Tuberculosis

Practical guide for clinicians, nurses, laboratory technicians and medical auxiliaries

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Tuberculosis drug information sheets
Amikacin (Am)
Amoxicillin/clavulanic acid ratio 4:1 (Amx/Clv)
Bedaquiline (Bdq)
Clofazimine (Cfz)
Cycloserine (Cs) or terizidone (Trd)
Delamanid (Dlm)
Ethambutol (E)
Ethionamide (Eto) or prothionamide (Pto)
Imipenem/cilastatin (Ipm/Cln)
Isoniazid - Standard dose (H)
Levofloxacin (Lfx)
Linezolid (Lzd)
Meropenem (Mpm)
Moxifloxacin (Mfx)
Para-aminosalicylate sodium (PAS)
Pretomanid (Pa)
Pyrazinamide (Z)
Rifabutin (Rfb)
Rifampicin (R)
Rifapentine (P)
Streptomycin (S)

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Patients on drug-resistant TB treatment
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Abdominal pain
Diarrhoea
Epigastric pain
Hepatotoxicity
Metallic taste
**Nausea and vomiting**
- Neurotoxicity

**Depression**

**Headache**

**Optic neuritis**

**Ototoxicity**

**Peripheral neuropathy**

**Psychosis**

**Seizures**
- Endocrine disorders

**Gynecomastia**

**Hypothyroidism**
- Dermatological disorders

**Alopecia**

**Fungal infection**

**Photosensitivity**

**Skin reactions**
- Musculoskeletal disorders

**Arthralgias**

**Tendinitis/tendon rupture**
- Miscellaneous

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**Haematologic disorders**

**Lactic acidosis**

**Nephrotoxicity**

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Introduction

Tuberculosis is an infectious disease caused by *Mycobacterium tuberculosis*. Tuberculosis typically attacks the lungs, but can also affect other parts of the body. The disease has become rare in high income countries, but is still a major public health problem in low- and middle-income countries.

It is estimated that between the years 2000 and 2010, eight to nine million new cases emerged each year. Approximately 1.5 million people die from the disease each year. In adults, tuberculosis is the second leading cause of death due to an infectious disease (after AIDS), with 95% of deaths occurring in low-income countries. Tuberculosis is a major problem of children in poor countries where it kills over 100,000 children each year.
The treatment of tuberculosis remains a constraint for patients and a heavy burden for the healthcare system. Drug-susceptible tuberculosis requires at least six months of therapy under close supervision. A treatment for multidrug-resistant tuberculosis requires nearly two years of treatment with poorly tolerated and less effective drugs. In most places the diagnosis still relies mainly on direct microscopy that is unable to detect a large proportion of patients. The BCG vaccine, developed almost a century ago, confers only partial protection.

After 40 years of minimal progress in the tools to fight tuberculosis there are some reasons for hope. A few new drugs are reaching the final phase of development; a new molecular test that can be decentralized to some extent and allows the rapid diagnosis of tuberculosis and of resistance to rifampicin has been introduced. Though this is undeniable progress, much will be needed to bring the new tools and drugs to the patients in need. Furthermore, a true “point of care” diagnostic test still does not exist and little progress has been made in research for a more effective vaccine.

Case management of patients does not necessarily have to involve a major, vertical programme. It should be incorporated into the framework of other medical activities in order to offer comprehensive and integrated treatment even if the number of patients being treated is relatively small.

This guide has been developed jointly by Médecins Sans Frontières and Partners In Health. It aims at providing useful information to the clinicians and health staff for the comprehensive management of tuberculosis. Forms of susceptible and resistant tuberculosis, tuberculosis in children, and HIV co-infection are all fully addressed.

As treatment protocols are constantly changing, medical staff are encouraged to check this website for updates.

**Abbreviations and acronyms**
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACH</td>
<td>Air change per hour</td>
</tr>
<tr>
<td>AFB</td>
<td>Acid-fast bacilli</td>
</tr>
<tr>
<td>AIDS</td>
<td>Acquired immunodeficiency syndrome</td>
</tr>
<tr>
<td>Amk</td>
<td>Amikacin</td>
</tr>
<tr>
<td>Amx/Clv</td>
<td>Amoxicillin/clavulanic acid</td>
</tr>
<tr>
<td>ARI</td>
<td>Annual risk of infection</td>
</tr>
<tr>
<td>ART</td>
<td>Antiretroviral therapy</td>
</tr>
<tr>
<td>ARV</td>
<td>Antiretroviral</td>
</tr>
<tr>
<td>BCG</td>
<td>Bacillus Calmette-Guérin</td>
</tr>
<tr>
<td>Bdq</td>
<td>Bedaquiline</td>
</tr>
<tr>
<td>CDC</td>
<td>United States Centers for Disease Control and Prevention</td>
</tr>
<tr>
<td>Cfz</td>
<td>Clofazimine</td>
</tr>
<tr>
<td>Cm</td>
<td>Capreomycin</td>
</tr>
<tr>
<td>CMX</td>
<td>Cotrimoxazole</td>
</tr>
<tr>
<td>CPC</td>
<td>Cetylpyridinium chloride</td>
</tr>
<tr>
<td>CPT</td>
<td>Cotrimoxazole preventive therapy</td>
</tr>
<tr>
<td>Cs</td>
<td>Cycloserine</td>
</tr>
<tr>
<td>CXR</td>
<td>Chest X-ray</td>
</tr>
<tr>
<td>DOT</td>
<td>Directly observed therapy</td>
</tr>
<tr>
<td>DR</td>
<td>Drug resistance</td>
</tr>
<tr>
<td>DR-TB</td>
<td>Drug-resistant tuberculosis</td>
</tr>
<tr>
<td>DST</td>
<td>Drug susceptibility test(ing)</td>
</tr>
<tr>
<td>E</td>
<td>Ethambutol</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiogram</td>
</tr>
<tr>
<td>EPTB</td>
<td>Extrapulmonary tuberculosis</td>
</tr>
<tr>
<td>Eto</td>
<td>Ethionamide</td>
</tr>
<tr>
<td>FDC</td>
<td>Fixed-dose combination</td>
</tr>
<tr>
<td>Acronym</td>
<td>Definition</td>
</tr>
<tr>
<td>---------</td>
<td>------------</td>
</tr>
<tr>
<td>FNAC</td>
<td>Fine needle aspiration cytology</td>
</tr>
<tr>
<td>FQ</td>
<td>Fluoroquinolone</td>
</tr>
<tr>
<td>GFR</td>
<td>Glomerular filtration rate</td>
</tr>
<tr>
<td>H</td>
<td>Isoniazid</td>
</tr>
<tr>
<td>HCW</td>
<td>Health care worker</td>
</tr>
<tr>
<td>HIV</td>
<td>Human immunodeficiency virus</td>
</tr>
<tr>
<td>HPF</td>
<td>High-power field</td>
</tr>
<tr>
<td>IC</td>
<td>Infection control</td>
</tr>
<tr>
<td>IM</td>
<td>Intramuscular</td>
</tr>
<tr>
<td>Imp/Cin</td>
<td>Imipenem/cilastatin</td>
</tr>
<tr>
<td>IPT</td>
<td>Isoniazid preventive therapy</td>
</tr>
<tr>
<td>IRIS</td>
<td>Immune reconstitution inflammatory syndrome</td>
</tr>
<tr>
<td>IUATLD</td>
<td>International Union against Tuberculosis and Lung Disease</td>
</tr>
<tr>
<td>Km</td>
<td>Kanamycin</td>
</tr>
<tr>
<td>LFT</td>
<td>Liver function test</td>
</tr>
<tr>
<td>Lfx</td>
<td>Levofloxacin</td>
</tr>
<tr>
<td>LPA</td>
<td>Line probe assay</td>
</tr>
<tr>
<td>Lzd</td>
<td>Linezolid</td>
</tr>
<tr>
<td>MDR</td>
<td>Multidrug resistance</td>
</tr>
<tr>
<td>MDR-TB</td>
<td>Multidrug-resistant tuberculosis</td>
</tr>
<tr>
<td>Mfx</td>
<td>Moxifloxacin</td>
</tr>
<tr>
<td>MGIT</td>
<td>Mycobacteria growth indicator tube</td>
</tr>
<tr>
<td>MODS</td>
<td>Microscopic observation of drug susceptibility</td>
</tr>
<tr>
<td>Mpm</td>
<td>Meropenem</td>
</tr>
<tr>
<td>NNRTI</td>
<td>Non-nucleoside reverse transcriptase inhibitor</td>
</tr>
<tr>
<td>NRTI</td>
<td>Nucleoside reverse transcriptase inhibitor</td>
</tr>
</tbody>
</table>
Chapter 1: Introduction and epidemiology

1.1 Characteristics of Mycobacterium tuberculosis bacillus

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NTM</td>
<td>Non tuberculous mycobacteria</td>
</tr>
<tr>
<td>Ofx</td>
<td>Ofloxacin</td>
</tr>
<tr>
<td>PAS</td>
<td>Para-aminosalicylic acid</td>
</tr>
<tr>
<td>PCP</td>
<td>Pneumocystosis</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PI</td>
<td>Protease inhibitor</td>
</tr>
<tr>
<td>PO</td>
<td>Orally (per os)</td>
</tr>
<tr>
<td>Pto</td>
<td>Prothionamide</td>
</tr>
<tr>
<td>R</td>
<td>Rifampicin</td>
</tr>
<tr>
<td>Rfb</td>
<td>Rifabutin</td>
</tr>
<tr>
<td>RR</td>
<td>Rifampicin resistance</td>
</tr>
<tr>
<td>S</td>
<td>Streptomycin</td>
</tr>
<tr>
<td>TAT</td>
<td>Turn around time</td>
</tr>
<tr>
<td>TB</td>
<td>Tuberculosis</td>
</tr>
<tr>
<td>Thz</td>
<td>Thioacetazone</td>
</tr>
<tr>
<td>TLA</td>
<td>Thin-layer agar</td>
</tr>
<tr>
<td>TSH</td>
<td>Thyroid-stimulating hormone</td>
</tr>
<tr>
<td>TST</td>
<td>Tuberculin skin test</td>
</tr>
<tr>
<td>UVGI</td>
<td>Ultraviolet germicidal irradiation</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
<tr>
<td>XDR</td>
<td>Extensive drug resistance</td>
</tr>
<tr>
<td>XDR-TB</td>
<td>Extensively drug-resistant tuberculosis</td>
</tr>
<tr>
<td>Z</td>
<td>Pyrazinamide</td>
</tr>
</tbody>
</table>
1.1 Characteristics of Mycobacterium tuberculosis bacillus

*Mycobacterium tuberculosis*, along with *M. bovis*, *M. africanum*, *M. microti* and others, make up the *Mycobacterium tuberculosis* complex, a group of bacteria that cause clinical tuberculosis (TB) in humans. Most TB cases are caused by *M. tuberculosis*. Cases due to other species are far less prevalent.

*M. tuberculosis* is a small, rod-shaped, strictly aerobic, acid-fast bacillus. Like other mycobacteria, it is slow growing, resulting in more gradual development of disease when compared with other bacterial infections.

**Notas**
(a) Acid-fast bacilli are bacilli, which once stained, resist discoloration by acid and alcohol.

1.2 Transmission

*M. tuberculosis* is transmitted from human-to-human and spread is mainly airborne. The source of infection is usually a patient with pulmonary or laryngeal TB. During coughing, speaking, or sneezing, the patient produces tiny infectious droplets. These particles, called droplet nuclei, are about 1 to 5 microns in diameter. Depending on the environment, they can remain suspended in the air for several hours.

Transmission may occur when these infectious droplets are inhaled. UV light (sunshine or artificial sources) and ventilation reduce the probability of transmission (Chapter 14).

Other modes of transmission are far less common. Inoculation of cutaneous or mucous membranes rarely occurs, although such cases have been observed in laboratory personnel. Congenital infection (by transplacental transmission or via aspiration or swallowing of infected amniotic fluid at birth) has been reported, but is very rare. Transmission through breast milk does not occur.

The infectiousness of a patient is associated with the quantity of bacilli contained in their sputum. Patients with smear-positive sputum on microscopy are by far the most infectious. Those with smear-negative/culture-positive results are less infectious, but still contribute to TB transmission due to more frequent delays in diagnosis.
Patients infected with *M. tuberculosis*, but who have not developed active TB (latent tuberculosis infection), are not infectious. Patients with extrapulmonary TB (EPTB) are only infectious in exceptional circumstances.

Children are generally much less infectious than adults. This may be due to weaker cough mechanics, less sputum production and lower bacillary load.

Not everyone who is exposed to an infectious TB patient becomes infected with *M. tuberculosis*. The probability that TB will be transmitted depends on several factors:

- **Infectiousness of the source** (the most important factor)
  - Bacteriological status: smear-positive patients are the most infectious.
  - Virulence of the bacilli: some strains are highly transmissible (and/or more likely to cause active TB).

- **Environment where the exposure occurred**
  - Outdoor environments or those with good ventilation and sunlight are less likely to lead to transmission. Small rooms or rooms with no ventilation are conditions most likely to lead to transmission.
  - The proximity of the person to the patient is also important (e.g. the risk is higher if the person sleeps next to the patient than if they sleep 20 metres away from the patient).

- **Duration of exposure**
  - People in close and prolonged contact with TB patients are at highest risk of becoming infected with *M. tuberculosis*. They may be family members, roommates, friends, co-workers or other people who spend several hours a day with the infectious patient.

The best way to stop transmission is to start effective TB treatment as soon as possible. It is estimated that a person with untreated smear-positive TB transmits the bacillus to 10 to 20 people a year (with variations according to living conditions and environment).

### 1.3 Evolution of tuberculosis infection and disease in humans

When a person inhales infectious droplets containing *M. tuberculosis*, most of the larger droplets become lodged in the upper respiratory tract (nose and throat) where infection is unlikely to develop. However, smaller droplet nuclei may reach the small air sacs of the lung (the alveoli) where infection can occur.

#### 1.3.1 Primary infection and latent tuberculosis infection

After transmission, *M. tuberculosis* multiplies slowly, in most cases in the terminal alveoli of the lungs (primary focus) and in the lymph nodes of corresponding drainage areas: this is the primary infection. The primary focus and related hilar lymphadenopathy form the primary complex.

In one to two months, due to the action of lymphocytes and macrophages (cellular immunity), the primary focus is contained and encapsulated, with a central zone of parenchymal necrosis (caseous lesions). It is not usually detectable on chest x-ray, unless it calcifies or grows substantially. Primary infection is usually asymptomatic. In most cases (90 to 95% of non-HIV infected patients), the pulmonary lesions gradually heal.

During the primary infection, specific immunity develops and a positive skin reaction to tuberculin is observed[^1]. This immune response may persist without clinical signs of TB. The patient is infected by *M. tuberculosis*, but does not develop the disease. This is referred to as latent tuberculosis infection (LTBI).

In 5 to 10% of infected people, primary infection and/or LTBI progresses to active TB over their lifetime. For HIV co-infected patients, this risk is much higher.

#### 1.3.2 Active tuberculosis

Before immunity is established, bacilli from the primary infectious focus or from a near-by lymph node can be transported and disseminated throughout the body via the lymph system or the bloodstream. Secondary foci can develop this way, particularly in the lungs, lymph nodes, serous membranes, meninges, bones and kidneys. As soon as an immune response is mounted, most of these foci resolve spontaneously. However, some bacilli may remain dormant in the secondary foci for months and sometimes years. Different factors can reduce the immune response (e.g. HIV infection) and lead to reactivation of the bacilli and their multiplication in one or more of these foci. This reactivation or progression of the primary or secondary foci results in active TB. An active TB lesion contains actively, slowly or sporadically multiplying bacilli as well as dormant bacilli. While active TB may occur months or years following primary infection, half of TB cases appear in the year following infection.

1.3.3 Risk factors for developing active tuberculosis

Certain factors increase the risk of developing active TB within the first two years of being infected. These factors include any factor that results in a weakened immune system, damaged lungs and the intensity and duration of exposure.

Host immune response factors:
- HIV infection
- Children under 5 years
- Malnutrition
- Persons over 60 years
- Diabetes mellitus
- Other risk factors: prolonged corticosteroid therapy (g. prednisolone) and other immunosuppressive therapies, severe kidney disease, alcoholism, substance abuse, certain types of cancer (e.g. leukaemia, Hodgkin’s lymphoma, cancer of the head and neck); pregnancy

Conditions that damage the lung:
- Tobacco smoking
- Silicosis
- Chronic obstructive pulmonary disease (COPD)

Intensity of exposure (high number of inhaled bacilli):
- Highly infectious source
- Poorly ventilated environment
- Proximity with infectious source, including residents and employees of institutions such as prisons, boarding schools and residential care facilities
- Long duration of exposure

Referencias


1.4 Prognosis

Without treatment, TB is a severe and potentially fatal disease. After 5 years without treatment, the outcome of smear-positive pulmonary TB (PTB) in non-HIV-infected patients is as follows\(^1\):

- 50 to 60% die (case fatality ratio (CFR) for untreated TB);
- 20 to 25% are cured (spontaneous cure);
- 20 to 25% continue to have symptoms.

In non-HIV infected patients, the CFR is estimated at 3%\(^2\). Untreated TB in HIV-infected patients (not on effective antiretroviral therapy) is almost always fatal. Even on antiretroviral therapy, the CFR is higher than in non-HIV infected patients\(^3\,\!^4\).

Risk factors for poor outcomes of TB treatment (death and relapse) include co-morbidities (e.g. HIV infection, diabetes, COPD), cavities on chest x-ray, high bacillary load and resistance to TB drugs.

### Referencias


1.5 Factors modifying tuberculosis epidemiology

Five major factors influence TB epidemiology: (1) socioeconomic conditions, (2) TB treatment, (3) HIV infection, (4) diabetes and (5) BCG vaccination.

### 1.5.1 Socioeconomic conditions

The principal factors leading to a reduction in TB cases are improved social and housing conditions. Most cases occur in low-income countries. In industrialised countries, TB generally affects the most disadvantaged social groups.

### 1.5.2 Tuberculosis treatment

Diagnosing and initiating effective treatment in a patient early during their TB disease before they can infect multiple people is considered the most effective preventive measure against TB. Once an effective TB treatment is started, there is a rapid reduction in transmission\(^1\,\!^2\).

Since the introduction of TB treatment, the risk of TB infection decreased by approximately 10% per year in industrialised countries\(^8\). This trend was observed in countries with a BCG vaccination programme as well as in those without one. Detection programmes, diagnosis and treatment of TB contributed to this reduction in the risk of TB infection.
1.5.3 HIV infection

Immunodeficiency induced by HIV infection is a major risk factor for progression to active TB and has a considerable impact on the epidemiology of TB. While the lifetime risk of developing active TB in the general population is 5 to 10% after infection with *M. tuberculosis*, this risk is approximately 10% per year in patients co-infected with HIV and *M. tuberculosis*. Approximately 8% of incident TB cases in the world are among HIV-infected patients (highest in the WHO African Region, more than 50% in parts of southern Africa)[4].

1.5.4 Diabetes

The risk of TB among people with diabetes is higher than among those without diabetes. It is estimated that diabetes contributes to 15% of TB cases worldwide[6]. Diabetes is also associated with poor absorption of TB drugs and therefore higher rates of drug resistant tuberculosis (DR-TB).

1.5.5 BCG vaccination

**Effectiveness of BCG at the individual level**

BCG vaccination, if given at birth, is highly effective against the severe forms of TB (miliary and meningitis) in children[6].

**Epidemiological impact of vaccination**

Despite some protection from the BCG vaccination, the impact of BCG vaccination on TB transmission and the TB epidemic is considered negligible[7].

1.5.6 Other factors

Other modifying factors include infection control measures (Chapter 14) and treatment of LTBI (Chapter 16). The degree to which in a given context the TB epidemiology is affected by these measures is not known.

**Referencias**


**1.6 Epidemiological indicators**
When a National TB Programme (NTP) functions well, indicators can be obtained from the local authorities and NTP. The WHO tuberculosis country profiles also provide an estimation of TB indicators by individual country.

**Box 1.1 - Most common indicators**

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Description</th>
<th>Numerator</th>
<th>Denominator</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Annual incidence rate of TB cases</strong>&lt;sup&gt;(a)&lt;/sup&gt;</td>
<td>The number of new TB cases (all forms) that occur in a population over one year</td>
<td>number of new TB cases</td>
<td>population at the start of the year</td>
</tr>
<tr>
<td><strong>Annual incidence rate of smear-positive PTB cases</strong>&lt;sup&gt;(a)&lt;/sup&gt;</td>
<td>The number of new smear-positive PTB cases that occur in a population over one year</td>
<td>number of new smear-positive PTB cases</td>
<td>population at the start of the year</td>
</tr>
<tr>
<td><strong>Prevalence of smear-positive PTB cases</strong></td>
<td>The number of smear-positive PTB cases over a given period of time, usually one year</td>
<td>number of smear-positive PTB cases</td>
<td>population at the start of the period of time</td>
</tr>
<tr>
<td><strong>Proportion of multidrug- and rifampicin-resistant TB cases among TB cases</strong></td>
<td>The number of multidrug- and rifampicin-resistant TB cases over a given period of time</td>
<td>number of multidrug- and rifampicin-resistant TB cases</td>
<td>denominator(s): Total number of TB cases, Number of new TB cases, Number of previously treated TB cases</td>
</tr>
<tr>
<td><strong>Proportion of extensively drug-resistant TB cases among TB cases</strong></td>
<td>The number of extensively drug-resistant cases over a given period of time</td>
<td>number of extensively drug-resistant cases</td>
<td>as for multidrug- and rifampicin-resistant TB cases</td>
</tr>
<tr>
<td><strong>Proportion of HIV-infected patients among new TB cases</strong></td>
<td>The number of HIV-infected patients over a given period of time</td>
<td>number of HIV-infected patients</td>
<td>number of new TB cases</td>
</tr>
</tbody>
</table>

(a) The rate is expressed as the number of new TB cases (or new smear-positive PTB cases) per 100,000 population.

(b) Prevalence is expressed as the number of smear-positive PTB cases per 100,000 population. It includes new and pre-existing cases. Prevalence represents approximately double the incidence rate.

(c) Proportion is expressed in %.

---

**1.7 Global burden of tuberculosis**

**1.7.1 Latent tuberculosis infection**

The global prevalence of LTBI is unknown due to difficulties in diagnosis. However, WHO estimates that one-quarter of the world population has LTBI<sup>(l)</sup>.
1.7.2 Active tuberculosis

Globally, active TB remains a leading cause of death from infectious disease. WHO estimates that each year there are approximately 10 million incident cases of TB and 1.5 million deaths due to TB, including 1.3 million among HIV-negative individuals and 214,000 among HIV-infected individuals[2]. Children under 15 years account for 11% of all estimated TB cases[6]. However, TB cases in children are frequently undiagnosed and unreported.

While the absolute number of global TB cases is stable, there are large individual country and regional differences in incidence and prevalence.

Most TB cases are in Southeast Asia (43%), Africa (25%) and the Western Pacific (18%), with lower percentages in the Eastern Mediterranean, the Americas and Europe[2].

1.7.3 Drug-resistant tuberculosis

Drug-resistant TB (DR-TB) is a growing worldwide problem, and no region is spared.

WHO estimates that annually worldwide there are[3]:
• More than one million rifampicin-susceptible and isoniazid-resistant TB (Hr-TB) cases (11% of all incident TB cases).
• 3.3% of new cases and 18% of previously treated cases, with multidrug-resistant TB (MDR-TB) and rifampicin-resistant TB (RR-TB)b representing 465,000 cases and 182,000 deaths.

In Eastern Europe and Central Asia, TB incidence is lower than in Southeast Asia and Africa, but up to 30% of new and 65% of retreatment cases exhibit rifampicin-resistance.

In China and India, there is a low proportion of rifampicin-resistant cases among all TB cases. However, because of their large populations, these two countries represent 41% of global MDR/RR-TB cases.

Resources for detecting drug resistance are limited in many parts of Africa. However, available data suggest that the MDR-TB burden is significant, especially in the south.

The prevalence of extensively drug-resistant TB (XDR-TB)c, according to the new WHO definition, is currently unknown.

Notas
(a) Multidrug-resistant: resistance to at least rifampicin and isoniazid.
(b) Rifampicin-resistant: resistance to rifampicin, with or without resistance to other TB drugs.
(c) Extensively drug-resistant: rifampicin-resistance with resistance to any fluoroquinolone, and at least either bedaquiline or linezolid.

Referencias

Chapter 2: Clinical presentation
2.1 Pulmonary tuberculosis

Prolonged cough (more than 2 weeks), with or without sputum production, is a common symptom in patients with pulmonary tuberculosis (PTB).

Other frequent, less specific, signs and symptoms include weight loss, anorexia, fatigue, haemoptysis (blood in sputum), shortness of breath, chest pain, moderate fever and night sweats.

Signs and symptoms may vary between individuals and generally evolve in a chronic, insidious manner. History-taking is therefore of the utmost importance.

Advanced forms and complications are common:

- Respiratory insufficiency due to extensive lesions and destroyed lungs;
- Massive haemoptysis due to large cavities with hyper-vascularisation and erosion of vessels;
- Pneumothorax due to the rupture of a cavity in the pleural space.

In endemic areas, the diagnosis of PTB should be considered in any patient consulting for respiratory symptoms lasting more than 2 weeks.

Table 2.1 provides a differential diagnosis of PTB for non-HIV infected patients.

Table 2.1 - Differential diagnosis for PTB (non-HIV infected patients)
Starting from a pulmonary localisation (primary infection), *M. tuberculosis* can spread to other organs during a silent phase, usually soon after primary infection (Chapter 1). Active TB can develop in many other parts of the body, particularly in lymph nodes, meninges, bones and joints, kidneys, genital organs and the abdominal cavity.

Extrapulmonary tuberculosis (EPTB) can develop at any age. Due to relative immunodeficiency, young children, HIV-infected and malnourished patients are more at risk of developing EPTB.

Approximately 15% of global TB cases are classified as EPTB, although this figure varies according to the local epidemiology. A patient with EPTB may also have pulmonary involvement, which should be searched for whenever EPTB is diagnosed or suspected.

Table 2.3 at the end of this chapter summarises the characteristics of EPTB.

### 2.2.1 Lymph node tuberculosis

Lymph node TB is common, particularly in certain areas of Africa and Asia, and especially in children and HIV-infected patients.
The presentation of lymph node TB is a non-inflammatory adenopathy. Nodes are cold and painless, multiple (usually bilateral) or single, evolving in a chronic mode towards softening and fistulisation. Cervical localisation is most frequent. Axillary and mediastinal localisations are also common. Other sites may be involved. Diagnosis may be clinical, but whenever possible, fine needle aspiration should be performed (Chapter 3 and Appendix 7). Adenopathy usually disappears within 3 months of treatment initiation. Paradoxical reactions may occur at the beginning of treatment (appearance of abscesses, fistulas or other lymph nodes), but a change in the treatment is not required. Differential diagnoses include malignancies (lymphoma, leukaemia, ear/nose/throat tumours, Kaposi sarcoma) and other infections (bacterial, viral, non-tuberculosis mycobacteria, toxoplasmosis, HIV infection, syphilis, African trypanosomiasis).

2.2.2 Tuberculous meningitis

TB meningitis is a serious form of TB that affects the meninges. It is most common in children under 2 years and in HIV-infected patients. It is a medical emergency. Any delay in diagnosis or treatment will result in irreversible neurological sequelae or death. TB meningitis typically has a subacute insidious course over days or weeks. Symptoms include headaches, irritability, fever, vomiting and altered mental status, which worsen if treatment is delayed. The meningeal syndrome (stiff neck, hypotonia in infants, photophobia and headache) is present in most cases. Third cranial nerve palsy (oculomotor paralysis) may occur. Diagnosis is assisted by examination of cerebrospinal fluid (Chapter 3).

The main differential diagnoses are other forms of meningitis.

2.2.3 Tuberculosis of bones and joints

Up to 40% of patients with TB of bones and joints have concurrent PTB.

**Spinal TB (spondyloisitis or Pott's disease)**

TB can affect vertebrae and intervertebral disks, causing destruction and deformation of the spine. The thoracic spine is the most frequently affected. Localised back pain may precede by several months the appearance of the first radiological anomalies (destruction of an intervertebral disk). A spinal prominence (gibbus) due to destruction and deformity of the vertebral bodies may be felt. Paravertebral cold abscesses and/or neurological complications can develop. A missed diagnosis of thoracic or cervical spinal TB can result in paralysis.

**Arthritis**

TB most frequently causes a chronic mono-arthritis, starting insidiously, with little or no pain and accompanied by joint destruction. The joints most often affected are the hips, knees, elbows and wrists.

**Osteitis**

Osteitis is the least common presentation of TB of the bones. It may be a primary osteitis or an osteitis secondary to TB arthritis. Typically, long bones are affected. Cold abscesses may occasionally occur. Like arthritis, it is distinguished from common bacterial infections by the presence of mild symptoms, despite bone and joint destruction.

The diagnosis is based on the patient's history, clinical examination and radiography, as biopsy and culture are difficult to perform in many settings. A history of prolonged and insidious osteitis or arthritis associated with a deterioration of the general physical condition favours TB aetiology, as opposed to bacterial osteomyelitis or brucellosis. The patient may have a history of non-response to antibiotics.

2.2.4 Urogenital tuberculosis

Renal involvement is frequent and may be asymptomatic for a long period, with a slow development of signs and symptoms: painful urination (dysuria), urinary urgency and frequency (pollakiuria), including during the night (nocturia); back/abdominal pain; tenderness/swelling of the testes or epididymitis or haematuria. General physical condition is generally preserved. Diagnosis is suspected in the presence of pyuria (white blood cells in the urine) and micro- or macroscopic haematuria, which does not respond to antibiotics. Examination of the urine helps with diagnosis (Chapter 3).

In men, genital localisation is secondary to renal involvement. Signs are most often epididymitis with scrotal pain.
In women, genital tract infection can also occur by a hematogenous path. Signs are non-specific: pelvic pain, leucorrhoea and abnormal vaginal bleeding. Infertility is often the reason leading women to seek medical attention. Extension may be found in the peritoneum, with resulting ascites.

### 2.2.5 Abdominal tuberculosis

Abdominal TB commonly presents as ascites resulting from the peritoneal localisation of the infection. Abdominal mass (often in the right lower quadrant), pain and diarrhoea may be present. The frequency of chronic ascites in tropical regions, with its many different causes, makes this relatively uncommon form of TB difficult to diagnose. Diagnosis is assisted by examination of the ascitic fluid via paracentesis. Constitutional symptoms (fever, night sweats, malaise and weight loss) may be present. Accumulation of ascites may mask weight loss.

### 2.2.6 Tuberculous pleural effusion

Tuberculous pleural effusion is one of the most common forms of EPTB. It is often asymptomatic, especially if less than 300 ml. Shortness of breath and chest pain (often unilateral) occur when the effusion is large. Sputum production and cough are present in the case of concurrent PTB, which is common. Constitutional symptoms such as fever, night sweats, malaise and weight loss may also be present. Effusion can progress to tuberculous empyema, characterised by purulent fluid containing large numbers of bacilli. Tuberculous empyema is often associated with thickened, scarred and calcified pleura. Diagnosis is assisted by examination of the pleural fluid via paracentesis and chest x-ray. See Chapter 3.

### 2.2.7 Tuberculous pericardial effusion

Clinical signs of a tuberculous pericardial effusion include chest pain, shortness of breath, oedema of the lower limbs and sometimes ascites. Clinical examination may show pericardial friction rub, raised jugular pressure and tachycardia. CXR and ultrasound are key elements for diagnosis. Pericardiocentesis may be necessary in the event of acute heart failure with haemodynamic compromise. It must be performed by experienced personnel in well-equipped hospitals, and when possible, under direct visualisation with ultrasound.

### 2.2.8 Cutaneous tuberculosis

The clinical presentation of cutaneous TB is chronic, painless, non-pathognomonic lesions, ranging from small papula and erythema to large tuberculomas. The diagnosis is based on culture from a biopsy.

### Referencias


2.3 Disseminated or miliary tuberculosis

Miliary TB is a generalised massive infection characterised by hematogenous diffusion of *M. tuberculosis* throughout the body. It is a medical emergency.

The disease may manifest as a miliary pattern, or very small nodular elements ('millet seeds') in the lungs.

The classic acute form is mostly found in children, young adults and HIV-infected patients. The presentation can be either abrupt or insidious, with progressive deterioration in the patient’s physical condition. The clinical picture is often completed within one to two weeks and is characterised by a profoundly altered physical condition, marked wasting, headache and constant high fever. Discrete dyspnoea and coughing suggest a pulmonary focus; however, lungs can often be clear on auscultation. A moderate hepatosplenomegaly is occasionally found. Certain forms of miliary TB evolve in a subacute manner over several months. Given this non-specific clinical picture, typhoid fever and septicaemia should be considered in the differential diagnosis.

Diagnosis of miliary TB is confirmed by CXR (Chapter 3).

When feasible, fundoscopy may reveal choroidal tubercles.

Sputum smear examination is usually negative.

When there is no possibility of obtaining CXR, the lack of response to antibiotics is an argument in favour of miliary TB.

The tuberculin skin test is more likely to be falsely negative than in any other form of TB.

In children, the risk of meningitis (20-40%) is high. Lumbar puncture should be routinely performed if miliary TB is suspected.

Referencias


2.4 Clinical presentation in HIV-infected patients

Among HIV-infected patients, TB is the most common opportunistic infection and the leading cause of morbidity and mortality. According to the WHO clinical staging system for HIV/AIDS, patients with PTB are in clinical stage 3 and patients with EPTB in clinical stage 4.

In the early stages of HIV infection, when the immune system is functioning relatively normally, the clinical signs of TB are similar to those in seronegative individuals.

As the immune system deteriorates in later stages of the disease, smear-negative PTB, disseminated TB and EPTB become more common. These cases are more difficult to diagnose, and have a higher fatality rate than smear-positive PTB cases. Patients may have difficulty expectorating, so more advanced sputum collection techniques may be necessary (Chapter 3 and Appendix 3).

Algorithms presented in Chapter 4 use clinical criteria combined with laboratory and other investigations to help diagnose TB in HIV-infected individuals.

Table 2.2 provides a differential diagnosis of PTB in HIV-infected patients.

**Table 2.2 - Differential diagnosis for PTB in HIV-infected patients**
The most common EPTB in HIV-infected patients are miliary TB, TB meningitis and diffuse lymphadenopathy in children, and lymph node TB, pleural effusion, pericarditis, TB meningitis and miliary TB in adults.

Immune reconstitution inflammatory syndrome (IRIS) is a clinical presentation of TB in patients starting antiretroviral therapy. See Chapter 12 for clinical presentation and management of IRIS.

### Referencias


   [https://www.who.int/hiv/pub/guidelines/HIVstaging150307.pdf](https://www.who.int/hiv/pub/guidelines/HIVstaging150307.pdf)

### 2.5 Summary of clinical presentations of tuberculosis

**Table 2.3 - Clinical presentations and considerations for HIV-infected patients**

<table>
<thead>
<tr>
<th>Diseases</th>
<th>Comments</th>
</tr>
</thead>
</table>
| Other pneumonia (bacterial, viral, atypical) | • Bacterial pneumonia (most often *S. pneumoniae, H. influenzae*) is common at all stages of HIV infection.  
• Atypical pneumonia (*M. pneumoniae, C. pneumoniae*) and viral pneumonia are possible at any CD4 count, except in the case of cytomegalovirus, which occurs at CD4 < 50. |
| Pneumocystosis (Pneumocystis jirovecii pneumonia or PCP or PJP) | • Pneumocystis has many characteristics in common with PTB (insidious onset, persistent cough, fever) but tends to occur in the advanced stages of HIV infection (CD4 < 200).  
• Pneumocystis is unlikely in patients taking co-trimoxazole prophylaxis.  
• It imparts a greater degree of dyspnoea, rarely produces effusions, and is not usually accompanied by haemoptysis. |
| Pulmonary Kaposi's sarcoma (KS) | • KS can resemble PTB, with slow onset of cough, fever, haemoptysis, night sweats and weight loss. It is a disease of advanced stage HIV, and in most cases, is preceded or accompanied by lesions involving the skin and mucus membranes. |
| Less common diseases | • Pulmonary cryptococcosis, histoplasmosis and other fungal infections.  
• Pulmonary nocardiosis: on direct smear, nocardia are weakly acid-fast, and similar in appearance to mycobacteria (although they are branching filamentous bacilli, particularly on Gram staining). |
<table>
<thead>
<tr>
<th>Sites</th>
<th>Clinical presentations</th>
<th>Considerations for HIV patients</th>
</tr>
</thead>
</table>
| Pulmonary TB          | • Prolonged cough (> 2 weeks), with or without sputum production.  
• Weight loss, anorexia, fatigue, shortness of breath, chest pain, moderate fever, night sweats, haemoptysis. | • Fever and weight loss are more common and pronounced.  
• Cough and haemoptysis may be less common (less inflammation and cavity formation).  
• See algorithms, Chapter 4.                                                                                                                                                                           |
| Disseminated miliary TB | • Non-specific symptoms: high fever, headache, weight loss.  
• Deterioration over days or weeks.  
• Simultaneous involvement of multiple organs.  
• High risk of meningitis in children.  
• Miliary findings CXR. | • May be confused with severe wasting in advanced HIV disease.  
• *M. tuberculosis* sometimes isolated from blood cultures.                                                                                                                                 | |
| Lymph nodes TB        | • Most often in cervical region.  
• Non-inflammatory, painless node > 2 cm, chronic (> 4 weeks); fistulisation possible. | • HIV infection can cause persistent generalised lymphadenopathy (PGL). PGL lymph nodes are painless, and symmetrical. Posterior cervical or epitrochlear nodes are often involved.  
• Other common causes of lymphadenopathy include lymphoma, carcinomatous metastases, Kaposi sarcoma.  |
| TB meningitis         | • Subacute, insidious.  
• Headaches, irritability, fever, altered mental status.  
• Meningeal syndrome usually present. | • Rule out cryptococcal meningitis: perform antigen test on serum and CSF.                                                                                                                                                                           |
| Bone and joint TB     | • Monoarthritis with joint destruction and little or no pain.  
• Deformity of the spine (Pott’s disease). | • Multifocal disease more common.                                                                                                                                                                                                                   |
| Urogenital TB         | • Renal: urinary symptoms, few constitutional symptoms; suspected when no response to antibiotics for urinary infection.  
• Non-specific gynaecological symptoms, infertility or epididymitis with scrotal pain. |                                                                                                                                                                                                                                                   |
| Abdominal TB          | • Ascites (may mask weight loss).  
• Abdominal mass, pain, diarrhoea. | • PTB is more frequently associated.                                                                                                                                                                                                             |
| Effusions             | • Pleural: pleuritic chest pain, dyspnoea.  
• Pericardial: chest pain, dyspnoea, lower limb oedema or ascites, pericardial friction rub. | • Serious effusions are common.  
• TB is the most likely aetiology in high TB-HIV prevalence settings.                                                                                                                                                                         |
TB is considered as non-severe if the following criteria are met:

- negative smear microscopy, and
- uncomplicated PTB with a small infiltrate confined to one lobe and no cavities, or
- uncomplicated extra-thoracic lymph node TB, or
- uncomplicated intrathoracic lymph node TB.

Chapter 3: Diagnosis and follow-up investigations

3.1 Active tuberculosis

3.2 Latent tuberculosis infection

3.3 Other investigations

Update: October 2022

3.1 Active tuberculosis

3.1.1 Introduction

Active tuberculosis (TB) is bacteriologically confirmed by the detection of *M. tuberculosis* complex through different bacteriological tests. These tests detect either the organism (smear microscopy and culture), or some of its genetic material (genotypic tests, including rapid molecular tests and genome sequencing).

Specimens used for bacteriological testing include respiratory specimens (sputum, nasopharyngeal aspirate and, in children, gastric aspirate) and extrapulmonary specimens (Table 3.6).

Drug susceptibility testing (DST) is indicated for all patients with confirmed TB. It can be performed using genotypic or phenotypic tests:

- Genotypic DST (gDST) can detect resistance to TB drugs by identifying specific gene mutations.
- Phenotypic DST (pDST) can detect resistance to TB drugs by measuring the growth of *M. tuberculosis* in the presence of the drug.

