




Identification of *Mycobacterium tuberculosis* Resistance to Common Antibiotics: An Overview of Current Methods and Techniques

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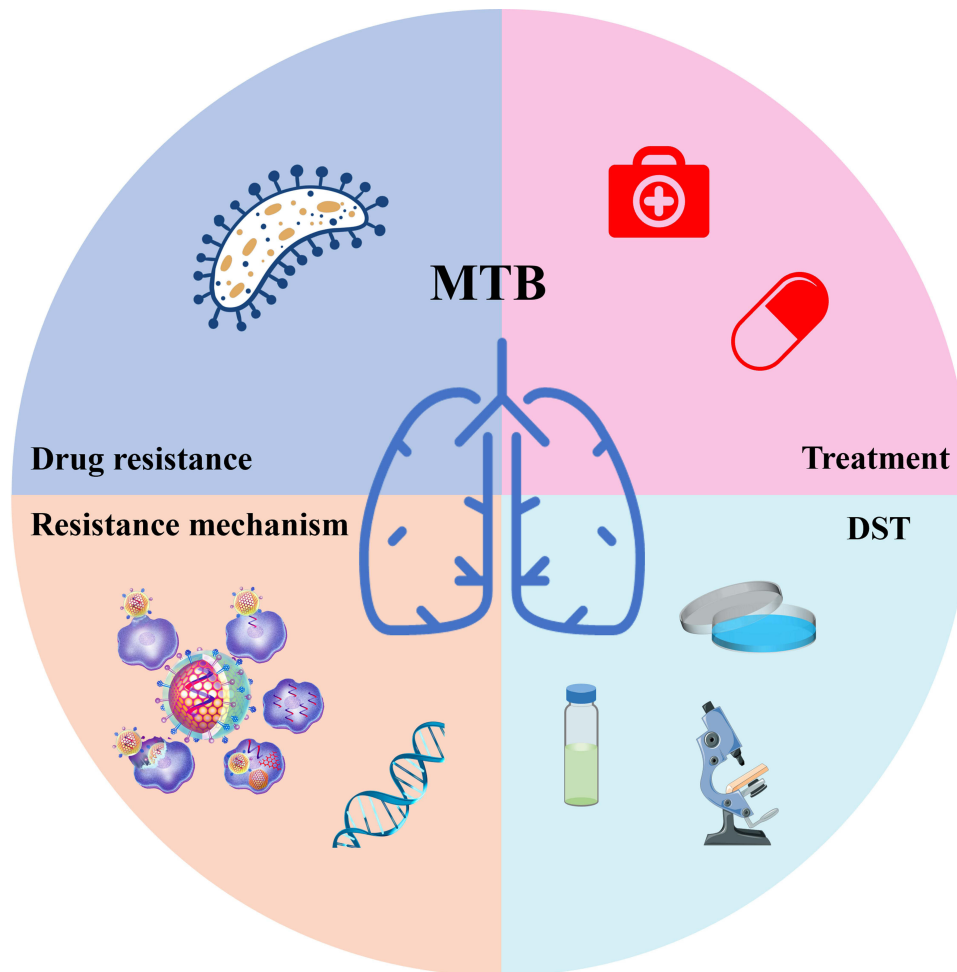
Abstract: Multidrug-resistant tuberculosis (MDR-TB) is an essential cause of tuberculosis treatment failure and death of tuberculosis patients. The rapid and reliable profiling of *Mycobacterium tuberculosis* (MTB) drug resistance in the early stage is a critical research area for public health. Then, most traditional approaches for detecting MTB are time-consuming and costly, leading to the inappropriate therapeutic schedule resting on the ambiguous information of MTB drug resistance, increasing patient economic burden, morbidity, and mortality. Therefore, novel diagnosis methods are frequently required to meet the emerging challenges of MTB drug resistance distinguish. Considering the difficulty in treating MDR-TB, it is urgently required for the development of rapid and accurate methods in the identification of drug resistance profiles of MTB in clinical diagnosis. This review discussed recent advances in MTB drug resistance detection, focusing on developing emerging approaches and their applications in tangled clinical situations. In particular, a brief overview of antibiotic resistance to MTB was present, referred to as intrinsic bacterial resistance, consisting of cell wall barriers and efflux pumping action and acquired resistance caused by genetic mutations. Then, different drug susceptibility test (DST) methods were described, including phenotype DST, genotype DST and novel DST methods. The phenotype DST includes nitrate reductase assay, Roche™ solid ratio method, and liquid culture method and genotype DST includes fluorescent PCR, GeneXpert, PCR reverse dot hybridization, ddPCR, next-generation sequencing and gene chips. Then, novel DST methods were described, including metabolism testing, cell-free DNA probe, CRISPR assay, and spectral analysis technique. The limitations, challenges, and perspectives of different techniques for drug resistance are also discussed. These methods significantly improve the detection sensitivity and accuracy of multidrug-resistant tuberculosis (MRT) and can effectively curb the incidence of drug-resistant tuberculosis and accelerate the process of tuberculosis eradication.

Keywords: MTB, antibiotic resistance, Raman spectroscopy, rapid detection

Introduction

Just as the 2023 global tuberculosis report released by the World Health Organization (WHO), 7.5 million tuberculosis patients were newly diagnosed globally in 2022, with an estimated 1.3 million deaths, nearly twice the rate of HIV/AIDS. At present, tuberculosis, a class A infectious disease aroused by *Mycobacterium tuberculosis* (MTB), has turned into the first cause of death of a single infectious disease after novel coronavirus pneumonia. Although the clinical detection rate of MTB has increased, over 90% of the pathogenic bacteria of human pulmonary tuberculosis are MTB. Its main transmission routes are air and contact, and it is easy to form an explosive epidemic. The main high-risk places include

