITS	ITS-F	TGGATCCGACGAAGTCGTAACAAG G	PAN-04R	ATGCTBCAABCACTATCCA	271	ITS_n16f: CTTTCTAAGGAGCACCACGA	11060072 ; 15814999
Mycobacteria	ITS	Ec16S.1390p	TTGTACACACGCCCGTCA	Mb23S.44n	TCTCGATGCCAAGGCATCCACC	480	Ec16S.1390p/Mb23S.44n complement	9431937
Mycobacteria	16S rRNA	B1 (8F)	TGGAGAGTTTGATCCTGGCTCAG	Primer 264	TGCACACAGGCCACAAGGGA	1029	B9: CGTGCTTAACACATGCAAGTC	30415449 ; 7505291
Mycobacteria	16S rRNA	4F	TTGGAGAGTTTGATCCTGGCTC	801R	GGCGTGGACTTCCAGGTATCT	786	4F/801R	25932452
Mycobacteria	16S rRNA	Seq1 (KY18)	CACATGCAAGTCGAACGGAAGG	Seq2 (modified KY75)	GCCCGTATCGCCGACGCT	582	Primer 244: CCCACTGCTGCCTCCCGTAG and 259: TTTCACGAACAACGCGACAA	9431937 ; 7505291
Mycobacteria	16S rRNA	Primer 285	GAGAGTTTGATCCTGGCTCAG	Primer 264	TGCACACAGGCCACAAGGGA	1026	Primer 244	7505291
Bacteria	16S rRNA full	pA = B1 - 3bp	AGAGTTTGATCCTGGCTCAG	pHr	AAGGAGGTGATCCAGCCGCA	1525	pA/pHr	2798131
Bacteria	16S rRNA	pA = B1 - 3bp	AGAGTTTGATCCTGGCTCAG	pDr	GTATTACCGCGGCTGCTG	515	pA/pDr	2798131
Bacteria	16S rRNA	B1 (8F)	TGGAGAGTTTGATCCTGGCTCAG	D = pDr + 2 bp	ACGTATTACCGCGGCTGCTG	520	B9: CGTGCTTAACACATGCAAGTC	10351948
Mycobacteria	hsp65 (Telenti fragment)	Tb11	ACCAACGATGGTGTGCCAT	Tb12	CTTGTGGAACCGCATAACCT	441	Restriction analysis	8381805
Mycobacteria	hsp65	HSPF3	ATCGCCAAGGAGATCGAGCT	HSPR4	AAGGTGCCGCGATCTTGTT	604	HSPF3/HSPR4	25932452
Mycobacteria	rpoB-V	MycF	GGCAAGGTCACCCGAAGGG	MycR	AGCGGCTGCTGGGTGATCATC	752	MycoseqF: GAAGGGTGAAGAC and MycoseqR: GCTGGG TGATCATCGAGTACGG CGAGCTGAC	14662964
Mycobacteria	argH	ARGHF	GACGAGGGCGACAGCTTC	ARGHS	GTGCGGAGCAGATGATG	661	ARGHF/ARGHS	36121509
Mycobacteria	cya	ACF	GTGAAGCGGGCCAAGAAG	ACFR1	AACTGGGAGGCCAGGAGC	647	ACF/ACF1	36121509