To diagnose TB and determine the appropriate regimen at baseline:

- All patients should be tested with a rapid molecular test (RMT) to detect *M. tuberculosis* and rifampicin resistance.
- Whatever the result of the rifampicin susceptibility test (resistance detected or not), all patients, if possible, should be tested with an RMT for isoniazid resistance and at least those with high risk of isoniazid resistance (for definition of high risk of resistance, see below).
- All patients with rifampicin resistance should be tested for resistance to fluoroquinolones and other TB drugs.
- All patients with isoniazid resistance and rifampicin susceptibility should be tested for resistance to fluoroquinolones.
- Culture, pDST and genome sequencing may be required.

In limited-resource settings, resistance to TB drugs should be investigated in priority in patients with:

- High risk of mortality: e.g. HIV-infected patients or patients with extensive disease.
- High risk of resistance: patients with previous TB treatment, or in contact with a TB case resistant to TB drug(s), or coming from an area of high prevalence of resistance to TB drug(s).

Notes:

- Negative bacteriological tests for *M. tuberculosis* does not rule out TB.
- A negative DST does not necessarily rule out drug resistance.
Other investigations can assist TB diagnosis. These investigations include: lateral flow urine lipoarabinomannan assay (LF-LAM) which detects an antigen of *M. tuberculosis* cell wall excreted in urine, medical imaging, and some biological tests.

### 3.1.2 Rapid molecular tests

RMTs are nucleic acid amplification tests (NAATs). They can detect *M. tuberculosis* and drug resistance by identifying resistance-conferring mutations in certain genes (Table 3.1). Other drug resistance-conferring mutations may be present, but not detected by RMTs. In areas where prevalence of these mutations is high, RMT sensitivity may be decreased[1].

#### Table 3.1 – Rapid molecular tests and detection of drug resistance

<table>
<thead>
<tr>
<th>Tests</th>
<th>TB drug resistance (targeted genes)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Low complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>Xpert MTB/RIF</td>
<td>Rifampicin (<em>rpoB</em>)</td>
</tr>
<tr>
<td>Xpert MTB/RIF Ultra</td>
<td></td>
</tr>
<tr>
<td>Truenat MTB-RIF Dx</td>
<td></td>
</tr>
<tr>
<td><strong>Moderate complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td><strong>High complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>GenoType MTBDRplus (V2.0)</td>
<td></td>
</tr>
<tr>
<td>Genoscholar NTM+MDRTB II</td>
<td></td>
</tr>
<tr>
<td><strong>Low complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>Xpert MTB/XDR</td>
<td>Isoniazid high-level resistance (<em>katG</em>)</td>
</tr>
<tr>
<td><strong>Moderate complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>GenoType MTBDRplus (V2.0)</td>
<td>Isoniazid low-level resistance, thionamides (<em>inhA promoter</em>)</td>
</tr>
<tr>
<td>Genoscholar NTM+MDRTB II</td>
<td></td>
</tr>
<tr>
<td><strong>High complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>GenoType MTBDRs/ (V2.0)</td>
<td>Fluoroquinolones (<em>gyrA, gyrB</em>)</td>
</tr>
<tr>
<td></td>
<td>Aminoglycosides (<em>rrs, eis</em>)</td>
</tr>
<tr>
<td><strong>High complexity NAATs</strong></td>
<td></td>
</tr>
<tr>
<td>Genoscholar PZA-TB II</td>
<td>Pyrazinamide (<em>pncA</em>)</td>
</tr>
</tbody>
</table>

(a) Mutations in other genes can result in resistance to thionamides. Consequently, absence of inhA mutation does not rule out resistance.

(b) Specific mutations in gyrA (e.g. mutations recognized by the probes MUT3B, 3C, 3D) are associated with high-level fluoroquinolones resistance.

RMTs have a good specificity, but are less sensitive than culture. Their various levels of complexity determine their use at different levels of health facilities. Low complexity RMTs are preferred in routine practice.

#### Low complexity nucleic acid amplification tests

1) **Xpert assays**

Xpert assays are almost fully automated. An uninterrupted power supply and a computer are required to perform and read assays.

Xpert assays can be performed on:
- Respiratory specimens
- Extrapulmonary (EP) specimens:
- Lymph node biopsy or aspirate: suspicion of lymph node TB or detection of rifampicin resistance in clinically diagnosed lymph node TB;
- Cerebrospinal fluid (CSF): suspicion of TB meningitis;
- Pleural fluid: suspicion of TB with pleural effusion;
- Stool: suspicion of PTB in children;
- Pericardial fluid: suspicion of TB with pericardial effusion (sampling to be performed only by experienced clinicians);
- Urine: suspicion of genitourinary TB; suspicion of disseminated TB in HIV-infected patients;
- Synovial fluid: suspicion of TB arthritis;
- Peritoneal fluid: suspicion of abdominal TB;
- Blood: suspicion of disseminated TB in HIV-infected patients.

Xpert MTB/RIF and Xpert MTB/RIF Ultra assays provide simultaneously results for *M. tuberculosis* detection and rifampicin resistance.

Sensitivity of Xpert MTB/RIF Ultra assay is higher than that of Xpert MTB/RIF assay. It provides a result “trace” corresponding to the lowest bacillary load for *M. tuberculosis* detection. It is preferred for HIV-infected patients, children, EP specimens, and sputum smear-negative specimens. Its specificity is lower in patients with a history of TB, as a “trace” result may indicate that the specimen contains fragments of dead bacilli.

WHO has validated their use on lymph node biopsy or aspirate, CSF, and pleural fluid, synovial fluid. Xpert assays can be performed on any biopsy specimens (lymph node, bone, skin, resection material, etc.) with good performance. Xpert assays have shown acceptable performances in various studies on other specimens (peritoneal fluids, stools, and urine). Xpert assays on blood have a low sensitivity compared to culture and are not routinely recommended.

Xpert MTB/XDR assay detects resistance to isoniazid (low- and high-level), fluoroquinolones (low- and high-level), aminoglycosides, and thionamides. It does not detect resistance to rifampicin.

Xpert MTB/XDR assay employ the same platform as other Xpert assays, but require a 10-colour module instead of the 6-colour module used for Xpert MTB/RIF and Xpert MTB/RIF Ultra assays. The 10-colour module can also read Xpert MTB/RIF and Xpert MTB/RIF Ultra assays.

Xpert MTB/XDR assay should be used:
- When resistance to rifampicin has been detected by Xpert MTB/RIF or Xpert MTB/RIF Ultra, to detect resistance to other drugs.
- When *M. tuberculosis* has been detected by Xpert MTB/RIF or Xpert MTB/RIF Ultra or culture, to detect resistance to isoniazid in all patients, if possible, and at least those with high risk of isoniazid resistance (Section 3.1.1).
- Before using a fluoroquinolone containing regimen in isoniazid-resistant TB (Hr-TB), multidrug-resistant (MDR-TB), rifampicin-resistant TB (RR-TB) or drug-susceptible TB treated with the regimen 2HPZ-Mfx/2HP-Mfx.
- Before treating drug-susceptible TB meningitis with the regimen 6HRZ-Eto.
- In patients with fluoroquinolone-susceptible TB, initially treated with a fluoroquinolone-containing regimen, and presenting a smear-positive microscopy at Month 2 or later.

**Table 3.2 – Main performances of Xpert assays**
For more information on specimen processing and Xpert instruments see Appendix 1.
For interpretation of Xpert assay results see Appendix 2.

2) Truenat assays

Truenat assays require:
- Several manual steps (pipetting).
- Sequential testing for *M. tuberculosis* detection (Truenat MTB Plus), then for rifampicin resistance detection (Truenat MTB-RIF Dx).
- Separate kits for specimen preparation, DNA extraction, DNA amplification, and detection of *M. tuberculosis* and rifampicin resistance.

Truenat MTB Plus can only be performed on sputum specimens (positive or negative smear microscopy). It is not recommended for other respiratory specimens or EP specimens\[2][7].
Specificity is high, i.e. a positive result is unlikely to be a false positive\[14].
Tests can be run at room temperatures of up to 40 °C and humidity of up to 80%. Truenat instruments are battery-operated and can be used in peripheral or mobile health facilities.
Interpretation of results is the same as for Xpert (Appendix 2).

**Table 3.3 – Main performances of Truenat assays**
3) TB-LAMP

Although validated by WHO, this test has major limitations:

- It does not detect rifampicin resistance.
- Its sensitivity is lower than that of other low complexity NAATs in HIV-infected or smear-negative patients.
- It cannot be used for the diagnosis of extrapulmonary TB (EPTB)[2].

Box 3.1 – Choice of low complexity NAATs

- **Xpert**: first line tests for the diagnosis of TB in children and adults.
- **Truenat**: if no power supply or operating temperature between 31 and 40 °C.
- **TB-LAMP**: not recommended.

**Moderate complexity nucleic acid amplification tests**

WHO recommends these tests for the simultaneous detection of *M. tuberculosis* and resistance to rifampicin and isoniazid, from smear-positive and negative respiratory specimens, in children and adults, including HIV-infected patients.

**Table 3.4 – Performances of moderate complexity NAATs**

<table>
<thead>
<tr>
<th>Tests</th>
<th>Performances</th>
</tr>
</thead>
</table>
| - Abbott Real Time MTB and MTB RIF/INH BD MAX MDR-TB  
- Hain FluoroType MTB and MTBDR  
- Roche cobas MTB and MTB-INH/RIF | Detection of MTB compared to culture:  
- Sensitivity 93%  
- Specificity 97.7%  
Detection of rifampicin resistance compared to pDST:  
- Sensitivity 96.7%  
- Specificity 98.9%  
Detection of isoniazid resistance compared to pDST:  
- Sensitivity 86.4%  
- Specificity 99.8% |

NAATs of moderate complexity have several limitations:

- Need for space, equipment, qualified staff; only feasible in regional laboratories.
- Their use does not eliminate the need for pDST, high complexity NAATs, or genome sequencing to:
  - test susceptibility to other TB drugs;
  - confirm a negative result in patients at high risk of drug resistance.
- Their use on EP specimens is not validated.
High complexity nucleic acid amplification tests

Line probe assays (LPA) can detect specific rifampicin, isoniazid, fluoroquinolones, aminoglycosides, and pyrazinamide resistance encoding mutations in *M. tuberculosis*.

These tests can be performed on isolates of *M. tuberculosis* (indirect testing). Some can be performed on sputum specimens (direct testing).

NAATs of high complexity have several limitations:
- Need for space, equipment, highly qualified staff; only feasible in reference and national laboratories.
- Risk of cross-contamination (tests are performed in an open system that can lead to the detection of DNA from sources other than the specimen).
- To benefit from the short turnaround time of these tests, efficient logistical support is required to ensure specimens are transported to the laboratory and the results are delivered in a timely manner.
- Their use does not eliminate the need for pDST or genome sequencing to:
  - test sensitivity to other TB drugs;
  - confirm a negative result in patients at high risk of drug resistance.
- Although direct test results can be obtained in 1 to 2 days, for indirect tests, it is necessary to wait the time required for bacterial growth (Appendix 5).
- Their use on respiratory (non-sputum) or EP specimens is not validated.

**Box 3.2 – WHO validated LPAs**

**First-line LPAs**
- GenoType MTBDRplus version 2 ("Hain first line test"): initial test to detect resistance to rifampicin and isoniazid on smear-positive sputum specimens and *M. tuberculosis* isolates. Compared to pDST, sensitivity is 98.2% for rifampicin, and 97.8% for isoniazid; specificity is 95.4% for rifampicin, and 98.8% for isoniazid[^15]. On smear-negative sputum specimens, sensitivity is low (44.4%), and its use is not recommended[^18].
- Genoscholar NTM+MDRTB II ("Nipro test"): performances comparable to GenoType MTBDRplus to detect resistance to rifampicin and isoniazid on smear-positive sputum specimens and *M. tuberculosis* isolates. Not recommended on smear-negative sputum specimens. Can differentiate *M. avium*, *M. intracellulare* and *M. kansasii* from other non-tuberculous mycobacteria.
- Genoscholar PZA-TB II: to detect resistance to pyrazinamide on *M. tuberculosis* isolates. Compared to pDST, sensitivity is 81%, and specificity is 97%[^18].

**Second-line LPA**
GenoType MTBDRsl/ version 2 ("Hain second line test"): in patients with confirmed MDR/RR-TB, to detect resistance to fluoroquinolones (high- and low-level) and aminoglycosides on smear-positive or smear-negative sputum specimens and *M. tuberculosis* isolates. The number of "indeterminate" results is higher for smear-negative than for smear-positive sputum specimens. For smear-positive sputum specimens, sensitivity is 93% for fluoroquinolones, and 88.9% for aminoglycosides; specificity is 98.3% for fluoroquinolones, and 91.7% for aminoglycosides[^16].

### 3.1.3 Genome sequencing

Genome sequencing can only be performed in highly specialized reference laboratories.

It can rapidly:
- Detect mutations associated with TB drug resistance. When available, it is particularly useful to identify:
  - resistance to TB drugs for which pDST is unreliable, or no RMTs are available;
  - mutations missed by RMTs (+ 20% of drug resistance detection compared to RMTs has been described[^17]).
- Detect mixed infection (infection with distinct *M. tuberculosis* strains).
- Identify heteroresistance (same strain, with different resistance profiles).
- Differentiate treatment relapse and reinfection with a different strain.

Genome sequencing methods include Sanger sequencing (reference method) and next generation sequencing (NGS). The advantage of NGS is that, unlike Sanger sequencing, it provides results for a large number of genes in a single reaction. NGS results are interpreted by reference laboratories using specific software and mutation databases[^9].
Some mutations associated with resistance to recently introduced drugs (e.g. bedaquiline and delamanid) and their therapeutic implications are still not well-known.

The two main NGS techniques are targeted NGS (tNGS) and whole genome sequencing (WGS):

- tNGS (on smear-positive sputum specimens or culture isolates): detection of resistance conferring mutations on 18 selected genes: first-line TB drugs, fluoroquinolones, aminoglycosides, linezolid, bedaquiline, clofazimine, ethionamide (Deeplex® Mycobacteria). Used in routine.
- WGS (on culture isolates): detection of resistance conferring mutations on whole genome (i.e. potentially all TB drugs). Used for research.

### 3.1.4 Smear microscopy

The purpose of smear microscopy is to detect acid-fast bacilli (AFB) in stained specimens.

Smear microscopy has several limitations:

- It has a sensitivity lower than RMTs and culture in respiratory specimens (65% compared to culture[^11]) and EP specimens (48% compared to culture[^18]).
- It has a low sensitivity in patients with low bacillary load in sputum (paucibacillary TB), e.g. children and HIV-infected patients.
- It cannot differentiate between *M. tuberculosis* and non-tuberculous mycobacteria. However, in areas with high TB prevalence, AFB detected on smear microscopy are most likely *M. tuberculosis*.
- It does not determine if bacilli are viable (alive) or non-viable (dead).
- It does not determine susceptibility of the bacilli to TB drugs.

Sputum smear microscopy is no longer the recommended initial diagnostic test for PTB. However, it still plays a role:

- When RMTs are not available.
- For assessing the infectiousness of PTB patients.
- For monitoring the response to TB treatment in patients with:
  - drug-susceptible PTB ([Chapter 9](#)).
  - drug-resistant PTB. However, culture is also required for monitoring treatment response in these patients ([Chapter 10](#)) and ([Chapter 11](#)).

For improving the sensitivity of smear microscopy:

1. Two sputum specimens should be examined. Approximately 86% of sputum smear-positive patients are identified during the first examination, and an additional 12% during the second. It is not necessary to carry out more than 2 examinations[^18].
2. Light-emitting diode (LED) fluorescent microscopy to examine auramine-stained smears is preferred to Ziehl-Neelsen microscopy, as it is more sensitive, and reading is more rapid.

Concentration techniques can also increase the sensitivity of smear microscopy[^20].

For sputum specimen collection, storage and shipment see [Appendix 3](#).

For sputum smear preparation and staining techniques see [Appendix 4](#).

### 3.1.5 Culture

Culture consists of growing *M. tuberculosis* in specific liquid or solid media.

Culture on liquid medium (automated or manual mycobacterial growth indicator tube, MGIT) is the reference method for the diagnosis of PTB and EPTB. Given the long turnaround time and equipment required, it is not used as initial diagnostic test. Culture on solid medium (Lowenstein-Jensen) is cheaper, less prone to contaminations than cultures on liquid media, but its turnaround time is longer.

Other culture techniques are less commonly used[^d].

Culture is necessary to:

- Confirm treatment failure.
- Assess treatment response in patients with drug-resistant PTB ([Chapter 10](#)) and ([Chapter 11](#)).
- Evaluate treatment outcome in patients with drug-resistant PTB ([Chapter 17](#)).
- Provide isolates for the following tests:
  - First-line LPAs on sputum smear-negative and EP specimens
  - Genoscholar PZA-TB II, regardless of sputum smear positivity
Culture may help to diagnose TB when other bacteriological tests are negative or inconclusive:
- In patients with signs and symptoms of TB and a negative RMT, particularly when resistance is suspected.
- In adults with history of TB in the previous 5 years and showing a “trace” result by Xpert MTB/RIF Ultra.

Culture has several limitations:
- Only specialized laboratories implementing systematic quality assurance procedures can be relied upon for culture (often national reference laboratories or supranational).
- *M. tuberculosis* is a slow-growing bacillus. Positive culture results are obtained after 2 to 4 weeks.

For sputum specimen collection, storage and shipment see Appendix 3.
For the time required to obtain the results see Appendix 5.

### 3.1.6 Phenotypic drug susceptibility testing

Phenotypic DST (pDST) determines if a strain is resistant to a TB drug by evaluating the growth in the presence of the drug. It can determine two levels of resistance (low and high) for isoniazid and fluoroquinolones.

The pDST is essential to detect resistance to drugs for which there are no reliable RMTs, and when genome sequencing is not available.

In addition, pDST may be necessary:
- If an RMT indicates *M. tuberculosis* “detected” and drug resistance “indeterminate”.
- If an RMT indicates drug susceptibility in a patient at high risk of resistance.
- In areas with a high prevalence of mutations not detected by RMTs.

Phenotypic DST is performed on culture isolates by specialized laboratories (often national reference laboratories or supranational).

The pDST is not reliable for all drugs, even when performed by a highly qualified laboratory\[^1\].

**Table 3.5** – Reliability of pDST for first- and second-line TB drugs

<table>
<thead>
<tr>
<th>Reliability of pDST</th>
<th>TB drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Highly reliable</td>
<td>IsoniazidDTV</td>
</tr>
<tr>
<td></td>
<td>Rifampicin</td>
</tr>
<tr>
<td></td>
<td>Fluoroquinolones</td>
</tr>
<tr>
<td></td>
<td>Aminoglycosides</td>
</tr>
<tr>
<td>Unreliable (should not be performed)</td>
<td>Ethambutol</td>
</tr>
<tr>
<td></td>
<td>Ethionamide</td>
</tr>
<tr>
<td></td>
<td>Cycloserine or terizidone</td>
</tr>
<tr>
<td></td>
<td>Para-aminosalicylic acid (or sodium)</td>
</tr>
<tr>
<td></td>
<td>Delamanid</td>
</tr>
<tr>
<td>Reliable, but limited access outside of supranational laboratories</td>
<td>Bedaquiline</td>
</tr>
<tr>
<td></td>
<td>Linezolid</td>
</tr>
<tr>
<td></td>
<td>Clofazimine</td>
</tr>
<tr>
<td>Reliable when performed in a high-quality laboratory (difficult to perform)</td>
<td>Pyrazinamide</td>
</tr>
</tbody>
</table>

[^1]: Reference or citation information is needed for the statement regarding the unreliability of pDST for all drugs.
3.1.7 Summary of bacteriological tests

The tables below provide an overview of the specimens that can be used for each test, and of the tests that can detect resistance to each TB drug.

**Table 3.6 – Specimens for bacteriological tests**

<table>
<thead>
<tr>
<th>Tests</th>
<th>Specimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xpert, microscopy, culture</td>
<td>Respiratory or EP specimens</td>
</tr>
<tr>
<td>Truenat</td>
<td>Sputum (smear-positive or negative)</td>
</tr>
<tr>
<td>Moderate complexity NAATs</td>
<td>Respiratory specimens</td>
</tr>
<tr>
<td>GenoType MTBDRplus version 2</td>
<td>Sputum (smear-positive only) M. tuberculosis isolate</td>
</tr>
<tr>
<td>Genoscholar NTM+MDRTB II</td>
<td></td>
</tr>
<tr>
<td>Genoscholar PZA-TB II</td>
<td>M. tuberculosis isolate</td>
</tr>
<tr>
<td>GenoType MTBDRs/ version 2</td>
<td>Sputum (smear-positive or negative) M. tuberculosis isolate</td>
</tr>
<tr>
<td>tNGS</td>
<td>Sputum (smear-positive only) M. tuberculosis isolate</td>
</tr>
<tr>
<td>WGS</td>
<td>M. tuberculosis isolate</td>
</tr>
</tbody>
</table>

**Table 3.7 – Tests to detect specific drug resistance**
### 3.1.8 Lateral flow urine lipoarabinomannan assay

TB lipoarabinomannan (LF-LAM) is a urine-based point-of-care test that detects lipoarabinomannan (LAM) antigen, which is a marker of active TB.

This test is easy to perform by trained staff, including in peripheral health facilities.

Advantages of LF-LAM over sputum-based tests include:
- Urine specimens easier to collect.
- No risk of staff contamination during specimen collection or processing.
The urine is applied to the test strip, left at room temperature for 25 minutes, then read by the naked eye by comparing the band for positivity to a grading scale provided by the manufacturer.[4]

This rapid test should be used in the diagnosis of PTB and EPTB in HIV-infected children and adults. Its rapidly obtained result can contribute to reducing TB mortality among these patients.[7]

Its performances depend on the individual level of immunodeficiency at the time of testing. Its sensitivity is low, but it has an acceptable specificity (see below).

The LF-LAM test is recommended for the following patient groups[22]:
- HIV-infected patients with signs and symptoms of TB or seriously ill[9], irrespective of CD4 count (sensitivity: 35%; specificity: 95%).
- Hospitalised patients with advanced HIV disease[9] (sensitivity: 64%; specificity: 82%).
- HIV-infected outpatients with CD4 count < 100 cells/mm³ (sensitivity: 40%; specificity: 87%).

If LF-LAM test is positive: TB treatment should be initiated[1].

Due to the low sensitivity of the LF-LAM test, a negative result does not rule out TB. The test does not provide information on drug susceptibility. Therefore, all above-mentioned patients should be tested with an RMT, regardless of whether the LF-LAM result is positive or negative.

### 3.1.9 Medical imaging

**Radiography**

Chest x-ray (CXR) is used to:
- Detect abnormalities suggestive of PTB and other intra-thoracic TB localisations (pleural, pericardial, miliary).
- Evaluate the severity of intra-thoracic lesions.

It is particularly useful in the diagnosis of PTB in children (Chapter 5).

For PTB, CXR has a higher sensitivity than TB symptoms[23]: a patient with a normal CXR is unlikely to have PTB. For this reason, it can also be used as a screening tool (Chapter 6) and a triaging tool to identify patients with respiratory symptoms eligible for an RMT.

CXR is also used to:
- Evaluate the response to TB treatment.
- Look for possible complications in case of worsening respiratory symptoms (pneumothorax, tracheal stenosis, etc.).

CXR has several limitations:
- Low specificity: except for cavities or miliary TB, which are specific to TB other abnormalities seen on CXR may be due to other pulmonary diseases.
- Variable quality, depending on several factors:
  - equipment and supply
  - positioning (obtaining quality CXR in children is challenging)
  - reader training and proficiency
- Difficulty distinguishing active from healed lesions
- Error rate of approximately 20%[24] (specialists’ under/over-reading of the film)

When available, digital CXR has advantages over x-ray films:
- Consistent quality
- Easier image archiving
- No need for reagents and films
- Rapid transmission for teleconsultation and specialist advice
- Immediate results; possibility to screen large numbers of people within a short timeframe
- Lower radiation exposure for staff and patients.

Interpretation of digital CXR can be assisted by computer-aided detection (CAD) software packages. CAD analyses CXR for the presence of PTB-compatible abnormalities, and divides images into “normal” and “abnormal”, thereby reducing the number of CXR that need to be read by a clinician. CAD is as sensitive as a radiologist[28].
Computer-aided CXR interpretation assists clinicians when all CXR cannot be read by a radiologist. However, a radiologist should be consulted locally or via telemedicine to interpret difficult CXR (e.g. in children).

Bone x-ray is used to diagnose and evaluate severity of bone and/or joint TB and assess treatment response.

**Ultrasound**

Ultrasound (including point-of-care ultrasound, POCUS) may be useful in:

- **PTB**: pulmonary consolidation can support the diagnosis of PTB.
- **EPTB**: if suspected pleural/pericardial effusion or abdominal TB in children and adults, particularly in immunocompromised patients (e.g. HIV-infection, malnutrition).

**Table 3.8 – Medical imaging findings suggestive of TB**

<table>
<thead>
<tr>
<th>Sites</th>
<th>Findings</th>
</tr>
</thead>
</table>
| **Pulmonary TB** | **Children**  
See [Chapter 5](#)  
**Adolescents and adults**  
CXR can show:  
- Infiltrates typically located in apical and posterior segment of upper lobes and superior segments of lower lobes.  
- Cavities (specific for TB), patchy, poorly defined consolidations.  
**Patients with TB/HIV**  
As above.  
- In advanced immunodeficiency, infiltrates tend to be more homogeneous, diffuse and located in the lower lungs.  
- Less cavities than in non-HIV-infected patients.  
- Mediastinal and hilar lymphadenopathy may be observed.  
- Miliary pattern. |
| **Miliary TB**   | CXR can show miliary nodules (1-3 mm in diameter) disseminated in both fields and uniformly distributed throughout the lung.                                                                                   |
| **Pleural effusion** | - CXR: effusion (even with minimal clinical signs):  
  - Mostly unilateral.  
  - Obliteration of costophrenic angle.  
  - Opacity with curved upper margin.  
  - Ultrasound: anechogenic fluid on the costophrenic angle (may be echogenic in empyema).  
                                                                                                                               |
| **Pericardial effusion** | - CXR: cardiac silhouette enlargement, “water bottle” silhouette (very large effusions).  
- Ultrasound: anechogenic fluid around the heart (may be echogenic if purulent).                                                                 |
| **Bone/joint TB**| X-ray can show:  
- Any bone/joint: osteopenia (demineralization), bone destruction with relative preservation of cartilage space.  
- Spine: destruction of an inter-vertebral disk, osteopenia, irregularity of bone margin, bone destruction, paravertebral abscesses. |
| **Abdominal TB** | Ultrasound can show enlarged lymph nodes consistent with TB (and other diseases, especially in HIV infection), bowel wall thickening (ileo-caecal region), hypoechoic micro-abscesses of liver and/or spleen, ascites. |

**Notes:**

- Radiographical and ultrasound findings of EPTB are non-specific. A differential diagnosis should always be considered.
In HIV-infected patients in settings of high TB prevalence, pleural/pericardial effusion, enlarged abdominal lymph nodes, splenic microabscesses, and ascites are highly suggestive of EPTB. Adolescents typically have CXR abnormalities similar to those found in adults, however, they may also have abnormalities commonly seen in children, such as enlarged hilar lymph nodes.

### 3.1.10 Other laboratory tests on tissues and body fluids

The diagnosis of TB can be supported by biological tests performed on tissues or body fluids.

**Table 3.9 – Findings suggestive of TB in tissues or body fluids**

<table>
<thead>
<tr>
<th>Tissues/fluids</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymph node</td>
<td>Cytology: granulomatous tissue, presence of giant Langhans cells, and/or caseous necrosis. AFBs are not always found by microscopy. See Lymph node fine needle aspiration, Appendix 7.</td>
</tr>
<tr>
<td>CSF</td>
<td>Clear, hyper-concentrated liquid.</td>
</tr>
<tr>
<td></td>
<td>High protein level &gt; 0.40 g/l (see Pandy test, Appendix 8).</td>
</tr>
<tr>
<td></td>
<td>Low glucose &lt; 60 mg/l.</td>
</tr>
<tr>
<td></td>
<td>Ratio CSF glucose/blood glucose &lt; 0.5.</td>
</tr>
<tr>
<td></td>
<td>Between 100 and 1,000 white cells/mm³, of which over 80% are lymphocytes. In HIV-infected patients, rule out cryptococcal meningitis.</td>
</tr>
<tr>
<td>Peritoneal fluid</td>
<td>Translucent, yellow-coloured liquid.</td>
</tr>
<tr>
<td></td>
<td>Exudate rich in lymphocytes, usually &gt; 300 white cells/mm³; Rivalta test positive (Appendix 8).</td>
</tr>
<tr>
<td></td>
<td>Serum-ascites albumin gradient (SAAG): &lt; 1.1 g/dl: consistent with TB (and many other conditions).</td>
</tr>
<tr>
<td></td>
<td>&gt; 1.1 g/dl: peritoneal TB unlikely.</td>
</tr>
<tr>
<td></td>
<td>Adenosine deaminase (ADA) &gt; 39 U/l, likely due to TB[27].</td>
</tr>
<tr>
<td>Pleural fluid</td>
<td>Straw-coloured fluid.</td>
</tr>
<tr>
<td></td>
<td>High protein level ≥ 30 g/l (Rivalta test, Appendix 8).</td>
</tr>
<tr>
<td></td>
<td>Rich in white cells (1,000-2,500/mm³), with predominant lymphocytes.</td>
</tr>
<tr>
<td></td>
<td>ADA typically &gt; 50 U/l. Pleural effusion with an ADA &lt; 40 U/l is much less likely due to TB. The specificity is increased when ADA is &gt; 50 U/l and the lymphocyte-neutrophil ratio is &gt; 0.75[28].</td>
</tr>
</tbody>
</table>

**Notes:**
- ADA levels increase in TB. ADA is therefore a surrogate marker for TB in pleural and peritoneal fluids. Although not widely available, kits can be purchased to perform the test if a spectrophotometer is available.
- The sensitivity of ADA in peritoneal fluid is lower in patients with cirrhosis.
- HIV-infected patients may have lower levels of ADA.

**Notas**

(a) When microscopy is the only diagnostic test available, specimens should be sent to a facility with capacity to perform RMTs.

(b) For more information, see: Global Laboratory Initiative. Line probe assays for drug resistant tuberculosis detection Interpretation and reporting guide for laboratory staff and clinicians. [http://stoptb.org/wg/gli/assets/documents/LPA_test_web_ready.pdf](http://stoptb.org/wg/gli/assets/documents/LPA_test_web_ready.pdf)

(c) For more information:
- WHO catalogue of mutations in *M. tuberculosis* complex and their association with drug resistance: [https://www.who.int/publications/i/item/9789240028173](https://www.who.int/publications/i/item/9789240028173)
(d) Microscopic observation of drug susceptibility (MODS), nitrate reductase assay (NRA), thin layer agar and colorimetric redox indicator (CRI).

(e) LAM antigen is a component of the mycobacterial cell walls released by M. tuberculosis then excreted by the kidneys.

(f) Alere Determine® TB LAM Ag (Alere Inc, Waltham, MA, USA).

(g) Seriously ill: respiratory rate > 30/minute, temperature > 39 °C, heart rate > 120/minute and unable to walk unaided.

(h) For children > 5 years and adults: CD4 count < 200 cells/mm³ or a WHO clinical stage 3 or 4. All children < 5 years are considered as having advanced HIV disease.

(i) HIV-infected patients diagnosed with TB using the LF-LAM should be recorded as bacteriologically confirmed TB cases.

References


3.2 Latent tuberculosis infection

Diagnosis is based on exclusion of active TB and demonstration of latent tuberculosis infection (LTBI).

For demonstrating LTBI, one of the following tests may be performed. However, these tests are not mandatory prior to initiating LTBI treatment in:

- Children under 5 years household contact of a TB case;
- HIV-infected children and adults[1].

3.2.1 Tuberculin skin test
A positive tuberculin skin test (TST) indicates that a mycobacterial infection has occurred. For interpretation of TST results, see Appendix 9.

TST has several limitations:
- It does not distinguish infection by *M. tuberculosis* from exposure to environmental mycobacteria.
- It does not distinguish latent/active TB.
- Prior BCG vaccination can result in a false positive TST.
- False negative TST is common, particularly in HIV-infected patients and malnourished children.

After having ruled out active TB, a positive TST is an indication for treatment of LTBI (Chapter 16).

Notes:
- TST is also used to check the absence of TB in neonates on isoniazid monotherapy (Chapter 16).
- Other skin tests are available, but have not yet been evaluated by WHO.

### 3.2.2 Interferon gamma release assays

The test is performed in vitro on blood to which *M. tuberculosis* antigens are added. This results in the rapid stimulation of memory T cells and release of interferon-gamma in patients previously exposed to the bacillus.

The following tests measure:
- QuantIFERON-TB Gold In-Tube: the amount of interferon-gamma released.
- T-SPOT.TB test: the number of interferon-gamma producing T cells[2].

The advantage of IGRAs over TST is the absence of cross-reaction with BCG vaccine and most environmental mycobacteria.

IGRAs have some limitations:
- They do not distinguish latent/active TB.
- They are more complex than TST (equipment and trained laboratory technicians) and are not widely available.

A positive test indicates that LTBI is likely; a negative test indicates that it is unlikely. After having ruled out active TB, a positive IGRA is an indication for treatment of LTBI (Chapter 16).

### Referencias


### 3.3 Other investigations

In addition to TB diagnosis tests, the following investigations should be performed at baseline and during treatment. The purpose is to identify common comorbidities, contra-indications, as well as adverse effects of TB drugs.

*Table 3.10 – Other investigations in TB treatment*
<table>
<thead>
<tr>
<th>Tests</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrocardiogram (ECG)</td>
<td>Patients on QT prolonging drugs</td>
</tr>
<tr>
<td>Brief peripheral neuropathy screen (BPNS)</td>
<td>Patients on linezolid</td>
</tr>
<tr>
<td>Visual function tests</td>
<td>Patients on MDR/RR-TB treatment including ethambutol, linezolid or thionamides</td>
</tr>
<tr>
<td>Audiometry (b)</td>
<td>Patients on aminoglycosides</td>
</tr>
<tr>
<td>Full blood count</td>
<td>Patients on linezolid (or rifabutin)</td>
</tr>
<tr>
<td>Liver function tests</td>
<td>Patients with pre-existing hepatic disease</td>
</tr>
<tr>
<td>Aspartate aminotransferase (AST)</td>
<td>Patients on MDR/RR-TB treatment</td>
</tr>
<tr>
<td>Alanine aminotransferase (ALT)</td>
<td></td>
</tr>
<tr>
<td>Creatinine</td>
<td>Patients with pre-existing renal disease</td>
</tr>
<tr>
<td>Serum creatinine</td>
<td>Patients on aminoglycosides</td>
</tr>
<tr>
<td>Creatinine clearance</td>
<td></td>
</tr>
<tr>
<td>Serum electrolytes (potassium)</td>
<td>Patients on aminoglycosides</td>
</tr>
<tr>
<td>Glycated haemoglobin (HbA1c), or Blood glucose level (fasting or random)</td>
<td>All patients</td>
</tr>
<tr>
<td>HIV, hepatitis B and C testing</td>
<td>Patients with undocumented HIV, hepatitis B and C status</td>
</tr>
<tr>
<td>CD4 count and viral load</td>
<td>Patients with TB/HIV coinfection</td>
</tr>
<tr>
<td>Thyroid stimulating hormone (TSH)</td>
<td>Patients on thionamides or PAS</td>
</tr>
<tr>
<td>Pregnancy test</td>
<td>Patients of childbearing age with MDR/RR-TB</td>
</tr>
</tbody>
</table>

(a) BPNS is a clinical examination for detecting peripheral neuropathy and grading the severity of symptoms (Appendix 16).
(b) For children under 5 years, a specialized equipment and consultation are required.

For more information see Chapter 9, Chapter 10 and Chapter 11.

Chapter 4: Diagnostic algorithms for pulmonary tuberculosis (PTB) in adults and adolescents

4.1 Guiding principles for the use of the algorithms

4.2 Adult and adolescent algorithms
4.1 Guiding principles for the use of the algorithms

The aim of algorithms is to assist the diagnostic process and minimize incorrect diagnosis. The following algorithms are for adults and adolescents. For diagnostic algorithms for children < 10 years, see Chapter 5.

4.1.1 Clinical assessment

– An assessment for danger signs is the first part of the clinical assessment. The adult or adolescent is classified as seriously ill if one or more of the following danger signs are present:
  • Respiratory rate > 30/minute;
  • Fever > 39°C;
  • Pulse rate > 120/minute;
  • Unable to walk unaided.

– In cases where there is no bacteriological confirmation of TB, the clinical (and radiological) assessment should determine if the patient needs broad-spectrum antibiotics and/or anti-TB drugs.

– HIV testing should be routinely offered to all individuals suspected of having TB. If testing is refused or unavailable, it might be assumed that a certain patient is likely to be HIV-positive (according to context and/or clinical presentation). In this event, follow the algorithm for HIV-infected patients.

4.1.2 Clinical response

For patients who are treated empirically for bacterial pneumonia or pneumocystosis (PCP), a “non-response to antibiotics” increases the likelihood of TB. The converse is not necessarily true, such that a response to antibiotics does not automatically exclude TB in a person suspected of having TB, particularly if respiratory symptoms persist after treatment. Pneumonia or PCP may occur in patients with underlying TB.

Antibiotic treatment is appropriate for HIV-infected patients with cough because bacterial infections are common both with and without TB. All seriously ill patients being started on anti-TB treatment should also be treated empirically, with broad-spectrum antibiotics for bacterial pneumonia because benefits outweigh the risks.

Referencias


4.2 Adult and adolescent algorithms

Diagnostic algorithm 1

PTB in HIV-negative patients with low risk of MDR-TB
When the patient’s serological status is unknown, this algorithm should be used in settings with HIV prevalence < 5%.

Patients are considered to be at low risk of multidrug-resistant TB (MDR-TB) if they do not meet one of the following criteria: 1) resident in areas with high MDR-TB prevalence; 2) all retreatment categories; 3) exposure to a known MDR-TB case; 4) patient remaining smear + at 2 months; 5) exposure to institutions with high risk of MDR-TB (e.g. prisons).

Danger signs: respiratory rate > 30/min and/or fever > 39°C and/or pulse rate > 120/min and/or unable to walk.

Smear microscopy: two sputum examinations performed on the same day.

Broad spectrum ATB:
- If no danger signs: amoxicillin for 7 days (NO fluoroquinolones);
- If danger signs: parenteral ATB (e.g. ceftriaxone).

Clinical response to a broad spectrum antibiotic does not rule out TB. Patient should be informed to return for reassessment if symptoms recur.

According to setting:
Xpert MTB/RIF available: two sputum smear microscopy on the same day and one Xpert MTB/RIF from one of the samples collected for smear microscopy;

Xpert MTB/RIF not available: two sputum smear microscopy on the same day.

h: In groups of patients with high level of resistance to isoniazid (> 10%) it is recommended to perform a conventional DST at baseline (and/or a line probe assay) in order to provide adequate treatment.

According to setting:

- In groups of patients with prevalence of MDR-TB < 10%, patients seriously ill should immediately be initiated under empiric MDR-TB treatment. H and R will be included in the regimen until confirmation of MDR-TB by conventional methods. If the patient is stable, the clinician may choose to wait for confirmation before initiating a MDR treatment.
- In groups of patients with prevalence of MDR-TB ≥ 10%, patients should be initiated under empiric MDR-TB treatment. Consider adding H in settings where mono-resistance to R is not uncommon.

Clinical signs and chest X-ray (CXR) findings tend to be more typical in those who are HIV-negative having active TB:

<table>
<thead>
<tr>
<th></th>
<th>TB</th>
<th>Bacterial pneumonia</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clinical signs</strong></td>
<td>Weight loss, productive cough, purulent sputum, haemoptysis, pleuritic chest pain</td>
<td>• Acute onset</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Fever</td>
</tr>
<tr>
<td><strong>CXR</strong></td>
<td>• Infiltrates, nodules with or without cavitation in the upper lobes and in the superior segments of the lower lobes.</td>
<td>• Lobar consolidation</td>
</tr>
<tr>
<td></td>
<td>• Pleural effusions</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Adenopathy in the mediastinum or hila (rare in TB in adults and adolescents)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Miliary disease</td>
<td></td>
</tr>
</tbody>
</table>

When clinical signs AND CXR are strongly suggestive of active TB, treatment should be initiated without waiting for diagnosis confirmation.