Graphical Abstract



schools, nursing homes, and other vital social places, which cause potential significant harm to public security. The treatment cycle of tuberculosis and multidrug-resistant tuberculosis is time-consuming and expensive, resulting in a substantial social and economic burden on countries and individual patients.¹ Then, the incidence data of multidrug-resistant tuberculosis (MRT) and rifampicin-resistant tuberculosis in China is the fourth in the world, and China is one of the high-burden countries with drug-resistant tuberculosis in the world. So, timely and effective identification of MTB infection diagnosis and its drug resistance can effectively realize the early detection, early control, and precise treatment of tuberculosis, an effective tuberculosis prevention and control measure. Therefore, rapid identification of tuberculosis infection and drug resistance types is crucial to curb the current high incidence rate, high mortality, and high economic burden of tuberculosis. In a word, the treatment situation is dire.²

Moreover, the treatment of tuberculosis is facing pain points such as long cycles, high drug resistance rate, and prolonged drug sensitivity reporting time, which brings great trouble to the diagnosis and treatment of tuberculosis. This study aims to explore an immediate method for rapid drug sensitivity diagnosis of MTB and to provide tuberculosis patients with an efficient and reliable laboratory basis for treatment. The diagnosis of tuberculosis mainly includes smear staining, gamma interferon test, bacterial culture, metagenome sequencing, targeted DNA sequencing technology, mass spectrometry technology, and other emerging technologies. The diversity of diagnostic methods provides more selectivity for diagnosing and treating tuberculosis, but the help for tuberculosis still needs to be improved. To realize the early

diagnosis and treatment of tuberculosis, rapid results of drug sensitivity diagnosis of tuberculosis must be required. Therefore, we have summarized the latest progress in drug resistance detection of MTB and attempted to find a fast, low-cost, and easy-to-operate drug sensitivity detection system for MTB.

Antibiotic Resistance of MTB

The drug resistance mechanism of MTB is complex. Currently, the partially elucidated mechanism is divided into intrinsic and acquired resistance. Inherent drug resistance mainly includes reduced cell wall permeability, efflux pump function, and cell metabolism. Mutations in target genes mainly cause acquired drug resistance and are the primary mechanism in MTB. The main resistance mechanisms of MTB are shown in Figure 1, and one of these acquired resistance is exemplified by the *inhA* gene mutations that caused resistance to isoniazid.

Inherent Drug Resistance

The inner layer of the cell wall of MTB was made up of peptidoglycan, arabinogalactan, and mycolic acid, which are jointed to form the mycolic acid–peptidoglycan–arabinogalactan (MA-AG-PG) complex, constituting a permeable barrier.³¹ Then, the outer cell wall is mainly covered by lipids and proteins. Therefore, the reduced permeability of the cell wall of MTB will interfere with the transport of nutrients or drugs. At the same time, the low permeability plays a protective role in the mycobacteria survival in the host, resulting in the drug resistance of MTB. The cell wall barrier makes MTB impermeable to most antibiotics.³² The survival and pathogenicity of MTB depend on the trehalose, the critical ingredient of glycolipids, with the help of type I ATP-binding cassette (ABC) transporter LpqY-SugABC, the only pathway for trehalose to enter MTB.^{33,34} Studies have shown that trehalose metabolism is related to drug tolerance in