Target species	Target Region	Fw name	Fw sequence 5'-3'	Rv name	Rv sequence 5'-3'	Length (bp)*	Sequence primer	Reference (PMID)
<i>M. avium complex</i>	ITS	MACF	CCCTGAGACAACACTCGGTC	MACR	GTTTCATCGAAATGTGTAATT	141	multiplex PCR/Gel electrophoresis	11060072
<i>M. fortuitum</i>	n.r	FORF	CCGTGAGGAACCGTTGCCCT	FORR	TAGCACGCAGAATCGTGTGG	223	multiplex PCR/Gel electrophoresis	11060072
<i>M. chelonae</i>	n.r	CHEF	GTTACTCGCTTGGTGAATAT	CHER	TCAATAGAATTGAAACGCTG	93	multiplex PCR/Gel electrophoresis	11060072
<i>M. goodnae</i>	n.r	GORF	CGACAACAAGCTAAGCCAGA	GORR	GCATCAAATGTATGCGTTG	152	multiplex PCR/Gel electrophoresis	11060072
<i>M. scrofulaceum</i>	n.r	SCOF	TCGGCTCGTTCTGAGTGGTG	SCOR	TAAACGGATGCGTGGCCGAA	99	multiplex PCR/Gel electrophoresis	11060072
<i>M. szulgai</i>	n.r	SZUF	AACACTCAGGCTTGCCAGA	SZUR	GAGGGCAGCGCATCCAATTG	105	multiplex PCR/Gel electrophoresis	11060072
<i>M. avium</i>	16S	M. avium FWD-01	CCTCAAGACGCATGTCTTC	MAC REV-02	ACCTACCGTCAATCCGAGAA	299	Real time PCR	19297596
<i>M. intracellulare</i>	16S	M. intracellulare FWD-02	GACCTTTAGRCGCATGTCTTT	MAC REV-02	ACCTACCGTCAATCCGAGAA	301	Real time PCR	19297596
Mycobacteria	16S	AFB genus FWD-06	CCGCAAGRCTAAAACCTCAA	AFB genus REV-01b = Primer 264	TGCACACAGGCCACAAGGGA	149	Real time PCR	19297596
M. chelonae/abscessus	ITS	MCAG FWD-01	TAAGGAGCACCATTCCCAG	MCAG REV-01	CGACGTTTTGCCGACTACC	128	Real time PCR	19297596
M. fortuitum group	16S	MFG FWD-01	CCACGCGCTTCATGGTGT	MFG REV-01	ACTTGCGCTTCGTCCTAT	286	Real time PCR	19297596
M. fortuitum group	16S	MFG FWD-02	CCGCGCTTCATGGGGT	MFG REV-01	ACTTGCGCTTCGTCCTAT	286	Real time PCR	19297596
M. fortuitum group	16S	MFG FWD-03	ACCACGCATTCATGGTGT	MFG REV-01	ACTTGCGCTTCGTCCTAT	287	Real time PCR	19297596
Mycobacteria	Ag85b	MT1	TTCCTGACCAGCGAGCTGCCG	MT2	CCCCAGTACTCCAGCTGTGC	500	multiplex PCR/Gel electrophoresis	8789008
M. avium complex	DnaJ	DnaJ-1	GACTTCTACAAGGAGCTGGG	DnaJ-2	GAGACCGCCTTGAATCGTTC	140	multiplex PCR/Gel electrophoresis	18037461
M. avium	16S rRNA	MYCGEN-F = pA	AGAGTTTGATCTTGCTCAG	MYCAV-R	ACCAGAAGACATGCGTCTTG	180	multiplex PCR/Gel electrophoresis	18037461
M. avium subsp. avium/silvaticum	IS901	IS901-1	GGATTGCTAACCACGTGGTG	IS901-2	GCGAGTTGCTTGATGAGCG	577	multiplex PCR/Gel electrophoresis	18037461
M. avium subsp. avium/silvaticum and hominissuis	IS1245	IS1245-1	GAGTTGACCGGTTTCATCG	IS1245-2	CGTCGAGGAAGACATACGG	385	multiplex PCR/Gel electrophoresis	18037461

* Length can vary depending on species. ** ITS = internal transcribed spacer. *** nr: not reported.