**Diagnostic algorithm 2**

**PTB in HIV-positive patients**
When the patient's serological status is unknown, this algorithm should be used in settings with HIV prevalence > 5%.

TB suspect is defined as: cough for more than 2 weeks or any cough with at least one of the following signs: loss of weight, night sweats, fever, and suspicion based on clinical judgment.

Danger signs: respiratory rate > 30/min and/or fever > 39°C and/or pulse rate > 120/min and/or unable to walk.

According to setting:
- Xpert MTB/RIF available: two sputum smear microscopy on the same day AND one Xpert MTB/RIF from one of the samples collected for smear microscopy;
- Xpert MTB/RIF not available: two sputum smear microscopy on the same day.

In patients groups with high level of resistance to isoniazid (> 10%) it is recommended to perform a conventional DST at baseline (and/or a line probe assay) in order to provide adequate treatment.
When possible a culture should be performed. A positive culture result at any point in time in the algorithm should lead to a full TB treatment.

TB treatment should be started when clinical signs AND chest X-ray (CXR) are suggestive of TB (Note k).

Broad spectrum ATB/PCP:
- If no danger signs: amoxicillin for 7 days (or recommended oral agent for community-acquired pneumonia in the area). Do NOT use fluoroquinolones;
- If danger signs: parenteral ATB (e.g. ceftriaxone) AND high dose cotrimoxazole.

If no danger signs: patient should be re-assessed after 7 days.
If danger signs: patient should be assessed daily and if no response, TB treatment should be considered after 3 to 5 days.
Clinical response to broad-spectrum ATB does not rule out TB. Patient should be informed to return for reassessment if symptoms recur.

Differential diagnosis of a coughing HIV-infected adult/adolescent: bacterial (including atypical) pneumonia, PCP, fungal infection, non-tuberculous mycobacteria, nocardiosis, Kaposi sarcoma and lymphoma.

The diagnosis should be based on clinical assessment, CXR and CD4 results, whether cotrimoxazole preventive therapy (CPT) was used, and other treatment already used in the patient. If the index of suspicion for active TB is high, empiric TB treatment should be initiated without waiting for diagnosis confirmation. Other treatments such as broad-spectrum ATB or therapy for PCP may be needed in addition to TB treatment.

<table>
<thead>
<tr>
<th>Clinical signs</th>
<th>TB</th>
<th>PCP (HIV+)</th>
<th>Bacterial pneumonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current cough</td>
<td>• Dry cough</td>
<td>• Acute onset</td>
<td></td>
</tr>
<tr>
<td>Weight loss</td>
<td>• Dyspnoea ++</td>
<td>• High fever</td>
<td></td>
</tr>
<tr>
<td>Purulent sputum and haemoptysis less likely if HIV-positive with low CD4 count</td>
<td>• Hypoxemia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fever</td>
<td>• Not on CPT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Night sweats</td>
<td>• More likely if low CD4 count</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pleuritic chest pain</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CXR</th>
<th>TB</th>
<th>PCP (HIV+)</th>
<th>Bacterial pneumonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper lobe infiltrates and cavitation only likely in HIV-positive adults with higher CD4 counts. Any lobe of the lung may be affected</td>
<td>• Bilateral interstitial infiltrate with reticulonodular markings that are more pronounced in the lower lobes</td>
<td>• Lobar consolidation</td>
<td></td>
</tr>
<tr>
<td>In HIV-positive adults with lower CD4 counts, the following 4 patterns are suggestive of TB:</td>
<td>• Findings lag behind symptoms and may be normal early in the disease</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. miliary pattern</td>
<td>2. pleural effusion without airspace (with straw-coloured liquid aspirate)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. hilar and mediastinal adenopathy</td>
<td>4. large heart (especially if symmetrical and rounded)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In the absence of any improvement of clinical signs (no weight gain, persistent cough, pain, etc.) AND no improvement on CXR after 2 months of a well conducted TB treatment, diagnosis and treatment should be reconsidered. MDR-TB should also be considered.

In addition to the differential diagnosis in Note k above, DR-TB should be considered.

Immediately start empiric MDR treatment, even if positive predictive value of Xpert MTB/RIF for R resistance is low (this is done to avoid the rapid and high mortality due to untreated MDR-TB in HIV patients). H and R should be included in the regimen until confirmation of MDR-TB by conventional methods if the patient comes from a group with less than a prevalence of MDR-TB < 10%. In groups of patients with prevalence of MDR-TB ≥ 10%, patients should be initiated under an empiric MDR treatment without H or R, although one can consider adding H in settings where mono-resistance to R is not uncommon.
Diagnostic algorithm 3 with Xpert MTB/RIF

PTB in patients with high risk of MDR-TB

The following patients are considered to be at high risk of MDR-TB: 1) resident in areas with high MDR-TB prevalence; 2) all retreatment categories; 3) exposure to a known MDR-TB case; 4) patient remaining smear-positive at 2 months; 5) exposure to institutions with high risk of MDR-TB (e.g. prisons).

Groups of patients at risk of MDR-TB are also at risk of other types of DR-TB as well. DST to the first-line should be performed in order to provide adequate treatment for possible mono- or poly-drug resistance.

In populations with a prevalence < 10% of MDR-TB, the resistance to R diagnosed by Xpert MTB/RIF must be confirmed by conventional methods. Drug sensitivity testing (DST) to both first-line drugs and secondline TB drugs should be performed if possible.

In groups of patients with prevalence of MDR-TB < 10%, the decision to start the MDR-TB treatment will be made on clinical presentation of the patient and immunological status. Patients seriously ill and/or HIV+ should be initiated immediately under empiric MDR-TB treatment. H and R will be included in the regimen until confirmation by conventional methods.

In groups of patients with prevalence of MDR-TB ≥ 10%, the patient should be initiated using an empiric MDR-TB treatment. Consider adding H in settings where mono-resistance to R is not uncommon.

Baseline sputum smear microscopy result on 1 specimen in order to: 1) allow patient follow-up with microscopy; 2) take immediate decisions related to TB infection control.

Chapter 5: Diagnosis of tuberculosis in children

5.1 Background

5.2 Characteristics of tuberculosis in children

5.3 Diagnostic approach

5.4 Key elements of the diagnosis

5.5 Collecting sputum specimens in children
5.1 Background

Tuberculosis (TB) is a significant cause of morbidity and mortality in children in settings of high TB prevalence. The lack of an accurate diagnostic method has contributed to a gross underestimation of its true burden. In high prevalence settings, children < 15 years old are expected to make up 10 to 20% of all TB cases.

The characteristics of TB disease and approach to diagnosis in children > 10 years old are similar to those for adults. This section focuses on children < 10 years old.

Referencias


5.2 Characteristics of tuberculosis in children

Children have a high risk of TB infection. In endemic areas, children are likely to be exposed to TB in their household or community. A careful contact history is extremely important in children with signs and symptoms suggestive of TB. In children with diagnosed TB, an effort should be made to detect the source case and any other undiagnosed cases in the household. The younger the child, the more likely it is that a contact can be identified. Equally, all TB cases, especially children or adults who are smear-positive, should be asked about close contact with children (Chapter 16).

Children have a high risk of progression to active TB disease and of developing severe forms: 90% of young children, who develop TB, do so within 12 months of infection. Children < 3 years and children with immune suppression (e.g. HIV, malnutrition, post-measles) are particularly vulnerable. TB screening with prompt treatment or prophylaxis is especially critical in these children.

Most cases are pulmonary TB (PTB), but smear positivity is rare because children generally have low bacillary loads. Furthermore, sputum samples can be difficult to obtain from children. As a result, smear-positive TB represents only an estimated 10% of all TB observed in the 0 to 14 age group.

Extrapulmonary TB (EPTB) is common in children. The site of EPTB disease is age related. Miliary and meningeal TB is more frequently seen in young children. TB lymphadenitis and osteoarticular TB are more common in older children.

BCG administered at birth offers partial protection against severe forms in young children. It offers little if any protection against pulmonary TB. A history of BCG vaccination does not exclude the possibility of TB in a child with suggestive signs and symptoms.

Referencias

5.3 Diagnostic approach

For the majority of children, careful history, clinical assessment and follow up alone are sufficient to make a diagnosis of TB, even if confirmation is not possible.

Bacteriology, chest x-ray and tuberculin skin test are useful but not essential in most cases.

A trial of treatment with anti-TB drugs is not recommended as a method to diagnose TB. The decision to treat a child should be carefully considered. Once such a decision is made, the child should be treated with a full course of therapy.

5.4 Key elements of the diagnosis

Recommendations of how to combine the following elements to arrive at a treatment decision are summarized in the diagnostic algorithms (Section 5.6).

5.4.1 Careful history

- Contact with a known or presumed TB case:
  - Timing of the exposure: greater risk if exposure occurred in the past 12 months;
  - Closeness of the contact: greater risk if living in same household or sleeping in the same room;
  - Type of TB of the source case: greater risk if smear-positive or cavities on x-ray; resistance pattern of the source case.
- Symptoms suggestive of TB:
  - Cough persistent for > 2 weeks and not improving;
  - Unexplained fever for > 1 week;
  - Unexplained weight loss or failure to thrive;
  - Unexplained fatigue, lethargy or reduced playfulness.

5.4.2 Clinical examination

- Vital signs: fever and increased respiratory rate may be present.
- Growth: weigh the child and compare with previous records. Weight loss or flattening of the growth curve can signal chronic disease.
- Respiratory examination: abnormal auscultation or percussion may be present. Signs of severe respiratory infection: tachypnoea, cyanosis, hypoxemia (SpO$_2$ < 90%), nasal flaring, chest indrawing, grunting and feeding difficulties in infants.
- Physical signs of EPTB (see also Chapter 2):
  - Highly suggestive, e.g.:
    - Angular deformity of the spine;
    - Cervical lymph node with fistula formation.
  - Non specific requiring further investigation, e.g.:
    - Sub-acute meningitis not responding to antibiotic therapy;
    - Distended abdomen with ascites;
    - Lymphadenopathy without fistula formation;
    - Non-painful enlarged joint.
- Other: certain physical findings may point to alternative diagnoses (e.g. asthma) or relevant co-morbidities (e.g. HIV).

5.4.3 Re-assessment and follow up

The diagnosis is rarely made at the first consultation, as the initial clinical presentation is usually non specific. Follow up is critical to assess if signs and symptoms persist despite a trial of well-monitored non-TB antibiotic treatment.
Particularly suggestive of TB disease are:
- Persistent pneumonia after appropriate, well-monitored antibiotic treatment;
- Measured or reliably reported fever of > 38 °C for > 1 week, after common causes such as malaria or pneumonia have been excluded;
- No weight gain despite appropriate nutritional support;
- Persistent or worsening fatigue.

### 5.4.4 HIV testing

HIV testing should be routinely offered to children with presumed or diagnosed TB.

### 5.4.5 Diagnostic investigations

The following investigations should be performed in children suspected of TB whenever possible. The unavailability of a test due to resource limitations should not delay the diagnosis of TB.

**Tuberculin skin test (TST)**
- A positive test can support a diagnosis of TB in a symptomatic child;
- A negative TST does not exclude TB;
- Causes of false positive: BCG, atypical mycobacteria (NTM);
- Causes of false negative: HIV infection, malnutrition.

**Chest x-ray**

Chest x-ray can be helpful for the diagnosis of intrathoracic TB[^1]. Hilar lymphadenopathy is the most common finding. However, obtaining quality films in children and accurately interpreting them may be difficult. This limits their utility in many settings.

**Bacteriology**

Try to confirm TB, although treatment should not be delayed if clinically indicated.

For EPTB, obtain specimens from the suspected sites for microscopy and, when possible, for culture, cytology or histopathological examination and molecular methods (e.g. Xpert MTB/RIF).

Bacterial yields are higher in older children, and in children of all ages with severe disease. Two sputum specimens should be obtained: an on-the-spot specimen (at first evaluation), and an early morning specimen. Alternatively, two specimens collected one hour apart are an acceptable option (see Appendix 3).

Xpert MTB/RIF is the initial test of choice in screening for multidrug-resistant TB (MDR-TB). When Xpert MTB/RIF is not available, conventional drug susceptibility test (DST) can be done. DST indications are the same as for adults.

### Referencias


### 5.5 Collecting sputum specimens in children
Given the importance of trying to obtain confirmation of disease in areas of high drug-resistant TB prevalence or in contact cases of MDR-TB, optimizing the collection of appropriate specimens is critical.

Children < 6 years old, and some as old as 10 years old, may be unable or unwilling to spontaneously expectorate. Explanation and encouragement are important. Chest clapping is a simple, yet, often effective measure to help expectoration.

If these measures fail, sputum specimen can be obtained by sputum induction or gastric aspiration (Appendix 3). Given the distress caused to the child and the generally low yield on smear microscopy, these procedures should only be done if culture or Xpert MTB/RIF is available.

**5.6 Paediatric diagnostic algorithms**

**Paediatric diagnostic algorithm 1**

**Contact of a TB case**

---

**a.** Contact: child living in the same household or in close and regular contact with any known or suspected TB case in the last 12 months.

**b.** Malnutrition or growth curve flattening.

**c.** Clinical assessment (including growth assessment), bacteriological tests, HIV testing (in high HIV prevalence areas), and when relevant and available: X-ray (CXR), investigations for EPTB, TST.

**d.** Examples of “obvious TB” may include cases of Pott’s disease, TB meningitis, lymph node TB with fistula formation, smear or Xpert MTB/RIF positive or highly suggestive chest X-ray (e.g. hilar lymphadenopathy, upper lobe infiltrates, miliary picture).

**e.** Broad spectrum ATB:

- If no danger signs: amoxicillin PO for 7 days;
Clinical response to a broad-spectrum antibiotic does not rule out TB. Carer should be informed to consult if symptoms re-occur.

Paediatric diagnostic algorithm 2

Symptomatic child

Day 1

Cough > 2 weeks or poor weight gain\(^a\) or fever\(^b\) > 1 week or suspicion of EPTB

Clinical assessment and other investigations\(^c\)

Antibiotics\(^d\), nutritional support or other treatment according to clinical findings

If obvious TB\(^d\), start TB treatment

Clinical assessment\(^d\) and other investigations after 1 week

Is the child still symptomatic?

NO

Day 7

Is the child HIV exposed or infected or is a contact of a TB case?

NO

Antibiotics\(^d\), nutritional support or other treatment according to clinical findings for one week

YES

Clinical assessment\(^d\):
- Poor weight gain
- Persistent cough
- Persistent fever
- Fatigue or lethargy
- CXR suggestive of TB

Day 7-12

None present\(^f\)

One present

≥ 2 present

TB unlikely

Start TB treatment

TB treatment in particular if child is HIV+ or < 3 years or presents severe malnutrition, or TST+.

---

\(a\). Malnutrition or growth curve flattening.

\(b\). Temperature > 38°C.
Clinical assessment (including growth assessment), bacteriological tests, HIV testing (in high HIV prevalence areas), and when relevant and available: X-ray (CXR), investigations for EPTB, TST.

- Smear microscopy positive or Xpert MTB/RIF positive, CXR showing suggestive lesions (e.g. hilar lymphadenopathy, upper lobe infiltrates, miliary picture), gibbus.

- Broad spectrum antibiotics:
  - If no signs of severity:
    - first-line: amoxicillin PO for 7 days (NO fluoroquinolones). Advise carer to return with the child if no improvement after 48 hours of antibiotics;
    - if a second course of antibiotic if needed: azithromycin PO for 5 days.
  - If signs of severity: parenteral antibiotics (ceftriaxone ± cloxacillin if *S. aureus* is suspected).

In addition: PCP treatment should be given presumptively to all HIV-exposed or HIV-infected children < 1 year of age, and any older child with severe immune suppression and not on CTX prophylaxis. For all other HIV-exposed or HIV-infected children, it should be considered if there is poor response to broad spectrum antibiotics after 48 hours.

Clinical response to a broad-spectrum antibiotic does not rule out TB. Carer should be informed to consult if symptoms re-occur.

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**Chapter 6: Intensive case finding in HIV-infected individuals**

**6.1 Routine screening**

**6.2 Purposes of screening**

---

**6.1 Routine screening**

Intensive case-finding (ICF) should be in place in all projects providing care to HIV-infected patients. Screening can be performed at multiple points in time by different levels of health care workers (e.g. counsellors during HIV testing, health care providers during clinical consultations).

All children and adults should be regularly screened for TB using the following criteria:

**Table 6.1 - Screening criteria/symptoms in children and adults[^1]**

<table>
<thead>
<tr>
<th>Children</th>
<th>Adults</th>
</tr>
</thead>
<tbody>
<tr>
<td>![current_cough][a]</td>
<td>![current_cough][a]</td>
</tr>
<tr>
<td>Fever</td>
<td>Fever</td>
</tr>
<tr>
<td>![poor_weight_gain][b]</td>
<td>Weight loss</td>
</tr>
<tr>
<td>Contact with a contagious person</td>
<td>Night sweats</td>
</tr>
</tbody>
</table>

[^1]: (a) Asking about “current cough”, rather than cough for 2 weeks, is more sensitive for TB disease in HIV-infected individuals
(b) Poor weight gain is defined as reported weight loss or underweight or confirmed weight loss > 5% since last visit, or growth curve flattening.
6.2 Purposes of screening

6.2.1 Early detection and treatment of active TB

Children and adults found to have one or more of the above symptoms/criteria during screening may have active TB, and they should be evaluated with an appropriate TB diagnostic algorithm in order to rapidly diagnose those who do have TB (see Diagnostic algorithms, Chapter 4 and Chapter 5).

6.2.2 Identification of patients eligible for isoniazid preventive therapy (IPT)

The significant proportion of asymptomatic active TB described in patients eligible for ART (15 to 20%) leads to use this screening method with caution. It should only be used for ruling out TB in patients not yet eligible for antiretroviral therapy (CD4 greater than 350 and no WHO stage 3 or 4 illnesses) or after three months of treatment in patients started under antiretroviral therapy in order to allow for possible unmasking of TB.

For IPT, see Chapter 16.

Referencias


Chapter 7: Case definitions for registration

7.1 Definition of a tuberculosis case

7.2 History of prior anti-TB treatment

7.3 Anatomical site of the disease

7.4 Bacteriological status

7.5 HIV status

7.6 Other co-morbidities

7.7 Summary of patient registration
7.1 Definition of a tuberculosis case

A tuberculosis (TB) case is a patient that has been diagnosed as such by a clinician, regardless if the diagnosis has been confirmed bacteriologically or not.

The elements necessary for defining a TB case are: the TB treatment history, the bacteriological status, the anatomical site of the disease and the patient’s HIV status.

**Note:** any person receiving treatment for TB should be recorded as a TB case.

7.2 History of prior anti-TB treatment

Patients who have interrupted or failed a previous anti-TB treatment have a higher risk of developing drug-resistance (DR). Therefore, it is important to question patients about their previous treatment prior to treatment initiation.

Case registration distinguishes between [1]:

- **New patients:** patients who have never been treated for TB or have taken anti-TB drugs for less than 1 month.
- **Previously treated patients:** patients who have received 1 month or more of anti-TB drugs in the past.
  - Previously treated patients are further sub-classified into relapse, failure and return after treatment interruption:
    - **Relapse:** patients who were cured or completed treatment on their last TB treatment;
    - **Failure:** patients who have failed their most recent treatment (see Chapter 17 for outcome definitions for failure);
    - **Treatment interruption:** patients who interrupted (see Chapter 17 for outcome definition of treatment interruption) their last treatment should be classified as “Return after treatment interruption”.
- **Others:** patients who cannot be included in one of the above categories (e.g. patients who have previously been treated via an erratic or unknown TB regimen).

**Referencias**


7.3 Anatomical site of the disease

**Pulmonary TB (PTB)**

Refers to a case of TB presenting with involvement of the lung parenchyma.

**Notes:**
- Miliary TB is also classified as PTB because there are lesions in the lungs.
- Any patient presenting with PTB and an EPTB form at the same time is classified as a PTB case for recording purposes.
Extrapulmonary TB (EPTB)

Refers to a case of TB involving organs other than the lungs. Diagnosis is based on clinical signs corresponding to extrapulmonary active TB and a decision by a clinician to treat with a full course of anti-TB drugs.

Notes:
- Sputum smear microscopy should always be done, and if possible culture and/or molecular test.
- Patients presenting with tuberculous pleural effusion, or mediastinal lymphadenopathy without evidence of parenchymal localization are classified in this category.

Notas

(a) If possible, obtain histological or bacteriological evidence (microscopy, culture or molecular test).

7.4 Bacteriological status

Bacteriological status refers to the detection of *M. tuberculosis* by smear, culture or molecular methods. The bacteriological status can be further sub-classified on the basis of drug sensitive and drug resistant cases.

7.4.1 Detection of *M. tuberculosis*

Every TB case should be classified into one of two categories:
- **Confirmed TB case**: a case with a positive bacteriological result (microscopy, culture or molecular method).
- **Non-confirmed TB case**: a case where investigations are negative (microscopy, culture or molecular method) and for whom a clinician prescribes anti-TB treatment.

Confirmed TB cases are further sub-classified as:
1. smear positive/negative/not done
2. culture positive/negative/not done
3. molecular test positive/negative/not done

7.4.2 Strain sensitivity/resistance

When possible, culture and DST should be done to determine if the strain presents resistance to some drugs:

- **Susceptible TB**: the strain is not resistant to any first-line anti-TB drugs.
- **Drug-resistant TB**:
  - **Monodrug-resistant TB**: resistance to one first-line anti-TB drug only;
  - **Polydrug-resistant TB** (PDR-TB): resistance to more than one first-line anti-TB drug, other than isoniazid and rifampicin;
  - **Multidrug-resistant TB** (MDR-TB): resistance to at least isoniazid and rifampicin;
  - **Extensively drug-resistant TB** (XDR-TB): MDR-TB resistant to at least one fluoroquinolone and at least one second-line injectable drug (Km, Amk, Cm).

Patients with DR-TB should be classified in the following manner:

- **Confirmed isoniazid resistance and rifampicin susceptible**: resistance to isoniazid but not rifampicin. Resistance to first and second-line anti-TB drugs may be present.
- **Confirmed rifampicin resistant TB** (RR-TB): resistance to rifampicin confirmed by phenotypic drug susceptibility test or line probe assay or Xpert MTB/RIF (isoniazid susceptible or unknown).
- **Confirmed MDR-TB**: resistance to isoniazid and rifampicin, with or without resistance to first and second-line anti-TB drugs.
- **Confirmed XDR-TB**: resistance to isoniazid and rifampicin, and to at least one fluoroquinolone, and one second-line injectable drug (Km, Amk, Cm).
- **Unconfirmed DR-TB**: patients treated as DR-TB but without DST results (e.g. children who are contacts of a known case, patients with clinical failure and for whom no DST was available for some reason).
7.5 HIV status

Determining and recording the patient’s HIV status is critical for treatment decisions, as well as for assessing programme performances. The TB treatment card and TB register, which should be treated as confidential documents, should include: dates and results of HIV tests, starting date of cotrimoxazole and antiretroviral therapy.

7.6 Other co-morbidities

Any other significant diseases, such as diabetes, hepatitis B or C, cancer and malnutrition, should be noted at registration.

7.7 Summary of patient registration

Table 7.1 summarizes the elements necessary for defining a TB case.

Table 7.1 - Patient registration by outcome of most recent TB treatment[1]

<table>
<thead>
<tr>
<th>Registration groups based on treatment history</th>
<th>Further classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>New</td>
<td></td>
</tr>
<tr>
<td>Relapse</td>
<td>1. PTB or EPTB? If EPTB, indicate site.</td>
</tr>
<tr>
<td>Failure</td>
<td>2. Bacteriologically confirmed or non-confirmed TB case?</td>
</tr>
<tr>
<td>Treatment interruption</td>
<td>3. Sub-category of bacteriological status:</td>
</tr>
<tr>
<td></td>
<td>▪ Smear positive/negative/not done</td>
</tr>
<tr>
<td></td>
<td>▪ Culture positive/negative/not done</td>
</tr>
<tr>
<td></td>
<td>▪ Molecular test positive/negative/not done</td>
</tr>
<tr>
<td>Previously treated</td>
<td>4. If previously treated:</td>
</tr>
<tr>
<td>Treatment interruption</td>
<td>▪ Document last regimen received</td>
</tr>
<tr>
<td></td>
<td>▪ History of second-line drug use</td>
</tr>
<tr>
<td>Other</td>
<td>5. DST pattern: susceptible to H and R, confirmed H resistance and R susceptible, RR-TB, MDR-TB or XDR-TB</td>
</tr>
<tr>
<td></td>
<td>6. HIV status (negative/positive/not done)</td>
</tr>
<tr>
<td></td>
<td>7. Other co-morbidities?</td>
</tr>
</tbody>
</table>

Referencias

Chapter 8: Tuberculosis drugs and treatment regimens

8.1 Introduction

A combination of several antituberculosis drugs is needed to treat tuberculosis (TB) and prevent the emergence of resistance. Each TB drug has a specific action on one or more bacillary populations, but none on dormant bacilli.

TB drugs are classified into two categories:
- Drugs for drug-susceptible TB (DS-TB), also referred to as “first-line TB drugs”.
- Drugs for drug-resistant TB (DR-TB), also referred to as “second-line TB drugs”. WHO has further classified DR-TB drugs in groups based on their effectiveness and safety profile.

Treatment regimens define the TB drug combinations used and the intended duration of TB treatment.

For more information on the TB drugs, see Appendix 10.

8.2 Standard code for treatment regimens

8.2.1 Tuberculosis drugs

Each TB drug has an abbreviation.

Table 8.1 - Categories and abbreviations of TB drugs
### 8.2.2 Treatment regimens

TB treatment regimens are expressed as follows:

- **Drugs** are designated by their abbreviation.
- For some regimens, the treatment is divided into two phases: initial (or intensive) phase, and continuation phase. The phases are separated by a slash `/`.
- **Letters in brackets ()** indicate fixed-dose combinations (FDCs).
- **Letters that are not in brackets** indicate individual drugs.
- **Second-line drugs** are separated by a hyphen.
- **Letters in square brackets [ ]** indicate that drugs are used, but not considered as likely effective (Chapter 10).
- A superscript h (h) indicates that the drug is administered in a high dose.
- **Numbers before letters** indicate the duration (in months) of the treatment or of each phase.
- **Numbers in subscript and angle brackets < >** after a drug indicate the duration (in months) of the treatment with this drug.

**Box 8.1 – Examples**

<table>
<thead>
<tr>
<th>Categories</th>
<th>TB drugs</th>
<th>Abbreviations</th>
</tr>
</thead>
</table>
| **Drug-susceptible TB** (first-line drugs) | Isoniazid (standard dose)  
Rifampicin  
Pyrazinamide  
Ethambutol  
Rifabutin  
Rifapentine | H  
R  
Z  
E  
Rfb  
P |
| **Drug-resistant TB** (second-line drugs) |  
**Group A**  
Levofloxacin or moxifloxacin  
Bedaquiline  
Linezolid | Lfx or Mfx  
Bdq  
Lzd |
| **Group B** | Clofazimine  
Cycloserine or terizidone | Cfz  
Cs or Trd |
| **Group C** | Delamanid  
Ethambutol  
Pyrazinamide  
Imipenem/cilastatin or meropenem  
Amikacin or streptomycin  
Ethionamide or prothionamide  
Para-aminosalicylate sodium or para-aminosalicylic acid  
Isoniazid (high-dose) | Dlm  
E  
Z  
Ipm/Cln or Mpm  
Am or S  
Eto or Pto  
PAS  
H<sub>h</sub> |
| **Ungrouped** | Pretomanid | Pa |

**Notes:**

- High-dose isoniazid, although not a Group C drug according to the WHO classification, is considered in this guide as a Group C drug as it is used as such when building a treatment regimen for DR-TB.
- Pretomanid is not categorized in the WHO classification and is only used in standard treatment regimens for DR-TB (Chapter 10).
8.3 Drugs for drug-susceptible tuberculosis

All drugs used for DS-TB treatment are taken 7 days a week.

8.3.1 First-line drugs

Table 8.2 – Main characteristics of first-line TB drugs

<table>
<thead>
<tr>
<th>TB drugs</th>
<th>Activity</th>
<th>Resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid</td>
<td>Bactericidal</td>
<td>• High level of resistance in some regions. • Cross-resistance with thionamides.</td>
</tr>
<tr>
<td>Rifampicin</td>
<td>Bactericidal</td>
<td>• High level of resistance to rifampicin in some regions. • High level of cross-resistance between rifamycins.</td>
</tr>
<tr>
<td>Rifabutin</td>
<td>Bactericidal</td>
<td></td>
</tr>
<tr>
<td>Rifapentine</td>
<td>Bacteriostatic</td>
<td>Unknown (no reliable drug susceptibility test for ethambutol).</td>
</tr>
<tr>
<td>Ethambutol</td>
<td>Bacteriostatic</td>
<td></td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>Weakly bactericidal</td>
<td>High level of resistance in regions where rifampicin resistance is frequent.</td>
</tr>
</tbody>
</table>

Isoniazid

Isoniazid is usually well tolerated at recommended doses. It may cause peripheral neuropathy, hepatotoxicity, and hypersensitivity reactions. Peripheral neuropathy can be prevented by administration of pyridoxine (vitamin B₆). See Appendix 17.

Rifamycins (rifampicin, rifabutin, rifapentine)

Rifamycins are usually well tolerated at recommended doses. They may cause hypersensitivity reactions, hepatotoxicity, and thrombocytopenia. They are strong inducers of cytochrome P450 and can affect the plasma concentrations of many drugs (Appendix 19). Rifampicin is the most used rifamycin in the treatment of DS-TB. Rifabutin is used instead of rifampicin in patients taking certain antiretrovirals (Appendix 19). Rifapentine is only used in the 4-month regimen 2HPZ-Mfx/2HP-Mfx.

Note: rifampicin and rifapentine are also used to treat latent TB infection (Chapter 16).

Ethambutol

Ethambutol is usually well tolerated, including in children, particularly with respect to ocular toxicity. Ocular toxicity is dose- and duration-dependent. It is uncommon when ethambutol is used at the recommended dose for 2 months.

Pyrazinamide
Pyrazinamide is usually well tolerated however, it may cause hepatotoxicity, gout, arthralgias and photosensitivity.

### 8.3.2 Other drugs

Two second-line drugs are also used in the treatment of DS-TB: moxifloxacin (Section 8.4.1) and ethionamide (Section 8.4.3).

### Referencias


### 8.4 Drugs for drug-resistant tuberculosis

Drugs used for DR-TB treatment (except bedaquiline) are taken 7 days a week.

#### 8.4.1 Group A drugs

*Table 8.3 – Main characteristics of TB drugs Group A*

<table>
<thead>
<tr>
<th>TB drugs</th>
<th>Classes</th>
<th>Activity</th>
<th>Resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levofloxacin</td>
<td>Fluoroquinolones (FQs)</td>
<td>Bactericidal</td>
<td>• Resistance common in some regions.</td>
</tr>
<tr>
<td>Moxifloxacin</td>
<td></td>
<td></td>
<td>• Cross-resistance between FQs.</td>
</tr>
<tr>
<td>Bedaquiline</td>
<td>Diarylquinolines</td>
<td>Bactericidal</td>
<td>• Partial cross-resistance with Cfz.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Growing resistance as use increases.</td>
</tr>
<tr>
<td>Linezolid</td>
<td>Oxazolidinones</td>
<td>Bactericidal</td>
<td>Resistance assumed to be rare due to its limited use.</td>
</tr>
</tbody>
</table>

**Fluoroquinolones (levofloxacin, moxifloxacin)**

FQs are usually well tolerated. They may cause tendinopathy and QT prolongation. Moxifloxacin is sometimes used at high dose (Mfx³) in the presence of low-level resistance to FQs.

**Bedaquiline**

Bedaquiline is usually well tolerated. It may cause hepatotoxicity and QT prolongation. Bedaquiline has a long half-life (5.5 months). Therefore, adverse effects can persist after the drug is stopped, and if TB is still active, resistance can develop. Bedaquiline is metabolized in the liver by the cytochrome P450 (CYP450) system enzymes. Drugs, which induce or inhibit CYP450, can affect bedaquiline plasma concentrations and should be avoided (Appendix 19). The extent of cross-resistance bedaquiline/clofazimine and the clinical implications are not fully understood[1][2][3].

**Linezolid**

Linezolid may cause myelosuppression, dose- and duration-dependent neuropathy and lactic acidosis.
Pyridoxine supplementation (vitamin B₆) is recommended for all patients on linezolid, although there is no evidence that pyridoxine can prevent linezolid-induced neuropathy. Adverse effects frequently lead to reducing the dose or discontinuing linezolid. The optimal dose and duration of treatment are not established.

Linezolid has many interactions and overlapping toxicities with other drugs (e.g. risk of serotonin syndrome when administered with serotonergic drugs[4]). However, it is not always possible to avoid concomitant use of these drugs (e.g. even on linezolid, a patient with depression may require an antidepressant).

### 8.4.2 Group B drugs

Table 8.4 – Main characteristics of TB drugs Group B

<table>
<thead>
<tr>
<th>TB drugs</th>
<th>Classes</th>
<th>Activity</th>
<th>Resistance</th>
</tr>
</thead>
</table>
| Clofazimine       | Riminophenazine (anti-leprosy drug)    | Probably bacteriostatic | • Partial cross-resistance with Bdq.  
|                   |                                       |                | • Growing resistance as use increases.                                      |
| Cycloserine/Terizidone  | Analogue of D-alanine                  | Bacteriostatic | • Resistance common in areas where it has been used extensively.  
|                   |                                       |                | • Full cross-resistance between the 2 drugs.                                |

**Clofazimine**

Clofazimine is a QT-prolonging drug. Orange-pink to brownish-black discoloration of the skin and body fluids occur in almost all patients. These changes are reversible and not harmful.

Clofazimine has a long half-life (approximately 70 days). Consequently, its adverse effects can persist for several weeks or months after the drug is stopped.

**Cycloserine or terizidone**

Cycloserine and terizidone are structural analogues used at the same dose. Both drugs may cause neurotoxicity including psychiatric adverse events.

To prevent neurotoxicity, pyridoxine (vitamin B₆) should be administered along with these drugs throughout the course of treatment ([Appendix 17](#)).

### 8.4.3 Group C drugs

Table 8.5 – Main characteristics of TB drugs Group C
**Table of TB drugs**

<table>
<thead>
<tr>
<th></th>
<th>Classes</th>
<th>Activity</th>
<th>Resistance</th>
</tr>
</thead>
</table>
| Delamanid        | Nitroimidazooxazines | Bactericidal | • Potential cross-resistance with pretomanid.  
• Resistance assumed to be rare due to its limited use. |
| Ethambutol       |               | Bacteriostatic | High prevalence of resistance among MDR/RR-TB patients (> 49% in some settings\(^6\)[\(^8\)]. |
| Pyrazinamide     |               | Bactericidal | High prevalence among MDR/RR-TB patients (> 80% in some areas\(^7\)[\(^8\)]. |
| Imipenem/cilastatin Meropenem | Carbapenems |               | Full cross-resistance between carbapenems. |
| Amikacin Streptomycin | Aminoglycosides | Bactericidal | Partial cross-resistance between the 2 drugs. |
| Ethionamide Prothionamide | Thionamides | Weak bacteriostatic | • Full cross-resistance between thionamides.  
• Cross-resistance with isoniazid if inhA mutation present.  
• High prevalence of resistance among MDR-TB patients in some areas\(^9\). |
| Para-aminosalicylate sodium Paraaminosalicylic acid | Weak bacteriostatic |               | Common in some regions. |
| Isoniazid high-dose |               |               | Cross-resistance with thionamides if inhA mutation present. |

**Delamanid**

Delamanid is usually well tolerated.  
It may cause QT prolongation\(^10\).  
It is particularly useful in patients with pre-existing hepatic disease (no reported hepatotoxicity) or HIV infection (no significant drug interactions or overlapping toxicities with antiretrovirals).  
It is also useful for replacing a Group A or B drug causing toxicity.

**Ethambutol**

See Section 8.3.1. Vision monitoring is required when ethambutol is administered for more than 2 months (risk of optic neuritis).

**Pyrazinamide**

See Section 8.3.1.

**Carbapenems (imipenem/cilastatin, meropenem)**

Imipenem is always combined with cilastatin. Cilastatin has no antibacterial activity, its role is to inhibit a renal enzyme that inactivates imipenem.  
Meropenem does not need to be combined with cilastatin, as it is metabolised through a different pathway.  
High cost and difficulty with administration limits the use of carbapenems.  
Carbapenems may cause gastrointestinal disturbances, neurotoxicity and hypersensitivity reactions.  
Meropenem should be used in children and adolescents under 15 years, and if possible, in epileptic patients and patients with TB meningitis (risk of seizures lower than with imipenem/cilastatin).  
The first dose is always administered in a health facility so that an eventual hypersensitivity reaction can be managed. If conditions permit, carbapenems can be continued as an outpatient.
Amoxicillin/clavulanic acid is routinely administered prior to carbapenems, as clavulanic acid prevents the development of carbapenem resistance.

**Aminoglycosides (amikacin, streptomycin)**

Aminoglycosides should only be used when no alternative is available. Most DR-TB patients can be treated without aminoglycosides, including some cases of extensively drug-resistant TB (XDR-TB).

Aminoglycosides are nephrotoxic and ototoxic drugs. Streptomycin is less nephrotoxic than other aminoglycosides, but causes vestibular toxicity more frequently. If an aminoglycoside is used, close monitoring is essential (audiometry, electrolytes and renal function). If close monitoring cannot be ensured, aminoglycosides should not be used.

**Note:** kanamycin and capreomycin are no longer recommended, as their use is associated with higher rates of treatment failure and death.

**Thionamides (ethionamide, prothionamide)**

Ethionamide and prothionamide are used at the same dose.

They may cause gastrointestinal disturbances, hypothyroidism (especially if co-administered with paraaminosalicylic acid), neuropathy and hepatotoxicity.

In diabetic patients, the dose of antidiabetics may need to be adjusted.

**Para-aminosalicylate sodium or para-aminosalicylic acid**

PAS often causes gastrointestinal disturbances and can decrease the absorption of other TB drugs.

It may also cause hypothyroidism, especially when co-administered with a thionamide.

**High-dose isoniazid**

See Section 8.3.1. There is limited evidence to support the use of high-dose isoniazid.

High-dose isoniazid may cause more adverse effects than the standard dose.

It has overlapping toxicity with linezolid (neuropathy) and hepatotoxic drugs.

To prevent peripheral neuropathy, pyridoxine (vitamin B₆) should be administered to all patients throughout the course of treatment (Appendix 17).

### 8.4.4 Ungrouped drugs

**Pretomanid**

Pretomanid belongs to the same class as delamanid and has bactericidal activity.

It is used only as part of standard regimens for DR-TB in the following combinations: BPaLM, BPaL (and BPaLC in operational research conditions only), see Chapter 10.

Regimens that include bedaquiline, pretomanid and linezolid may cause hepatotoxicity, lactic acidosis, myelosuppression, neuropathy and QT prolongation.

Pretomanid/delamanid cross-resistance is likely.

### 8.4.5 Other drugs

**Amoxicillin/clavulanic acid**

Amoxicillin/clavulanic acid is administered before each dose of carbapenem.

The clavulanic acid component prevents the development of carbapenem resistance.

Only formulations with a ratio of 4:1 (e.g. 500/125 mg) or 2:1 (e.g. 250/125 mg) are suitable for this indication. Do not use formulations with a ratio of 8:1 or 7:1.

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**Referencias**

8.5 Tuberculosis drug formulations

Only quality-assured drugs should be used. Several internationally recognized mechanisms ensure the quality of TB drugs a, b. 

8.5.1 Fixed-dose combinations

FDC formulations combine several TB drugs (2, 3 or 4) in the same tablet. FDCs are only available for first-line TB drugs. FDCs improve adherence (decreased pill burden, decreased risk of omission of one or more drugs).

Table 8.6 – Quality-assured FDC formulations

---


### 8.5.2 Individual drugs

Quality-assured single drug formulations are available for all first-line TB drugs. It may be necessary to use them when FDCs cannot be used due to adverse effects or drug interactions. There are no quality-assured FDCs for second-line TB drugs. The treatment of DR-TB is based on a combination of individual drugs.