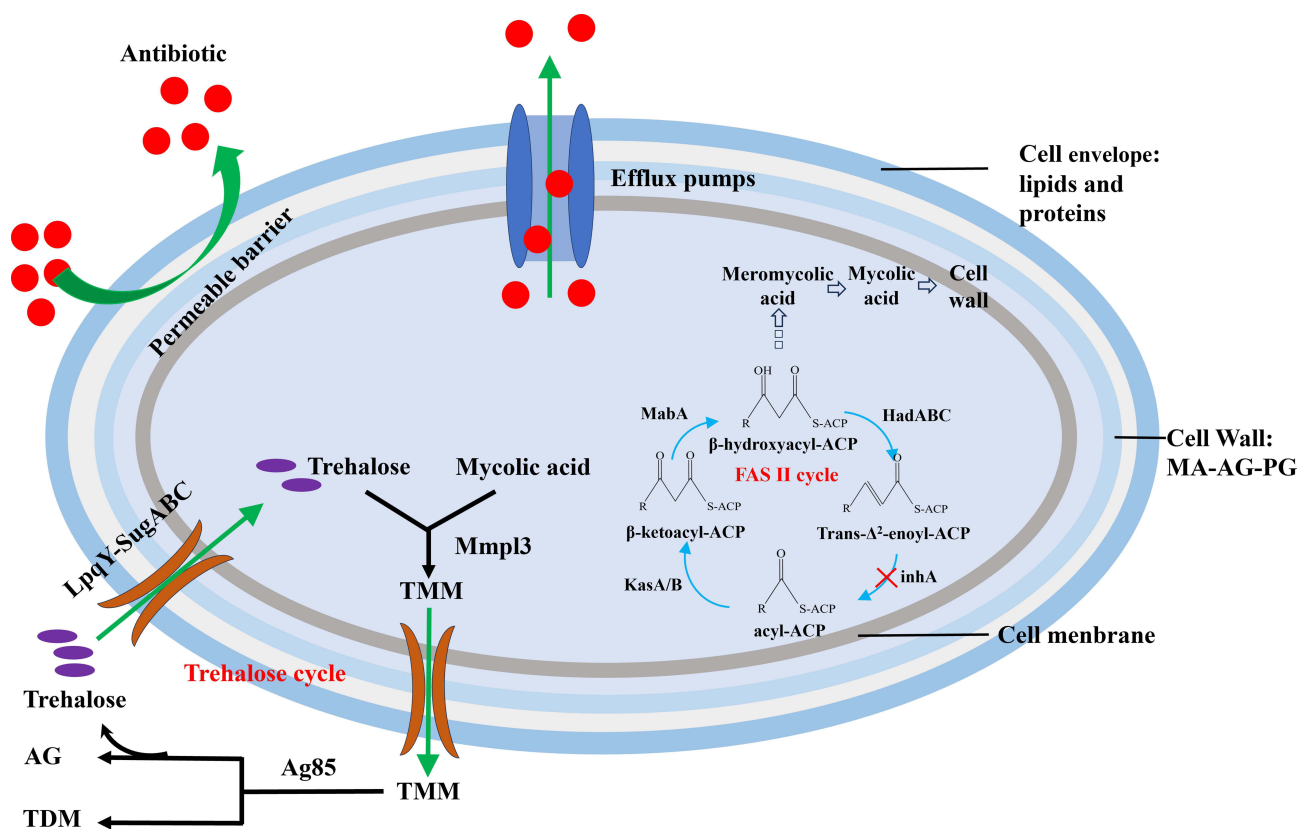


Figure 1 Schematic illustration of the molecular mechanism of MTB antibiotic resistance. There are four ways to show the molecular mechanism: cell wall permeability, efflux pump function, cell metabolism (trehalose cycle), and resistance mutations (taking the *inhA* gene for example). MA-AG-PG: mycolic acid–arabinogalactan–peptidoglycan; AG: arabinogalactan; TMM: trehalose monomycolate; TDM: trehalose dimycolate; Ag85: mycolyltransferase antigen 85; MmpL3: Mammalian membrane protein large 3; LpqY-SugABC: The type I ATP-binding cassette (ABC) transporter; ACP: enoyl-acyl carrier protein; FAS II cycle: fatty acid synthase II cycle systems; Maba, KasA/B, HadABC, InhA: four enzymes of the FAS-II cycle.

MTB.³⁵ Meantime, cellular metabolism is also an extremely important mechanism of drug resistance in MTB. Then, the efflux pump system also plays a specific role in developing drug resistance in MTB. The efflux pump is a transport protein located in the bacterial plasma membrane. When antibiotics enter the bacteria to act, the gene-associated efflux pump can be overexpressed so that the expression of efflux pump-related proteins is up-regulated, and the antibiotics in the body will be actively removed so that the reduction of drug concentration in the bacterium, resulting in the formation of drug resistance. There are currently 6 types of efflux pumps discovered in bacteria, including major facilitator superfamily (MFS), ATP-binding cassette family (ABC), protein bacterial antibacterial compound efflux family (proteobacterial antimicrobial compound efflux family (PACE), small multidrug resistant family (SMR), resistance–nodulation–division family (RND), and multidrug-resistant and toxic compound efflux Family (Multidrug and toxic compound extrusion, MATE).³⁶ Previous studies showed verapamil, one of the efflux pump inhibitors, could effectively inhibit drug efflux.³⁷ At the same time, studies have also demonstrated that combining verapamil with isoniazid or rifampicin (RIF) can cause the MIC of isoniazid (INH) or RIF reduction and even reverse the resistance of MTB.³⁸ About 20 genes in the genus MTB have been identified as related genes encoding efflux pumps, mainly concentrated in MFS, RND, and SMR.³⁹ It was shown that most of the exocytosis pump gene expression levels were upregulated in isoniazid-resistant MTB strains exposed to low concentrations of antibiotics.

Acquired Drug Resistance

Acquired resistance may occur through mutations or horizontal gene transfer. However, studies on the horizontal transfer of resistance genes through plasmids or removable genetic elements in MTB have yet to be reported. All currently known acquired resistance to MTB is mediated by chromosome mutations in the presence of antibiotics.⁴⁰ RIF resistance in MTB occurs due to mutations in the *rpoB* gene encoding the β subunit of RNA polymerase with 1178 amino acids. Mutations in the gene cause a change in the amino acids, resulting in a conformational change in the RIF binding site on the RNA polymerase molecule, which results in a loss of the ability to bind RIF, leading to the development of resistance.³ Mutations usually occur at codons 531, 526, and 516. Then, MIC assays have shown that mutations in codons 531 and 526 are associated with high levels of resistance to RIF. In contrast, mutations in codons 511, 516, 518, and 522 can only lead to low resistance levels to RIF.³ The *inhA* gene encodes an enoyl lipid acyl carrier protein reductase that binds to the NADH and fatty acid synthase II (FAS-II) complexes and forms a covalent 4-subunit complex with the β -ketolipid acyl carrier protein synthase (*kasA*) and the acyl carrier protein AcpM. The activated INH attacks the target site, prevents the production of mature mycolic acid, and depletes the mature mycolic acid.^{41,42}

It has been shown that *Ald* gene deletion increases MTB resistance to cycloserine by testing and comparing MTB MICs of sensitive MTB strains, resistant strains, and *Ald* knockout strains.¹⁰ The *Ald* gene encodes L-alanine dehydrogenase, which catalyzes the conversion of pyruvate to L-alanine with the participation of NADH, hindering the biosynthesis of peptidoglycan. Thereby, the cell wall of MTB is damaged.⁴³ Moreover, the mechanism of resistance to capreomycin (CPM) is mainly due to mutations in the *tea* and *rrs* genes, leading to changes in the target of drug action. Mutations in the *tlyA* gene result in a lack of methylation of bacterial ribosomes, leading to the development of drug resistance.^{12,13} And the *tlyA* mutations primarily cause low-level resistance to CPM.⁴⁴ The *rrs* gene encodes 16S rRNA, and mutations in this gene lead to alterations in the CPM action target, resulting in drug resistance.^{12,14} In addition, mutations in the *eis* gene can lead to CPM resistance.¹⁵ The standard mutant gene segments and functions of the main anti-tuberculosis drugs are shown below (Table 1).

As is known to all, antibiotics work by binding to the target protein on the cell wall, cell membrane, or intracellular to inhibit the synthesis of nucleic acid, protein, and the associated metabolic pathway (Figure 2).