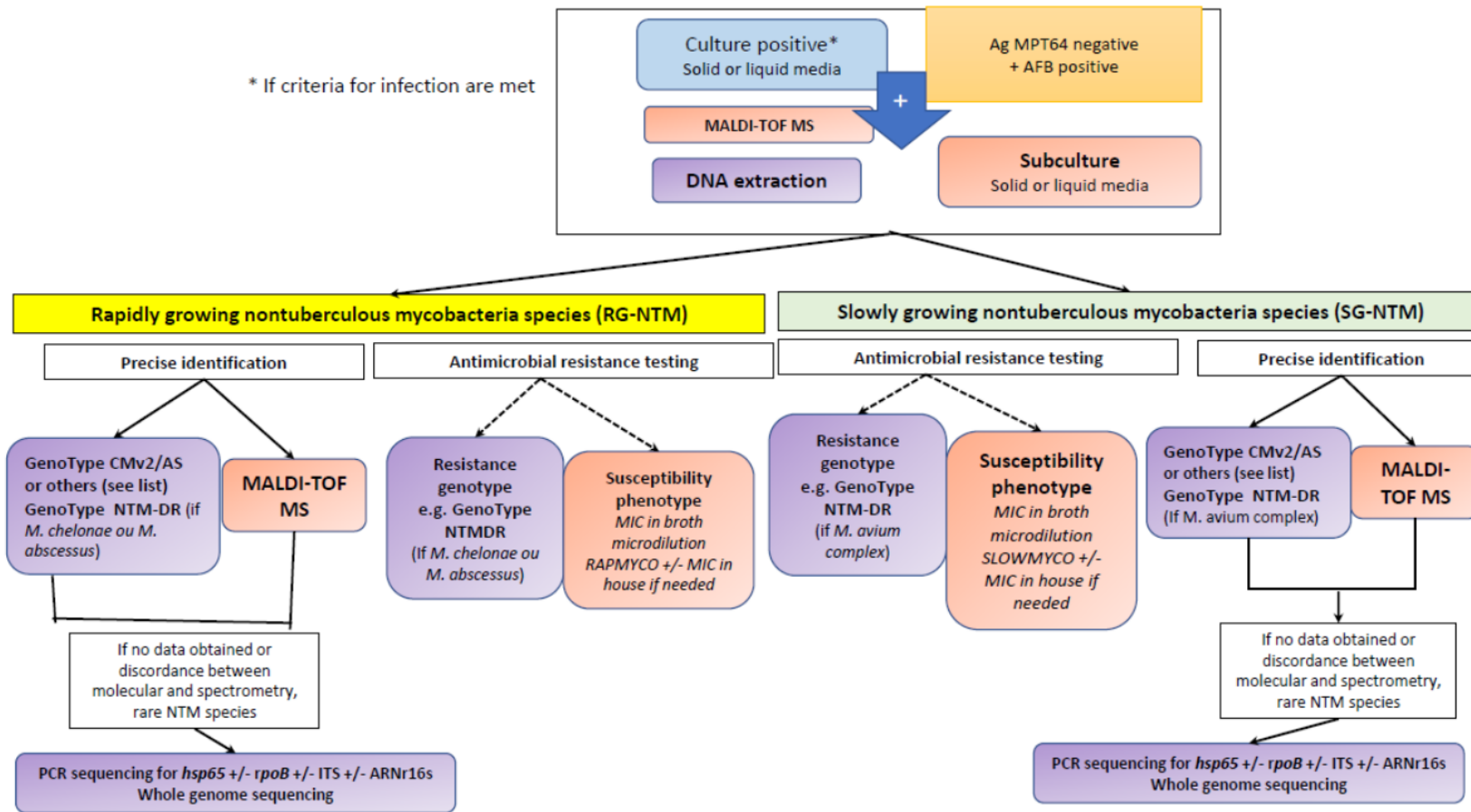
Supplementary Table 2. Commercial molecular tests for the identification and species differentiation of nontuberculous mycobacteria

Company	GenoType CMdirect V1.0	Genotype CM V2.0	GenoType AS V1.0	Genotype NTM-DR V1.0	INNO-LiPA MYCOBACTERIA V2	Genoscholar™ NTM+MDR TB	Fluorotype Mycobacteria V1	Anyplex MTB/NTM Me	NeoPlexTM TB/NTM
Type	Bruker	Bruker	Bruker	Bruker	Fujirebio	Nipro	Bruker	Seegene	Genematrix
	LPA	LPA	LPA	LPA	LPA	LPA	RT PCR with fluorescence	RT PCR with fluorescence	RT PCR with fluorescence
Status	CE-IVD	CE-IVD	CE-IVD	CE-IVD	CE-IVD	CE-IVD; WHO approved	CE-IVD	CE-IVD	CE-IVD
Input	Primary specimen	DNA from culture	DNA from culture	DNA from culture	DNA from culture	DNA from culture	DNA from culture	Primary specimen	Primary specimen
<i>M. abscessus</i>	✓	✓		✓			✓		
<i>M. abscessus subsp. abscessus</i>				✓			✓		✓
<i>M. abscessus subsp. bolletii</i>				✓			✓		
<i>M. abscessus subsp. massiliense</i>				✓			✓		✓
<i>M. asiaticum</i>			✓				✓		
<i>M. avium</i>	✓	✓		✓	✓	✓	✓		✓
<i>M. celatum</i>			✓		✓		✓		
<i>M. chelonae</i>	✓	✓		✓	✓		✓		
<i>M. chimaera</i>				✓			✓		
<i>M. fortuitum group</i>	✓	✓			✓		✓		
<i>M. gastri</i>			✓				✓		
<i>M. genavense</i>			✓		✓		✓		
<i>M. goodii</i>			✓				✓		
<i>M. goodii</i>			✓				✓		
<i>M. goodii</i>			✓				✓		
<i>M. gordonae</i>	✓	✓			✓		✓		