### 8.5.3 Paediatric formulations

Paediatric formulations should be used whenever possible. However, they are not available for all TB drugs. When the only option is to manipulate the adult formulations:

- Preferably use scored tablets.
- Ensure that tablets/capsules can be split, crushed or opened (e.g. active ingredients may be protected from gastric acidity by an enteric coating).
- If tablets must be crushed (or capsules opened), a fraction of the powder corresponding to the required dose is mixed with food or liquids. Such manipulations should be done immediately before administering the drug. Any remaining powder should be discarded.
- The preparation of extemporaneous formulations using adult formulations is an alternative, however, this can only be considered if there are qualified personnel to ensure preparation in compliance with the appropriate compounding procedures.

### Notes

(a) Quality assurance:
- WHO Prequalification Scheme: [http://apps.who.int/prequal/](http://apps.who.int/prequal/)
- Stringent Regulatory Authorities (SRA): [https://www.who.int/initiatives/who-listed-authority-reg-authorities/SRAs](https://www.who.int/initiatives/who-listed-authority-reg-authorities/SRAs)

(b) Supply:

### Chapter 9: Treatment of drug-susceptible tuberculosis
9.1 Introduction

Drug-susceptible tuberculosis (DS-TB) treatment is indicated:
- When susceptibility to rifampicin and isoniazid is confirmed by drug susceptibility testing (DST), or
- If the probability of resistance to rifampicin and isoniazid is low:
  - while waiting for DST results for rifampicin and/or isoniazid,
  - when susceptibility to rifampicin is confirmed and susceptibility to isoniazid cannot be tested.

The probability of resistance is considered low in the following situations:
- No previous TB treatment;
- No contact with a drug-resistant TB (DR-TB) patient;
- The patient comes from an area of low prevalence of resistance according to drug resistance surveys.

Patients with DS-TB should start a conventional regimen based on first-line drugs (Table 9.1) or, if eligible, an alternative regimen (Table 9.2).

All regimens for DS-TB are standard regimens.

For dosages of fixed-dose combinations see Appendix 13.
For dosages of individual drugs see Appendix 10.

9.2 Conventional treatment regimens

Table 9.1 – Conventional DS-TB regimens according to the infection site
If bacteriological testing and/or CXR are not available, children meeting the following criteria are eligible for the 4-month regimen 2(HRZE)/2(HR):

- **Pulmonary TB (PTB)**
  - microscopy smear-negative or Xpert result “negative”, “trace”, “very low” and “low” or
  - clinically diagnosed with TB lesions confined to one lobe and no cavities on chest x-ray (CXR)

- **Extrapulmonary TB (EPTB) non severe, i.e.**:
  - pleural effusion without complications (e.g. no empyema, pneumothorax or fistula)
  - extra- or intra-thoracic lymph node TB with no airway obstruction

If after one month of treatment symptoms have completely resolved, continue treatment until the end. If symptoms have not completely resolved, further investigations are needed.

If after 4 months of treatment symptoms have not completely resolved and/or there is no weight gain, further investigation is needed. The treatment can be extended to 6 months if causes of non-response to treatment (including DR-TB, non-adherence and non-TB disease) are ruled out or unlikely.

Ethambutol can be removed from the 4- and 6-month regimens in non-HIV-infected children living in areas where the prevalence of HIV and/or isoniazid resistance is low with:

- **PTB and EPTB (except miliary TB, TB meningitis and bone and joint TB)**

For spinal TB, rest and back support bracing are indicated in addition to drug therapy. For patients with neurological deficit or unstable spine lesion, surgery can also be considered.

### Regimens and Duration

<table>
<thead>
<tr>
<th>Regimens</th>
<th>Eligibility</th>
</tr>
</thead>
</table>
| 2(HRZE)/2(HR)       | Children > 3 months and adolescents < 16 years with:
| 4 months            | Pulmonary TB (PTB)                                                          |
|                     | • microscopy smear-negative or Xpert result “negative”, “trace”, “very low” |
|                     | • clinically diagnosed with TB lesions confined to one lobe and no cavities |
|                     | Extrapulmonary TB (EPTB) non severe, i.e.:                                  |
|                     | • pleural effusion without complications (e.g. no empyema, pneumothorax     |
|                     | • extra- or intra-thoracic lymph node TB with no airway obstruction         |
| 2(HRZE)/4(HR)       | PTB and EPTB (except miliary TB, TB meningitis and bone and joint TB)       |
| 6 months            | Adolescents ≥ 16 years and adults                                           |
|                     | Children and adolescents < 16 years not eligible for the 4-month regimen    |
|                     | or when the national protocol does not include the 4-month regimen.         |
| 2(HRZE)/10(HR)      | Miliary TB and TB meningitis                                               |
| 12 months           | All children, adolescents and adults.                                       |
| 2(HRZE)/7-10(HR)    | Bone and joint TB                                                         |
| 9-12 months         | All children, adolescents and adults.                                       |

If bacteriological testing and/or CXR are not available, children meeting the following criteria are eligible for the 4-month regimen 2(HRZE)/2(HR):

- **Signs and symptoms not requiring hospitalisation**:
- Extra-thoracic lymph node TB without involvement of other EP sites.

If after one month of treatment symptoms have completely resolved, continue treatment until the end. If symptoms have not completely resolved, further investigations are needed.

If after 4 months of treatment symptoms have not completely resolved and/or there is no weight gain, further investigation is needed. The treatment can be extended to 6 months if causes of non-response to treatment (including DR-TB, non-adherence and non-TB disease) are ruled out or unlikely.

Ethambutol can be removed from the 4- and 6-month regimens in non-HIV-infected children living in areas where the prevalence of HIV and/or isoniazid resistance is low with:

- PTB microscopy smear-negative, or
- Extra- or intra-thoracic lymph node TB

For spinal TB, rest and back support bracing are indicated in addition to drug therapy. For patients with neurological deficit or unstable spine lesion, surgery can also be considered.

### Notas

(a) Symptoms requiring hospitalisation: signs of severe respiratory disease or distress, severe acute malnutrition, fever > 39 °C, severe pallor, restlessness, irritability or lethargy, etc.

### Referencias

   [https://apps.who.int/iris/rest/bitstreams/1414333/retrieve](https://apps.who.int/iris/rest/bitstreams/1414333/retrieve)
9.3 Alternative treatment regimens

Table 9.2 – Alternative DS-TB regimens according to the infection site

<table>
<thead>
<tr>
<th>Regimen</th>
<th>Duration</th>
<th>Eligibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>2HPZ-Mfx/2HP-Mfx</td>
<td>4 months</td>
<td>PTB and non-severe EPTB&lt;sup&gt;1,2&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Adolescents ≥ 12 years and adults meeting all the following criteria:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Weight ≥ 40 kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• CD4 ≥ 100 if HIV-infected</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• No resistance to fluoroquinolones (FQs) or living in areas where the prevalence of FQs resistance is low.</td>
</tr>
<tr>
<td>6HRZ-Eto</td>
<td>6 months</td>
<td>TB meningitis&lt;sup&gt;3&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Non-HIV infected children and adolescents under 20 years without inhA mutation detected.</td>
</tr>
</tbody>
</table>

Regimen 2HPZ-Mfx/2HP-Mfx  
- This regimen is an alternative to the conventional regimens for PTB and EPTB in eligible patients.  
- Implementation requires DST to FQs and supply of rifapentine.  
- There are no fixed-dose combinations (FDC) for this regimen which makes treatment adherence more difficult.

Regimen 6HRZ-Eto  
- Small studies have shown lower mortality, but more neurological sequelae with the 6HRZ-Eto regimen compared to the 12-month conventional regimen. However, no clinical trials have been conducted to compare the two regimens<sup>3</sup>.  
- The advantages of this regimen are short duration and better central nervous system penetration of ethionamide compared to ethambutol.  
- Implementation requires supply of ethionamide.  
- There are no FDC for this regimen which makes treatment adherence more difficult.  
- The daily doses of TB drugs in this regimen are higher than those of other regimens:  
  - isoniazid 20 mg/kg daily (max. 400 mg)  
  - rifampicin 20 mg/kg daily (max. 600 mg)  
  - pyrazinamide 40 mg/kg daily (max. 2 g)  
  - ethionamide 20 mg/kg daily (max. 750 mg)
9.4 Special situations

9.4.1 Women (pregnant or breastfeeding or of childbearing age)

Pregnant or breastfeeding women

- All first-line TB drugs, except rifabutin and rifapentine, can be used during pregnancy and breastfeeding\(^1\).
- Isoniazid may cause peripheral neuropathy due to vitamin B\(_6\) (pyridoxine) deficiency:
  - Pregnant and breastfeeding women should receive pyridoxine PO (10 mg once daily) throughout the course of TB treatment.
  - Breast-fed neonates or infants should receive pyridoxine PO (5 mg once daily).
- Rifampicin may cause clotting disorders due to increased vitamin K (phytomenadione) metabolism:
  - Women in late pregnancy on rifampicin (or rifabutin) should receive phytomenadione PO (10 mg once daily) for 2 weeks prior to expected date of delivery.
  - Neonates should also receive phytomenadione IM at birth (1 mg single dose) to prevent haemorrhagic disease of the newborn.
- Regimens containing rifapentine, moxifloxacin\(^2\) and ethionamide cannot be used to treat DS-TB in pregnant and breastfeeding women.

Women of childbearing age

Women on contraception should use an intra-uterine device or a progestogen-only injectable throughout the courses of TB treatment, as rifamycins reduce the effectiveness of implants and oral contraceptives.

9.4.2 Malnutrition or risk of malnutrition

- For patients with malnutrition, therapeutic feeding should be initiated.
- For children with severe acute malnutrition, a 6-month regimen is preferred over a 4-month regimen until more data on the efficacy of the 4-month regimen in these patients become available.
- For at-risk populations, such as children, pregnant and breastfeeding women and the elderly, nutritional supplementation with a standard food package or ready-to-use food may be considered during the first 2 months of treatment.

9.4.3 Diabetes

TB can impair glycaemic control in patients with diabetes\(^3\). It is necessary to increase blood glucose monitoring in these patients. TB drugs can exacerbate complications of diabetes (e.g. peripheral neuropathy). Avoid prescribing ethambutol in patients with pre-existing diabetic retinopathy. Rifampicin can reduce the effect of sulfonylureas (e.g. glibenclamide, gliclazide). In contrast, first-line TB drugs have no interactions with metformin.

If diabetes is diagnosed, treat and monitor according to standard protocols.

References


[1] Isoniazid may cause peripheral neuropathy due to vitamin B\(_6\) (pyridoxine) deficiency.

[2] Regimens containing rifapentine, moxifloxacin and ethionamide cannot be used to treat DS-TB in pregnant and breastfeeding women.

At the end of TB treatment, it is recommended to schedule a specialist consultation for a complete evaluation and, if necessary, adjust antidiabetic treatment.

### 9.4.4 Renal insufficiency

In patients with renal insufficiency, creatinine clearance should be calculated. If it is less than 30 ml/minute, doses of certain TB drugs should be adjusted.

For the formula to estimate the creatinine clearance and dose adjustments in renal insufficiency see [Appendix 12](#).

---

**Referencias**


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### 9.5 Adjunctive therapy

#### 9.5.1 Pyridoxine prophylaxis

Pyridoxine (vitamin B<sub>6</sub>) prophylaxis is indicated for all patients at risk of peripheral neuropathy, i.e. pregnant or breastfeeding women and patients with HIV infection, chronic alcohol use, malnutrition, diabetes, chronic hepatic disease or renal impairment (see [Appendix 17](#)).

#### 9.5.2 Corticosteroid therapy

Corticosteroid therapy is indicated for:

- Treatment and prevention of TB-associated immune reconstitution inflammatory syndrome (TB-IRIS). See [Chapter 12](#).

There is insufficient evidence regarding the use of corticosteroids in other indications<sup>[3][4]</sup>.

**Table 9.3 – Corticosteroid treatment**

<table>
<thead>
<tr>
<th>Indications</th>
<th>Dosage and duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>TB meningitis</td>
<td><strong>Dexamethasone</strong> IV: 0.4 mg/kg once daily for 7 days then, <strong>Prednisolone</strong> PO: 2 mg/kg once daily, tapered off over 6 to 8 weeks</td>
</tr>
<tr>
<td>TB pericarditis</td>
<td><strong>Prednisolone</strong> PO&lt;br&gt;Child: 1.5 mg/kg once daily for 4 weeks, tapered off over 6 weeks&lt;br&gt;Adul: 60 mg once daily for 4 weeks, tapered off over 6 weeks</td>
</tr>
</tbody>
</table>
9.6 Patient monitoring

Patients should be assessed at baseline, then, regardless of the regimen prescribed, monitored throughout the course of treatment.

Monitoring includes:
- Assessment of treatment response
- Detection of adverse effects and adherence issues.

For the schedule of follow-up examinations see Appendix 14.

Baseline and follow-up findings should be noted in the patient file to enable the detection and interpretation of potential changes.

9.6.1 Clinical visits

Baseline assessment

Assessment includes:
- Symptoms of TB and their severity (cough, fever, night sweats, weight loss, shortness of breath, ability to perform daily activities).
- Vital signs and weight.
- Comorbidities and other risk factors for adverse effects requiring monitoring adaptation.
- Psychological assessment.

Other investigations may be needed depending on the drugs used in the regimen prescribed (Section 9.6.3).

Clinical assessment should be performed by a clinician. Psychological assessment should be performed whenever possible by personnel with appropriate training.

All patients starting treatment should be given the information they need to understand the disease and its treatment (Appendix 21).

Follow-up visits

Each follow-up visit, assessment includes:
- Clinical progress, vital signs and weight. Dosages should be adjusted to the weight if necessary.
- Occurrence of adverse effects.
- Adherence to treatment (Appendix 22).
- Psychological condition.

Frequency of visits depends on the patient’s clinical condition and evolution:
- A visit every other week for the first month, then once a month if there is no particular problem.
- Additional visits may be required in case of comorbidities, severe or multiple adverse effects, pregnancy, etc.

Visits should coincide with bacteriological examinations and other investigations when possible.

References


The clinician should take into account any information and concerns regarding treatment tolerance and adherence reported by the patient or the team responsible for the patient’s follow-up and support.

### 9.6.2 Bacteriological tests

To assess treatment response in patients with:
- PTB: bacteriological tests are essential.
- EPTB: evaluation is based on clinical evolution. However, bacteriological tests are required if patients also develop PTB.

#### Baseline tests

Baseline tests are those performed on specimens collected just prior to treatment initiation. They include:
- Rapid molecular tests (RMTs) for detection of *M. tuberculosis* and rifampicin and isoniazid resistance.
- Smear microscopy to monitor treatment progress.
- Culture and phenotypic DST (pDST) when indicated.

For more information see Chapter 3.

#### Follow-up tests

- **Smear microscopy**
  - Microscopy should be performed every 2 months until treatment completion.
  - If treatment is effective, microscopy at Month 2, 4 and 6 should be negative.

  **Notes:**
  - Patients with high bacillary load at baseline may have dead bacilli in their sputum for several months.
  - As microscopy cannot distinguish dead from live bacilli, a positive result does not necessarily indicate that the treatment has failed.

- **Rapid molecular tests**
  - RMTs cannot be used to monitor treatment progress. However, if microscopy or culture is positive at Month 2 or later, RMTs should be performed to detect the emergence of new drug resistance not present at baseline (Chapter 3).

- **Culture and pDST**
  - Culture and pDST should be performed:
    - at Month 2 or later, if RMTs show a new resistance to rifampicin or isoniazid;
    - at Month 4, if microscopy is positive.
  - Full pDST (for first- and second-line drugs) should be performed on any positive culture.

  **Note:** bacteriological tests are performed at the end of the month (e.g. Month 2 means the end of the 2\textsuperscript{nd} month of treatment).

Regardless of the above schedule, RMTs, culture and pDST should be performed if the patient’s clinical condition deteriorates.

#### End of treatment test

Microscopy should be performed at end of treatment to confirm the end of treatment outcome (Chapter 17).

### 9.6.3 Other investigations

#### Radiography

- **CXR:** for children with presumptive PTB, patients with non-bacteriologically confirmed PTB, suspicion of other intra-thoracic TB at baseline, then if indicated (e.g. worsening respiratory symptoms, non-response to TB treatment).
- **Bone x-ray:** for patients with osteoarticular and spinal TB at baseline, then every 6 months.

#### Biological tests

**Table 9.4** – Blood tests at baseline and during treatment
9.7 Adverse effects

Rapid management of adverse effects is essential to increase tolerance and improve outcomes.
In the event of minor adverse effects, drugs should not be stopped. Providing support and using ancillary medicines is all that is necessary.
In the event of major adverse effects, the regimen may need to be adapted.

Table 9.5 – Main adverse effects and likely responsible drugs

<table>
<thead>
<tr>
<th>Tests</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full blood count[a]</td>
<td>HIV-infected patients on rifabutin or zidovudine (AZT), at baseline, then once a month for the first 2 months, then if indicated.</td>
</tr>
<tr>
<td>Liver function tests[b]</td>
<td>Patients with pre-existing hepatic disease, at baseline, then once a month.</td>
</tr>
<tr>
<td>Serum creatinine[c]</td>
<td>Patients with renal insufficiency at baseline, then if indicated.</td>
</tr>
<tr>
<td>HbA1C and/or blood glucose level</td>
<td>All patients, at baseline, to detect diabetes. If diabetes is detected, monitor according to standard protocols.</td>
</tr>
<tr>
<td>HIV, hepatitis B and C</td>
<td>For patients with undocumented HIV, hepatitis B and C status; HIV test every 6 months in high HIV prevalence areas. Tests can be repeated in case of recent exposure.</td>
</tr>
<tr>
<td>CD4 count and viral load</td>
<td>HIV-infected patients: at baseline, then every 6 months.</td>
</tr>
</tbody>
</table>

(a) Haemoglobin, red and white blood cells, platelets.
(b) Aspartate aminotransferase (AST) and alanine aminotransferase (ALT). Bilirubin if elevated liver enzymes.
(c) For estimation of creatinine clearance see Appendix 12.
9.8 Treatment adaptation and change of treatment

9.8.1 Treatment adaptation

The whole treatment or individual drug(s) may be temporarily interrupted by the clinician in case of severe adverse effects (Appendix 17).

This is considered as treatment adaptation, as long as it does not meet the definition of “treatment failure” (Chapter 17).

9.8.2 Change of treatment

The clinician should replace the DS-TB treatment with:

- A treatment for isoniazid-resistant TB when RMT or pDST show:
  - the development of isoniazid resistance (Chapter 11) after treatment initiation, or
  - undetected isoniazid resistance at baseline, for any reason.
- A treatment for multidrug-resistant or rifampicin-resistant TB (MDR/RR-TB, see Chapter 10) in the following circumstances[1]:
  - Development of rifampicin resistance after treatment initiation.
  - Rifampicin resistance not detected at baseline, for any reason.
  - No bacteriological conversion or bacteriological reversion (Chapter 17).
  - Insufficient clinical response to treatment in patients:
    - with non-bacteriologically confirmed TB (e.g. miliary TB, some forms of EPTB, TB in children).
    - with bacteriologically confirmed TB, when the bacteriological response cannot be assessed, or the result is inconclusive.

The above treatment changes meet the outcome definition of “treatment failure” except when the reason for change is a resistance undetected at baseline[1] (Chapter 17).

### Adverse effects

<table>
<thead>
<tr>
<th>Minor</th>
<th>Drug(s) likely responsible</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea, vomiting</td>
<td>Eto, Z</td>
<td>Appendix 17</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>Z</td>
<td>Appendix 17</td>
</tr>
<tr>
<td>Peripheral neuropathy</td>
<td>H, Eto</td>
<td>Appendix 17</td>
</tr>
<tr>
<td>Orange/red urine, tears, etc.</td>
<td>R, P</td>
<td>Patients should be told that this is normal before starting treatment.</td>
</tr>
</tbody>
</table>

### Major

| Skin reactions               | E, Z, R, H, P, Mfx, Eto   | Appendix 17                                     |
| Hepatotoxicity               | Z, H, R, P, Eto           | Appendix 17                                     |
| Optic neuritis               | E                          | Appendix 17                                     |
| Haematologic disorders       | R, P, H, E                | Appendix 17                                     |

For more information on adverse effects see Appendix 10.
9.9 Treatment interruption

Treatment interruption can lead to the emergence of new resistances. Problems of treatment interruption by the patient (e.g. discontinuation of certain drugs, recurrent treatment interruptions) should be identified and addressed (reinforcement of patient support and management of adverse effects if necessary).

Table 9.6 – Management of patients who interrupt treatment

<table>
<thead>
<tr>
<th>Length of treatment before interruption</th>
<th>Length of interruption</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 1 month</td>
<td>&lt; 2 weeks</td>
<td>Continue treatment at the point it was stopped. Doses missed during interruption must be made up to complete the treatment.</td>
</tr>
<tr>
<td></td>
<td>2-7 weeks</td>
<td>Restart treatment or perform RMTs (see below) depending on patient’s clinical evolution.</td>
</tr>
</tbody>
</table>
|                                        | ≥ 8 weeks              | Perform RMTs:  
|                                        |                        | • if no resistance, restart treatment.  
|                                        |                        | • if resistance, start DR-TB treatment. |
| ≥ 1 month                              | < 2 weeks              | Continue treatment at the point it was stopped. Doses missed during interruption must be made up to complete treatment. |
|                                        | ≥ 4 weeks              | Perform RMTs:  
|                                        |                        | • if no resistance, restart treatment.  
|                                        |                        | • if resistance, start DR-TB treatment. |

For patients on 6-month regimen who have received adequate treatment for 4 months or more, who return smear negative, are in good clinical condition and with no resistance detected, the decision to re-start a treatment is considered on a case-by-case basis.

When a DST is not feasible (e.g. miliary TB, some forms of EPTB, TB in children), clinical and radiological evaluation should guide the decision to either restart DS-TB treatment or switch to an DR-TB treatment.

Chapter 10: Treatment of multidrug-resistant TB (MDR-TB)

10.1 Design of therapeutic regimens in MDR-TB

10.2 Selection of anti-TB drugs in MDR-TB regimens

10.3 Building a treatment regimen for MDR-TB
10.1 Design of therapeutic regimens in MDR-TB

The following are the basic principles involved in MDR-TB regimen design[1]:

- The intensive phase includes at least four core Group 2 to 4 anti-TB drugs likely to be effective, including an injectable agent – plus pyrazinamide (Z).

- In the case of unclear evidence about the effectiveness of a certain drug, it can be part of the regimen but it should not be counted as one of the four core second-line anti-TB drugs.

- An anti-TB drug is considered “likely to be effective” when:
  1. The drug has not been used in a regimen that failed for the individual patient;
  2. Drug susceptibility testing (DST) performed on the patient’s strain indicates that the strain is susceptible. Only DST for isoniazid, rifampicin, Group 2 and 3 drugs is considered reliable;
  3. No known resistance to drugs with high cross-resistance;
  4. No known close contacts with a patient infected with a strain resistant to the drug;
  5. In the absence of DST or for drugs in which individual DST is not reliable, a drug resistance survey demonstrates that resistance to the drug is rare in patients with similar TB history.

- It is not always possible that all five criteria can be ascertained and clinical judgment is often necessary on whether to count a drug as “likely effective”.

- An important pitfall in designing MDR-TB regimens is due to the turnaround time necessary for DST, the patient may have already received months of a treatment by the time DST results become available from the laboratory. The possibility of further acquired resistance during this time must be considered. If there is a high probability of acquired resistance to a drug after the specimen for DST was collected, this drug should not be counted as one of the four second-line anti-TB drugs in the core regimen, but can be included as an adjunctive agent.

- The most effective regimens for MDR-TB include at least a fluoroquinolone (preferably a third-generation), an injectable agent, ethionamide (or prothionamide), either cycloserine or para-aminosalicylic acid, and pyrazinamide.

- There are conditions when more than five drugs may be started, as is the case if the susceptibility pattern is unknown or the effectiveness is questionable for a drug(s).

- A drug should not be used when patient is known to have a major contraindication of usage (e.g. known major drug-drug interactions, history of allergic reaction, pregnancy).

- Each dose is given under directly observed therapy (DOT) throughout the treatment. A treatment card is marked for each observed dose. DOT can be performed either facility-based or home-based (often referred to as community-based). See Chapter 13.

- Treatment is given six or seven days a week. Six days a week is chosen for those patients managed in outpatient settings where DOT cannot be done everyday.
10.2 Selection of anti-TB drugs in MDR-TB regimens

See reference[1]

Group 1 (Oral first-line agents)

Pyrazinamide is routinely added to MDR regimens if susceptibility (by DST) is documented or if DST is unknown. If well tolerated it is used for the entire treatment, although patients doing well and with minimal lung disease can have it stopped with the injectable agent and continue with at least three likely effective drugs.

Ethambutol is not routinely added to MDR regimens, however it can be added if the criteria of it being a likely effective drug are met.

For patients with strains resistant to low concentrations of isoniazid, but susceptible to higher concentrations, the use of high-dose isoniazid may have some benefit (see Group 5).

The newer rifamycins, such as rifabutin have very high cross-resistance to rifampicin and are not used in MDR regimens.

Group 2 (Injectable agents)

All patients should receive a Group 2 injectable agent if susceptibility is documented or the drug is considered likely to be effective.

Kanamycin or amikacin are the first choice injectable agent. Both are low cost, and have been used extensively for the treatment of MDR-TB. They are considered to be very similar and have a high frequency of cross-resistance.

Given the high rates of resistance to streptomycin in patients with MDR-TB, streptomycin is not used in MDR-TB treatment regimens.

If the strain is susceptible to capreomycin or if resistance is rare in the patient population and if aminoglycosides are contra-indicated or poorly tolerated or ineffective on the patient’s strain, capreomycin should be used. Capreomycin should also be used while waiting for the DST results in places where resistance to kanamycin and amikacin is common.

Group 3 (Fluoroquinolones)

The most potent available fluoroquinolones in descending order based on in vitro activity and animal studies are: moxifloxacin > levofloxacin > ofloxacin[2][3].

This guide recommends not using ofloxacin (second-generation fluoroquinolone) as it has inferior performance against TB compared to the other Group 3 fluoroquinolones. In addition, resistance may develop more easily to the fluoroquinolone group when ofloxacin is used in a multidrug regimen. Ciprofloxacin (second-generation fluoroquinolone) is not included in Group 3 and should never be used to treat drug-susceptible or DR-TB because of its low efficacy against TB bacilli[4].

Third-generation fluoroquinolones (moxifloxacin and levofloxacin) may have some efficacy against ofloxacin-resistant strains[5].

Referencias

Mostly based on cost and availability, levofloxacin is often the fluoroquinolone used in most MDR-TB regimens, whereas moxifloxacin is reserved for cases of high resistance (resistance to ofloxacin, injectable agents, or other second-line anti-TB drugs).

In case of resistance to fluoroquinolones, the use of bedaquiline should be considered (see below).

**Group 4 (Oral bacteriostatic second-line anti-TB drugs)**

Ethionamide and prothionamide are considered the most potent Group 4 drugs\(^{[6]}\). However it should be noted that these drugs do have some cross-resistance with isoniazid. Ethionamide and prothionamide can be included in the regimen if inhA gene is detected but should not be counted as a likely effective drug.

Cycloserine and/or para-aminosalicylic acid should be included in MDR-TB regimens. Both share no cross-resistance to other anti-TB drugs. Since the combination of ethionamide or prothionamide and para-aminosalicylic acid often causes a high incidence of gastrointestinal disturbances and hypothyroidism, these agents are usually used together only when three Group 4 agents are needed.

The drugs in Group 4 may be started at a low dose and escalated over 1 to 2 weeks to improve tolerance.

**Group 5 (Drugs with limited data on efficacy and/or long-term safety)**

Group 5 drugs are recommended in cases where adequate regimens are impossible to design with the drugs from Groups 1 to 4. Compared to other drugs in this group bedaquiline is the only one with proven efficacy against TB. While there is no clear evidence for the hierarchy of use of Group 5 drugs, these guidelines propose that the three most attractive agents from this group in order of preference are: bedaquiline, linezolid, clofazimine.

**Bedaquiline**\(^{[7][8][9]}\) : Bedaquiline is a diarylquinoline with bactericidal anti-mycobacterial activity. This new drug was registered by the US FDA in December 2012\(^{[8]}\) for MDR-TB patients with no other therapeutic options. It is recommended in case of resistance to fluoroquinolones or when it is not possible to have four effective anti-TB drugs from Group 2 to 4 in the regimen. The dosage in adult is 400 mg once daily for 2 weeks followed by 200 mg 3 times per week for 22 weeks.

The drug is not yet recommended for children or pregnant women. The main adverse effects are nausea, arthralgia, headache and QT prolongation. QT prolongation can result in cardiac arrhythmia and sudden death. Baseline and regular electrocardiogram (ECG) monitoring should be performed. QT prolongation is more pronounced when combined with clofazimine. Combination with other QT prolonging drugs (moxifloxacin, ondansetron, etc.) should be avoided or closely monitored. Bedaquiline must not be combined with rifamycins and some antiretrovirals (see Chapter 12). Bedaquiline is not registered in most high burden countries and only available through compassionate use (see also Appendix 11).

For situations that require the use of Group 5 drugs other than bedaquiline (or when bedaquiline is not available), use at least two other drugs from Group 5 given the limited knowledge of their efficacy.

**Linezolid** : Linezolid has good activity in vitro and in animal studies. There are also a number of reports and case series in MDR-TB and XDR-TB\(^{[10][11][12][13][14][15][16][17]}\) and a recent study showing efficacy in XDR-TB\(^{[18]}\). It has numerous severe adverse effects including myelo-supression and irreversible peripheral neuropathy. It is presently very expensive.

**Clofazimine** : There is a moderate amount of experience with clofazimine in MDR-TB treatment but no clear in vivo data on efficacy against TB. It is usually added to regimens for XDR-TB.

**Amoxicillin/clavulanic acid** : Generally the B-lactam antibiotics are not regarded as very useful drugs in TB. However, the addition of the B-lactamase inhibitor makes them active in vitro against TB. There is one in vivo study that showed good early bactericidal activity. While amoxicillin/clavulanic acid is probably a relatively weak anti-TB drug, it is often included because it is available, inexpensive and causes only minor adverse effects.
**High-dose isoniazid:** High-dose isoniazid (16-20 mg/kg/day) can be used as a Group 5 drug in the presence of resistance to low concentrations of isoniazid[^19] (> 1% of bacilli resistant to 0.2 mcg/ml but susceptible to 1 mcg/ml of isoniazid). Isoniazid is not recommended for high-dose resistance (> 1% of bacilli resistant to 1 mcg/ml of isoniazid[^20]) or in presence of katG gene mutation (see LPA, Chapter 3, **Section 3.4.2**).

**Notes:**
- **Gatifloxacin** (Group 3): Although gatifloxacin is similar to moxifloxacin in efficacy against TB, it is associated with serious hypo/hyperglycaemia, and new onset diabetes. Thus, its use is not recommended.
- **Terizidone** (Group 4): It is unknown whether this drug is equally efficacious as cycloserine, therefore these guidelines recommends the use of cycloserine over terizidone.
- **Imipenem/cilastatin and meropenem** (Group 5): These beta-lactam/carbapenems are only given intravenously. Given the cost and difficulty of the twice-daily intravenous administration, it is not commonly used in resource-constrained settings. Meropenem is preferred in children as there is more experience with its use. Meropenem can be combined with oral doses of clavulanate. These drugs are commonly used for a duration of two months past conversion.
- **Clarithromycin** (Group 5): This drug is included in various TB manuals[^21] yet evidence to support its efficacy in MDR-TB is minimal. It may have a synergistic effect on first-line anti-TB drugs with enhanced intracellular effectiveness against the TB bacilli. However, until more information on effectiveness in TB and MDR-TB, its use is not recommended.
- **Thioacetazone** (Group 5): While thioacetazone is known to be active against TB bacilli, it is placed in Group 5 because its role in DR-TB treatment is not well established. Thioacetazone has cross-resistance with some of the other anti-TB agents (Chapter 8, **Section 8.5**) and overall is a weakly bacteriostatic drug. It is contraindicated in HIV-infected individuals[^22] due to a risk of serious adverse reactions (Stevens-Johnson syndrome and death). Persons of Asian descent also have a higher incidence of Stevens-Johnson syndrome. For these reasons, thioacetazone is rarely added as a Group 5 drug. Until there is more information in its role in MDR-TB therapy, its use is not recommended.

**Notas**
(a) Bedaquiline is relatively well tolerated. Data and experience on its use is very limited. In a blinded randomized placebo-control study there were an increase number of deaths in the study arm that received bedaquiline. While none of the deaths were considered directly related to the drug, the possibility that the use of the drug carries an increase risk of death cannot be ruled out. The risks and benefits of receiving this drug should be fully explained to the patient.

(b) [http://www.accessdata.fda.gov/drugsatfdadocs/label/2012/204384s000lbl.pdf](http://www.accessdata.fda.gov/drugsatfdadocs/label/2012/204384s000lbl.pdf)

**Referencias**


10.3 Building a treatment regimen for MDR-TB

Individual regimens are designed based on DST of the infecting strain, history of TB treatment, and contact history. Figure 10.1 describes the steps to build a regimen for MDR-TB treatment.

Figure 10.1 - Building a regimen for MDR-TB
Box 10.1 - Examples of how to initiate and design MDR-TB regimens
Example 1 - Patient doing poorly on first-line treatment

A patient receiving first-line treatment for new patients (2 HRZE/4 HR) continues to be smear positive at Month 3 with symptoms including weight loss, fever, shortness of breath and cough. The patient feels the shortness of breath is getting severe and he spends more than 50% of the day in bed. No DST was performed at the start of treatment. Xpert MTB/RIF performed at Month 3 shows MTB+ and rifampicin resistance. What should be done?

**Answer:** A positive Xpert MTB/RIF at Month 3 in a patient doing poorly on a first-line regimen that shows R resistance is highly likely to be a true positive. This patient should be placed on MDR-TB therapy. A confirmatory DST with conventional methods to at least H and R and if possible to injectable agents and fluoroquinolones should be performed.

If a rapid molecular test was not available, this patient should be placed on an MDR-TB regimen while waiting conventional DST results.

- If there is low second-line drug resistance in patient’s strains with MDR-TB in the area then a common regimen is: Km-Lfx-Eto (or Pto)-Cs-ZE.
- If there is moderate to high second-line drug resistance in MDR-TB strains in the area or if the level of resistance to second-line drugs is not known: Cm-Mfx-Eto (or Pto)-Cs-PASZE. Once DST becomes available the regimen can be adjusted. In this case, the infecting strain was determined to be resistant to H-R-S and susceptible to Km-Cm-Ofx-E; resistance to Z was unknown. Given the DST results, it is recommended to continue with Km-Lfx-Eto (or Pto)-Cs-ZE and drop the PAS if it was used in the initial regimen.

Example 2 - Xpert RIF positive in a patient with low probability of MDR-TB

A HIV-negative smear-negative TB suspect is referred to Xpert MTB/RIF to establish the diagnosis of TB. The result of the Xpert is MTB+ and rifampicin resistance. The patient has never been diagnosed with TB. The MDR-TB prevalence for new patients in the area is 1%. The patient only complains of a mild cough for 3 weeks and X-ray shows minimal lesions. What should be done?

**Answer:** The RIF resistance positive predictive value (PPV) for the Xpert MTB/RIF in the setting of 1% rifampicin resistance prevalence is 32% ([Appendix 3](#)). Because of the relatively low PPV of the Xpert MTB/RIF under these circumstances and the fact that patient is HIV-negative and not seriously ill, he can be placed on a first-line drug regimen while waiting confirmation DST. If possible, DST confirmation should be done through a rapid phenotypic method or using LPA on culture (indirect method). If the patient deteriorates clinically at any time while waiting confirmation DST, an empirical MDR-TB regimen should be started. When the DST returns, the regimen should be adjusted if the resistance to rifampicin is confirmed.

**Note:**

An alternative shorter 9 month standard regimen (4 Km-Gfx-Pto-Cfz-high dose H-ZE/5 Gfx- Cfz-ZE) has shown good effectiveness in a study in Bangladesh. Adaptations are made in some countries in Western Africa with moxifloxacin replacing gatifloxacin and extension of the regimen to 12 months. At present, this regimen is still considered experimental.

Given the limited evidence supporting this regimen these guidelines recommend the following:

- Obtain country-level and institutional ethical approval before implementation.
- Implement it under operational research conditions following good practices.
- Consider this regimen on a case-by-case basis for programmes with proper follow-up and outcome documentation in unstable settings where a 2 year-treatment is not an option.
- Perform DST to the fluoroquinolones in a liquid medium and do not use in any patient with documented fluoroquinolone resistance (the third-generation fluoroquinolones are the backbone of the regimen and the regimen does not perform well against strains resistant to fluoroquinolones).
- Use only in HIV-negative patients until more information is published on the regimen and its use in HIV-positive patients.
- Do not use in areas with a high prevalence of resistance to second-line anti-TB drugs until more information is published.

**Notas**

10.4 Duration of MDR-TB regimens

10.4.1 Intensive phase

Duration of intensive phase is guided by culture. The injectable agent should be continued for at least 8 months\(^1\) and at least 4 months after the patient becomes culture negative – which ever is longer.

The use of an individualized approach which reviews the cultures, smears, X-rays and the patient’s clinical status may also aid in deciding whether or not to continue an injectable agent longer than the above recommendation, particularly in the case of patients for whom the susceptibility pattern is unknown, effectiveness is questionable for an agent(s) or extensive or bilateral pulmonary disease is present.

A change to intermittent therapy with the injectable agent (3 times weekly) is done when signs of toxicity are noticed. Three times a week therapy is recommended in patients after 6 months of an injectable agent and who have had culture conversion, as toxicity becomes a greater risk to patients with longer periods of the injectable agent.

10.4.2 Length of treatment

The duration of treatment is guided by culture. It is recommended continuing therapy for a minimum of 20 months\(^1\) and at least 18 months after the patient becomes culture negative.

Extension of therapy to 24 months may be indicated in chronic cases with extensive pulmonary damage.

Referencias

   http://ajrccm.atsjournals.org/content/182/5/684.full.pdf


10.5 Follow-up for patients treated for MDR-TB

Table 10.1 - Routine patient monitoring
<table>
<thead>
<tr>
<th>Evaluation</th>
<th>Frequency</th>
</tr>
</thead>
</table>
| **Assessment by a clinician**                  | During intensive phase: every day during the first weeks if hospitalized and at least every week if treated as outpatient, until the treatment is well tolerated. Once stable, the patient is seen once or twice monthly.  
During continuation phase: monthly assessment unless there is a medical necessity to see the patient more often.  
The DOT supporter sees the patient daily and signals any concerns to the clinician. |
| **Treatment adherence and tolerance**          | Daily at every DOT encounters by the DOT supporter.                        |
| **Sputum smear and cultures**                  | Monthly until the end of treatment.  
*Note: programmes with very limited culture capacity may consider doing smears monthly but cultures every other month for the continuation phase.* |
| **Weight**                                     | At baseline and then monthly.                                              |
| **DST**                                        | At baseline and for any positive culture during treatment.                 |
| **Chest X-rays**                               | At baseline and then every three to six months.                           |
| **Serum creatinine**                           | At baseline, then twice a month for the first two months, then monthly while receiving an injectable agent.  
Every one to three weeks in HIV-infected patients, diabetics throughout the course of the injectable agent. |
| **Serum potassium (K+)**                       | At baseline, then twice a month for the first two months, then monthly while receiving an injectable agent.  
Every one to three weeks in HIV-infected patients, diabetics throughout the course of the injectable agent. |
| **Thyroid stimulating hormone (TSH)**          | Every six months if receiving Eto/Pto and/or PAS (every three months in HIV positive patients) and whenever signs/symptoms of hypothyroidism are present. TSH is sufficient for screening for hypothyroidism and it is not necessary to measure hormone thyroid levels. |
| **Liver serum enzymes**                        | At baseline then monthly during the intensive phase. Every 3 months thereafter. Monthly monitoring for HIV-infected.  
In patients with viral hepatitis: once weekly for the first month, then every one to four weeks.  
Monthly for patients taking Bdq. |
| **Bilirubine**                                 | Monthly for patients taking Bdq.                                           |
| **HIV screening**                              | At baseline then repeat when clinically indicated or every 6 months in high HIV prevalence settings. |
| **Pregnancy tests**                            | At baseline for women of childbearing age, and repeat if indicated.        |
| **Haemoglobin**                                | If on Lzd, weekly during the first month, then monthly or as needed based on symptoms; there is little clinical experience with prolonged use of Lzd.  
For HIV-infected patients on AZT: monthly initially and then as needed based on symptoms.  
If patient not on Lzd or AZT, routine monitoring is not indicated. |
| **White blood count**                          | Baseline audiogram, then monthly during intensive phase (and whenever clinically indicated). Ask patient about changes in hearing at every clinic visit and evaluate their ability to participate in normal conversation. |
| **Hearing tests**                              |                                                                           |
10.6 Management of adverse effects in patients on second-line regimens

Treating rapidly and aggressively adverse reactions is an important means to increase tolerance and is critical to improve outcomes.