Phenotypic Drug Sensitivity Test

Currently, chemotherapy for MTB follows a “one dose fits all” principle. Yet, the individual differences of the anti-tuberculosis drug were neglected, referring to absorption, distribution, metabolism, and drug excretion. High-performance liquid chromatography mass was used to monitor the first- and second-line antibiotic concentrations in plasma or serum and was successfully applied in two clinical patients.⁴⁵ The drug resistance of MTB is mainly about the peripheral lipids of the MTB cell envelope and the increased amounts of nonpolar lipids.⁴⁶ The classic

Table I Summary of Antibiotic Resistance Genes in Drug-Resistant MTB

Drugs	Genes	Functionality	Reference
Rifampicin	<i>ropB</i>	Encodes the β subunit of RNA polymerase	[3]
Isoniazid	<i>katG</i>	Encodes catalase-peroxidase	[4]
	<i>inhA</i>	Encodes NADH-enoyl-dependent acetyl carrier protein reductase	[4]
Streptomycin	<i>rpsL</i>	Encodes S12	[5]
	<i>rrs</i>	Encodes 16S RNA	[5]
	<i>gidB</i>	Encodes 7-methylguanosine-methyltransferase	[6]
Ethambutol	<i>embB</i>	Encodes arabinofuranosyltransferase, related to arabinogalactan synthesis	[7–9]
	<i>embC</i>	Encodes arabinofuranosyltransferase, related to arabinoglycan lipid synthesis	[7–9]
	<i>embA</i>	Encodes arabinofuranosyltransferase, related to arabinogalactan synthesis.	[7–9]
Cycloserine	<i>Ald</i>	Encodes L-alanine dehydrogenase	[10]
	<i>Alr</i>	Encodes alanine racemase	[11]
	<i>ddlA</i>	Encodes D-ala-D-ala ligase	[11]
Capreomycin	<i>tlyA</i>	Regulate methylation of bacterial ribosomes	[12,13]
	<i>rrs</i>	Encodes 16S rRNA	[12,14]
	<i>eis</i>	Encodes aminoglycoside acetyltransferase	[15]
Linezolid	<i>rrl</i>	Encodes 23S rRNA	[16]
	<i>rplC</i>	Encodes the ribosomal L3 protein	[17,18]
Pyrazinamide	<i>PncA</i>	Encodes pyrazinamidase	[19]
	<i>rpsA</i>	Encodes the 30S ribosomal protein S1	[19]
	<i>panD</i>	Encodes aspartate decarboxylase	[19]
Bedaquiline	<i>Rv0678</i>	Enhances the expression of MTB membrane exocytosis MmpS5/MmpL5 membrane proteins	[20]
	<i>atpE</i>	Encodes the c subunit of F1/F0-ATP synthase	[21]
	<i>pepQ</i> (<i>Rv2535c</i>)	Encodes a protein with two structural domains, a 100-aa N-terminal α/β structural domain and 250 KDA C-terminal peptidase structural domain	[22]
	<i>rv1979c</i>	Encodes a permease involved in amino acid transport	[23]
Clofazimine	<i>Rv0678</i>	The encoded product is a regulatory protein	[24,25]
	<i>Rv1979c</i>	Efflux pump expression	[25]
	<i>Rv2535c</i>	Permease, Peptidase	[25]
Delamanid	<i>ddn</i>	Encodes F420-cofactor-dependent nitroreductase	[26]
	<i>fbIA</i>	Encodes 2-phospholactate transferase	[27]
	<i>fbIB</i>	Encodes L-glutamate ligase	[28]
	<i>fbIC</i>	Encodes FO synthase	[29]
	<i>fgdI</i>	Encodes 6-phosphogluconate dehydrogenase	[30]

Notes: The italicized fonts represent the mutation genes associated with drug resistance.

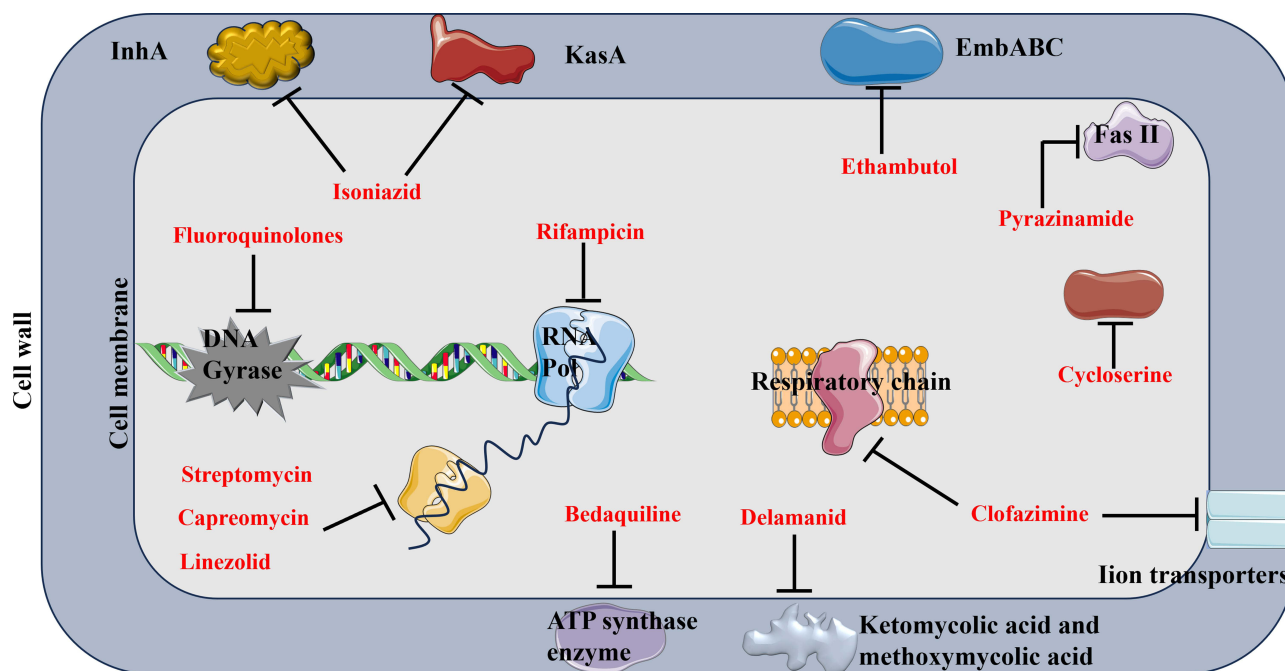


Figure 2 Schematic diagram of MTB drug resistance mechanism. Four major modes of anti-MTB drug mechanism, as follows: work on cell wall proteins (isoniazid, ethambutol, bedaquiline, delamanid, and clofazimine); work on DNA replication machinery (fluoroquinolones); work on translational machinery (streptomycin, capreomycin, linezolid) and work on metabolic pathways (pyrazinamide and cycloserine). InhA, KasA, EmbABC: the target protein on the cell wall to corresponding antibiotics. Note: Reprinted from Life Sciences, 274, Aditi Chauhan, Manoj Kumar, Awanish Kumar, Kajal Kanchan, Comprehensive review on mechanism of action, resistance and evolution of antimycobacterial drugs, 119301, Copyright 2021, with permission from Elsevier.²⁵