Company	GenoType CMdirect V1.0	Genotype CM V2.0	GenoType AS V1.0	Genotype NTM-DR V1.0	INNO-LiPA MYCOBACTE RIA V2	Genoschola ⁺ ™ NTM+MDR TB	Fluorotype Mycobacteria V1	Anyplex MTB/NT Me	NeoPlexT M TB/NTM
<i>M. haemophilum</i>		(✓)	✓		✓		✓		
<i>M. heckeshomense</i>			✓				✓		
<i>M. intermedium</i>			✓				✓		
<i>M. interjectum</i>	✓	✓					✓		
<i>M. intracellulare</i>	✓	✓		✓	✓	✓	✓		✓
<i>M. kansasii</i>	✓	✓	✓		✓	✓	✓		✓
<i>M. lentiflavum</i>			✓				✓		
<i>M. malmoense</i>	✓	✓			✓		✓		
<i>M. marinum</i>	✓	✓			✓		✓		
<i>M. mucogenicum</i>			✓				✓		
<i>M. peregrinum</i>							✓		
<i>M. phlei</i>			✓				✓		
<i>M. scrofulaceum</i>	✓	✓			✓		✓		
<i>M. shimoidei</i>			✓				✓		
<i>M. simiae</i>			✓		✓		✓		
<i>M. smegmatis</i>			✓		✓		✓		
<i>M. szulgai</i>	✓	✓	✓				✓		

Company	GenoType CMdirect V1.0	Genotype CM V2.0	GenoType AS V1.0	Genotype NTM-DR V1.0	INNO-LiPA MYCOBACTERIA V2	Genoscholar™ NTM+MDR TB	Fluorotype Mycobacteria V1	Anyplex MTB/NTM Me	NeoPlexTM TB/NTM
<i>M. ulcerans</i>	✓	✓	✓		✓		✓		
<i>M. xenopi</i>	✓	✓			✓		✓		
TB vs NTM	✓	✓			✓	✓		✓	✓
Resistance prediction MAC*				✓					
Resistance prediction <i>M. abscessus</i>*				✓					
Comment	Cannot differentiate <i>M. ulcerans</i> / <i>M. marinum</i>	Cannot differentiate <i>M. ulcerans</i> / <i>M. marinum</i> ; <i>M. abscessus</i> and <i>M. chelonae</i> / <i>M. immunogenum</i> ; <i>M. (para)scrofulaceum</i> / <i>M. paraffinicum</i> ; <i>M. malmoense</i> / <i>M. haemophilum</i> / <i>M. palustre</i> / <i>M. nebraskense</i>	Cannot differentiate <i>M. genavense</i> / <i>M. triplex</i> ; <i>M. szulgai</i> / <i>M. intermedium</i> ; <i>M. haemophilum</i> / <i>M. nebraskense</i>	Cannot differentiate <i>M. chelonae</i> / <i>M. immunogenum</i> ; Rare cases of <i>M. abscessus</i> subsp. <i>abscessus</i> or <i>bolletii</i> with truncated <i>erm(41)</i> genes might be misclassified as subsp. <i>massiliense</i>	Cannot differentiate <i>M. ulcerans</i> / <i>M. marinum</i>		Cannot differentiate <i>M. heckeshornense</i> / <i>M. xenopi</i>		

Supplementary Figure 1. Example of laboratory strategy when culture is positive with nontuberculous mycobacteria



Source: ECDC

Annex 3. Network partners

This list, which was revised by Yen Holicka in 2022, includes both former and current contributors to the project.

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