All patients should be informed that they are likely to experience adverse effects. Adverse effects appear most commonly at the start of therapy, especially during the first few weeks of treatment where the patient can feel quite lousy – with nausea and vomiting being the most common adverse effect. Patients should be informed that many of the common minor adverse effects will improve with time and medical treatment.

Patients are monitored for general toxicities and drug-specific toxicity at every DOT encounter. They should be educated that if serious adverse effects appear (e.g. hearing loss, dizziness, ringing in the ears, jaundice, edema, decreased urine output, skin rash or burning in the legs), they must inform the health care worker immediately.

It is often difficult to ascertain whether a given adverse effect is due to a single drug or is the result of several drugs given simultaneously. If after management of adverse effects the patient remains intolerably symptomatic, a dose reduction or elimination of one of the drugs may be necessary. Permanent dose reduction or definitive elimination of a drug should be considered only after all other possibilities have been exhausted i.e., in cases of significant organ dysfunction or intractable intolerance. Ideally, any drug eliminated from a treatment regimen should be replaced with an equally effective drug, as to not compromise the overall effectiveness of the regimen.

Dose reduction can be done in a systematic manner by starting with the most likely offending drug for one week to see whether the symptoms diminish or disappear. If symptoms persist, the drug is returned to its original dose and the same process repeated for the other drugs, until all potentially responsible drugs have been tested. Systematic dose reduction of multiple drugs simultaneously would be the next option.

Whenever reducing or holding a drug to determine the cause of an adverse effect, tell the patient that this is a test to determine which drug is involved and that the drug dose will be increased back to therapeutic dose in a manner that will be better tolerated. Returning back to therapeutic doses gradually (over one to two weeks) while implementing strategies to decrease a specific adverse effect can often allow the patient to better tolerate it.

Treatment supporters and nurses working with TB programmes should report an adverse event to the physician at the earliest appropriate time. Only the managing physician should do dose changes or eliminate a specific anti-TB drug.

Often, if an adverse effect cannot be completely eliminated, patients may be asked to tolerate symptoms until they subside. Often reassurance and emotional support can result in the avoidance of adding yet another medication to the high burden of medications the patient is already receiving.

For specific management of common adverse effects, see Appendix 17.

Ancillary medicines (anti-emetics, potassium replacement, thyroid hormone, medicines for psychiatric conditions, etc.) should be provided free of charge to the patient.

10.7 Surgery as an adjunctive treatment measure

<table>
<thead>
<tr>
<th>Vision tests</th>
<th>For patients on long-term E or Lzd, use the Ishihara test (test for changes in the vision of colour). Perform at baseline as a certain percentage of the population has colour blindness. Then monthly in patients taking Lzd.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psycho-social consultation</td>
<td>At baseline by trained personnel in the skills of psycho-social management, during treatment and repeat as indicated. Refer to psychiatrist when indicated.</td>
</tr>
<tr>
<td>ECG</td>
<td>Patients taking Bdq: at baseline then, after 2 weeks then, monthly.</td>
</tr>
</tbody>
</table>
Surgery can be considered only in optimal surgical facilities with trained thoracic surgeons. Specialized surgical facilities should include stringent infection control measures since infectious substances and aerosols are generated in large quantities during surgery, during mechanical ventilation and post-operative pulmonary hygiene manoeuvres.

General indications for surgery for programmes with limited access to surgery include patients with resistance to a large number of drugs and localized pulmonary disease. Computerized tomography, pulmonary function testing, and quantitative lung perfusion/ventilation are recommended as part of the preoperative work-up.

The most common operative procedure in patients with MDR-TB is resection of part or all of a lung. Large case series analysis suggest surgical resection can be effective and safe under appropriate surgical conditions. It is considered an adjunct to chemotherapy and appears to be beneficial for patients when skilled thoracic surgeons and excellent postoperative care are available. It is not indicated in patients with extensive bilateral disease.

Resection surgery should be timed so as to offer the patient the best possible chances of cure with the least morbidity. Thus, the timing of surgery may be earlier in the course of the disease when the patient’s risk of morbidity and mortality are lower, for example, when the disease is still localized to one lung or one lobe. Furthermore, bacilli excretion during treatment has a “window” when the bacilli load decreases under pressure of anti-TB drugs and it can be registered by decreasing or even disappearing of mycobacteria in smear and/or culture. This “window” is the best time for surgery. It is critical to operate before the mycobacterial count begins to rise. The best time for surgery is usually considered to be between two and six months after initiation of treatment. Surgery should not be considered a last resort.

Even with successful resection, an additional 12 to 24 months of chemotherapy should be given.

Referencias


10.8 Management of patients whose treatment failed and palliative care

When it has been determined a patient is failing therapy for DR-TB, the first priority is to design a new regimen using the principles described in Sections 10.1 to Section 10.3. The new regimen should contain at least two new effective drugs.

The employment of newly developed TB drugs available for compassionate use is encouraged. For some of these drugs (delamanid), approval is expect in 2013.
When no therapeutic option or new regimen is possible, the patient can be continued on an anti-TB regimen that is reasonably tolerated (and if the patient desires) or the regimen can be completely stopped. The decision to stop therapy should be made after careful evaluation and consultation with the patient, the family and the MDR-TB treatment team.

Palliative/supportive care should be continued. Supportive measures for minimizing suffering due to the disease or the therapy should be implemented according to the patient needs.

Supportive measures may include:

- Relief of respiratory symptoms: oxygen should be used to alleviate shortness of breath; corticosteroids (prednisolone) are beneficial in severe respiratory insufficiency; codeine helps control cough.
- Identification, assessment and treatment of pain: according to the standard recommendations (non opioids/mild opioids/strong opioids adapted to the level of pain).
- All necessary ancillary medications needed should be used.
- Patients with poor nutritional status should receive nutritional support.
- In debilitated patients, important measures for making patients comfortable and preventing complications must be taken. Regular scheduled movement of the bedridden patients prevents bedsores. Bathing and oral care assistance keeps patients clean and comfortable, while preventing skin infections.
- Disorders such as anxiety or depression due to prolonged sickness, separation from family, difficult living conditions, etc. should be addressed when present. The patient as well as the family may need support.
- Potential social problems should also be addressed. When necessary, hospice-like care should be offered to families who want to keep the patient at home. Inpatient end-of-life care should be available to those for whom home care is not available.

Note: the above palliative/supportive measures should be implemented to all DR-TB patients if indicated whether or not they are failing treatment. Some measures may even need to be continued after a patient’s TB has cured, but the patient still remains with significant respiratory damage.

### 10.9 Special situations

#### 10.9.1 Pregnant women

Pregnant women should be carefully evaluated, such that the risks and benefits of treatment considered according to gestational age and severity of disease.

- The primary goal is culture conversion to protect the health of the mother and child, both before and after birth.
- If the patient is very stable with minimum disease, treatment may be delayed and started in the second trimester with 3 or 4 drugs known to be safe in pregnancy and active on the infecting strain. In most cases of moderate to severe disease the treatment should be started right away with the risks and benefits explained to the mother.
- Aminoglycosides are contraindicated. If an injectable agent is required, capreomycin is the only option as there are case reports of safe use in pregnancy.
- Ethionamide and prothiomanide should be avoided due to data suggesting teratogenicity in animals.
- Fluoroquinolones are considered acceptable to use despite limited data.
- Moxifloxacin, para-aminosalicylic acid, cycloserine and amoxicillin/clavulanic acid is an appropriate initial regimen with a consideration of capreomycin in cases of advanced disease (extensive parenchymal damage or life-threatening condition).
- If some drugs were withheld because of the pregnancy, they can be added back postpartum if needed to make a more complete regimen.

The child should receive BCG at birth.

#### 10.9.2 Breastfeeding women

Most anti-TB drugs will be found in the breast milk in concentrations that would equal only a small fraction of the therapeutic dose used in an infant. Effects on infants of such exposure during the full course of DR-TB treatment have not been established. Therefore, when resources and training are available, it is recommended to provide infant formula as an alternative to breastfeeding. If infant formula is used, the infant formula, clean water, fuel for boiling water and the apparatus (stove, heating pans and bottles) must be provided to the mother, as well as training on how to prepare and use the infant formula. If infant formula cannot be provided regularly and used safely, the child should be breastfed and the risks/benefits explained to the mother.

Treatment administered timely and properly is the best way to prevent transmission of tubercle bacilli to the breastfed infant.
If a mother is smear-positive and there is a possibility the mother is failing treatment, the care of the infant should be entrusted to family members until she becomes smear-negative, if feasible. Otherwise, nursing mothers with DR-TB should not be separated from their infants.

### 10.9.3 Women of child-bearing age

A pregnancy test should be performed before starting anti-TB therapy (to be repeated if indicated).

Women of child-bearing age should be provided contraception in addition to MDR-TB treatment.

Patients should be advised to take their oral contraceptives at times well away from when they may experience vomiting caused by the anti-TB drugs. Patients who vomit within the first two hours of taking the contraceptive tablet should use a barrier method of contraception for the duration of symptoms and for seven days after recovery.

**Note:** for patients with mono- and poly-drug resistant TB susceptible to rifampicin ([Chapter 11](#)), rifampicin interacts with hormonal contraceptives and decreases their efficacy. Patients may choose between these options, throughout the course of anti-TB treatment: medroxyprogesterone IM or barrier methods (diaphragm, condom, UID) or, as a last resort, oral contraceptive containing a high dose of estrogen (50 micrograms/tablet).

### 10.9.4 Children

Children with DR-TB generally have primary resistance transmitted from an adult contact with DR-TB.

Culture and DST, if available, should be used to guide therapy. In other cases, the child should be treated empirically, guided by the DST pattern of the index case. However, every effort should be made to obtain a sample from the child for culture and DST.

Given the severity of DR-TB, there are no drugs that are absolutely contraindicated in children.

Children generally tolerate well second-line anti-TB drugs.

The administration of second-line drugs can be problematic due to the lack of commercially available paediatric formulations.

### 10.9.5 Extrapulmonary drug-resistant TB

Regimen construction and duration for extrapulmonary DR-TB is the same as for pulmonary DR-TB.

If a patient with DR-TB has symptoms suggestive of central nervous system involvement, the regimen should include drugs with good cerebrospinal fluid (CSF) penetration:\[1\][2]:

- Ethionamide or prothionamide and cycloserine have good penetration into the CSF.
- Kanamycin, amikacin, and capreomycin do so only in the presence of meningeal inflammation.
- Para-aminosalicylic acid and ethambutol have little or no penetration.
- Fluoroquinolones have variable CSF penetration, with better penetration seen in the higher generations.

### 10.9.6 Renal insufficiency

Renal insufficiency may be due an injectable anti-TB drug or other aetiologies including longstanding TB infection.

In patients with renal insufficiency, the creatinine clearance should be calculated. If less < 30 ml/min, anti-TB drugs should be adjusted. The formula to estimate the creatinine clearance, and the dose of anti-TB drugs in renal insufficiency are presented in Appendix 12.

### Referencias

10.10 Treatment of extensively drug-resistant TB (XDR-TB)

XDR-TB is much more difficult to treat than other MDR-TB and extremely difficult to treat in HIV-infected patients\(^1\)\(^2\)\(^3\)\(^4\). While reports of HIV-infected patients being promptly diagnosed with XDR-TB and placed on adequate regimen are non-existent to date, a few reports of cohorts of HIV-negative patients have been shown to have cure rates that exceed 50\%(\(^1\)\(^5\)\(^6\)\(^7\)\(^8\)\(^9\)\(^10\)\(^11\)\(^12\)\(^13\)\(^14\)\(^15\)\(^16\)\(^17\)\(^18\)\(^19\)\(^20\)\(^21\)\(^22\)).

There is very limited data on different clinical approaches to XDR-TB. Management of a patient with documented, or almost certain, XDR-TB should be as follows\(^1\)\(^2\)\(^3\):  

1 - Consider a longer duration of use for the injectable agent (12 months or possibly the whole treatment). If the patient’s strain is resistant to all injectable agents, use one the patient has never used before\(^3\)\(^4\)\(^5\)\(^6\).  
2 - Use a third-generation fluoroquinolone such as moxifloxacin. The potential benefit of moxifloxacin should be weighed against the increased risk of QT prolongation when combined with bedaquiline.  
3 - Use all Group 4 agents that have not been used extensively in a previous regimen or any that are likely to be effective.  
4 - Use two or more agents from Group 5. Add bedaquiline. Consider high-dose H if low-level resistance is documented or no katG mutation is detected.  
5 - Use any likely effective Group 1 drugs.  
6 - Consider adjuvant surgery if there is localized disease.  
7 - Consider compassionate use of new agents (Appendix 11).  

Extension of therapy to 24 months is the suggested minimum length of treatment for XDR-TB.

**Box 10.2 - A case of XDR-TB and example regimen**

**Example:**

A patient is receiving Km-Ofx-Eto-Cs-Z and remains smear-positive and culture-positive after 8 months of treatment. In addition the patient is not improving clinically. The DST performed on a sputum collected 2 months ago reveals resistance to H, R, Z, E, S, Km, Cm and Ofx. This patient has XDR-TB.

The regimen should be designed based on the principles described in Section 10.1. Bdq should be considered. A higher generation FQ may have some effect.  

The recommended regimen to be considered in this patient would be:  
Lfx-Cs-PAS-Bdq-Lzd-plus two Group 5 drugs (Cfz-Amx/Clv).  
- Lfx causes less QT prolongation than Mfx.  
- Cfz has an additive effect to the QT prolongation when used with Bdq.  
- ECG monitoring is required.  
- The risk of sudden death versus the benefits of Bdq should be fully explained to the patient.  
- Consider also compassionate use of new anti-TB agents under development.

**Notas**

\(a\) While the reproducibility and reliability of DST to injectables is good, there is little data on clinical relevance of the test. Options with XDR-TB are very limited and some strains may be affected *in vivo* by an injectable agent even though they are testing resistant *in vitro*.  


Chapter 11: Treatment of mono- and poly-drug resistant tuberculosis (PDR-TB)

11.1 Treatment schemes

11.1.1 Choice of the treatment scheme

Mono- and poly-drug resistant tuberculosis (PDR-TB) management is based on the PDR treatment schemes presented in Table 11.1.

Table 11.1 - Resistance pattern and recommended treatment schemes
Sus. = susceptible; Res. = resistant.
The treatment schemes of mono/PDR-TB are based on the assumption that a full baseline drug susceptibility testing (DST) is performed before or at the start of treatment with first line anti-TB drugs. There is little published evidence to determine the best treatment for mono/PDR-TB. The treatment schemes are therefore based on the principles of TB treatment and expert opinion.

At least 3, ideally 4, likely effective drugs are included in the regimen. DST results at baseline and previous treatment history are used to choose the appropriate scheme.

The use of Xpert MTB/RIF can greatly aid in getting patients on the proper regimens when isoniazid resistance is present and amplification of resistance to rifampicin is a possibility.

Perform second-line DST if patients come from a region of high second-line resistance and if there is a history of second-line anti-TB drug use. Resistance to second-line anti-TB drugs will impact the choice of regimen.

### 11.1.2 PDR Scheme A for cases with H or HS resistance

For new patients, the treatment regimen is 9 RZE. However, the combination HRZE can be used if more convenient since it can be given as fixed-dose combination.

At Month 2, perform smear, Xpert MTB/RIF, and culture:

<table>
<thead>
<tr>
<th>Resistance category</th>
<th>H</th>
<th>R</th>
<th>E</th>
<th>S</th>
<th>Treatment scheme</th>
</tr>
</thead>
<tbody>
<tr>
<td>H resistance</td>
<td>Res.</td>
<td>Sus.</td>
<td>Sus.</td>
<td>Sus.</td>
<td>PDR Scheme A(a)</td>
</tr>
<tr>
<td></td>
<td>Res.</td>
<td>Sus.</td>
<td>Sus.</td>
<td>Res.</td>
<td>PDR Scheme A(a)</td>
</tr>
<tr>
<td></td>
<td>Res.</td>
<td>Sus.</td>
<td>Res.</td>
<td>Sus.</td>
<td>PDR Scheme B</td>
</tr>
<tr>
<td></td>
<td>Res.</td>
<td>Sus.</td>
<td>Res.</td>
<td>Res.</td>
<td>PDR Scheme B</td>
</tr>
<tr>
<td>R resistance</td>
<td>Sus.</td>
<td>Res.</td>
<td>Sus.</td>
<td>Sus.</td>
<td>PDR Scheme C</td>
</tr>
<tr>
<td></td>
<td>Sus.</td>
<td>Res.</td>
<td>Sus.</td>
<td>Res.</td>
<td>PDR Scheme C</td>
</tr>
<tr>
<td></td>
<td>Sus.</td>
<td>Res.</td>
<td>Res.</td>
<td>Sus.</td>
<td>PDR Scheme C</td>
</tr>
</tbody>
</table>

(a) Except previously treated patients, for whom PDR Scheme B + ethambutol is preferred.
| Xpert available | **Xpert RIF+**: switch to empiric MDR regimen while waiting for full DST results then, adapt treatment accordingly.  
| **Xpert RIF−**: continue PDR Scheme A. |
|---|---|
| Xpert not available | **Culture+**: switch to empiric MDR regimen with the inclusion of R while waiting for full DST results.  
- DST is unchanged (H or HS resistance only): stop the MDR regimen, and resume PDR Scheme B;  
- DST has changed: adapt treatment accordingly.  
**Culture−**: continue PDR Scheme A. |

Perform smear and culture every other month. If cultures or smears are positive, switch to MDR regimen while waiting for full DST results then, adapt treatment accordingly.

For previously treated patients, it is safer to use Scheme B plus ethambutol, as DST to this drug should not be relied upon if the patient has already received it.

### 11.1.3 PDR Scheme B for cases with HE or HES resistance

Start patients on 3 Cm (or Km)-Lfx-RZ/7 Lfx-RZ regardless of smear status at the time of diagnosis.

At Month 2, perform smear, Xpert MTB/RIF and culture:

| Xpert available | **Xpert RIF+**: switch to empiric MDR regimen while waiting for full DST results then, adapt treatment accordingly.  
| **Xpert RIF−**: continue PDR Scheme B. |
|---|---|
| Xpert not available | **Culture+**: switch to empiric MDR regimen with the inclusion of R while waiting for full DST results.  
- DST is unchanged (HE or HES resistance only): stop the MDR regimen, and resume PDR Scheme B;  
- DST has changed: adapt treatment accordingly.  
**Culture−**: continue PDR Scheme B. |

At Month 3, perform smear, Xpert MTB/RIF, and culture. If Xpert shows RIF+ or if the culture is still positive, this regimen is declared “failure”. Switch to MDR treatment.

Even if found susceptible, streptomycin should not be used given the high rates of resistance to this drug in patients with DR-TB and the poor reliability of the DST.

### 11.1.4 PDR Scheme C for cases with R or RS or RE or RES resistance

Start patient on MDR regimen until confirmation that the strain is susceptible to fluoroquinolones and injectable agents.

When DST results confirm resistance to R, RS, RE or RES and susceptibility to H, fluoroquinolones and an injectable agent, there are two options:

1 - Continue the full course of MDR-TB treatment plus isoniazid. This is a reasonable consideration given that DST reliability is not 100%. This is recommended if the suspicion for MDR-TB is high (i.e. a contact of an MDR-TB patient or failure of a first-line regimen).

2 - Start PDR Scheme C: 3 Cm (or Km)-Lfx-HZ (+/- E)/12 Lfx-HZ (+/- E). Ethambutol is added if it is likely to be effective.

Even if found susceptible, streptomycin should not be used given the high rates of resistance to this drug in patients with DR-TB and the poor reliability of the DST.

At Month 2, perform smear and culture:  
**Culture+**: start empiric MDR regimen and repeat DST.
Culture--: complete PDR Scheme C.

At Month 3, perform smear and culture. If the culture is still positive, this regimen is declared “failure.” Switch to MDR treatment.

Note: if the baseline DST is performed by LPA (Hain® test), only DST for R and H are available. In order to avoid possible resistance amplification, the worst scenario should be assumed:

- If only resistance to H is detected, treat with Scheme B, even new patients while waiting for full DST.
- If only resistance to R is detected, treat as MDR-TB as sensitivity of Hain® test for H resistance is low.

Referencias


   [http://www.currytbcenter.ucsf.edu/drtb/](http://www.currytbcenter.ucsf.edu/drtb/)


11.2 Treatment algorithms for PDR-TB

PDR scheme A
Note: for previously treated patients it is safer to use Scheme B + ethambutol.

PDR scheme B
PDR scheme C

PDR SCHEME B
HE (+/- S) resistance

Start adapted regimen
3 Cm (or Km)-Rf-RZ

At Month 2:
Xpert Rif+ or culture+

Yes
Start empiric MDR-TB treatment
while waiting for DST results.

No
Complete intensive phase

If DST unchanged: complete contin-
uation phase to a total of 7 Lfx-RZ
after culture negativation. Perform
smear and culture every other month.
If DST changed: adapt treatment
accordingly.

At Month 3:
Xpert Rif+ or culture+

Yes

No
Start continuation phase: 7 Lfx-RZ
Perform smear and culture every
other month.

Any smear+/culture+

Yes
Failure
Resume MDR-TB treatment.

No
Cured or treatment completed

PDR scheme C
Chapter 12: Co-management and treatment of HIV in TB disease

12.1 HIV testing and counselling for patients known or suspected to have TB
12.2 Prophylaxis against opportunistic infections
12.3 Anti-TB regimens in HIV patients
12.4 Concomitant treatment TB and HIV
12.5 Drug interactions
12.6 Overlapping toxicities with anti-TB drugs and antiretrovirals
12.7 Immune reconstitution inflammatory syndrome (IRIS)
12.8 HIV-infected children with TB
12.9 HIV-infected pregnant women with TB
12.10 HIV-infected patients with DR-TB
12.1 HIV testing and counselling for patients known or suspected to have TB

HIV testing is recommended for all patients with signs and symptoms of tuberculosis (TB), whether TB is suspected or already confirmed. HIV testing should be offered as part of an “opt out” approach, which means the patients will have to specifically decline the HIV test after receiving the pre-test counselling if they do not want the test performed.

HIV-infected TB patients may have household members who are also living with HIV. Testing for HIV is recommended in immediate family members where horizontal or vertical transmission may have occurred.

12.2 Prophylaxis against opportunistic infections

It is recommended that cotrimoxazole preventive therapy (CPT) be initiated or continued during TB treatment, as it is associated with a reduced risk of death. CPT prevents a number of infections such as pneumocystosis, toxoplasmosis, some diarrhoea and other bacterial infections (respiratory, urinary tract, etc.) and malaria.

If the patient is receiving prophylaxis against other opportunistic infections, the prophylaxis should continue during TB therapy.

12.3 Anti-TB regimens in HIV patients

HIV patients follow the usual first-line or second-line TB regimens. Intermittent regimens should not be used in HIV-positive TB patients.

Case definitions, treatment categories, sputum examination follow-up, and treatment outcomes are equally applicable for HIV-infected TB patients.

12.4 Concomitant treatment TB and HIV

Antiretroviral therapy (ART) dramatically improves survival in HIV-infected patients. In addition, ART reduces TB rates greatly both at individual and population levels.

ART must be initiated in all HIV positive patients with active TB irrespective of the CD4 cell count. Start the anti-TB treatment first, followed by ART as soon as possible and within eight weeks of starting TB treatment [1][2][3].

For the following patients, at high risk of mortality, consider starting ART within the first two weeks:
- Patients with low CD4 count (especially CD4 < 50);
- Young children (especially < 1 year of age);
- Patients with drug-resistant TB (DR-TB).
The first-line ART regimen should contain two nucleoside reverse transcriptase inhibitors (NRTIs) plus one non-nucleoside reverse transcriptase inhibitor (NNRTI). The preferred NNRTI in patients starting ART while on TB treatment is efavirenz (EFV), since there is less interaction between EFV and rifamycins compared to other NNRTIs. The preferred NRTI in the first-line ART regimen is tenofovir (TDF), combined with either lamivudine (3TC) or emtricitabine (FTC). If TDF is not available, then zidovudine (AZT) is preferred over stavudine (d4T) due to the long-term adverse effects.

In summary, for adults and adolescents:

- Tenofovir/lamivudine/efavirenz (TDF/3TC/EFV) is the preferred first-line ART regimen because of the once daily dosing and the availability of a fixed-dose combination.
- In the event of severe central nervous system intolerance to EFV:
  a) Give triple NRTI ART regimen: zidovudine/lamivudine/abacavir (AZT/3TC/ABC);
  or
  b) Replace rifampicin (R) with rifabutin (Rfb, 300 mg daily) and start nevirapine (NVP) based ART regimen with lead-in dosing of NVP for 2 weeks;
  or
  c) Give NVP based ART regimen tenofovir/lamivudine/nevirapine (TDF/3TC/NVP) but without lead-in dose when used with rifampicin containing first-line TB treatment. In patients with CD4 > 250, close clinical and ALT monitoring at 4, 8 and 12 weeks is recommended.

Referencias


12.5 Drug interactions

12.5.1 Antituberculous and antiretrovirals

Rifamycins and antiretrovirals

Interactions between rifamycins and 2 groups of antiretrovirals (ARVs) — NNRTIs and protease inhibitors (PIs) — must be expected due to liver enzyme induction of the rifamycins. For possible combinations of ARVs and rifamycins, see Table 12.1.

Patients receiving NVP when TB is diagnosed:

- If rifabutin is available, give 2 months of HZE-Rfb followed by 4 months of H-Rfb.
- If rifabutin is not available, replace NVP with EFV 600 mg. When the TB treatment is completed, NVP may be resumed.
- If rifabutin is not available and EFV is contraindicated, see options in previous section.

Patients receiving protease inhibitors (PI):

- When PIs and rifamycins are given to the same patient, PI serum levels can decrease to sub-therapeutic levels, while the serum levels of rifamycins could rise to toxic levels.
- Rifabutin is a less potent enzyme inducer than rifampicin; rifabutin is the preferred drug in patients using PIs.
- If rifabutin is not available, dosages of lopinavir and ritonavir (LPV/r) must be significantly increased in patients taking both LPV/r and rifampicin (see Table 12.1). Liver enzymes should be monitored.
### Table 12.1 Possible combinations of ARVs and rifamycins

<table>
<thead>
<tr>
<th></th>
<th>Rifampicin</th>
<th>Rifabutin</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>NNRTIs</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nevirapine (NVP)</td>
<td>Do not combine unless Rfb is not available and there are no other options.</td>
<td>Rfb: 300 mg/day NVP: usual dose</td>
</tr>
<tr>
<td>Efavirenz (EFV)</td>
<td>May be combined. R: usual dose EFV: 600 mg/day</td>
<td>–</td>
</tr>
<tr>
<td><strong>NRTIs</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abacavir (ABC)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Didanosine (ddI)</td>
<td>May be combined without dose adjustments.</td>
<td>–</td>
</tr>
<tr>
<td>Lamivudine (3TC)</td>
<td></td>
<td></td>
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<tr>
<td>Stavudine (d4T)</td>
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<td>Zidovudine (AZT)</td>
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<tr>
<td>Tenofovir (TDF)</td>
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<tr>
<td><strong>PIs</strong></td>
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<tr>
<td>Indinavir (IDV)</td>
<td>Do not combine</td>
<td>Rfb: 300 mg/day IDV: 1 g every 8 hours</td>
</tr>
<tr>
<td>Nelfinavir (NFV)</td>
<td>Do not combine</td>
<td>Rfb: 300 mg/day NFV: usual dose</td>
</tr>
<tr>
<td>Lopinavir/ritonavir (LPV/r)</td>
<td>May be combined if Rfb is not available. LPV/r: double dose of LPV/r (800 mg/200 mg twice daily) R: usual dosage</td>
<td>Rfb: 150 mg/day[1] LPV/r: usual dose</td>
</tr>
<tr>
<td>Atazanavir/ritonavir (ATZ/r)</td>
<td>Do not combine</td>
<td>Rfb: 150 mg/3 times a week[2] ATZ/r: usual dose</td>
</tr>
</tbody>
</table>

**Bedaquiline and antiretrovirals**[3][4]

Data from drug-drug interaction studies with bedaquiline and ARVs to date are extremely limited.

- **NNRTI**: EFV (enzyme inducer) is estimated to decrease bedaquiline concentrations by 50%. Nevirapine does not significantly affect bedaquiline concentrations.
- **NRTI** are unlikely to affect bedaquiline concentrations.
- **PI**: ritonavir is an enzyme inhibitor. The use of ritonavir-boosted lopinavir (LPV/r) with bedaquiline may result in a significant accumulation of bedaquiline and its metabolites. This combination is therefore not recommended.
The following ART regimens can therefore be considered in association with bedaquiline:
1) 2 NRTIs + nevirapine: e.g. AZT/3TC or FTC/NVP or TDF-3TC-NVP;
or
2) 3 NRTIs: e.g. AZT/3TC/ABC.

**Fluoroquinolones and didanosine**

Buffered didanosine contains an aluminium/magnesium-based antacid that, if given with a fluoroquinolone, can result in its decreased absorption. If it is not possible to avoid prescribing these drugs together, didanosine should be given 2 hours apart fluoroquinolone administration. The enteric-coated formulation of didanosine can be given without such precaution.

Other drug-drug interactions can occur between anti-TB drugs and ARVs, further complicating treatment. Most of the drugs used in the treatment of DR-TB have not had drug-drug interaction studies performed with ARVs.

**12.5.2 Other interactions**

Rifampicin can interact with drugs commonly used in opportunistic infections.

Interaction occurs with fluconazole. Rifampicin may decrease blood levels of fluconazole by as much as 25-50%. The two drugs can be taken 12 hours apart (i.e. rifampicin in the morning, fluconazole in the evening) without dosage adaptation. However, the patient’s clinical condition should be carefully monitored, as the dosage of fluconazole may need to be increased if clinical improvement is suboptimal.

For the treatment oral candidiasis, miconazole mucoadhesive tablets (gum patches) can be used (no interaction with rifampicin).

**Notas**

(a) If a patient is changed from EFV back to NVP on completion of TB treatment, no lead-in dosing of NVP is necessary.

**Referencias**


3. Sundari Mase, Terence Chorba, Philip Lobue, Kenneth Castro. Provisional CDC Guidelines for the Use and Safety Monitoring of Bedaquiline Fumarate (Sirturo) for the Treatment of Multidrug-Resistant Tuberculosis. MMWR Recommendations and Reports, October 25, 2013/62(rr09);1-12.


**12.6 Overlapping toxicities with anti-TB drugs and antiretrovirals**

The main potential overlapping toxicities between anti-TB drugs and ARVs are:
- Hepatic reactions;
The use of agents with shared adverse effect profiles should be avoided if possible. Often, however, the benefit of using drugs that have overlapping toxicities outweighs the risk. Thus, if two drugs with overlapping toxicities are essential in a regimen, increased monitoring for potential adverse effects is recommended rather than avoidance of a certain combination.

Important points:
- HIV patients are more likely to develop isoniazid-related peripheral neuropathy. Thus, all patients on isoniazid should receive pyridoxine PO (vitamin B₆): 10 mg daily or 25 mg twice a week.
- The use of thiacetazone is contraindicated in HIV patients due to the high frequency of Stevens-Johnson syndrome and corresponding risk of mortality.
- Due to reports of increased renal toxicity during concurrent use of TDF and injectable agents (kanamycin, amikacin, and capreomycin), the use of TDF is not recommended during the intensive (i.e. injectable) phase of DR-TB treatment. If TDF is absolutely necessary, serum creatinine and creatinine clearance, and electrolytes should be monitored frequently.

For potential overlapping toxicities of antiretrovirals and anti-TB drugs, see Appendix 19.

12.7 Immune reconstitution inflammatory syndrome (IRIS)

Immune reconstitution inflammatory syndrome (IRIS) occurs after initiation of ART, in the presence of a previously unrecognized, often subclinical, opportunistic infection. Patients present with paradoxical worsening of their clinical status as the immune system recovers on ART.

Symptoms vary according to the infection, but can include fever, enlarging lymph nodes, worsening pulmonary infiltrates, respiratory distress, neurologic signs, or exacerbation of inflammatory changes at other sites.

IRIS can occur anytime between 10 and 180 days after ART initiation (usually 2-4 weeks) and is more common with CD4 count < 50. Mild to moderate forms of IRIS are relatively common in TB patients who are started on ART (seen in up to one third of patients in some studies); however, severe IRIS is relatively rare.

IRIS occurs with respect to TB in two circumstances:
1 - Paradoxical TB IRIS: A patient is diagnosed with TB, starts TB treatment, followed by ART after a few weeks, and then develops IRIS.
2 - Unmasking TB IRIS: A patient is screened for TB before initiation of ART and no TB is found. The patient then starts ART, followed by onset of TB symptoms and signs.

It is important to note that IRIS with respect to TB is a diagnosis of exclusion. Patients with advanced HIV infection may show clinical deterioration for a number of other reasons (all of which should be ruled out before giving the diagnosis of IRIS):
- Clinical worsening due to new opportunistic infections;
- Other subclinical infections unmasked following immune reconstitution with ART initiation;
- TB treatment failure due to DR-TB.

The management of IRIS depends on the clinical status of the patient and the site and extent of involvement. Nonsteroidal anti-inflammatory drugs are used in mild to moderate IRIS cases and corticosteroids in severe IRIS cases (Chapter 9, Section 9.3). The use of corticosteroids can be dangerous if IRIS is misdiagnosed and the clinical deterioration is in fact DR-TB or a different opportunistic infection.

Most cases of IRIS can be treated without interruption of ART; in very severe forms of IRIS, ART may need to be suspended.

12.8 HIV-infected children with TB

- Cutaneous reactions;
- Neuropathy;
- Nephrotoxicity.

HIV patients are more likely to develop isoniazid-related peripheral neuropathy. Thus, all patients on isoniazid should receive pyridoxine PO (vitamin B₆): 10 mg daily or 25 mg twice a week.

The use of thiacetazone is contraindicated in HIV patients due to the high frequency of Stevens-Johnson syndrome and corresponding risk of mortality.

Due to reports of increased renal toxicity during concurrent use of TDF and injectable agents (kanamycin, amikacin, and capreomycin), the use of TDF is not recommended during the intensive (i.e. injectable) phase of DR-TB treatment. If TDF is absolutely necessary, serum creatinine and creatinine clearance, and electrolytes should be monitored frequently.
Most HIV-positive children with TB respond well to the 6-month TB regimen, similar to HIV-uninfected children. If the clinical response is slow, other causes should be considered such as poor adherence to therapy, inadequate drug absorption, DR-TB, and other infections.

The following ARV regimens are preferred in children on TB treatment:

- Child < 3 years old or < 10 kg: AZT preferred or D4T/3TC + ABC;
- Child > 3 years and > 10 kg: AZT preferred or ABC or D4T/3TC + EFV.

TDF is considered safe in children above 3 years of age.

### 12.9 HIV-infected pregnant women with TB

TB in HIV-positive pregnant/postpartum women is associated with significant maternal and infant mortality. ART in pregnant women with TB is summarized below:

- TDF is the preferred NRTI and is safe to use throughout pregnancy.
- Safety of EFV is considered acceptable during pregnancy.[1]

**Referencias**


### 12.10 HIV-infected patients with DR-TB

DR-TB does not appear to be more prevalent in HIV-infected patients compared to HIV-uninfected patients. However, high mortality rates have been reported in patients coinfected with HIV and DR-TB.

Prompt initiation of appropriate DR-TB therapy (and subsequent initiation of ART) can help to reduce mortality.

**Chapter 13: Adherence to tuberculosis treatment**

13.1 Introduction

13.2 Treatment delivery model

13.3 Factors that influence adherence

13.4 Patient education and support

**Update: January 2022**
13.1 Introduction

Good adherence is when the patient follows the treatment as prescribed. Patient understanding, acceptance and motivation to start and complete TB treatment are essential to maximise chances of cure. Good knowledge of drug dosing, length of treatment, required clinical follow-up and common adverse effects help patients to follow the prescribed therapy.

Failure to take tuberculosis (TB) drugs consistently, or in an inappropriate manner, or stopping the treatment too soon, can lead to treatment failure or relapse. It may also contribute to the development of resistance, which can complicate subsequent treatment, thereby decreasing the chances of a successful outcome.

13.2 Treatment delivery model

13.2.1 Self-administered treatment

Self-administered treatment (SAT) is taken autonomously by the patient without daily supervision. The patient is seen at a health facility at regular intervals (e.g. monthly) to receive drugs, support and treatment education. SMS telephone reminders may be considered to reinforce adherence.

13.2.2 Directly observed therapy

Drugs are sometimes provided daily to the patient and the treatment is taken under direct observation (DOT) by a third party.

DOT may be provided:
- In health facilities (facility-based DOT): in this model, DOT is implemented in a centralised setting and treatment is administered by healthcare workers.
- Outside of health facilities (community or home-based DOT): in this model, DOT is implemented in a decentralised setting and is usually provided by supervised, trained and remunerated treatment supporters.
  For the roles and responsibilities of treatment supporters see Appendix 20.
- Remotely (video-observed therapy or VOT): VOT uses secure Internet connections via a smart phone or computer application to remotely supervise patients taking their treatment.

DOT is labour-intensive to implement and can be inconvenient for patients. Community and home-based DOT and VOT require fewer resources (personnel and transport) than facility-based DOT and may be more convenient for patients.

Box 13.1 – Recommended treatment delivery models
13.3 Factors that influence adherence

Several factors can influence adherence, including barriers related to the patient, the treatment or the therapeutic environment. While it is not always feasible to address all these factors, at the very least it is possible to control the treatment and therapeutic environment-related factors.

13.3.1 Patient-related factors

A discussion should be held with the patient prior to treatment initiation and then during every contact they have with the healthcare team. The objective is to identify and anticipate barriers to treatment adherence. Barriers may include:

- Socioeconomic factors (work and home responsibilities, treatment-related costs, decreased income, etc.).
- Psychological factors (feelings of shame, fear of stigma or marginalisation, uncertainty about the future, conceptions about the disease and its treatment, etc.).
- Physical or mental disability.
- Lack of knowledge about the disease and treatment.
- Perception of the disease and treatment (a patient might abandon treatment due to improvement or absence of improvement, a negative experience with a previous treatment, etc.).

Solutions depend on the context and the patient’s problem, and therefore should be identified on a case-by-case basis.

13.3.2 Treatment-related factors

Drug-susceptible TB (DS-TB)

- DOT has not been proven to improve treatment outcomes for DS-TB when compared to SAT in controlled trials[1].
- When there is no factor to complicate adherence, and provided the patient receives appropriate support, treatment should be self-administered.
- There are some situations in which DOT may be preferred:
  - Patients with mental health issues or serious socioeconomic problems (e.g. the homeless) and all patients incapable of taking drugs on their own.
  - Prisoners (risk of drugs being sold or stolen).

Drug-resistant TB (DR-TB)

- Due to the lack of fixed-dose combinations (FDC), length of treatment, adverse effects of TB drugs and lack of therapeutic alternatives if treatment fails, patients usually require reinforced support.
- If DOT is considered useful, home-based DOT[2] or VOT are preferred to facility-based DOT. A combination of approaches may be required for some patients.

Latent TB infection (LTBI)

- LTBI treatments can be self-administered.
- DOT may be preferred with the 3HP regimen, as it may cause serious hypersensitivity reactions. However, SAT can be considered if the patient well informed and is able to seek rapid medical attention if adverse effects develop.