laboratory examination method for MTB includes an E-test, nitrate reductase assay, bacterial culture method, and molecular diagnostic technique, with different features and limitations. Although E-tests could successfully rapidly determination of drug resistance to streptomycin (STM), INH, RIF, and ethambutol (EMB), it existed many limitations, such as the high risk of cross-contamination, laboratory infection, expense, high false-positive rate.⁴⁷

Roche™ Solid Ratio Method

The Roche™ solid ratio method is a classic method for detecting MTB susceptibility. It is to directly inoculate sputum, alveolar lavage fluid, hydrothorax, and ascites of tuberculosis patients and other samples onto Löwenstein-Jensena (L-J) medium, which contains different anti-tuberculosis drugs. The classic culture method of MTB is the gold standard for drug sensitivity detection. It can detect resistance levels to more than a dozen first and second-line antituberculosis drugs simultaneously, has well-established commercial reagents, and is inexpensive to test. However, it requires a long culture cycle, specific instruments, and a laboratory environment (P3 laboratory), which significantly inconveniences the treatment of tuberculosis.^{48–50}

Nitrate Reductase and Indirect Proportion Methods

The nitrate reductase test is used to determine the growth of MTBs by taking advantage of the fact that most MTBs produce an active nitrate reductase enzyme that breaks down nitrate during growth. The test operation has the advantages of simplicity, rapidity, and low cost. Studies have shown that nitrate reductase assay analyses MTB sensitivity testing to six common anti-tuberculosis drugs, which achieved high specificity and sensitivity.^{51,52} In addition, drug sensitivity testing can be performed directly on microscopy-positive clinical specimens^{53,54}, with a shorter cycle time compared to other phenotyping methods. DST by indirect proportion methods on L-J media was performed at the final drug concentrations. In brief, two appropriate dilutions of the bacilli, 10^{-2} and 10^{-4} dilutions (undiluted = 10^6 – 10^8 colony-forming units/mL) were inoculated on a drug-containing and drug-free media, to obtain countable colonies on both media. The drug resistance was taken 28–42 days.⁵⁵

Liquid Culture Method

The BACTEC MGIT 960 system is BD's fully automated, continuously monitorable, non-invasive liquid culture system that employs a fluorescent substance as a mycobacterial growth indicator and hence is known as the Mycobacteria growth indicator tube (MGIT) method. The main liquid susceptibility test method used in tuberculosis laboratories today is the modified proportional susceptibility test of BD MGIT960. MGIT960 system may test many tuberculosis drugs, such as isoniazid, rifampicin, ethambutol, streptomycin, levofloxacin, moxifloxacin, bedaquiline, and linezolid.⁵⁶ The turnaround time with the BACTEC MGIT 960 system (7.5 ± 1.8 days) was significantly shorter than the indirect proportion method (28 days or 42 days).⁵⁷ But both liquid- and solid-based media and indirect proportion methods have fair sensitivity and specificity, compared to the Xpert MTB/RIF and PCR and other molecular methods.⁵⁸

Genotype Drug Sensitivity Test

Fluorescence Quantitative Real-Time PCR

Multi-Fluorescence quantitative Real-Time PCR (MF-qRT-PCR) assay was an efficient, accurate, reliable, and easy-to-operate method for detecting drug resistance in RIF and INH. It can be used to distinguish MTC and NTM from clinical isolates. Compared with DNA sequencing, the sensitivity and specificity of MF-qRT-PCR to RIF resistance were 97.2% and 100%, and the sensitivity and specificity to INH resistance were 97.9% and 96.4%, respectively. The limitation of this study is that other anti-TB drug genes or regions could not be tested on time.⁵⁹ Another study showed genotypic predictions of the susceptibility of MTB to first-line drugs were found to be associated with phenotypic susceptibility to these drugs.⁶⁰ However, there were still many aspects to improve.

Fluorescent PCR Probe Melting Curve Method

Some studies have used the fluorescent PCR probe melting curve method to detect RIF, isoniazid, and fluoroquinolone drug sensitivity in tissue samples of tuberculosis patients, which has high sensitivity and specificity compared to phenotypic drug sensitivity testing results.⁶¹ High-resolution melt curve analysis assay showed a high level of concordance with DNA sequencing, without phenotypic drug susceptibility testing data, and could directly detect fluoroquinolone resistance in sputum samples, including direct smear-negative samples.⁶²

GeneXpert MTB/RIF and GeneXpert-Ultra

The Xpert MTB/RIF Assay System developed by Cepheid is a stand-alone assay that uses integrated microfluidics technology in combination with fully automated nucleic acid analysis to detect the *rpoB* gene fragment of MTB in samples. It is highly integrated and automated, significantly reducing the potential for contamination and human error in the assay.⁶³ Studies have shown the Xpert MTB/RIF Assay to be 95% of sensitivity and 98% of specificity.⁶⁴ GeneXpert Ultra uses fully closed nested multiplexed real-time fluorescence PCR to detect the "core" region of the RIF resistance-associated *rpoB* gene (RRDR) and two additional target genes (IS1081 and IS6110). The GeneXpert MTB/RIF Ultra system is more sensitive than the first generation because it dramatically increases the sensitivity of MTB strains from 112.6 to 15.6 CFU/mL, optimizes the interpretation of silent mutations, and reduces the false-positive rate of the RFP resistance assay at low levels of MTB in samples and mixed infections.⁶⁵

PCR Reverse Dot Hybridization Method

The main principle of the PCR reverse dot hybridization method consists of both amplification and hybridization. The unlabeled oligonucleotide probe is first immobilized on a solid-phase membrane, and then the labeled PCR product is hybridized with the solidified probe.⁶⁶ It is a gene diagnostic technique that combines probe, nucleic acid hybridization, and enzyme-linked chromatography. It seems superior in sensitivity and specificity, easy operation, fast results, and large detection throughput. Line Probe Assays (LPA), a kind of PCR-single-stranded probe reverse hybridization assays, were used to detect isoniazid and rifampicin resistance, based on the mutations in *rpoB*, *inhA*, and *katG* genes. It could turn around and detect samples within a few hours. LPAs apply biological amplification of target nucleic acids (DNA) by applying specific primers of resistance genes. Then, the product of LPA is denatured and hybridized to a specific

oligonucleotide probe immobilized on a nylon membrane, and multiple target sequences can be detected in one hybridization. For the diagnosis of rifampicin mono-resistance, LPA had a perfect agreement with a sensitivity of 90.0% and a specificity of 99.1%. Considering the advantage of LPA, WHO approved the method for the diagnosis of MDR-TB and RIF-resistant TB in smear-positive TB.^{67–69}

Compared with the phenotypic drug sensitivity assay using this technique by Guo et al,⁷⁰ the sensitivity and specificity of RIF resistance were 91.2% and 98.3%, respectively. It detects whether MTB is drug-resistant and resistant to which drugs and provides insight into the specific mutation sites or regions of the drug-resistant genes. This method is expensive in terms of reagents and instruments and has low detection of mutant loci in other gene fragments not designed on the membrane. Moreover, the interpretation of the spots on the membrane after color development is affected by subjectivity, causing false negatives of MTB.

Digital PCR

Digital PCR combines microfluidic technology with PCR to quantify individual DNA copies accurately and achieve accurate quantification of target DNA with high sensitivity and specificity. It does not require the construction of a standard curve. Zhang et al used digital PCR to detect the susceptibility of MTB to RIF, isoniazid, and streptomycin. The existence of subgroups with different sensitivities to drugs is known as hetero-resistance. Studies have shown that 8.5% and 14.2% of tuberculosis patients are hetero-resistant to INH and RIF,⁷¹ respectively. MeltPro TB/INH only detects 20% to 40% of the INH resistance heterogeneity.⁷² ddPCR can see mutant sequences as low as 0.01%.⁷³ So, digital PCR offers significant advantages in detecting and quantifying heterologous drug resistance in MTB populations.^{74,75}

Next-Generation Sequencing

Next-generation sequencing (NGS), also termed high-throughput or massively parallel sequencing, is used for pathogen detection and monitoring the hospital microbiome and its drug resistance. It is a technology genre that allows thousands to billions of DNA/RNA fragments to be tested and independently sequenced.^{76,77} Metagenomic next-generation sequencing (NGS) detected MTB with high specificity and sensitivity but overlooked the drug resistance information.^{78,79} Then, targeted next-generation sequencing (NGS) may be an early detection method for drug resistance directly from MTBC-positive specimens, with the resistance mutations by 100%. The turnaround time of the tNGS assay showed less than culture and Whole Genome Sequencing (WGS) workflows with a two-week and a similar cost.⁸⁰ The other studies also showed that tNGS could detect various antibiotic resistance genes and related mutations and has great potential in predicting MTB drug resistance.⁸¹ The performance of nanopore-targeted sequencing in bronchoalveolar lavage fluid and metagenomic next-generation sequencing for diagnosing pneumonia pathogens and both methods show higher sensitivity than conventional microbial testing. Nanopore-targeted sequencing can be considered a reliable method for diagnosing pneumonia pathogens.^{82,83} However, its application in MTB research remains limited because of less experience and the lack of sequence data in the bioinformatics analysis of MTB.⁸⁴ Thus, next-generation sequencing is a comfortable tool to predict MTB drug resistance, but the limitations make its implementation in monitoring drug resistance of clinical pathogens premature.

Gene Chips

Gene chips work by applying known nucleic acid sequences as probes immobilized on a substrate, such as glass, and then hybridizing with the DNA or RNA complementary target nucleotide sequences of the sample to be tested to obtain information about the sample. Gene chip technology can detect and analyze many sequences of a sample simultaneously due to the simultaneous immobilization of many probes on the support. Thus, it solves the shortcomings of traditional nucleic acid blot hybridization technology, such as cumbersome operation, low automation, a small number of operated sequences, and low detection efficiency. Moreover, by designing different probe arrays, the use of specific analysis methods can give the technology a variety of other applications. It was shown that gene microarrays have high sensitivity and specificity in the detection of resistance to RIF, isoniazid, fluoroquinolones, and streptomycin with the advantages of rapidity, accuracy, and high throughput but lower sensitivity in the detection of ethambutol.⁸⁵ Current gene-core technology can detect wild-type and 13 mutations at six sites of *rpoB*, wild-type at one site of *katG*, and wild-type

and mutations in the promoter region of the *inhA* gene.^{86,87} Therefore, it can be seen that the gene chip technology cannot detect all the drug-resistant genotypes of MTB, so it cannot completely replace the traditional drug sensitivity test. Still, its sensitivity and efficiency can complement the conventional drug sensitivity test.