References


• Simplicity of treatment improves adherence. The use of FDC simplifies the treatment by reducing the number of tablets. In addition, FDC prevents omission of one or more prescribed TB drugs.
• Adverse effects may lead patients to interrupt their treatment, so these should be detected and managed promptly.

13.3.3 Factors related to the therapeutic environment

• To ensure the widest possible access to treatment, TB diagnosis, monitoring and treatment (including TB drugs and drugs for adverse effects and co-morbidities) should be provided free of charge.
• The relationship between patients and healthcare workers influences if patients have confidence in healthcare workers, they are more likely to follow recommendations and engage with the treatment process. Patients are also more likely to bring questions and concerns to the attention of healthcare workers. The same applies to the relationship with treatment supporters.
• In health facilities, the way in which patients are received is Waiting times for diagnosis or follow-up visits should be reasonable.
• Drug supply management must be rigorous. Shortages can lead to treatment interruption and negatively impact adherence (patients waste time in unnecessary travel and lose confidence in the health facility).
• The proximity of drug distribution sites limits the number of patients who abandon due to transportation problems. To anticipate potential problems, give the patients a few extra days of treatment in case they are unable to come to get their drugs on the scheduled.
• For the co-management of TB and HIV infection, patients should receive TB and HIV treatment at the same time and in the same place (“one-stop service”). This reduces the number of visits and decreases waiting times, which results in greater patient satisfaction and improved treatment outcomes. Co-management of other co-morbidities (e.g. diabetes, hypertension) should, when possible, use the same approach.
• Hospitalisation should be limited to patients with clinical conditions requiring hospital level care. If hospitalisation is necessary, accommodation (comfort, food, heating, etc.) should be adequate. The duration of stay should be as short as possible and patients should be discharged as soon as their clinical condition allows.

13.4 Patient education and support

Patient education and support require the involvement of the entire healthcare team (clinicians, nurses, treatment supporters, social workers, etc.). In large-scale programmes, the healthcare team sometimes includes trained counsellors who provide information and support.

Treatment education and support may be provided through various channels: organising educational sessions during in-facility or home visits, video and telephone contacts.

Patient education and support are required throughout treatment, as adherence may vary over time and patients may experience phases of treatment acceptance and rejection.

Due to the toxicity and long duration of treatment, patients on DR-TB treatment usually require substantial support.

13.4.1 Patient education

Patient education consists of:
• Helping patients to understand the disease and treatment.
• Enabling patients to acquire and maintain skills that allow them to manage their treatment and disease in their everyday lives.
• Answering patients’ questions throughout the treatment.

For more information see Appendix 21.

13.4.2 Emotional support

Listen to patients and give them encouragement, so that they feel comfortable saying they have forgotten or have made a mistake with their treatment. This is common, and it is important to know so that solutions can be found.

Psychological problems, such as depression and anxiety are frequent, and may have a negative impact on adherence. The healthcare team should be sensitised to their early detection and management.
13.4.3 Social support

Implement social support measures for patients with limited resources. Depending on the situation and specific needs of patients:

- Social workers can help to obtain disability allowances, housing assistance, shelter for the homeless, etc.
- The programme can provide meals or food, vouchers or money for transportation or reimburse the cost, etc.

Chapter 14: Tuberculosis infection control

14.1 Introduction

14.2 Implementation of TB IC strategies

14.3 Administrative controls

14.4 Environmental controls

14.5 Personal protective measures

14.6 Hospital hygiene

14.7 Patients’ homes

14.1 Introduction

The largest source of *M. tuberculosis* transmission is the contagious patients with respiratory tuberculosis not yet diagnosed and put on treatment. Therefore, tuberculosis infection control (TB IC) relies, above all, on:

- Early diagnosis (including in clinics and any non-tuberculosis medical wards, whereby active case finding through cough surveillance of all admissions should avoid days or weeks of transmission from unsuspected TB cases);

AND

- Prompt implementation of effective treatment. With effective treatment, contagiousness decreases even after a few days and may be considered nil after 2 to 3 weeks of treatment[1][2][3][4]. It is essential the treatment is “effective,” as multidrug-resistant TB (MDR-TB) patients that are placed on first-line anti-TB drugs are likely to remain contagious.

However, in health care facilities where TB patients or persons suspected of having TB congregate, additional measures are needed to reduce the risk of transmission between patients, to health care staff and to vulnerable (particularly immunocompromised) patients/visitors[6].

TB infection control (IC)* consists in different strategies for preventing transmission of TB in health care facilities.

Notas

(a) This chapter reviews the basic TB IC strategies. More in depth information can be found from the Tuberculosis Coalition for Technical Assistance which has published a framework and developed a website [http://www.tbcta.org/Library](http://www.tbcta.org/Library) that provides a comprehensive set of examples.

Referencias

14.2 Implementation of TB IC strategies

There is a trio of infection control levels, which include (1) administrative, (2) environmental and (3) personal protective controls. The implementation of these measures requires a dedicated staff and an IC plan.

14.2.1 Infection control practitioner

A person should be clearly identified and designated as responsible for TB IC. This person should have the support and authority to conduct, apply and evaluate TB IC policies. This person in some settings is also known as IC officer.

14.2.2 Infection control committee

The IC practitioner would evaluate the need to create an infection control committee (ICC). The ICC might include doctors, nurses, laboratory technicians, logisticians and administration staff (including representation from the maintenance and housekeeping services). According to the context and degree of risk, experts in IC may be needed.

14.2.3 Infection control plan

All facilities should have a detailed written IC plan that is at least annually updated and distributed to healthcare staff. A simplified version of the plan must be accessible to all healthcare workers including staff not directly involved in TB patients’ management, such as cleaners, kitchen staff, etc.

The first step in developing an IC plan is assessing the health care facility’s risk for TB transmission. This should be performed by the IC practitioner. The plan must be specific to each facility. An example of risk assessment tool is given in Appendix 16.

The IC plan should include the different types of measures—administrative, environmental and personal. Information on specific precautions and procedures for high-risk areas should be detailed.

It is recommended to draw a floor plan of the facility with the different areas, including the patient flow and identifying areas of high risk.

Listed below from highest to lowest level of risk:

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https://apps.who.int/iris/bitstream/handle/10665/44148/9789241598323_eng.pdf?sequence=1
14.3 Administrative controls

The administrative controls aim at preventing the exposure to infectious droplet nuclei.

14.3.1 Patients triage

Upon entry into the health facility, a member of the medical staff should identify patients with a cough as soon as possible. Patients with a cough over two weeks should be sent to a separate waiting room if possible.

All patients with cough (including patients with less than two weeks of cough) should receive tissues or face masks, and they should be requested to cover their mouth and nose when they cough.

14.3.2 Patient, visitors and attendants' flow

Inside the TB department, circulation of patients and attendants is controlled:

- Encourage patients/attendants to spend as much time as possible outdoors if weather permits or in areas that are open on three or four sides.
- Have visible signage on entry doors to TB wards that forbid visitors to enter.
- Limit visitation duration, particularly for contagious patients.
- Encourage visits outside the building, especially for contagious patients.
- Have visiting areas well identified with signage.

### Referencias


### Tables

<table>
<thead>
<tr>
<th>Highest risk</th>
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<tbody>
<tr>
<td>- Smear-positive inpatient unit</td>
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<tr>
<td>- Diagnosis department</td>
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<tr>
<td>- Culture/drug susceptibility test (DST) and sputum smear preparation area (laboratory)</td>
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<tr>
<td>- Sputum collection area</td>
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<tr>
<td>- Radiology department</td>
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<td>- Waiting area</td>
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<tr>
<th>Limited risk</th>
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<tbody>
<tr>
<td>- Children inpatient ward</td>
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<tr>
<td>- Extrapulmonary TB (EPTB) and smear-negative unit</td>
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<tr>
<td>- Sputum reception and smear reading area (laboratory)</td>
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<td>- Waste management area</td>
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<tr>
<th>Lowest risk (non-TB zone)</th>
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<tr>
<td>- Kitchen area</td>
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<tr>
<td>- Administration</td>
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</table>
14.3.3 Segregation of hospitalized patients

Patients should preferably be treated in ambulatory care. Hospitalisation should be limited and reserved for clinically unwell patients.

TB wards must be separated from the others wards in the health structure compound.

Ideally, within the TB department, patients should be placed in single rooms. If this is not possible, cohort isolation must be implemented and different sections should be labelled according to the degree of contagiousness (smear/culture status) and risk of resistance.

The following is one scheme of separation. It does involve the use of some single isolation rooms (all TB inpatient facilities should have some isolation rooms. If none exist, a very high priority is to add some).

- Smear-positive patients with proven or suspected DR-TB, including chronic cases and retreatment cases that are likely to have MDR-TB. MDR-TB cases should have single isolation rooms (place in 2 to 4 person rooms with other MDR-TB patients if there are no single rooms and try to match DST patterns). It is particularly important not to mix MDRTB patients with extensively drug-resistant TB (XDR-TB) patients.
- Smear-positive patients with fully susceptible TB.
- Smear-negative patients (or patients who have converted), with proven or suspected DRTB (once patients are on effective treatment, they rapidly become non-contagious).
- Less or non-contagious TB: patients with smear-negative pulmonary TB (PTB), EPTB, patients having converted their sputum/culture and most children.
- Patients who are undergoing diagnosis as suspected cases: when possible do not hospitalize patients for diagnosis. If hospitalization is necessary, these patients need isolation rooms. Never put a patient who is not receiving TB medications in a TB ward.

If women and men are to be separated, this scheme requires at least 8 different wards and enough single rooms for suspect cases and MDR-TB patients.

14.3.4 TB IC training

All healthcare personnel should receive initial training on TB transmission, information on high-risk areas in the facility and on protective measures. Continuing education should be offered annually.

The training should also include how staff can teach patients, visitors and attendants about the risk of TB transmission and how to avoid it (cough etiquette, use of masks and respirators).

14.4 Environmental controls

The environmental measures aim at reducing the concentration of infectious droplet nuclei in the air.

14.4.1 Ventilation

Ventilation (replacement of inside air with outside air) is the most effective means for reducing the concentration of *M. tuberculosis* in the air, and as a result, the risk of transmission.

The WHO recommends that in areas where TB transmission might occur, a minimum ventilation rate of 12 air changes per hour (ACH) should be achieved. See Appendix 17 for recommendations on ACH measurement.

Effective ventilation can be obtained by natural (assisted or not) or mechanical means.

**Natural ventilation**
Natural ventilation, especially cross-ventilation (windows/doors in opposite sides of the room), has the best cost-effective ratio. It should be done with the windows and outside doors open (as much as weather conditions permit). Inside doors should be closed so that the flow of air is directed outside and not toward the corridors. Create shady spaces so that patients, attendants and visitors can stay outside during the day. Wind-driven roof turbines (whirly birds) or chimneys can also be used to improve natural ventilation, in that they can keep the principle of directing room air towards the exterior. In addition, fans can be used when the natural ventilation flow rate is too low (assisted natural ventilation).

**Mechanical ventilation**

When natural ventilation cannot reach adequate rates, centralised mechanical ventilation should be considered in some settings, such as within cold climates. Centralised mechanical ventilation relies on the use of mechanical equipment to maintain an air pressure difference between two areas in order to draw air into a room and vent it to the outside. It requires continuous and meticulous maintenance, which renders it costly and difficult to implement and operate.

Advantages and disadvantages of each ventilation technique are presented in Appendix 18.

### 14.4.2 Architectural considerations

Airborne infection control should be always considered during the planning/construction stages of new health facilities and those being modified. It is important to achieve the following:

- Building layout and design with maximised natural ventilation (assisted or not) and sunlight. Waiting areas should be open on three sides. Design of TB wards should avoid internal hallways with doors from the rooms and wards opening into them. Instead, doors should open to outside hallways that are open to air (this may not be feasible in cold climates).
- Specific areas (open air, sputum collection booth, etc.) should be reserved for procedures with a high risk of *M. tuberculosis* transmission (e.g. sputum collection, sputum induction, etc.).
- Allow patient flow that reduces exposure of patients at risk to patients that are infectious (e.g. separate waiting rooms for different cohorts, one patient per room in a hospital). If designing a new TB ward, incorporate plenty of single rooms or at least small rooms with 2 to 4 beds for easier separation of the different cohorts of patients. General hospitals should also have isolation rooms available for TB suspects and contagious patients.

Rehabilitation of existing structures in order to maximise natural ventilation could be a viable economical option instead of building expensive systems, like centralised mechanical ventilation.

### 14.4.3 Ultra-violet germicidal irradiation

Ultra-violet germicidal irradiation (UVGI) lamps® may be used when adequate ventilation cannot be achieved in high-risk areas. When properly installed, designed, maintained and operated, an UVGI system, in addition to 6-12 ACH ventilation, could be the equivalent of 10-25 ACH[7]. For technical information on upper room UVGI, see Appendix 19.

- Main requirements and constraints in UV lamps usage include:
  - Expertise in installation and testing;
  - Rigorous monitoring and maintenance;
  - Electricity, relative humidity less than 70, good air mixing.
- Potential hazards include: Transient eye and skin injuries from overexposure, mercury poisoning (broken or mishandled lamp).

### 14.4.4 Areas requiring specific measures

**Sputum collection areas**
These areas must be settled, wherever possible, outside in open air where bacilli will naturally be dispersed by wind rather than in a closed room where the concentration of bacilli will be high.

In cold regions, sputum collection should be performed in very well ventilated indoor rooms (at least 20 ACH) or in well ventilated rooms (at least 12 ACH) equipped with a UVGI system.

Another option for sputum collection areas in cold climate regions is to assign a specific room of small size (1 m²) with one single glass door opening outside. Keep the door largely open for 5 minutes between each patient. The small volume of air in this room facilitates rapid ventilation.

**Laboratory**

All laboratories should undergo a risk assessment, and IC measures should be adapted accordingly. In any case, limit the access to all TB laboratories.

The use of ventilated workstation (Appendix 7) is strongly recommended for smear preparation (microscopy and test Xpert). In laboratories where culture are carried out, biological safety cabinets type II must be used.

Laboratories must have easy to clean working surfaces (avoid wood) to allow proper disinfection. They should also have large windows to let in sunlight and allow natural ventilation if the laboratory has no mechanical ventilation.

Water-filters should be used to avoid contamination by saprophyte mycobacteria that are sometimes present in the water.

**Notas**

(a) UVGI inactivate bacilli. Natural light dries the droplet but does not inactivate bacilli.

**Referencias**


**14.5 Personal protective measures**

Personal protective measures aim at minimising the risk of bacillus transmission by providing barriers to inhaling or exhaling infectious droplet nuclei.

**14.5.1 Respirators (or high-filtration masks or anti-inhalation masks)**

A respirator is personal protective equipment that prevents inhalation of infectious droplet nuclei by the person who wears it.

**Exposed staff**

Staff must wear a respirator, regardless if they are the caregiver or not. Respirators should be worn:

- When in contact with contagious patients (suspect or confirmed TB case);
- When collecting sputum samples;
- When collecting and disposing of sputum containers;
- In areas where droplet nuclei could be present (i.e. a room that has been occupied by a TB case, prior to the time required for air cleaning).

Using respirators needs proper training, fit testing and continuous supervision. This also applies to home-based DOT supervisors.

**Visitors/attendants**

Visitors and attendants must wear a respirator when entering a contagious TB patient’s room.
14.5.2 Face or surgical masks

Face masks are medical devices that prevent patients from spreading infectious droplets when talking, coughing or sneezing. They should be worn by contagious patients (suspect or confirmed) when they leave their rooms to go to another department or any other enclosed area. They should not be worn when the patient is alone in his/her room and outdoors.

For more information on surgical masks, see Appendix 28.

Using a mask in public areas could be stigmatizing. Patients can use a cloth scarf to achieve the same purpose.

14.6 Hospital hygiene

14.6.1 Hygiene and disinfection

Sputum containers

Patients with pulmonary TB produce sputum that may contain tubercle bacilli.

- In the wards, patients' sputum containers should be large (about 200-ml), non-sterile, and sealable. They are to be replaced daily and cannot be re-used.
- In the laboratories, containers for sample collection are smaller (25-35 ml), with hermetic screw cap, non-sterile and for single use.

Environmental cleaning

Sterilization or the use of disinfecting chemicals in a TB patient’s room is not necessary. Ordinary cleaning of rooms and objects (linens, dishes, etc.) used by TB patients is sufficient. After the patient is discharged, air the empty room well according to the calculated ACH.

Reusable medical items

Standard operating procedures for reprocessing items should be followed. There are no specific measures for TB services.

Standard precautions

Standard precautions (hand hygiene, gowns, etc.) apply in TB wards, as they do in any other hospital department.

14.6.2 Waste management

Standard operating procedures for handling and the disposal of healthcare waste (including soft, sharp, etc.) should be followed. There are no specific measures for TB services.

Note: used sputum containers should be collected in a leak proof trash bag and incinerated without filling the containers with chlorine solution before incineration (this can produce toxic gases).

14.7 Patients’ homes

In settings where DR-TB (and HIV) is highly prevalent, systematic TB IC evaluations on patients’ homes are recommended.

TB IC at patients’ homes follows the same principles and measures as in healthcare facilities. Administrative, environmental and personal measures should be followed at least until patient’s smear is negative, ideally until culture conversion.

Administrative measures

- Assess the risk of TB transmission: gather information on the number of people that live in the house, number of rooms, etc.
Chapter 15: Follow-up of staff exposed to tuberculosis

15.1 Introduction

The following recommendations apply to staff who work in health facilities and are in contact with tuberculosis (TB) patients and/or infectious laboratory specimens.

They provide general guidance, but should be adapted to the context and regulations of each country.

15.2 Baseline assessment

New staff should undergo a baseline assessment. This includes:

- BCG status (BCG scar check)
- Tuberculin skin test (TST) or interferon gamma release assay (IGRA)
In addition, the following information should be provided:

- Risk of occupational transmission of *M. tuberculosis*
- Infection prevention and control (IPC) measures to reduce the risk of transmission
- Higher risk of active TB in immunocompromised individuals (e.g. HIV-infected, diabetics) and in pregnant women
- Vigilance required for, and self-reporting of, signs and symptoms suggestive of TB

Immunocompromised staff and pregnant women should not work in TB departments or areas where the risk of exposure to *M. tuberculosis* is high (Chapter 14).

### 15.3 BCG vaccination

Recommendations vary between countries, with some requiring staff to be BCG vaccinated if never vaccinated and TST negative.

There is limited evidence regarding the benefits of BCG vaccination in adults who have not previously had BCG vaccination[1]. Vaccination should be considered on a case-by-case basis in the following situations[2]:

- Significant exposure to multidrug-resistant TB (MDR-TB): facilities treating MDR-TB, prisons, or areas with high MDR-TB prevalence.
- While corrective actions are implemented:
  - when transmission of MDR-TB to staff has occurred;
  - when IPC measures are inadequate or poorly applied.

The following information should be provided to staff considered for BCG vaccination:

- Benefits and risks of BCG vaccination.
- Impact of BCG on the interpretation of TST results in diagnosing a potential latent TB infection (LTBI).
- No complete protection conferred by the vaccine: TB may still occur if other IPC measures are not used.

BCG vaccine should only be administered if the person:

- Is HIV-negative.
- Is not pregnant.
- Has never had a BCG vaccination.
- Has never had active TB.
- Has a TST negative result.

For more information on BCG vaccine see Appendix 29.

### Referencias


### 15.4 Follow-up
Follow-up of routinely exposed staff includes:

- An annual clinical evaluation.
- Assessment for TB (including CXR) and HIV, if symptomatic.

For staff who were TST or IGRA negative at baseline, TST may be performed once a year.

Staff working in a TB department and presenting with a recent immunodepression (e.g. HIV infection, immunosuppressive treatment) or a pregnancy, should be transferred to another department or to an area within the TB department where the risk of exposure to *M. tuberculosis* is low (Chapter 14).

LTBI treatment (Chapter 16) should be offered, after exclusion of active TB:

- Once to staff who become TST or IGRA-positive.
- To all HIV-infected staff.

## Chapter 16: Treatment of latent tuberculosis infection

### 16.1 Introduction

### 16.2 Target populations

### 16.3 Latent tuberculosis infection treatment regimes

### 16.4 Latent tuberculosis infection in HIV-infected patients

### 16.5 Latent tuberculosis infection in household contacts

### 16.6 Latent tuberculosis infection in other individuals at risk

### 16.7 Latent tuberculosis infection and multidrug-resistant tuberculosis

### 16.8 Follow-up for patients treated for latent tuberculosis infection

Update: January 2022

16.1 Introduction

Exposure to *M. tuberculosis* may result in latent tuberculosis infection (LTBI). WHO defines LTBI as a state of persistent immune response to stimulation by *M. tuberculosis* antigens with no evidence of clinically manifest active tuberculosis (TB)\(^n\). This is also referred to as “tuberculosis infection”.

Identification and treatment of LTBI can reduce TB morbidity and mortality, as well as TB transmission.

Tuberculin skin test (TST) or interferon-gamma release assay (IGRA) can be used to detect LTBI (Chapter 3).

The goal of LTBI treatment is to reduce the risk of progression to active TB. It must be initiated only once active TB has been ruled out by appropriate evaluation.

If a patient develops signs and symptoms of active TB while on LTBI treatment, a specimen should be taken for diagnosis and detection of drug resistance (Xpert MTB/RIF, Xpert MTB/XDR, culture and drug susceptibility test, DST) and according to the results, TB treatment should be initiated.
16.2 Target populations

TST or IGRA cannot predict which patients with LTBI are likely to develop active TB. Therefore, widespread LTBI testing and treatment are not recommended.

However, in certain populations, the risk of progression to active TB significantly exceeds that of the general population. For these at-risk populations, the benefits of LTBI treatment of preventing active TB and TB transmission outweigh the potential risks.

Populations who benefit most from LTBI treatment include:

- HIV-infected individuals.
- Household contacts of patients with bacteriologically confirmed pulmonary TB (PTB), in particular children under 5 years.
- Other individuals or populations at risk (e.g. health staff, prisoners).

16.3 Latent tuberculosis infection treatment regimens

There are 3 recommended LTBI treatment regimens and 2 alternative treatment regimens[^1]. The decision to prescribe one regimen rather than the other should take into consideration:

- Drug-susceptibility of the strain of the presumed source patient, if known.
- Co-morbidities (e.g. HIV infection, pre-existing hepatic disease or neuropathy).
- Risk of drug interactions (especially with antiretrovirals), tolerability, length of treatment and likelihood of adherence.
- Individual characteristics (e.g. age, pregnancy, living conditions, individual preference).
- Epidemiological and programmatic aspects (e.g. HIV prevalence, available drugs, national recommendations).

Table 16.1 - LTBI treatment regimens

---

### Recommended regimens

<table>
<thead>
<tr>
<th>Regimen</th>
<th>Isoniazid PO once daily:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid daily for 6 months (6H) or 36 months (36H)</td>
<td>Isoniazid PO once daily: &lt; 30 kg: 10 mg/kg (7 to 15 mg/kg)</td>
</tr>
<tr>
<td></td>
<td>≥ 30 kg: 5 mg/kg (4 to 6 mg/kg)</td>
</tr>
<tr>
<td></td>
<td>(max. dose 300 mg daily)</td>
</tr>
<tr>
<td>OR</td>
<td></td>
</tr>
<tr>
<td>Isoniazid + rifapentine weekly for 3 months (3HP)</td>
<td>Isoniazid PO once weekly: &lt; 30 kg and ≥ 2 years: 20 to 30 mg/kg</td>
</tr>
<tr>
<td></td>
<td>≥ 30 kg: 900 mg</td>
</tr>
<tr>
<td></td>
<td>+ rifapentine PO once weekly: 10 to 14 kg and ≥ 2 years: 300 mg</td>
</tr>
<tr>
<td></td>
<td>14.1 to 25 kg and ≥ 2 years: 450 mg</td>
</tr>
<tr>
<td></td>
<td>25.1 to 32 kg: 600 mg</td>
</tr>
<tr>
<td></td>
<td>32.1 to 49.9 kg: 750 mg</td>
</tr>
<tr>
<td></td>
<td>≥ 50 kg: 900 mg max.</td>
</tr>
<tr>
<td>OR</td>
<td></td>
</tr>
<tr>
<td>Isoniazid + rifampicin daily for 3 months (3HR)</td>
<td>Isoniazid PO once daily: &lt; 30 kg: 10 mg/kg (7 to 15 mg/kg)</td>
</tr>
<tr>
<td></td>
<td>≥ 30 kg: 5 mg/kg (4 to 6 mg/kg)</td>
</tr>
<tr>
<td></td>
<td>(max. dose 300 mg daily)</td>
</tr>
<tr>
<td></td>
<td>+ rifampicin PO once daily: &lt; 30 kg: 15 mg/kg</td>
</tr>
<tr>
<td></td>
<td>≥ 30 kg: 10 mg/kg</td>
</tr>
<tr>
<td></td>
<td>(max. dose 600 mg daily)</td>
</tr>
</tbody>
</table>

### Alternative regimens

<table>
<thead>
<tr>
<th>Regimen</th>
<th>Isoniazid PO once daily:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid + rifapentine daily for 1 month (1HP)</td>
<td>Isoniazid PO once daily: ≥ 13 years: 300 mg</td>
</tr>
<tr>
<td></td>
<td>+ rifapentine PO once daily: ≥ 13 years: 600 mg</td>
</tr>
<tr>
<td>OR</td>
<td></td>
</tr>
<tr>
<td>Rifampicin daily for 4 months (4R)</td>
<td>Rifampicin PO once daily: &lt; 30 kg: 15 mg/kg</td>
</tr>
<tr>
<td></td>
<td>≥ 30 kg: 10 mg/kg</td>
</tr>
<tr>
<td></td>
<td>(max. dose 600 mg daily)</td>
</tr>
</tbody>
</table>

### 16.3.1 Isoniazid monotherapy
Isoniazid monotherapy (or isoniazid preventive therapy, IPT) is the treatment currently most often used for LTBI. This treatment has proven to be effective in preventing active TB in both HIV-infected and non-HIV-infected patients. WHO recommends this treatment in all patients regardless of their HIV status, including children of any age and pregnant women. The main disadvantage of isoniazid monotherapy is the length of treatment. Patients are usually healthy and may not be motivated to complete a 6-month therapy. Adverse effects (e.g., peripheral neuropathy, hepatotoxicity) can also lead to treatment interruption. All patients at risk of peripheral neuropathy should receive pyridoxine (vitamin B\textsubscript{6}) for the entire duration of treatment to prevent this risk (for doses see Appendix 17). In HIV-infected patients, the treatment may be difficult due to additive adverse effects of antiretrovirals and isoniazid, the extending of the duration of treatment to 36 months in some adolescents and adults (Section 16.4.2) and the high number of tablets to be taken daily. The number of tablets can be reduced using a fixed-dose combination (FDC) of isoniazid/cotrimoxazole/pyridoxine.

### 16.3.2 Rifapentine-containing regimens

#### Combination isoniazid-rifapentine once weekly for 3 months (3HP)

This treatment has proven to be effective in preventing active TB in both HIV-infected and non-HIV-infected patients. WHO recommends this treatment in children 2 years and over, adolescents and adults, regardless of their HIV status. It is short, requires few doses, has a high completion rate and the risk of hepatotoxicity is low\textsuperscript{[5][8]}. The disadvantages of this regimen are the lack of FDC and the development of hypersensitivity reaction in almost 4\% of patients\textsuperscript{[4]} (Section 16.8.3).

#### Combination isoniazid-rifapentine once daily for 1 month (1HP)

This treatment has proven to be effective in preventing active TB in HIV-infected patients. WHO recommends this treatment as an alternative regimen in patients 13 years and over, regardless of their weight and HIV status. The treatment is short, has a high completion rate and the risk of hepatotoxicity is low\textsuperscript{[7]}. However, cutaneous reactions (rash, itching) are common.

Rifapentine containing regimens are not currently recommended for pregnant women. Despite some reassuring data\textsuperscript{[8]}, safety is not definitively established.

### 16.3.3 Rifampicin-containing regimens

#### Combination isoniazid-rifampicin once daily for 3 months (3HR)

This treatment has proven to be effective in preventing active TB in both HIV-infected and non-HIV-infected patients. WHO recommends this treatment in all patients regardless of their HIV status, including children of any age and pregnant women. It is short, safe, has a good completion rate\textsuperscript{[3]} and FDC are available for children and adults. Hypersensitivity reaction may occur in approximately 2\% of patients\textsuperscript{[9]}.

#### Rifampicin monotherapy once daily for 4 months (4R)

This treatment has proven to be effective in preventing active TB in non-HIV-infected patients of all ages. WHO recommends this regimen as an alternative regimen in all patients regardless of their HIV status, including children of any age and pregnant women. The advantages of this regimen (better safety profile and completion rate compared to 6H)\textsuperscript{[10]} should be weighed against the risk associated with use of rifampicin in monotherapy (development of resistance to rifampicin in patients with undiagnosed active TB).

**Notes** on rifamycin-containing regimens:
- Rifapentine and rifampicin have interactions with many drugs, particularly antiretrovirals (Appendix 19) and contraceptives (Chapter 9).
- For pregnant women taking rifampicin, administer phytonamenadione (vitamin K) in the last few weeks of pregnancy (Chapter 9).
- Rifapentine and rifampicin are not interchangeable.
- Rifabutin can replace rifampicin if rifampicin cannot be used due to drug interactions\textsuperscript{[2]}. 

16.4 Latent tuberculosis infection in HIV-infected patients

Treatment of LTBI reduces the risk of active TB by 33-64%.[1]

For patients not yet on antiretroviral treatment (ART), ART initiation should take priority over initiation of LTBI treatment. Among these patients, there is a high proportion of undiagnosed, asymptomatic TB cases and it is important to use all existing diagnostic means to rule out active TB.

Note: a treatment programme for LTBI should be combined with a screening programme for active TB in HIV-infected patients (Chapter 6).

16.4.1 Children

HIV-exposed childrena and HIV-infected children and who do not have active TB (for evaluation, see Chapter 5) should receive LTBI treatment:

- After contact with a TB case, including smear-positive, smear-negative and extrapulmonary TB (EPTB), regardless of their age;
- In high TB transmission areas: if aged 12 months and over, regardless of their contact history.

References


In addition, for children treated for active TB and living in high TB transmission areas, LTBI treatment may also be prescribed immediately after the successful completion of TB treatment to reduce the risk of reinfection.

### 16.4.2 Adolescents and adults

HIV-infected adolescents and adults who do not have active TB should receive LTBI treatment, regardless of contact history and TB prevalence in the area.

In areas with high TB transmission, HIV-infected adolescents and adults with a LTBI test positive or unknown and who are unlikely to have active TB (no cough, no fever, no weight loss, no night sweats) should receive the treatment for at least 36 months (long-term regimen).

This regimen is more effective in preventing TB in HIV-infected adults with a positive TST than those with a negative TST [2].

If TST is not feasible, or where the national guidelines do not recommend long-term isoniazid monotherapy, HIV-infected adolescents and adults without any TB symptoms should receive another LTBI treatment (6H or a rifapentine- or rifampicin-containing regimen).

**Table 16.2 – LTBI treatments for HIV-infected patients**

<table>
<thead>
<tr>
<th>Age</th>
<th>Recommended regimens</th>
<th>Alternative regimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Child &lt; 2 years</td>
<td>6H or 3RH</td>
<td>4R</td>
</tr>
<tr>
<td>Child ≥ 2 years</td>
<td>6H or 3HP or 3RH</td>
<td>4R</td>
</tr>
<tr>
<td>Adolescent and adult</td>
<td>6H or 3HP or 3RH or 36H</td>
<td>1HP (if ≥ 13 years) or 4R</td>
</tr>
</tbody>
</table>

**Notas**

(a) HIV-exposed children are children born to HIV-infected women whose HIV status has not been established and/or are still at risk of infection (e.g. still breastfed).

**Referencias**


### 16.5 Latent tuberculosis infection in household contacts

A household contact is a person who has shared the same enclosed living space as the index case for one or more nights or for frequent or extended daytime periods during 3 months before the start of the current treatment [1].

#### 16.5.1 Neonates of mothers with active pulmonary tuberculosis
All neonates born to mothers with active PTB should receive treatment for LTBI, after exclusion of active TB, if the mother:

- Has been treated for PTB less than 2 weeks at the time of birth, or
- Has a positive smear microscopy result on a sputum sample collected at birth or close to the time of birth[2].

A test Xpert MTB/RIF and Xpert MTB/XDR should be performed to rule out resistance to rifampicin and isoniazid before starting LTBI treatment.

The recommended regimens are 3HR or 6H. For HIV-exposed neonates receiving nevirapine, only 6H is recommended.

BCG vaccine should be administered just after LTBI treatment completion (not during the treatment).

If a TST is feasible and the regimen chosen is 6H:

- Administer isoniazid for 3 months, then perform a TST.
- If the TST is positive, complete isoniazid monotherapy.
- If the TST is negative, stop isoniazid and administer the BCG vaccine.

Notes:

- A neonate should not be separated from its mother unless severely ill.
- Breastfeeding should continue, and breastfed neonates should receive pyridoxine (vitamin B₆).

### 16.5.2 Other household contacts

**Children under 5 years**

It is not mandatory to perform TST or IGRA prior to LTBI treatment. All children < 5 years in contact with a confirmed PTB case and who do not have active TB (for evaluation, see Chapter 5) should receive LTBI treatment, regardless of their HIV and BCG vaccination status.

If LTBI treatment is contra-indicated or in case of parental refusal, monitor the child closely for one year to enable the early detection of active TB.

**Children 5 years and older, adolescents and adults**

A TST or IGRA should be performed prior to LTBI treatment. If this is not feasible, LTBI treatment may be considered, weighing benefits and risks.

- Children 5 years and over in contact with a confirmed PTB case and who do not have active TB (for evaluation, see Chapter 5) may receive LTBI treatment, regardless of their HIV status.
- Adolescents and adults in contact with a confirmed PTB case and who do not have active TB (no TB symptoms and no abnormality on CXR) may receive LTBI treatment, regardless of their HIV status.

### Table 16.3 - LTBI regimens for household contacts

<table>
<thead>
<tr>
<th>Age</th>
<th>Recommended regimens</th>
<th>Alternative regimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Child &lt; 2 years</td>
<td>6H or 3RH</td>
<td>4R</td>
</tr>
<tr>
<td>Child ≥ 2 years and &lt; 5 years</td>
<td>6H or 3HP or 3RH</td>
<td>4R</td>
</tr>
<tr>
<td>Child ≥ 5 years, adolescent, adult</td>
<td>6H or 3HP or 3RH</td>
<td>1HP (if ≥ 13 years) or 4R</td>
</tr>
</tbody>
</table>

### Referencias

16.6 Latent tuberculosis infection in other individuals at risk

Routine LTBI testing (TST or IGRA) and treatment after exclusion of active TB:
- Are recommended for patients with silicosis, on dialysis or taking long-term immunosuppressive therapy.
- Can be considered for health staff, populations in congregate living settings (e.g. prisoners, refugees), migrants from countries with a high TB prevalence, homeless people and drug users.

LTBI testing should be performed periodically (e.g. once a year).

Routine LTBI testing and treatment is not recommended for diabetic, malnourished or alcoholic patients, unless they belong to the above-mentioned risk groups.

16.7 Latent tuberculosis infection and multidrug-resistant tuberculosis

Due to limited evidence, routine LTBI treatment for all household contacts of multidrug-resistant TB (MDR-TB) patients cannot be recommended at this time.

However, treatment of LTBI should be considered in certain high-risk household contacts based on an individual risk-benefit assessment.

Individual assessment includes:
- High risk of progression to active TB: children under 5 years, individuals with HIV infection or on immunosuppressive therapy.
- Resistance pattern of the source case: the LTBI treatment regimen must be individually tailored as contacts of MDR-TB patients are often infected with the same strain.[1]
- Intensity of exposure.
- Contra-indication or risk of adverse drug reactions.

A TST or IGRA should be performed prior to LTBI treatment. If not feasible, LTBI treatment may be considered, weighing benefits and risks.

16.7.1 Household contacts of multidrug-resistant tuberculosis cases eligible for treatment

Evidence is lacking on the choice of treatment to prevent disease in MDR-TB contacts. Few observational studies, primarily using a fluoroquinolone (FQ) for 6 months, reported promising results[2][3]. Randomized clinical trials are ongoing[4][5].

For contacts of FQ-susceptible MDR-TB patients, levofloxacin PO for 6 months can be proposed at the following doses:

<table>
<thead>
<tr>
<th>Weight</th>
<th>5 to 9 kg</th>
<th>10 to 15 kg</th>
<th>16 to 23 kg</th>
<th>24 to 34 kg</th>
<th>35 to 45 kg</th>
<th>&gt; 45 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily dose</td>
<td>150 mg</td>
<td>200 to 300 mg</td>
<td>300 to 400 mg</td>
<td>500 to 750 mg</td>
<td>750 mg</td>
<td>1 g</td>
</tr>
</tbody>
</table>
If active TB develops during LTBI treatment, DST including resistance to FQs is necessary due to the potential risk associated with use of FQs in monotherapy (development of resistance to FQs in patients with undiagnosed active TB). Independent of LTBI treatment, monitor these patients for 2 years for the development of active TB.

16.7.2 Household contacts of multidrug-resistant tuberculosis cases not eligible for treatment

If the contact is not eligible for LTBI treatment, closely monitor for signs and symptoms of active TB every 3 months for the next 2 years.

If active TB develops, initiate TB treatment promptly with a regimen designed according to the DST. If DST is not feasible, a regimen can be designed according to the resistance profile of the source case.

Referencias


16.8 Follow-up for patients treated for latent tuberculosis infection

For the modality of administration of LTBI treatments see Chapter 13.

16.8.1 Baseline assessment of liver function

Before initiating LTBI treatment, look for clinical signs of hepatic disease and specific risks of hepatotoxicity.

For patients with hepatic disease, baseline liver function tests (LFTs), i.e. aspartate aminotransferase (AST), alanine aminotransferase (ALT) and bilirubin should be performed.

The benefit of LTBI treatment should be weighed against the potential risk of aggravation of existing hepatic disease. LTBI treatment is contra-indicated in patients with end-stage hepatic disease or LFTs > 5 times the upper limit of normal (ULN) and should be used with caution in patients with LFTs > 3 times ULN[1].

Depending on available resources, baseline LFTs can be performed in groups at risk for hepatotoxicity (e.g. patients with HIV infection, women during pregnancy and post-partum period, chronic alcohol consumption, age > 35 years, concomitant use of hepatotoxic drugs, history of hepatic disease).

16.8.2 Follow-up

All patients should be evaluated monthly for signs and symptoms of active TB, adverse effects and adherence.
TST or IGRA should not be repeated.

In patients with pre-existing hepatic disease:
- Baseline LFTs are normal: monitor LFTs monthly.
- Baseline LFTs are elevated or LFTs increase during LTBI treatment: monitor LFTs once a week[2].

Other patients should be tested if they develop symptoms of hepatotoxicity.

Any problems with adherence should be addressed with the patient.

If signs and symptoms of active TB develop, the patient should undergo full evaluation (Chapter 3, Chapter 4 and Chapter 5).

16.8.3 Management of adverse effects

Hepatotoxicity

Clinical features resemble that of viral hepatitis. Early symptoms include malaise, fatigue, loss of appetite, muscle and joint pain. Nausea, vomiting and abdominal pain are common in severe disease. Jaundice, scleral icterus, dark (tea-coloured) urine and discoloured stool are signs of clinical worsening. Clinical hepatitis can be fatal, so action should be taken immediately.

- Patient with symptoms of hepatitis:
  - Stop all TB drugs and perform LFTs:
    - a) AST or ALT or bilirubin ≥ 3 times ULN or severe symptoms: do not re-initiate LTBI treatment.
    - b) AST, ALT, and bilirubin < 3 times ULN and mild symptoms (no jaundice): after discussion with the patient on benefits and risk, treatment may be re-initiated. Closely monitor the patient and perform LFTs once a week. Continue treatment as long as LFTs levels remain < 3 ULN and there are no signs of worsening hepatitis.
    - c) If LFTs are not available, do not re-initiate LTBI treatment.