Novel Drug Sensitivity Test Metabolism Testing Assay

In recent years, the assay based on bacterial metabolism to the drug or exterior inhibitor provides a new direction for detecting MTB drug resistance. JHU083 is a glutamine (Gln) metabolic antagonist, when administered in a TB mouse model, leading to a decrease in immunosuppressive bone marrow cell levels, an increase in effector T cell levels, and an increase in citrulline and NO production levels.⁸⁸ According to another study, about 150 MTB metabolomics of ethambutol (ETH) and ethionamide (ETO) phenotypically resistant subgroups, associated with 54 pre-XDR, 63 XDR-TB, and 33 pan-susceptible, two metabolites (meso-hydroxyheme and itaconic anhydride) were found to distinguish the pre-XDR and XDR-TB groups from the pan-susceptible group with 100% sensitivity and 100% specificity.⁸⁹ Pyrazinamide (PZA) is a first-line anti-MTB regimen, which could be converted to pyrazinoic acid (POA) by the bacterial enzyme pyrazinamidase. Then, POA is excreted into the extracellular medium, with a cycle of protonation, cell influx, and efflux established. Thus, the presence of POA can be used as a marker to assess the PZA resistance.⁹⁰ RIF, a first-line anti-tubercular agent, has been widely used to treat TB for many years. It inhibits the MTB transcription, restraining protein synthesis and mediating bacterial killing. Many studies have previously reported that the RNA polymerase gene *rpoB* could lead to MTB resistance against RIF. The RIF-mediated metabolic changes associated with pyrimidine, purine, arginine, phenylalanine, tyrosine, and tryptophan metabolic pathways could exploit potential treatment agents of MTB.^{91,92} In summary, metabolism analysis of anti-MTB drug or drug-resistant tuberculosis may provide a new orientation for detecting MTB drug resistance and effective treatment references for clinical doctors.

Cell-Free DNA Probe Assay

Cell-free DNA (cfDNA) is a cell-free nucleic acid fragment found in human blood and other body fluids.⁹³ It often exists as DNA-protein complexes in 70–200 bp short fragments or 21 kb long fragments.⁹⁴ The cfDNA was released into the blood during the decomposition of dead cells and microorganisms.⁹⁵ Alternatively, DNA breaks during apoptosis and nuclear consolidation and is released into the bloodstream.⁹⁶ The cfDNA enters the body fluids with the blood circulation in various fractions and can be detected and analyzed by different techniques such as PCR or sequencing. CfDNA can be detected in a wide range of body fluid samples and is of great significance for children and adults who have difficulty retaining respiratory specimens and for extra-pulmonary tuberculosis such as osteoarticular tuberculosis. It has been shown that the growth of bacterial populations can be estimated by measuring the skewness of the bacterial genome coverage of cfDNA and that drug resistance can be assessed by sequencing cfDNA against a defined spectrum of drug-resistance genes. The detection of MTB-cfDNA in the blood of people with latent tuberculosis infection (LTBI) suggests that the detection of MTB-cfDNA in the blood can be used to screen people with LTBI, which is a breakthrough in LTBI detection.⁹⁷ The cfDNA in the patient's body is quickly cleared by immune cells after treatment.⁹⁸ Therefore, the detected cfDNA is more likely to be newly released into the bloodstream, which reflects the current disease and facilitates monitoring of the condition. Previous studies demonstrated that the intracellular pathogens' genomic DNA fragments could be tested in human plasma and could be potential biomarkers for finding the presence of pathogenic organisms.⁹⁹ Wu et al showed that cell-free DNA of bronchoalveolar lavage fluid could diagnose MTB and resistance to the first-line anti-MTB drug (RIF and INH).¹⁰⁰ Despite significant progress, the analysis of cfDNAs remains challenging, mainly due to contamination from cells degrading after sampling and releasing more nucleic acids (NAs) into the sampled body fluids. cfDNAs have a short half-life, low concentration, and high degree of fragmentation. Therefore, standard pre-treatment of samples is essential to improve specificity and sensitivity. In addition, large sample sizes are needed for clinical validation.

CRISPR Assay

Clustered Regularly Interspaced Short Palindromic repeats (CRISPR) and CRISPR-associated proteins (Cas) are immune defense systems derived from bacteria and archaea to cope with the invasion of exogenous nucleic acids.¹⁰¹ CRISPR-Cas systems are next-generation pathogen methods that detect single nucleotide polymorphisms with high sensitivity and specificity. With its superiority, it has now been developed as a strong tool for genome editing and is widely used in various research fields such as biology, medicine, and the environment. A study was conducted to detect streptomycin-resistant strains of MTB in one hour by a rapid fluorescent assay based on the CRISPR-CAS12a system, with 100% sensitivity and specificity compared with sequencing results.¹⁰² Bai et al realized a significant distinction between fluoroquinolone-resistant and sensitive strains of MTB using the CRISPR-CAS13a system, with specificity and sensitivity of 100% and 91.4%.¹⁰³ In addition, CRISPR assay can be used to explore the mechanisms of drug resistance. Li et al developed a CRISPR-interfering chemical genetics platform to discover multiple drug resistance mechanisms in MTB, such as *ettA* mutations causing low-level resistance to various drugs, and find that MTB in the Southeast Asian lineage with loss-of-function *whiB7* allele is highly sensitive to macrolides, which provides theoretical support for clarithromycin treatment of this sub-lineage.¹⁰⁴ Mei-Yi Yan group used CRISPR assay to show the antitubercular drug bedaquiline (BDQ) mechanisms and the possibility of optimizing tuberculosis therapy.¹⁰⁵ Additionally, the CRISPR assay could successfully prognosis the coordination action of cyclomarin A with isoniazid or RIF.¹⁰⁶ CRISPRi assay could provide a new thinking for anti-MTB drug development. Another group found an application of the Next-Generation CRISPR for Finding Low Abundance Sequences by Hybridization (FLASH), a technique for directly identifying MTB drug resistance from sputum samples.¹⁰⁷ CRISPR assay requires a small sample volume and short operational time compared to the Xpert method.