- Patient without symptoms of hepatitis, but elevated LFTs:
  - a) AST or ALT ≥ 5 times ULN or bilirubin ≥ 3 ULN: stop and do not re-initiate LTBI treatment.
  - b) AST and ALT < 5 times ULN and bilirubin < 3 ULN: stop LTBI treatment. Perform LFTs once a week. If LFTs return to normal, after discussion with the patient on benefits and risk, treatment may be re-initiated. Closely monitor the patient and perform LFTs once a week.

Note: 10-20% of patients taking isoniazid alone may have a mild, transient, asymptomatic elevation of LFTs (AST and/or ALT). In most cases, this does not require treatment interruption.

Hypersensitivity reaction

Approximately 2% of patients on 3HR regimen and 4% of patients on 3HP regimen have hypersensitivity reaction, typically after the first 3 to 4 doses[3]. Symptoms may include fever, headache, dizziness, nausea and vomiting, muscle and bone pain, rash, itching, red eyes, angioedema, shortness of breath and, more rarely, hypotension and altered consciousness.

In case of hypersensitivity reaction, treatment should be stopped immediately. Symptoms usually resolve within 24 hours after TB drug withdrawal. In case of mild reaction (fever, rash, itching), consider re-initiating the treatment. In this case, the patient should be observed at least 4 hours after each dose is administered to detect first signs of hypersensitivity reaction.

Other adverse effects

See Appendix 17.

Referencias

   [https://apps.who.int/iris/rest/bitstreams/1272664/retrieve](https://apps.who.int/iris/rest/bitstreams/1272664/retrieve)
Chapter 17: Monitoring and evaluation

17.1 Introduction

Monitoring and evaluation rely on both quantitative and qualitative information in order to provide information on the following:

- Programme performance (e.g. number of patients started on anti-TB treatment, treatment results, number of patients tested for MDR-TB, etc.);
- Planning for human resources, patient support, diagnostic tests and drug orders, etc.;
- Evaluation of the functioning of the programme (quality of drugs, diagnostics, patient support, etc.).

Recording and reporting are based on a set of standard case and outcome definitions.

Case definitions are presented in Chapter 7.

17.2 Definitions of treatment outcomes

For all forms of TB, outcome definitions have many similarities. These are:

- Outcome assignment is standardized, as to permit comparisons across clinicians, time and sites.
- Outcome assignment relies heavily, but not exclusively, on bacteriologic endpoints (smear or culture*).
- Outcomes are mutually exclusive and exhaustive.

For all forms of TB, definitions exist for:

- Interim outcomes (intended to have an indication on how the programme is functioning before final outcomes are available);
- Final outcomes (cure, completion, failure, treatment interruption, death or not evaluated).

17.2.1 Interim outcomes for drug-susceptible TB and MDR-TB

Given that TB treatment is long (6 to 18 months or more), interim outcomes provide early indicators of programme results. Table 17.1 provides a summary on interim outcomes.
17.2.2 Final outcomes for drug-susceptible TB and DR-TB

Table 17.2 provides definitions for the final outcomes.

Table 17.2 - Summary table of final outcome definitions[1][2]
<table>
<thead>
<tr>
<th>Outcomes</th>
<th>TB</th>
<th>Definitions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DS TB</td>
<td>Patient initially bacteriologically confirmed (microscopy, culture or molecular test) who completed treatment AND shows no signs of continued active disease AND has at least 2 negative smears or cultures: one at 4-5 months and the other at the end of treatment AND does not meet the definition of failure.</td>
</tr>
<tr>
<td>Cured</td>
<td>PDR-TB</td>
<td>Patient initially bacteriologically confirmed (culture or molecular test), who completed treatment AND has been consistently culture-negative with at least 3 results on sputum tested at least one month apart for the final 6 months of treatment AND does not meet the definition of failure.</td>
</tr>
<tr>
<td></td>
<td>MDR-TB</td>
<td>Patient initially bacteriologically confirmed (culture or molecular test), who completed treatment AND with at least 3 negative cultures in the last 8 months of treatment AND does not meet the definition of failure. If there is a lone positive culture or smear reported during that time, and no concomitant clinical evidence of deterioration, a patient may still be considered cured, provided that this positive culture is followed by a minimum of 3 consecutive negative cultures taken at least 30 days apart.</td>
</tr>
<tr>
<td>Completed</td>
<td>All</td>
<td>Patient who completed treatment AND has no signs of continued active disease AND does not meet the bacteriological criteria for cure.</td>
</tr>
<tr>
<td>Failure</td>
<td>DS TB</td>
<td>Patient with signs of continued active disease or deterioration requiring a treatment change: • Any patient with positive smear or culture at 4-5 months of treatment or thereafter. • Any patient with no significant clinical improvement, no significant gain of weight after 4-5 months of treatment and for whom the diagnosis of failure is established by a clinician.</td>
</tr>
<tr>
<td></td>
<td>DR-TB (a)(b)</td>
<td>Treatment terminated or need for permanent treatment change of at least 2 classes of anti-TB drugs because of one or more of the following: • Lack of monitoring cultures converting to negative by 6 months for MDR-TB (3 months for PDR-TB), and/or • Resistance amplification to rifampicin or isoniazid (PDR-TB) or to Group 2 or Group 3 drugs (MDR-TB), and/or • Bacteriological reversion (at least two positive smears or cultures at least 7 days apart after monitoring smears or cultures have become negative), or • A clinical decision has been made to terminate treatment early due to poor response or adverse events. These latter failures can be indicated separately in order to do sub-analysis.</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>Patient who interrupted treatment for 2 months or more.</td>
</tr>
<tr>
<td>Death</td>
<td>All</td>
<td>Patient who died on TB treatment or while awaiting TB treatment, irrespective of the cause of death. The cause of death should be recorded.</td>
</tr>
<tr>
<td>Treatment adapted (d)(e)</td>
<td>DS TB</td>
<td>Patient initially treated with a standard regimen and for whom the treatment is secondarily adapted according to the results of DST (and not because of a treatment failure).</td>
</tr>
</tbody>
</table>
If treatment is continuing at the time of a cohort analysis, an outcome of “still on treatment” may be provisionally assigned.

(a) A patient registered as “failure” can be re-registered as DR-TB “previously treated 2nd line” and started again on a new regimen if possible.

(b) This category does not include the changing of one drug due to an adverse effect or a temporary cessation of drugs in order to manage severe adverse event.

(c) If a patient was defined as a “failure”, and no appropriate treatment was possible, but the treatment was continued and the patient subsequently interrupted the treatment or died, the outcome is “failure” (the first outcome is recorded).

(d) For programmes that report using the WHO’s mutually exclusive six outcomes, the “treatment adapted” outcome can be added to failures for reporting purposes, but should also be kept track of separately for good programmatic monitoring and evaluation.

(e) Not applicable for DR-TB.

17.3 Recording tools

Forms used in recording and reporting can be found in the appendices. They are intended to be examples that programmes or country can use to produce their own forms.

17.3.1 Drug-susceptible TB treatment card and drug-susceptible TB register

Drug-susceptible TB treatment card and drug-susceptible TB register (Appendix 23 and Appendix 24) are used for all new patients or previously treated patients treated by standard first-line regimens (with or without confirmation of the drug susceptibility by a DST).

17.3.2 DR-TB treatment card and DR-TB register

DR-TB treatment card (Appendix 25) tracks, in particular, each dose of each drug taken during the full course of treatment. The number of actually observed doses and the number of expected observed doses are reported each month. In addition to the treatment card, it is recommended to keep a medical chart with a full admission note at the time of enrolment and a progress note at each medical encounter.

DR-TB register (Appendix 26) includes data on case definition, bacteriological exams (indicate date of specimen collection and not the date of result), type of treatment and treatment outcome. It is a separate register from the drug-susceptible TB register.

Each DR-TB patient detected should be registered, including patients who refuse treatment.
Transfer of patients from the drug-susceptible TB register to the DR-TB register is done usually while on treatment when DST results are available. Patient’s outcome is reported as ‘treatment adapted’ in the comment row of the drug-susceptible TB register (Appendix 24).

### 17.3.3 Laboratory request form(s) and register(s)

- Request form for microscopy and Xpert MTB/RIF (Appendix 27);
- Request form for sputum culture, LPA and DST (Appendix 28);
- Sputum smear microscopy register (Appendix 29);
- Xpert MTB/RIF register (Appendix 30).

### 17.3.4 Drug-O-Gram

The Drug-O-Gram is a summary of the patient’s treatment history. It includes consecutive DST and treatment changes presented in a chronological order and gives a short summary of the patient status (Appendix 31).

### 17.4 Reporting

The key evaluation tool for all forms of TB is the periodic report. It must be presented in a standardized manner in two parts: case enrolment and treatment outcomes. The data presented in the report comes from the TB register. It is generally completed by quarter for drug-susceptible TB and by semester for DR-TB.

Evaluation of interim and final treatment outcomes is a fundamental stage in the evaluation. This evaluation is done through a cohort analysis. A “cohort” is a group of individuals presenting certain common characteristics and undergoing the same events. In respect to the evaluation of TB patients, a cohort is represented by patients all put under treatment within a given period of time (usually a quarter for drug-susceptible TB and a semester for DR-TB). At the end of treatment, a final outcome is assigned to each patient (Table 17.1).

**Notes:**
- The number of patients in each group should, in principle, be identical to those registered for the same interval in the case enrolment part of the corresponding periodic report. If it is different, an explanation should be given (e.g., patients “interrupting before treatment” can be excluded from the outcome analysis).
- The outcomes of patients “transferred in” should not be included in the outcomes of the facility to which they were transferred. Their outcome results should be recorded in the facility that initially enrolled the patient in TB treatment.

### 17.4.1 Case detection and enrolment report for TB

The elements necessary for defining a TB case (treatment history, bacteriological status, anatomical site of the disease, and HIV status) are defined in Chapter 7.

See Quarterly report for case enrolment, Appendix 32.

**Main indicators**

- **Proportion of confirmed pulmonary TB (PTB)**
  \[ \text{Proportion of confirmed PTB} = \frac{\text{Number of PTB cases confirmed enrolled}}{\text{Total number of TB cases enrolled for the period}} \]
  
  With the introduction of automated molecular tests and rapid cultures, it is expected that the proportion of confirmed PTB cases will increase as compared to programmes where only smear microscopy is available.

- **Proportion of smear-negative PTB**
  \[ \text{Proportion of smear-negative PTB} = \frac{\text{Number of smear-negative PTB cases enrolled}}{\text{Total number of TB cases enrolled for the period}} \]
  
  This indicator essentially depends on the following: the quality of microscopy, the number of children under treatment (children are rarely smear-positive), the prevalence of HIV infection within the population (these patients present more smear-negative PTB), and the other diagnostics used (culture, Xpert MTB/RIF, etc).
  The proportion of smear-negative PTB is about 20% when HIV prevalence is low. It is 40 to 60% when HIV prevalence is high. Proportions that differ significantly from these should make one consider the possibility of under- or over-diagnosis of smear-negative forms.
17.4.2 Case detection and enrolment report for DR-TB

See standard DR-TB case detection and enrolment reports in Appendix 33.

Early detection of resistance is intended to ensure that an appropriate treatment is initiated from the start. DST is usually performed for patients at risk of DR-TB. Target groups vary according to local situation, but should at a minimum always include patients who have been previously treated and contacts of confirmed MDR-TB patients.

The indicators for detection aim at measuring the access of TB patients to DST. The frequency of MDR-TB among individuals in different risk groups is also evaluated.

All patients in whom DR-TB is highly suspected or detected should be started on appropriate treatment in the shortest time possible.

A comparison of enrolled patients under treatment to detected DR-TB cases gives an indication of access to care, though some patients started on treatment may have been detected prior to the period of assessment.

The period of assessment is six calendar months. This is usually counted from January to the end of June and July to the end of December. Indicators are measured three months after the end of the six-month period. All data can be extracted from the DR-TB register (Appendix 26), the laboratory register for culture and DST and the Xpert register (Appendix 30).

Each indicator should be calculated for all patients and for each risk group of patients, including: all cases, previously treated cases, failures, household contacts and other local risk groups according to the strategy.

Case detection indicators

- **Proportion of TB patients detected with DST result for isoniazid and rifampicin (for each risk group during the period)**
  \[ \frac{\text{Number of TB cases detected with DST result for both isoniazid and rifampicin}}{\text{Total number of TB cases detected}} \]

- **Proportion of TB patients detected with Xpert MTB/RIF result (for each risk group during the period)**
  \[ \frac{\text{Number of TB cases detected with Xpert MTB/RIF result}}{\text{Total number of TB cases detected}} \]
Interim treatment outcomes for drug-susceptible TB and DR-TB

Interim analysis should be completed approximately 3 months after all patients who were registered during a particular interval completed the intensive phase of treatment (three months should allow culture results for all those patients).

Interim results at Month 2 or 3 should be evaluated for all patients treated as new or previously treated patients by standard first-line regimens (with or without confirmation of the drug susceptibility by a DST). These results may be disaggregated by treatment history (new, previously treated, and by type of previous treatment).

At the beginning of a programme, when it is not yet possible to do cohort analysis, the conversion rate at Month 2-3 is a proxy indicator of the effectiveness of treatment, and it allows early detection of potential problems. The smear conversion rate of new smear-positive patients is the proportion of new smear-positive patients who are smear-negative at Month 2. The smear conversion rate of previously treated smear-positive patients is the proportion of previously treated smear-positive patients who are smear-negative at Month 3.

Interim treatment outcomes for DR-TB

The period of assessment is six calendar months, usually counted from January to end June, July to end December. All patients registered and starting treatment during the period of assessment are included in the calculation. The interim report form should be completed 9 months after the closing day of the cohort. This allows culture information at 6 months of treatment to be included for all patients in the cohort. For instance, interim results of TB patients who started treatment during the first semester of a year (1 January to 30 June), should be calculated at the beginning of April of the following year.

Culture conversion (for confirmed DR-TB cases) and death by six months are used as proxies for final outcomes. Information on treatment interruption by six months is helpful. It is also useful to know how many patients started on second-line drugs for MDR-TB turned out not to be MDR.

All data can be extracted from the DR-TB register (Appendix 26).

At six months:

- Proportion of death
  \[ = \frac{\text{Number of confirmed MDR-TB cases registered and started on MDR-TB treatment who died of any cause by the end of Month 6}}{\text{Total number of confirmed MDR-TB cases started on treatment for MDR-TB during the period}} \]

- Proportion of treatment interrupted
  \[ = \frac{\text{Number of confirmed MDR-TB cases started on MDR-TB treatment who interrupted by the end of Month 6}}{\text{Total number of confirmed MDR-TB cases started on treatment for MDR-TB during the period}} \]
17.4.4 Final treatment outcomes for TB

The final outcome is the most important direct measurement of the effectiveness of a TB programme in terms of patient care. All patients entered on the TB register should be assigned one of six mutually exclusive outcomes at the end of their therapy. All patients should be assigned the first outcome they experience for the treatment being evaluated.

Final treatment outcome cohort analysis could be carried out when all patients admitted in a given period of time had a chance to complete their treatment. In practice:

- For drug-susceptible TB (and all patients treated by standard first-line regimens) cohort results are analysed quarterly, one year after inclusion of the last patient of the cohort (e.g. cohort of patients admitted during the first quarter 2014 will be evaluated at the end of the first quarter 2015).
- For DR-TB, evaluation occurs 27 months after inclusion of the last patient in the cohort in order to have the results of cultures performed at 24 months. The period of assessment is six calendar months, usually counted from January to the end of June and July to the end of December. All patients starting treatment during this period are included in the calculation. Indicators are measured 24 months after the end of the semester of assessment. All data can be extracted from the DR-TB register.

Although the timing of the analysis is different for drug-susceptible TB and DR-TB, the indicators are the same. Indicators should be calculated for patients treated by standard first-line regimens (with or without confirmation of drug-susceptible TB by a DST), and for patients with PDR-TB and MDR-TB.

The most important indicators are:

- **Proportion with negative culture**
  \[ \text{Number of bacteriologically confirmed pulmonary MDR-TB cases registered and started on MDR-TB treatment with negative culture at Month 6} \div \text{Total number of bacteriologically confirmed pulmonary MDR-TB cases registered and started on treatment for MDR-TB during the period} \]

- **Proportion with positive culture**
  \[ \text{Number of bacteriologically confirmed pulmonary MDR-TB cases registered and started on MDR-TB treatment with positive culture at Month 6} \div \text{Total number of bacteriologically confirmed pulmonary MDR-TB cases registered and started on treatment for MDR-TB during the period} \]

- **Proportion found not to have MDR-TB**
  \[ \text{Number of patients started on MDR-TB treatment during the period and later found not to be MDR} \div \text{Total number of patients started on MDR-TB treatment during the period} \]

The final outcome is the most important direct measurement of the effectiveness of a TB programme in terms of patient care. All patients entered on the TB register should be assigned one of six mutually exclusive outcomes at the end of their therapy. All patients should be assigned the first outcome they experience for the treatment being evaluated. Although the timing of the analysis is different for drug-susceptible TB and DR-TB, the indicators are the same. Indicators should be calculated for patients treated by standard first-line regimens (with or without confirmation of drug-susceptible TB by a DST), and for patients with PDR-TB and MDR-TB.

The most important indicators are:

- **Proportion cured**
  \[ \text{Number of confirmed TB cases declared “cured”} \div \text{Total number of confirmed TB cases put under treatment during the period} \]

- **Proportion of treatment completed**
  \[ \text{Number of patients registered as “treatment completed”} \div \text{Total number of patients put under treatment for the period} \]

- **Proportion with successful outcome**
  \[ \text{Number of patients registered as “cured” or “treatment completed”} \div \text{Total number of patients put under treatment during the period} \]

- **Proportion of treatment interrupted**
  \[ \text{Number of patients registered as “treatment interrupted”} \div \text{Total number of patients put under treatment during the period} \]
17.5 Programme assessment

To be complete, evaluation should look at how well the programme functions, particularly with respect to three aspects: organization of care, established procedures and human resources. A set of quality criteria is evaluated for each of these aspects. The criteria may be either qualitative (description) or quantitative (indicators). The following tables can be used as a rough guide.

17.5.1 Organization

- Proportion of death
  \[ \text{Proportion of death} = \frac{\text{Number of patients registered as "death"}}{\text{Total number of patients put under treatment during the period}} \]
  This ratio usually does not exceed 5% for drug-susceptible TB. Over-mortality may be related to the poor functioning of a programme. It may also be due to a high prevalence of HIV infection among cases or late referrals.

- Proportion of failure
  \[ \text{Proportion of failure} = \frac{\text{Number of patients registered as "failures"}}{\text{Total number of patients put under treatment during the period}} \]
  A high failure rate in new cases can be related to poor treatment adherence, high rate of primary resistance or poor quality of anti-TB drugs. The failure rate should not be over 2% in new cases under treatment.

- Proportion of patients for whom HIV status is known
  \[ \text{Proportion of patients for whom HIV status is known} = \frac{\text{Number of patients for whom HIV status is known by the end of treatment}}{\text{Total number of patients put under treatment during the period}} \]
  This is one of the indicators that help evaluate the integration of TB and HIV services.

- TB-HIV co-infection rate
  \[ \text{TB-HIV co-infection rate} = \frac{\text{Number of HIV-infected TB patients}}{\text{Total number of TB patients put under treatment during the period and for whom HIV status is known at the end of treatment}} \]
  In high HIV-prevalence regions, co-infection rate may exceed 80%. This information is important in assessing other indicators, in particular the proportion of death.

Referencias

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Indicators</th>
<th>Goals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Access to care</td>
<td>• Accessibility of treatment facilities, decentralization, etc.</td>
<td>Easy access to care during the intensive/continuation phases</td>
</tr>
<tr>
<td></td>
<td>• Home-based treatment available when appropriate.</td>
<td></td>
</tr>
<tr>
<td>Patient comfort</td>
<td>• Patient welcome</td>
<td>• According to needs</td>
</tr>
<tr>
<td></td>
<td>• Condition of the facility, heating (or cooling), overall organization and</td>
<td>• Bed occupancy rate ≤ 100%</td>
</tr>
<tr>
<td></td>
<td>cleanliness.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Food during hospitalization and/or for outpatients (supplemental rations,</td>
<td></td>
</tr>
<tr>
<td></td>
<td>quantities, organization in charge).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Bed occupancy rate of the TB ward.</td>
<td></td>
</tr>
<tr>
<td>Information and therapeutic</td>
<td>Patient interviews conducted.</td>
<td>Patient understanding of treatment</td>
</tr>
<tr>
<td>education</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hospital hygiene</td>
<td>• Equipment (respirators, masks, gloves, gowns, autoclaves, cleaning supplies,</td>
<td>All necessary equipment is available and used.</td>
</tr>
<tr>
<td></td>
<td>etc.)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Waste management (sorting, incinerator, etc.)</td>
<td></td>
</tr>
<tr>
<td>Constant supply of lab</td>
<td>• Supplied by (government, agency or facility, other)</td>
<td>• 3-month buffer stock</td>
</tr>
<tr>
<td>materials</td>
<td>• Buffer stock</td>
<td>• No shortages</td>
</tr>
<tr>
<td></td>
<td>• Number and duration of shortages</td>
<td></td>
</tr>
<tr>
<td>Constant supply of quality-assured</td>
<td>• Stock card maintenance</td>
<td>Stock cards up-to-dated</td>
</tr>
<tr>
<td>anti-TB drugs</td>
<td>• Order frequency, delivery time, buffer stock</td>
<td>• One person in charge of the pharmacy</td>
</tr>
<tr>
<td></td>
<td>• Shortage(s)</td>
<td>• All adequate</td>
</tr>
<tr>
<td></td>
<td>• Drug sources</td>
<td>• No shortages</td>
</tr>
<tr>
<td></td>
<td>• Institution in charge of supply</td>
<td>• WHO-prequalified sources (or equivalent)</td>
</tr>
<tr>
<td></td>
<td>• Use of FDCs first-line drugs</td>
<td>• Use of FDCs</td>
</tr>
<tr>
<td></td>
<td>• Storage conditions</td>
<td>• Appropriate storage conditions</td>
</tr>
<tr>
<td></td>
<td>• Organization of supply for peripheral facilities</td>
<td>• Regular supply</td>
</tr>
<tr>
<td>Case detection</td>
<td>• Type of case detection (active or passive)</td>
<td>Know the type, in order to interpret the quantitative results of case</td>
</tr>
<tr>
<td></td>
<td>• Contacts screening</td>
<td>detection</td>
</tr>
<tr>
<td></td>
<td>• Detection rate of new smear-positive cases</td>
<td>• 100%</td>
</tr>
<tr>
<td></td>
<td>• Percentage of smear-positive patients out of the total number of patients</td>
<td>• Depends on the context</td>
</tr>
<tr>
<td></td>
<td>who had a sputum smear.</td>
<td>• &lt; 20%</td>
</tr>
<tr>
<td></td>
<td>• Detection rate of MDR-TB</td>
<td>• Depends on the context</td>
</tr>
<tr>
<td>Diagnosis of smear-negative</td>
<td>• Automated molecular test</td>
<td></td>
</tr>
<tr>
<td>PTB and EP forms</td>
<td>• Culture or molecular techniques</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• X-rays</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Others (e.g. ADA, Pandy, Rivalta, FNAC)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Algorithms used</td>
<td></td>
</tr>
<tr>
<td>DST</td>
<td>DST possible (methods, quality control)</td>
<td>Detection of DR-TB</td>
</tr>
<tr>
<td>Treatment support</td>
<td>Number of patients receiving treatment support/month</td>
<td>100% of those eligible for support</td>
</tr>
<tr>
<td>Identification of non-adherent</td>
<td>• System for identifying and looking for non-adherent</td>
<td>Yes</td>
</tr>
</tbody>
</table>
### 17.5.2 Procedures

<table>
<thead>
<tr>
<th>adherent patients</th>
<th>patients</th>
<th>Percentage of patients who resumed treatment among those missing for less than 2 months who had to be looked for</th>
<th>&gt; 90%</th>
</tr>
</thead>
</table>

<p>| Integrated TB/HIV care | Access to voluntary counselling and testing (VCT) | Access to ART | Access to cotrimoxazole prophylaxis | HIV treatment integrated in the TB service (or TB treatment in the HIV service) | Yes | Yes | Yes | Yes |</p>
<table>
<thead>
<tr>
<th>Criteria</th>
<th>Indicators</th>
<th>Goals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Registers/records</td>
<td>Description of the documents</td>
<td>Records reliable</td>
</tr>
<tr>
<td></td>
<td>• Consistency between TB registers and treatment cards</td>
<td>• 100%</td>
</tr>
<tr>
<td></td>
<td>• Consistency between TB register and lab registers</td>
<td>• 100%</td>
</tr>
<tr>
<td>Standard case definitions</td>
<td>Percentage of patients with exact case definition out of a randomized sample of patients</td>
<td>100%</td>
</tr>
<tr>
<td>Adequate standard treatment regimens and follow-up</td>
<td>• Percentage of new cases correctly treated (combinations, dosage, duration) out of a randomized sample of patients</td>
<td>&gt; 95%</td>
</tr>
<tr>
<td></td>
<td>• Percentage of patients who did not have bacteriological follow-up according to schedule out of a randomized sample of patients</td>
<td>&lt; 10%</td>
</tr>
<tr>
<td></td>
<td>• Percentage of MDR-TB patients who did not have biochemistry tests according to schedule out of a randomized sample of patients</td>
<td>&lt; 10%</td>
</tr>
<tr>
<td>HIV testing</td>
<td>Percentage of new cases tested for HIV</td>
<td>100%</td>
</tr>
<tr>
<td>ART</td>
<td>Percentage of HIV-positive TB cases started on ART</td>
<td>100%</td>
</tr>
<tr>
<td></td>
<td>ART started within:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• &lt; 2 weeks; 2 weeks-&lt; 2 months; ≥ 2 months</td>
<td></td>
</tr>
<tr>
<td>Criteria for cure</td>
<td>Percentage of confirmed cases declared cured who actually met the definition of cure out of a randomized sample of patients</td>
<td>&gt; 90%</td>
</tr>
<tr>
<td>Regular monitoring of drug-susceptible TB and DR-TB</td>
<td>• Quarterly report and cohort analysis for drug-susceptible TB</td>
<td>Quantitative data on inclusions and results collected</td>
</tr>
<tr>
<td></td>
<td>• Bi-annual report and cohort analysis for DR-TB</td>
<td>Rapid detection of potential problems</td>
</tr>
<tr>
<td>Adherence monitoring</td>
<td>• Percentage of patients coming in for their appointment out of number of patients expected</td>
<td>&gt; 90% in both the intensive and continuation phases</td>
</tr>
<tr>
<td></td>
<td>• Percentage of doses given under DOT for DR-TB treatment in a randomized sample of patients</td>
<td>100%</td>
</tr>
<tr>
<td>Prevention of <em>M. tuberculosis</em> airborne transmission in TB facilities</td>
<td>• Isolation</td>
<td>Isolation of smear positive patients</td>
</tr>
<tr>
<td></td>
<td>• Building ventilation, lights, UV lamps (hospital wards, outpatient clinics, laboratory); respirators for staff and visitors in contact with contagious</td>
<td>Isolation of DR smear positive patients</td>
</tr>
</tbody>
</table>
17.5.3 Human resources

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Indicators</th>
<th>Goals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staff</td>
<td>• Job descriptions (doctors, nurses, lab technicians, cleaning staff, etc.)</td>
<td>On average:</td>
</tr>
<tr>
<td></td>
<td>• Medical staff-to-patient ratio</td>
<td>• One nurse for 10-15 patients</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• One doctor for 40-50 patients</td>
</tr>
<tr>
<td>Training</td>
<td>Refer to training programme evaluation criteria</td>
<td>Competent staff</td>
</tr>
<tr>
<td>Other contributors</td>
<td>Description: other NGOs, local associations, etc.</td>
<td></td>
</tr>
</tbody>
</table>

A grid for evaluating TB clinic operations can be found in Appendix 35. Each criterion is rated either "satisfactory" or "unsatisfactory".

Appendices

Appendix 1. Xpert assays
Appendix 2. Interpretation of Xpert assay results
Appendix 3. Sputum specimen: collection, storage and shipment
Appendix 4. Sputum smear microscopy
Appendix 5. Time required for diagnostic test results
Appendix 6. Ventilated work station (VWS) and bio-safety cabinet (BSC)
Appendix 7. Lymph node fine needle aspiration
Appendix 8. Protein estimation
Appendix 9. Tuberculin skin test
Appendix 10. Drug information sheets and patient instructions for the treatment of tuberculosis
Tuberculosis drug information sheets

Amikacin (Am)
Amoxicillin/clavulanic acid ratio 4:1 (Amx/Clv)
Bedaquiline (Bdq)
Clofazimine (Cfz)
Cycloserine (Cs) or terizidone (Trd)
Delamanid (Dlm)
Ethambutol (E)
Ethionamide (Eto) or prothionamide (Pto)
Imipenem/cilastatin (Ipm/Cln)
Isoniazid - Standard dose (H)
Levofloxacin (Lfx)
Linezolid (Lzd)
Meropenem (Mpm)
Moxifloxacin (Mfx)
Para-aminosalicylate sodium (PAS)
Pretomanid (Pa)
Pyrazinamide (Z)
Rifabutin (Rfb)
Rifampicin (R)
Rifapentine (P)
Streptomycin (S)

Patient instructions

Patients on drug-susceptible TB treatment

Patients on drug-resistant TB treatment

Appendix 11. Use of tuberculosis drugs in pregnant or breastfeeding women

Appendix 12. Dose adjustments in renal insufficiency

Appendix 13. Daily dose of tuberculosis drugs using fixed-dose combinations

Appendix 14. Monitoring of patients on drug-susceptible tuberculosis treatment

Appendix 15. Monitoring of patients on drug-resistant tuberculosis treatment

Appendix 16. Additional investigations in drug-resistant tuberculosis

Appendix 16. Basic TB infection control risk assessment tool

Appendix 17. Management of adverse effects

Gastrointestinal disorders

Abdominal pain
Diarrhoea
Epigastric pain
Hepatotoxicity
Metallic taste
Nausea and vomiting
  Neurotoxicity
Depression
Headache
Optic neuritis
Ototoxicity
Peripheral neuropathy
Psychosis
Seizures
  Endocrine disorders
Gynecomastia
Hypothyroidism
  Dermatological disorders
Alopecia
Fungal infection
Photosensitivity
Skin reactions
  Musculoskeletal disorders
Arthralgias
Tendinitis/tendon rupture
  Miscellaneous
Electrolyte disorders
Haematologic disorders
Lactic acidosis
Nephrotoxicity
QT prolongation

Appendix 17. Air change per hour (ACH) measurement recommendations
Appendix 18. Compassionate use
Appendix 18. Advantages and disadvantages of ventilation techniques
Appendix 19. Drug interactions and overlapping toxicities
Appendix 19. Upper room ultraviolet germicidal irradiation (UVGI) system
Appendix 20. Treatment supporters
Appendix 21. Informing the patient
Appendix 23. Treatment card for patients on first-line anti-TB therapy
Appendix 24. Tuberculosis register for patients on first-line anti-TB therapy
Appendix 1. Xpert assays

Update: October 2022

1.1 Specimen processing

Staff members present during specimen preparation should wear a respirator (FFP2 or N95) to prevent the inhalation of bacilli. A biosafety cabinet should be used to protect staff from aerosols when centrifugation is needed (Appendix 6).

1.1.1 Sputum specimens

See Xpert MTB/RIF package insert:

See Xpert MTB/XDR package insert:

1.1.2 Lymph node and other tissue specimens

Adapted from WHO[1]

- Cut the tissue specimen in small pieces in a sterile mortar (or grinder).
- Add 2 ml of sterile phosphate buffer saline (PBS).
- Grind solution of tissue and PBS to obtain a homogeneous mixture.
- Transfer 0.7 ml of mixture into a centrifuge tube using a transfer pipette. Avoid transferring clumps that are not well homogenized.
- Add 1.4 ml of Xpert Sample Reagent (XSR) using a transfer pipette.
1.1.3 Cerebrospinal fluid specimens

Adapted from WHO[1]

The processing method for cerebrospinal fluid (CSF) depends on the volume available for testing.

<table>
<thead>
<tr>
<th>Volume of CSF</th>
<th>Procedure</th>
</tr>
</thead>
</table>
| 0.1 to 1 ml   | - Add XSR to the CSF to obtain a final volume of 2 ml.  
- Transfer 2 ml of the mixture into the Xpert cartridge.  
- Load the cartridge into the Xpert instrument as per the manufacturer's instructions. |
| 1 to 5 ml     | - Add an equal volume of XSR to the CSF.  
- Add 2 ml of the mixture directly into the Xpert cartridge.  
- Load the cartridge into the Xpert instrument as per the manufacturer's instructions. |
| > 5 ml        | - Centrifuge the CSF at 3,000g for 15 minutes.  
- Pour the supernatant and add XSR to the sediment to obtain a final volume of 2 ml.  
- Transfer 2 ml of the mixture into the Xpert cartridge.  
- Load the cartridge into the Xpert instrument as per the manufacturer's instructions. |

Note: a volume of CSF less than 0.1 ml is insufficient for testing.

1.1.4 Stool specimens[2]

Stool specimens can be used within 3 hours if kept at room temperature.
- Add 0.8 to 1 g of stool into the 8 ml XSR bottle.  
- Shake vigorously for 30 seconds.  
- Keep at room temperature for 10 minutes.  
- Shake vigorously for 30 seconds.  
- Sediment at room temperature for 10 minutes.  
- Without disturbing the sediment, transfer 2 ml of the supernatant into the Xpert cartridge.  
- Load the cartridge into the Xpert instrument as per the manufacturer's instructions.

1.1.5 Urine specimens[3]

Urine specimens can be used within 3 hours if kept at room temperature.
- Centrifuge 4 ml of urine at 3,000g for 5 minutes.  
- Pour the supernatant and add 2 ml of XSR to the sediment.  
- Shake vigorously.  
- Transfer 2 ml of the mixture into the Xpert cartridge.  
- Load the cartridge into the Xpert instrument as per the manufacturer's instructions.

1.2 Diagnostic accuracy of Xpert in specimens other than sputum
### 1.3 Logistic requirements

All Xpert assays are performed with the same instrument. The 10-colour module can read all Xpert cartridges. The 6-colour module can read Xpert MTB/RIF and Xpert MTB/RIF Ultra cartridges.

#### 1.3.1 Power supply

The instrument requires a constant and stable power supply. If power cuts are short (less than 10 minutes), use a 1500VA "on line" UPS. If power cuts are long, the system must be able to sustain a full cycle (approximately 45 minutes). Use a battery charger, a stationary battery, and a voltage stabilizer.

#### 1.3.2 Storage and operating temperatures

Storage of cartridges and reagents: between 2 and 28 °C for 12 months from date of manufacture.

Operating temperature for the Xpert instrument: between 15 and 30 °C. According to climate conditions, air conditioning may be required.

### Performances of Xpert MTB/RIF compared to culture[^4]

<table>
<thead>
<tr>
<th>Specimens</th>
<th>Performances of Xpert MTB/RIF compared to culture[^4]</th>
</tr>
</thead>
</table>
| Lymph node biopsy or aspirate                                            | Biopsy: sensitivity: 82%; specificity: 79%  
Aspirate: sensitivity: 89%; specificity: 86%                              |
| CSF                                                                        | Sensitivity: 70%; specificity: 97%                                                                                       |
| Pleural fluid                                                            | Sensitivity: 50%; specificity: 99%                                                                                       |
| Pericardial fluid                                                        | Sensitivity: 67.6%; specificity: 99.4%                                                                                    |
| Nasopharyngeal aspirate (children with suspected PTB)                    | Sensitivity: 46%; specificity: 100%                                                                                      |
| Gastric aspirate (children with suspected PTB)                           | Sensitivity: 73%; specificity: 98%                                                                                       |
| Stool (children with suspected PTB)                                      | Compared to respiratory specimens' culture:  
  - No HIV infection: sensitivity: 61%; specificity: 98%  
  - HIV infection: sensitivity: 70%; specificity: 98%                                                                     |
| Urine (suspected genitourinary TB)                                       | Sensitivity: 85%; specificity: 97%                                                                                       |
| Urine (HIV patients with suspected disseminated TB)                      | Sensitivity: 40%; specificity: 98%[^5]                                                                                   |
| Synovial fluid                                                           | Sensitivity: 97%; specificity: 94%                                                                                       |
| Peritoneal fluid                                                         | Sensitivity: 59%; specificity: 97%                                                                                       |
|                                                                            | Adult: sensitivity: 56%; specificity: 94%                                                                                 |

[^4]: the performances of Xpert MTB/XDR in non-sputum specimens are considered similar to those of Xpert MTB/RIF as the tests are based on similar technologies.

[^5]:

[^6]:
1.3.3 Calibration

The Xpert modules require annual calibration performed by an authorized service provider or carried out by swapping out the modules. A detailed contract with the supplier should guarantee regular maintenance, calibration, repair, and replacement as and when needed.

1.3.4 Required space

The dimensions of the Xpert IV instrument (4 modules enabling the processing of 4 specimens at the same time) are:
Width: 29.8 cm; height 35.6 cm; depth 31.1 cm; weight: 12 kg.

The instrument is designed for indoor use only. Provide at least 5 cm of clearance on each side to ensure adequate ventilation. Do not place the instrument close to the vents of other instruments or air-handling units.

The dimensions of the kits containing cartridges and reagents are:
Xpert MTB/RIF kit 50 tests: 31 cm x 28 cm x 20 cm
Xpert MTB/XDR kit 10 tests: 24 cm x 16 cm x 7 cm

1.3.5 Waste disposal

Same procedure as for sputum containers.
Xpert assays generate large volumes of waste.

Referencias


Appendix 2. Interpretation of Xpert assay results

Update: October 2022

2.1 Xpert MTB/RIF and Xpert MTB/RIF Ultra

MTB: M. tuberculosis; RIF: rifampicin
<table>
<thead>
<tr>
<th>Results</th>
<th>Interpretation and decisions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invalid/Error/No result</td>
<td>Perform a 2&lt;sup&gt;nd&lt;/sup&gt; test on a new specimen.</td>
</tr>
</tbody>
</table>
| MTB not detected                    | • Child with suspected PTB: perform a 2<sup>nd</sup> test on a new (respiratory or stool) specimen.  
   • Adult: re-evaluate clinically, perform x-ray if indicated, perform a 2<sup>nd</sup> test and/or a culture on a new specimen.                                        |
| MTB detected                        | **MTB detected**  
   No RIF resistance detected       | • Treat for DS-TB.  
   • Perform Xpert MTB/XDR, LPA or pDST to detect H resistance<sup>(a)</sup>; adjust treatment according to DST.                                                                                                            |
| MTB detected                        | **MTB detected**  
   RIF resistance detected         | Evaluate risk factors for rifampicin resistance (RR):  
   • High risk of RR<sup>(b)</sup>: treat for MDR/RR-TB.  
   • Low risk of RR<sup>(c)</sup>: perform a 2<sup>nd</sup> test on a new specimen<sup>(d)</sup>. If 2<sup>nd</sup> test shows:  
     • R susceptibility: treat for DS-TB.  
     • R resistance: treat for MDR/RR-TB.  
   For patients with MDR/RR-TB, perform:  
     • Xpert MTB/XDR or LPA and pDST or genome sequencing for resistance to other TB drugs.  
     • Culture and pDST for treatment monitoring.  
     • If discordant results with pDST (R resistance with Xpert, R susceptibility with pDST): treat for MDR/RR-TB<sup>(e)</sup>. |
| MTB detected                        | **MTB detected**  
   RIF resistance indeterminate     | • Xpert MTB/RIF:  
     • Perform a 2<sup>nd</sup> test on a new specimen. If still “indeterminate”, treat for DS-TB while investigating RR.  
     • Perform pDST or other gDST to confirm or rule out RR.  
     • Perform Xpert MTB/XDR, LPA or pDST to detect H resistance<sup>(a)</sup>.  
     • Xpert MTB/RIF Ultra:  
       • Send an extraction of the raw results (gxx file) to a reference laboratory for identification of possible mutations (interpretation of melting curves).  
       • If not feasible or still “indeterminate”: proceed as for Xpert MTB/RIF. |
| MTB detected                        | **MTB detected**  
   “trace” RIF resistance indeterminate<sup>(Xpert Ultra)</sup> | • HIV-infected patients, children and EP specimens: a “trace” result should be considered as positive.  
   • Adults with history of TB in the previous 5 years: a “trace” result cannot be interpreted, culture should be performed.  
   • No interpretation of RR is possible.  
   • If suspected resistance to R or other TB drugs: perform pDST or other gDST. Adjust treatment according to DST.  
   • Do not test the specimen with Xpert MTB/XDR as the Xpert MTB/XDR has a higher detection limit than Ultra. |

<sup>(a)</sup> For all patients if possible, and at least those with high risk of H resistance (patients with previous TB treatment with H, or contact with a TB case resistant to H, or from an area with a prevalence of resistance to H ≥ 3%).