Spectral Analysis Technique

Spectral analysis is a statistical method used to analyze a time series dataset. It identifies statistically meaningful frequencies in a time series to see whether they contain periodic or cyclical components. Spectral analysis is a powerful tool with many applications in numerous fields, such as environmental monitoring and traditional Chinese medicine. Matrix-assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF) is a rapidly developing and new type of soft ionized biomass spectrum, which was first used for MTB identification in 2004. Compared with traditional identification methods, MALDI-TOF MS simplifies the identification process, reduces the identification time, and has the advantages of accuracy, speed, sensitivity, and high throughput. MALDI-TOF MS can also identify different drug resistance types of MTB with other mutations and be a rapid and effective method to scan the drug resistance of MTB. It shows acceptable clinical application values in patients with relapsed tuberculosis.^{108–110}

Raman scattering is an inelastic scattering phenomenon triggered by light irradiation on the surface of a substance. The enhancement effect associated with rough metal surfaces such as gold, silver, and copper was known as the surface-enhanced Raman scattering effect, and the resulting spectra were known as surface-enhanced Raman spectroscopy (SERS). Nucleic acids, proteins, lipids, and carbohydrates in bacteria exhibit different spectral peaks on Raman spectra due to their different molecular vibration frequencies, which provides a basis for applying SERS detection technology in the field of bacteria.¹¹¹ With no need for labeling, non-invasive, easy operation, and short detection time, SERS has recently become a research hotspot in microbiology. Tang et al¹¹² used two deep learning methods (CNN and LSTM) to detect 117 staphylococcal strains of nine staphylococcal genera with SERS, and the accuracies reached 98.21% and 94.33%. SERS technology also shows excellent potential in the detection of MTB. Perumal et al¹¹³ used SERS technology to detect Mycobacterium acid for rapid and effective detection of tuberculosis. Dastgir et al used the SERS technique combined with PCR to obtain spectra from the PCR products of MTB to differentiate RFP resistance using principal component analysis and least discriminant analysis.¹¹³ Raman spectroscopy was combined with deep learning for the early diagnosis and to distinguish accurately in the three situations: 1) smear-positive and smear-negative sputum samples to MTB; 2) pulmonary and extra-pulmonary origin of MTB strains; and 3) different antibiotic resistance groups to RIF and INH for pulmonary MTB strains.¹¹⁴ The classification of MTB with different resistance to RFP and INH was predicted by applying SERS technology; its accuracy was 99.59%, precision was 99.64%, recall was 99.61%,

F1 score was 99.67%, and the accuracy of 5-fold cross-validation was 99.59%. Raman signals are closely related to the vibrations of multiple components and groups of bacteria. The collected Raman spectra are highly similar, and the data are complex.¹¹⁴ Thus, efficient data processing analysis methods are required to process and analyze Raman spectra. Deep learning is to extract learning features from large-scale raw data and build predictive models to make predictions.¹¹⁵ Many deep learning algorithms have emerged, including convolutional neural networks, fully connected neural networks, residual neural networks, etc.¹¹⁶ Deep learning performance is evaluated by concentrating on the data's local features and obtaining the data's global features. Classifying and identifying the data is more efficient than the traditional multivariate statistical analysis algorithms.

Limitations, Challenges, and Perspectives

From traditional testing techniques to emerging methods, all have demonstrated distinct advantages in detecting MTB drug resistance. Phenotypic drug sensitivity testing methods such as tuberculosis culture are accurate and inexpensive. They can test multiple first and second-line anti-tuberculosis drugs simultaneously. However, the operation is complicated, and the culture cycle is long. Many live bacterial detectors have high technical requirements for experimental sites, personnel, and laboratory biosafety risks. Traditional molecular detection techniques are mature and sensitive, with short detection time and low biosafety risk. However, there are areas for improvement in the following aspects: 1) challenging and complex to detect heterogeneous resistance; 2) synonymous and silent mutations that do not affect the phenotypes; 3) all phenotypically resistant strains cannot be detected. Traditional molecular diagnostic techniques are widely used in the clinic as complementary to phenotypic drug-sensitivity testing techniques.

Compared with classical detection methods, emerging detection technologies show great potential for application and need to be more mature and sophisticated in many aspects. The emerging digital PCR does not rely on standard curves and has higher sensitivity. No pre-amplification DNA treatment is required.¹¹⁷ However, amplifying the RNA gene may contain an incompletely reverse-transcribed template, causing off-target phenomena; false positives exist in a few reactions units.^{118,119} NGS enables a more comprehensive analysis of genomic drug resistance-related variants. Meanwhile, NGS can be used for epidemiological surveillance and exploration of drug resistance mechanisms. At present, genome sequencing technology needs a unified testing procedure. Laboratory operations are prone to exogenous nucleic acid contamination, producing false positive results. Nucleic acid molecules from non-pathogenic pathogens with low sequence counts or residual nucleic acid molecules from dead pathogens can be detected at the same time. These can easily interfere with proper clinical diagnosis and treatment. So, there are still challenges in establishing a unified quality control process and interpretation guidelines, addressing interference from background organisms, and reducing the cost of testing.

The majority of TB deaths occur in developing countries, and the 30 high-burden TB countries in the world are also mainly developing countries. Studies have shown that the economic burden on multidrug-resistant TB patients has been high and increasing in recent years. SERS technology is inexpensive, simple, and easy to disseminate, significantly reducing the economic pressure on developing countries for outbreak prevention and control and treating individuals with the disease. However, SERS requires a large number of samples for validation. In addition, Raman spectroscopy relies on complex algorithms. There are still significant challenges in the improvement and preparation of substrates to improve the sensitivity of spectra and the quality of spectral data, the establishment of a database of standard maps, the establishment of mature standard operating procedures, the promotion of mutual recognition of results, and the interpretation of consensus, among others.

Conclusion

Currently, there are many techniques for drug sensitivity testing of MTB, and all of them have shown their unique superiority. Under the pressure of emerging technologies, the classical drug-sensitivity testing techniques are still irreplaceable. While emerging technologies offer excellent application prospects, they also face many challenges. Combined drug resistance assays should be performed on a case-by-case basis to provide the clinic with more timely and accurate information, considering the methods' sensitivity and specificity, the reagents' cost, the reporting period,

and the limitations. Researchers should actively promote the research of emerging technologies and continuously improve them to provide more choices for clinical applications.

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