<sup>(b)</sup> Patients with previous TB treatment with R, or contact with a TB case resistant to R, or from an area of high prevalence of resistance to R.

<sup>(c)</sup> Patients with no previous TB treatment with R, or contact with a TB case resistant to R, and from an area of low prevalence of resistance to R.

<sup>(d)</sup> A 2<sup>nd</sup> test is necessary because in a population with a prevalence of resistance to rifampicin < 5%, the positive predictive value of one test is < 80%, i.e. > 20% of rifampicin resistant results are false positive.
### 2.2 Xpert MTB/XDR

MTB: *M. tuberculosis*; RIF: rifampicin; INH: isoniazid; FLQ: fluoroquinolones; ETH: ethionamide; AMK: amikacin; KAN: kanamycin; CAP: capreomycin

<table>
<thead>
<tr>
<th>Results</th>
<th>Interpretation and decisions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invalid/Error/No result</td>
<td>Perform a 2nd test on a new specimen.</td>
</tr>
<tr>
<td>MTB detected</td>
<td>After a positive Xpert MTB/RIF, an “MTB detected” result is expected because Xpert MTB/XDR and Xpert MTB/RIF have similar detection limit.</td>
</tr>
<tr>
<td>MTB not detected</td>
<td>After a positive Xpert MTB/RIF: perform a 2nd test on a new specimen. If the 2nd test is negative, it can be performed on culture isolates. After a “trace” result with Ultra, a negative result is expected because the Xpert MTB/XDR has a higher detection limit than Ultra.</td>
</tr>
<tr>
<td>MTB detected No resistance detected</td>
<td>• Treat according to the result of Xpert MTB/RIF or Ultra. Resistance cannot be ruled out because other resistance-conferring mutations are not detected by Xpert MTB/XDR (e.g. only 30% of Eto resistance conferring mutations are detected). • Perform pDST for resistance to other TB drugs and monitor treatment.</td>
</tr>
</tbody>
</table>
| MTB detected Low INH resistance detected | Evaluate risk factors of resistance for each drug: • High risk of resistance: consider as resistant to the drug. • InhA mutation and no katG mutation: H can be used, but not counted as a likely effective drug. • If low-level H resistance detected: Mfx can be used, but not counted as a likely effective drug. ● Resistance to Eto can be detected (inhA mutation). However, a negative result does not rule out resistance. Perform pDST for resistance to other TB drugs and monitor treatment. • Low risk of resistance: perform a 2nd test on a new specimen. If the 2nd test shows: ● Drug susceptibility: treat with the drug. ● Drug resistance: consider as resistant (see above for “High risk of resistance to the drug."
| MTB detected Drug resistance indeterminate | Perform a 2nd test on a new specimen. If still “indeterminate”: treat with likely effective drug(s) while investigating resistance with pDST or other gDST (second-line LPA, genome sequencing). |

(e) Patients with previous TB treatment with the drug or contact with a TB case resistant to the drug, or from an area of high prevalence of resistance to the drug.

(f) Patients with no previous TB treatment with the drug or contact with a TB case resistant to the drug and from an area of low prevalence of resistance to the drug.

(g) A 2nd test is necessary because in a population with a prevalence of resistance < 5%, the positive predictive value of one test is < 80% (i.e. > 20% of resistant results are false positive).

(h) No “indeterminate” result is given for Eto.

### Referencias

Appendix 3. Sputum specimen: collection, storage and shipment

3.1 Sputum collection techniques

Regardless the collection technique used, staff member present during sputum collection should wear a respirator to prevent bacilli inhalation.

3.1.1 Sputum obtained spontaneously

Two specimens are to be collected. When possible, specimens should be collected outside in the open air and far away from other people.

The first sample is collected on the spot, at the consultation, when the patient is identified as suspected TB case. If the patient has recently eaten, ask him/her to rinse his/her mouth with water in order to avoid the presence of food in the sample.

The second sample is collected the day after, in the early morning, right after the patient wakes up and before eating. The second sample may be collected at home then the patient brings it to the health facility. Alternatively, two sputum specimens can be collected one hour apart (frontloaded microscopy).

Collection technique:
- The patient must be given a labelled sputum container (or a Falcon® tube, if the sample is to be shipped by air).
- Have the patient take a deep breath, hold for a few seconds, exhale, repeat two or three times, then cough: sputum is material brought up from the lungs after a productive cough. One or two minutes of chest clapping are of benefit.
- Collect at least 3 ml and close the container hermetically.

The quality of sample determines the reliability of the result. Always check that the sample contains solid or purulent material and not only saliva. Take a new sample if unsatisfactory.

If the sample is collected at home, make sure that the patient has understood the technique, including closing the container hermetically after collecting the sputum.

3.1.2 Sputum induction

Sputum induction is sometimes used in children when sputa cannot be spontaneously expectorated, and only in order to perform cultures or Xpert MTB/RIF.

Sputum induction must be performed under close medical supervision. The child should be observed for respiratory distress during, and for 15 minutes after, the procedure. Bronchospasm may occur. Salbutamol spray and oxygen must be ready at hand.

Equipment
- Gloves and respirator
- Suction catheter (6, 7, 8F)
- Sputum container
- 50 ml syringe, needle
- Mask and tubing for nebulizer
- Holding chamber with child’s mask (to be sterilized between each patient)
- Sterile hypertonic solution of 5% sodium chloride (to be kept refrigerated)
Sterile solution of 0.9% sodium chloride (for the specimen)
Salbutamol spray
Oxygen

Procedure

The child should fast for at least 2 hours before the procedure.

- Prior to nebulization:
  - Explain the procedure to the child and/or the person accompanying him/her (this person must wear a respirator).
  - Place the child in a sitting position in the adult’s arms.
  - Administer 2 puffs of salbutamol via a holding chamber, 10 minutes before nebulization.
  - Prepare a sputum container.

- Nebulization:
  - Fill the nebulizer with 5 ml of 5% hypertonic saline solution (sputum inducer).
  - Place the nebulizer mask over the child’s mouth.
  - Leave the child to inhale until the reservoir is empty.

- Nasopharyngeal suction:
  - Do 1 to 2 minutes of clapping.
  - Clean out the nasal cavity.
  - During suction, the child is laid on his/her side, back to the operator, who is behind him/her.
  - Fit a suction catheter to a 50 ml syringe. Lubricate the end of the catheter.
  - Measure the distance from the tip of the nose to the angle of the jaw. Insert the suction catheter to that depth.
  - When inserting and withdrawing the tube, pull on the plunger of the syringe to create suction.
  - Once the syringe is filled with air and mucus, disconnect it from the suction catheter and purge the air (tip facing upward), so that only mucus is left in the syringe.
  - To collect the mucus: draw 2 ml of 0.9% sodium chloride into the syringe to rinse, then empty contents into the sample container.

3.1.3 Gastric aspiration

Gastric aspiration is sometimes used in children when sputa cannot be spontaneously expectorated nor induced using hypertonic saline, and only in order to perform cultures or Xpert MTB/RIF.

Equipment

- Gloves and respirator
- Suction catheter (6, 7, 8F)
- Sputum container
- 50 ml syringe
- Sterile water

Procedure

- Prior to inserting the suction catheter:
  - Explain the procedure to the child and/or the person accompanying him/her (this person must wear a respirator);
  - Place the child in a half-sitting or sitting position in the adult’s arms.
- Insert a nasogastric tube and check that it is correctly placed.
- First suction to collect the gastric fluid and place it in the sputum container, then rinse the stomach with 30 ml of sterile water and suction again. Add the suctioned fluid to the first sample.
- Start culture within 4 hours of collecting the sample. If there will be more than four hours’ delay, neutralize with 100 mg of sodium bicarbonate.

3.2 Sputum specimen storage

When examinations are not performed on the site of collection:

Specimen for smear microscopy
Smears should be performed within three-four days of collection and in the meanwhile stored refrigerated (2 to 8 °C) and protected from light. Contamination does not affect microscopy but heat make specimen liquefy, with selection of mucopurulent part of the sample more difficult.

**Specimen for culture in liquid medium**

Keep the specimen refrigerated (2 to 8 °C), protected from light. Do not use cetylpyrodinium chloride (CPC) as it is not compatible with MGIT.

The specimen should be processed as soon as possible.

**Specimen for culture on Lowenstein-Jensen medium (LJ)**

- Specimens that can be cultured in less than 3 days after collection:
  Keep refrigerated (2 to 8 °C) and protected from light until transport OR immediately transport to the laboratory for processing.

- Specimens that will be cultured more than 3 days after collection:
  Use Falcon tubes and add 1% CPC to preserve the specimen for up to 2 weeks. Specimens with CPC should not be refrigerated, as the CPC will crystallize and be ineffective.
  Samples with CPC can be inoculated on LJ. For inoculation on agar, they require prior neutralization by neutralizing buffer (Difco®).
  CPC can be used for specimens tested by Xpert MTB/RIF.

### 3.3 Sputum specimen shipment

**To a local laboratory**

- Without CPC transport medium: between 2 and 8 °C and protected from light;
- With CPC transport medium: should not be refrigerated because at low temperatures the CPC will crystallize and ruin the sample. Specimens should be kept at room temperature, protected from heat and light.

**By air to a reference laboratory for culture**

Samples are collected and shipped in 50 ml Falcon® conical tubes with screw caps. The tubes are labelled UN 3373, corresponding to Category B infectious substances. If transport times are less than 12 hours, even specimens without CPC can be transported at room temperature.

Samples are triple-packaged, in accordance with IATA packing instruction 650:
1. Primary container holding the sputum sample: tube tightly closed and placed into a latex glove;
2. Secondary container intended to protect the primary container: leak-proof box with enough absorbent material to absorb the entire sample, should the primary container break;
3. Outer packaging intended to protect the secondary container, with UN 3373 labelling.

Information to be provided:
- Primary container: label with the patient’s name or identification number and the sample collection date and location;
- Outer package: indicate the name of the receiving laboratory, the complete address (name, street, postal code, locality, country), and telephone number.
- All samples must be accompanied by the corresponding laboratory test request form (including clinical information).

**Notes:**
- Procedures for shipping bacterial strains obtained after culture are different, more complicated, and rarely feasible in practice. Cultures are classified as Category A infectious substances (UN 2814).
- For a detailed description of the shipment procedures, see MSF Medical catalogue, volume 4.

### Appendix 4. Sputum smear microscopy

**Update: January 2022**
4.1 Sputum smear preparation

Staff members present during sputum smear preparation should wear a respirator to prevent the inhalation of bacilli. Sputum smears should be prepared promptly after sputum collection.

**Equipment**
- Gloves
- Respirator (FFP2 or N95)
- New, clean glass slides (never re-use sputum smear slides)
- Wooden applicator sticks

**Technique**
- Label one end of the slide with the date of sputum collection and laboratory serial number.
- Select a mucopurulent or blood-stained portion of the sputum sample.
- Use an applicator stick to transfer to the slide.
- Smear the specimen over an area of 1.5 to 2 cm x 2 to 3 cm. Make it thin enough to be able to read through it.
- Allow the smear to air dry for 15 minutes. Do not dry the smear in direct sunlight or over a flame.
- Fix the smear by passing the underside of the slide through a flame for 2 to 3 seconds. Repeat 3 or 4 times.
- Allow to cool before staining.

4.2 Ziehl-Neelsen staining

**Equipment**
- Gloves
- Distilled or filtered water
- 0.3% carbol fuchsin
- 3% acid-alcohol
- 0.3% methylene blue
- Binocular microscope with oil immersion objective (100x magnification)

**Technique**
- Flood the slide with 0.3% carbol fuchsin (after filtering the carbol fuchsin).
- Gently heat the underside of the slide. Begin timing as soon as steam appears. Let it steam for 5 minutes. Do not let the stain boil or dry.
- Gently rinse the slide until the water runs clear, then drain off excess water.
- Flood the slide with 3% acid-alcohol for 2 to 3 minutes, then drain. Repeat this operation if the slide is not completely decolourised.
- Gently rinse the slide, then drain off excess water.
- Flood the slide with 0.3% methylene blue for one minute, then drain.
- Gently rinse the slide until the water runs clear, then drain off excess water.
- Allow to air dry. Do not wipe or blot.

**Reading**
- The slides should be examined by an experienced technician. Technicians must be given sufficient time to accurately read slides.
- Before reading the slide, apply a drop of immersion oil to the left edge of the stained smear. Do not touch the slide with the immersion oil applicator (risk of AFB transfer into the oil bottle and onto another slide).
- Examine at least one length (100 high power fields, HPF) before giving a negative result (this should take at least 5 minutes).
- AFB are red, straight or slightly curved rods. They may be found singly or in small groups. The background stains blue.

**Reporting**

Table 4.1 - Grading AFB scale (WHO-IUATLD)[1]
Note: 1-9 AFB per 100 HPF is a positive result. Note that 1-9 AFB per 100 HPF was previously reported as “scanty” followed by the number of AFB seen in 100 HPF (e.g. “scanty 3” meant there were 3 AFB in 100 HPF). Do not confuse "scanty 3" (3 AFB in 100 HPF) with AFB 3+ (more than 10 AFB per HPF).

### 4.3 Auramine O or auramine/rhodamine staining

#### Equipment
- Gloves
- Distilled or filtered (not chlorinated) water
- 0.1% auramine O or auramine/rhodamine solution
- 0.5% acid alcohol
- 0.5% potassium permanganate or 0.3% methylene blue
- Fluorescence microscope (or a LED device that can be attached to a standard light microscope)

#### Technique
- Flood the slide with auramine O or auramine/rhodamine solution for 15 minutes. Ensure that the staining solution remains on the smear.
- Gently rinse, then drain off excess water. Do not use chlorinated water to avoid disturbing the fluorescence reading.
- Flood the slide with 0.5% acid-alcohol for one minute, then drain.
- Gently rinse, then drain off excess water.
- Flood the slide with 0.5% potassium permanganate solution or 0.3% methylene blue for one minute, then drain.
- Gently rinse, then drain off excess water.
- Allow to air dry. Do not wipe or blot.

Note: to control the quality of the colouration include at least one known positive smear in the batch.

#### Reading
- The slides should be examined by an experienced technician (artefacts are frequent). Technicians must be given sufficient time to read slides.
- Use a 20x objective to screen the smear.
- Examine one length before giving a negative result.
- Always read the positive control smear first. If the positive control is not positive do not continue with the patient smears, but re-stain the batch.
- AFB are bright yellow, straight or slightly curved rods. They may be found singly or in small groups. The background is dark. Non-specific debris stains pale yellow.

#### Reporting

<table>
<thead>
<tr>
<th>Number of AFB (1000x magnification: one length = 100 HPF)</th>
<th>Reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zero AFB/one length</td>
<td>No AFB</td>
</tr>
<tr>
<td>1-9 AFB/one length or 100 HPF</td>
<td>Report exact number of AFB</td>
</tr>
<tr>
<td>10-99 AFB/one length or 100 HPF</td>
<td>1+</td>
</tr>
<tr>
<td>1-10 AFB/one HPF in at least 50 fields</td>
<td>2+</td>
</tr>
<tr>
<td>&gt; 10 AFB/one HPF in at least 20 fields</td>
<td>3+</td>
</tr>
</tbody>
</table>

**Note:**

Table 4.2 - Grading AFB scale (WHO-IUATLD)\(^\text{[1]}\)
**Number of AFB**  
(200-250x magnification: one length = 300 HPF)  

<table>
<thead>
<tr>
<th>Number of AFB/one length</th>
<th>Reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zero AFB/one length</td>
<td>No AFB</td>
</tr>
<tr>
<td>1-29 AFB/one length</td>
<td>Report exact number of AFB</td>
</tr>
<tr>
<td>30-299 AFB/one length</td>
<td>1+</td>
</tr>
<tr>
<td>10-100 AFB/one field on average</td>
<td>2+</td>
</tr>
<tr>
<td>&gt; 100 AFB/one field on average</td>
<td>3+</td>
</tr>
</tbody>
</table>

**Note:** The fluorescence stain remains stable when sheltered from light for only 3 days. Quality control should be done within this time.

**Referencias**


**Appendix 5. Time required for diagnostic test results**

Update: October 2022
Appendix 6. Ventilated work station (VWS) and bio-safety cabinet (BSC)

6.1 Ventilated workstation (VWS)

The VWS provides a safe work environment while preparing sputum smears for AFB staining and Xpert MTB/RIF. It is used when adequate natural ventilation cannot be achieved. Designed to be placed over a bench, it is constituted by a rectangular box ducted to the outside, where the duct is connected to an extraction fan.

VWS are used without filter and do not provide product protection. They should never be used for preparing cultures.

6.2 Class II BSC

Note: to provide negative results, cultures need to be incubated for 6 to 7 weeks on liquid media and 8 weeks on solid media.
A Class II BSC protects not only the operator and the environment, but also the material being manipulated inside the cabinet. The room air and the air circulating within the cabinet are drawn by a downward flowing current through a grate then, through a HEPA filter, which protects both the operator and the product. The air exiting the cabinet is filtered through a HEPA filter to protect the environment.

Class II BSCs are required for performing cultures.

Notes:
- Class I BSCs cannot be used for performing cultures and VWS are sufficient for preparing samples for microscopy and tests Xpert MTB/RIF. Therefore, their use is not recommended in this manual.
- Class III BSCs are generally not used for TB.

Appendix 7. Lymph node fine needle aspiration

FNAC is used to obtain material from lymph nodes. The material is expressed onto slides and prepared for examination. Two smears will be prepared with Giemsa stain to look for caseum, granuloma, giants cells, and epithelioid cells or histocytes and 1 or 2 will be prepared with Ziehl-Neelsen (ZN) stain to look for acid-fast bacilli (AFB).

Equipment
- Needle 23G (in very few cases, it would be possible to use 19G)
- 10 ml syringe
- 2 slides for Giemsa + one or 2 slides for ZN stain
- 10% povidone iodine, sterile gauze, gloves

Technique
- Disinfect the area.
- With the needle attached to the syringe, insert the needle deep into the lymph node.
- After the needle has entered the mass, pull back on the syringe plunger to create a vacuum.
- Rapidly move the needle in a to-and-fro fashion to allow material entering the needle.
- When blood or material appears in the needle hub the aspiration should be stopped. Try to aspirate as much as possible of materials, the amount of materials that has been aspirated would have effect on the specificity and sensitivity of diagnosis.
- Release the negative pressure before to take out the needle from the lymph node. Do not continue sucking while taking out the needle, this will avoid aspiration of materiel into the barrel of the syringe and avoid mixing the sample with the possible peripheral blood in the skin.
- Detach the needle from the syringe immediately after the aspiration.
- Fill the syringe with air (needle is still detached).

Slide preparation
- Slide should be identified prior to the aspiration and prepared immediately after the aspiration.
- Detach the needle from the syringe immediately after the aspiration.
- Fill the syringe with air (needle is still detached).

 Prepare the smear as follow:
- Giemsa
  - Reattach the needle to the syringe and carefully release one small drop of sample onto one end of the slide by pushing down the plunger of the syringe (if the drop is placed in the middle of slide it would be difficult to make smear afterwards).
  - Put another slide over the sample.
  - Slide the two slides against each other, in opposite directions, to spread the sample out completely between them. Do not press the slides together forcefully, to avoid crushing the cells.
  - Allow to air dry.
  - Fix the smears by methanol when they are completely dry.
  - Proceed to Giemsa staining.
- Ziehl-Neelsen
  - Place a small drop of ganglion aspirate on the slide.
- Make a smear that is neither to thin or too thick.
- Allow to air dry.
- Fix the smear by flame when it is completely dry.
- Proceed to ZN staining.

Reading after Giemsa staining

On each slide, one or several of the following aspects can be found:
- Caseation necrosis (caseum): a uniform, acellular, pinkish substance.
- Granuloma: cluster of epithelioid cells and lymphocytes scattered through out smear with or without caseous necrosis.
- Epithelioid cells: elongated, often semi-lunar cells with a fine granular nuclear chromatin surrounded by pink cytoplasm.
- Giant cells: huge multinuclear cells.

Notes:
- It would be better to look for granuloma and necrosis with the 10x and 40x power of microscope then to look for epithelioid cells and giant cells with 100x power.
- Observation of smear requires a competent reader with skills in cytology. Slides have to be sent to a referral cytopathology laboratory for quality control or confirmation.
- The quality of the specimen and the preparation are essential. The smear is to be done by skilled technicians.

Notas
(a) The golden standard of diagnosis for TB on tissue samples is hematoxylin-eosin stain, but Giemsa stain can be used as an alternative in remote areas with limited equipment.

Appendix 8. Protein estimation

Update: January 2022

8.1 Pandy test

Pandy test is used to detect an increase of protein in the cerebrospinal fluid (CSF).
The normal range of protein in CSF is 0.20 to 0.45 g/litre.
The Pandy test is positive when protein is superior to 0.45 g/litre.

Equipment
- Disposable gloves
- Pandy reagent
- Pasteur pipettes
- Conical centrifuge glass tube or test tube
- 1 ml pipettes

Preparation of 500 ml of Pandy reagent

Pandy is a saturated phenol solution.
- Weigh 30 g of phenol and transfer it into a 1000 ml bottle.
- Add 500 ml of distilled water and shake vigorously.
- Leave to stand for one 24 hours.
- Check that some phenol remains undissolved:
  - If so, filter: the solution is ready.
  - If all the phenol has dissolved, add a further 10 g of phenol and wait another 24 hours before filtering.
Pandy reagent is a highly corrosive and toxic solution:
- Label the bottle and mark it corrosive and poisonous.
- Wash hands after preparation.

**Technique**
- Place 1 ml of Pandy reagent in a centrifuge tube.
- Add 3 drops of CSF, drop by drop.
- After each drop, look for a white cloud in the tube.
- To facilitate the reading, place a black surface behind the tube.

**Results**
- Presence of a white precipitate: Pandy test
- Absence of a white precipitate: Pandy test

### 8.2 Rivalta test

The Rivalta test is used to detect an increase of protein in the body fluid (pleural fluid, ascites). The test is positive when the proteins are superior to 30 g/litre.

**Equipment**
- Disposable gloves
- Rivalta reagent
- Pasteur pipettes
- Conical centrifuge glass tube or test tube
- 5 ml pipette

**Preparation of 100 ml of Rivalta reagent**
- Place 50 ml of distilled water in a 100 ml measuring cylinder.
- With a 5 ml pipette, add 3 ml of glacial acetic acid and make up to the 100 ml mark with the remaining 50 ml of distilled water.
- Transfer the solution into a bottle.

**Technique**
- Place 2 ml of Rivalta reagent in a centrifuge tube.
- Add 3 drops of pleural fluid/ascites, drop by drop.
- After each drop, look for a white cloud in the tube.
- To facilitate the reading, place a dark surface behind the tube.

**Results**
- Presence of a white precipitate: Rivalta test positive.
- Absence of a white precipitate: Rivalta test

### Appendix 9. Tuberculin skin test

**Update:** January 2022

#### 9.1 Introduction

A delayed hypersensitivity reaction occurs after an intradermal injection of tuberculin (tuberculin skin test, TST) in persons infected by *M. tuberculosis* or vaccinated with BCG.
The test is performed by injecting 5 international units of tuberculin (purified protein derivative, PPD) intradermally on the ventral surface of the forearm (side of forearm exposed with palm facing up)\(^a\).

The test, which should be performed by a trained healthcare worker, requires 2 visits. The reading is done on the second visit, 48 to 72 hours after the tuberculin injection.
If the patient does not return within 72 hours, another TST should be performed.

The result is determined by the diameter of the reaction and individual characteristics of the person being tested (Table 9.1). It should be recorded in millimetres, not as “positive” or “negative”.

The reaction is the area of induration (swelling that can be felt) around the injection site.
Using a ruler, the diameter of induration is measured transversely. The erythema (redness) around the indurated area is not the reaction and should not be measured.
A reaction that appears several minutes, hours or even 24 hours after injection, but disappears on the day after its appearance, is of no significance.

There is no correlation between the diameter of the induration and:
- likelihood of active TB,
- risk of developing active TB,
- protection against TB disease in vaccinated people.

### 9.2 Positive TST

A positive TST signifies that a *M. tuberculosis* infection has occurred.
However, TST cannot differentiate between active and latent infection.
A positive test supports the diagnosis of latent TB when other diagnostic tools have been used to rule out active TB.
In children, a positive TST may be one element among many to establish the diagnosis of active TB.

#### Table 9.1 - Positive TST results

<table>
<thead>
<tr>
<th>Individual characteristics</th>
<th>Diameter of induration</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIV-infected persons</td>
<td>≥ 5 mm</td>
</tr>
<tr>
<td>Severely malnourished children</td>
<td></td>
</tr>
<tr>
<td>Patients taking corticosteroids (e.g. prednisolone ≥ 15 mg daily ≥ 1 month) or immunosuppressants</td>
<td></td>
</tr>
<tr>
<td>Recent contacts of TB patients</td>
<td></td>
</tr>
<tr>
<td>Persons with fibrotic changes on CXR consistent with prior TB</td>
<td></td>
</tr>
<tr>
<td>Persons from countries with high TB prevalence</td>
<td>≥ 10 mm</td>
</tr>
<tr>
<td>Mycobacteriology laboratory personnel</td>
<td></td>
</tr>
<tr>
<td>Persons working and/or living in congregate settings, including healthcare facilities, prisons, homeless shelters, etc.</td>
<td></td>
</tr>
<tr>
<td>Children &lt; 5 years</td>
<td></td>
</tr>
<tr>
<td>Children &gt; 5 years and adolescents exposed to adults at risk of TB</td>
<td></td>
</tr>
<tr>
<td>Other at-risk categories (e.g. diabetes, injecting drug users, end-stage renal disease, leukemia, low body mass index)</td>
<td></td>
</tr>
<tr>
<td>All other children and adults with no other risk factors or exposure to TB</td>
<td>≥ 15 mm</td>
</tr>
</tbody>
</table>

A highly positive (induration diameter > 20 mm) or phlyctenular reaction should be considered as an argument in favour of active TB but is not enough to decide on treatment.

Some persons may have a positive TST result even if they have not been infected with *M. tuberculosis*.
Causes of false positive results include:
- Errors in tuberculin administration
- Previous BCG vaccination
BCG is given at birth so previous BCG vaccination has limited impact on the interpretation of TST results, except in small children. The average diameter of the TST reaction 1 year after BCG vaccination is 10 mm, with extremes ranging from 4 to 20 mm. The reaction becomes weaker over time and disappears 5 to 10 years post-vaccination.

### 9.3 Negative TST

Usually, a negative TST result signifies that no *M. tuberculosis* infection has occurred. However, a negative TST result does not rule out TB infection.

Causes of false negative results include:

- Errors in tuberculin administration
- Recent viral illness or live virus vaccination (e.g. measles)
- Severe TB disease (e.g. TB meningitis or miliary TB)
- Recent (< 12 weeks) or very old (many years) TB infection
- Immunodepression or a weak immune response (e.g. the very elderly, children < 5 years, malnutrition, patients taking corticosteroids or immunosuppressants)
- Persons with diseases that result in anergy (e.g. AIDS, haemopathy, sarcoidosis)
- Natural extinction of post-vaccination reaction from the 5th year following BCG

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**Appendix 10. Drug information sheets and patient instructions for the treatment of tuberculosis**

**Update: October 2022**

- Tuberculosis drug information sheets
  - Amikacin (Am)
  - Amoxicillin/clavulanic acid ratio 4:1 (Amx/Clv)
  - Bedaquiline (Bdq)
  - Clofazimine (Cfz)
  - Cycloserine (Cs) or terizidone (Trd)
  - Delamanid (Dlm)
  - Ethambutol (E)
  - Ethionamide (Eto) or prothionamide (Pto)
  - Imipenem/cilastatin (Ipm/Cln)
  - Isoniazid - Standard dose (H)
  - Levofloxacin (Lfx)
  - Linezolid (Lzd)
  - Meropenem (Mpm)
  - Moxifloxacin (Mfx)
  - Para-aminosalicylate sodium (PAS)
  - Pretomanid (Pa)
Tuberculosis drug information sheets

Update: October 2022

- Amikacin (Am)
- Amoxicillin/clavulanic acid ratio 4:1 (Amx/Clv)
- Bedaquiline (Bdq)
- Clofazimine (Cfz)
- Cycloserine (Cs) or terizidone (Trd)
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- Levofoxicin (Lfx)
- Linezolid (Lzd)
- Meropenem (Mpm)
- Moxifloxacin (Mfx)
- Para-aminosalicylate sodium (PAS)
- Pretomanid (Pa)
- Pyrazinamide (Z)
- Rifabutin (Rfb)
- Rifampicin (R)
- Rifapentine (P)
- Streptomycin (S)

Amikacin (Am)

Update: January 2022

Forms, strengths and route of administration

- 500 mg amikacin base in 2 ml ampoule (250 mg/ml), for IM injection

Dosage

- Child and adult: 15 to 20 mg/kg once daily
- Patient 60 years and over: 15 mg/kg 3 times a week
- Maximum dose: 1000 mg daily
- Renal insufficiency: 12 to 15 mg/kg 2 or 3 times a week

Patient instructions

- Patients on drug-susceptible TB treatment
- Patients on drug-resistant TB treatment
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>Daily dose (ml) - IM injection&lt;sup&gt;(b)&lt;/sup&gt; (500 mg in 2 ml = 250 mg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75-100</td>
<td>0.4 ml</td>
</tr>
<tr>
<td>6</td>
<td>90-120</td>
<td>0.4 ml</td>
</tr>
<tr>
<td>7</td>
<td>105-140</td>
<td>0.6 ml</td>
</tr>
<tr>
<td>8</td>
<td>120-160</td>
<td>0.6 ml</td>
</tr>
<tr>
<td>9</td>
<td>135-180</td>
<td>0.6 ml</td>
</tr>
<tr>
<td>10</td>
<td>150-200</td>
<td>0.8 ml</td>
</tr>
<tr>
<td>11</td>
<td>165-220</td>
<td>0.8 ml</td>
</tr>
<tr>
<td>12</td>
<td>180-240</td>
<td>0.8 ml</td>
</tr>
<tr>
<td>13</td>
<td>195-260</td>
<td>1 ml</td>
</tr>
<tr>
<td>14</td>
<td>210-280</td>
<td>1 ml</td>
</tr>
<tr>
<td>15</td>
<td>225-300</td>
<td>1 ml</td>
</tr>
<tr>
<td>16</td>
<td>240-320</td>
<td>1.2 ml</td>
</tr>
<tr>
<td>17</td>
<td>255-340</td>
<td>1.2 ml</td>
</tr>
<tr>
<td>18</td>
<td>270-360</td>
<td>1.2 ml</td>
</tr>
<tr>
<td>19</td>
<td>285-380</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>20</td>
<td>300-400</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>21</td>
<td>315-420</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>22</td>
<td>330-440</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>23</td>
<td>345-460</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>24</td>
<td>360-480</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>25</td>
<td>375-500</td>
<td>2 ml</td>
</tr>
<tr>
<td>26</td>
<td>390-520</td>
<td>2 ml</td>
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<tr>
<td>27</td>
<td>405-540</td>
<td>2 ml</td>
</tr>
<tr>
<td>28</td>
<td>420-560</td>
<td>2 ml</td>
</tr>
<tr>
<td>29</td>
<td>435-580</td>
<td>2 ml</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to aminoglycosides.
- Administer with caution to patients 60 years and over or patients with pre-existing renal, vestibular, auditory or severe hepatic impairment.
- May cause:
  - nephrotoxicity, ototoxicity, electrolyte disturbances; rarely, hypersensitivity reactions;
  - local pain after injection.
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with other ototoxic and/or nephrotoxic drugs (furosemide, amphotericin B, tenofovir, etc.).
- Pregnancy: CONTRA-INDICATED
- Breastfeeding: no contra-indication

Monitoring

- Symptomatic monitoring.
- Audiometry, serum creatinine and electrolytes (K, Ca, Mg).

Patient instructions

- Maintain a good fluid intake to limit renal problems.

Remarks

- Use a different site for each injection (absorption may be delayed if the same site is used repeatedly).

Storage

- Below 25 °C
  Solution may darken from colourless to a pale yellow, but this does not indicate a loss of potency.

Amoxicillin/clavulanic acid ratio 4:1 (Amx/Clv)

Update: January 2022

Forms and strengths

- 500 mg amoxicillin/125 mg clavulanic acid tablet
Dosage (expressed in clavulanic acid)

- Child under 30 kg: 3 mg (0.25 ml)/kg of clavulanic acid 3 times daily, 60 minutes before each dose of meropenem
- Adolescent ≥ 15 years and ≥ 30 kg and adult: 125 mg of clavulanic acid 2 times daily, 60 minutes before each dose of carbapenem
- Maximum dose: 250 mg daily

- 250 mg amoxicillin/62.5 mg clavulanic acid per 5 ml, powder for oral suspension
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>500 mg/125 mg tablet</th>
<th>250 mg/62.5 mg per 5 ml oral suspension</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>50</td>
<td>–</td>
<td>1.3 ml x 3</td>
</tr>
<tr>
<td>6</td>
<td>60</td>
<td>–</td>
<td>1.5 ml x 3</td>
</tr>
<tr>
<td>7</td>
<td>70</td>
<td>–</td>
<td>2 ml x 3</td>
</tr>
<tr>
<td>8</td>
<td>80</td>
<td>–</td>
<td>2 ml x 3</td>
</tr>
<tr>
<td>9</td>
<td>90</td>
<td>–</td>
<td>2.5 ml x 3</td>
</tr>
<tr>
<td>10</td>
<td>100</td>
<td>–</td>
<td>2.5 ml x 3</td>
</tr>
<tr>
<td>11</td>
<td>110</td>
<td>–</td>
<td>3 ml x 3</td>
</tr>
<tr>
<td>12</td>
<td>120</td>
<td>–</td>
<td>3 ml x 3</td>
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<tr>
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<td>–</td>
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<td>14</td>
<td>140</td>
<td>–</td>
<td>3.5 ml x 3</td>
</tr>
<tr>
<td>15</td>
<td>150</td>
<td>–</td>
<td>4 ml x 3</td>
</tr>
<tr>
<td>16</td>
<td>160</td>
<td>–</td>
<td>4.5 ml x 3</td>
</tr>
<tr>
<td>17</td>
<td>170</td>
<td>–</td>
<td>4.5 ml x 3</td>
</tr>
<tr>
<td>18</td>
<td>180</td>
<td>–</td>
<td>5 ml x 3</td>
</tr>
<tr>
<td>19</td>
<td>190</td>
<td>–</td>
<td>5 ml x 3</td>
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<td>–</td>
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<td>210</td>
<td>–</td>
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<tr>
<td>22</td>
<td>220</td>
<td>–</td>
<td>6 ml x 3</td>
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<tr>
<td>23</td>
<td>230</td>
<td>–</td>
<td>6 ml x 3</td>
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<tr>
<td>24</td>
<td>240</td>
<td>–</td>
<td>6.5 ml x 3</td>
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<tr>
<td>25</td>
<td>250</td>
<td>–</td>
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<td>–</td>
<td>6.5 ml x 3</td>
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<td>250</td>
<td>–</td>
<td>6.5 ml x 3</td>
</tr>
<tr>
<td>28</td>
<td>250</td>
<td>–</td>
<td>6.5 ml x 3</td>
</tr>
<tr>
<td>29</td>
<td>250</td>
<td>–</td>
<td>6.5 ml x 3</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to penicillin-allergic patients and patients with history of hepatic disorders during a previous treatment with amoxicillin/clavulanic acid.
- Administer with caution to patients allergic to betalactams (cross-hypersensitivity may occur) and to patients with hepatic impairment.
- May cause: gastrointestinal disturbances (mainly diarrhoea), hypersensitivity reactions, hepatotoxicity.
- For the management of adverse effects see Appendix 17.
- Pregnancy: no contra-indication
- Breastfeeding: no contra-indication

Monitoring

- Symptomatic monitoring

Patient instructions

- Take with food.

Storage

- Powder for oral suspension: between 15 °C and 25 °C
- Once reconstituted, the oral suspension must be kept refrigerated (between 2 °C and 8 °C) and may be used for up to 7 days.

Bedaquiline (Bdq)

Update: September 2022

Forms and strengths

- 100 mg tablet
- 20 mg dispersible tablet

Dosage

- Child up to 15 kg: according to weight and age
- Child 16 to 29 kg: 200 mg once daily for 2 weeks, then 100 mg 3 times a week
- Child 30 kg and over and adult: 400 mg once daily for 2 weeks, then 200 mg 3 times a week

When administered 3 times a week, keep an interval of 48 hours between doses (Monday, Wednesday, Friday = M/W/F).
Alternatively, for children 16 to 29 kg: 10 dispersible tablets of 20 mg (200 mg) once daily on Weeks 1 and 2, then 5 dispersible tablets of 20 mg (100 mg) 3 times a week.

If 20 mg dispersible tablets are not available, 100 mg tablets can be crushed and suspended in 10 ml of water or fruit juice to obtain a solution containing 10 mg of bedaquiline per ml, then administered as follows:

### Contra-indications, adverse effects, precautions

- Do not administer (or discontinue) to patients with severe hepatic impairment, QTcF > 500 ms or clinically significant ventricular arrhythmia.
- Avoid or use with caution and under close monitoring in patients with:
  - history of syncopal episodes, *torsades de pointes*, congenital QT prolongation;
  - uncompensated heart failure, severe coronary artery disease, bradycardia;
  - electrolyte disturbances (correct first K, Ca, Mg), hypothyroidism (provide thyroxine);
  - severe renal impairment, end-stage renal disease (optimal dosing not established).
May cause:
- hepatotoxicity, moderate QT prolongation;
- nausea, vomiting, arthralgia, headache, increased amylase level.

For the management of adverse effects see Appendix 17.

Avoid or use with caution and under close monitoring in patients taking CYP450 inducers/inhibitors, some ARVs, or other QT prolonging drugs (Appendix 19).

**Pregnancy**: use if benefits outweigh the risks (safety not established).

**Breastfeeding**: avoid breastfeeding during treatment (safety not established).

Monitoring

- Symptomatic monitoring.
- Liver function, ECG, electrolytes (K, Ca, Mg).

Patient instructions

- Take with food.
- 100 mg tablets can be crushed and mixed with water or fruit juice.
- 20 mg tablets should be dispersed in water, juice, milk, yogurt, porridge, etc.
- Avoid alcohol during treatment.

Storage

- Below 25 °C

**Clofazimine (Cfz)**

Update: August 2022

**Forms and strengths**

- 50 mg and 100 mg soft capsules or tablets

**Dosage**

- Child under 10 kg: doses are administered 3 times a week (Monday, Wednesday, Friday = M/W/F)
- Child 10 to 29 kg: 2 to 5 mg/kg once daily
- Child 30 kg and over and adult: 100 mg once daily
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>100 mg capsule(a)</th>
<th>50 mg capsule(a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>–</td>
<td>–</td>
<td>1 caps (M/W/F)</td>
</tr>
<tr>
<td>6</td>
<td>–</td>
<td>–</td>
<td>1 caps (M/W/F)</td>
</tr>
<tr>
<td>7</td>
<td>–</td>
<td>–</td>
<td>1 caps (M/W/F)</td>
</tr>
<tr>
<td>8</td>
<td>–</td>
<td>–</td>
<td>1 caps (M/W/F)</td>
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<td>9</td>
<td>–</td>
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<td>1 caps (M/W/F)</td>
</tr>
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<td>10</td>
<td>20-50</td>
<td>–</td>
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</tr>
<tr>
<td>11</td>
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<td>12</td>
<td>24-60</td>
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<td>15</td>
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<td>16</td>
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</tr>
<tr>
<td>17</td>
<td>34-85</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>18</td>
<td>36-90</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>19</td>
<td>38-95</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>20</td>
<td>40-100</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>21</td>
<td>42-105</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>22</td>
<td>44-110</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>23</td>
<td>46-115</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>24</td>
<td>48-120</td>
<td>1 caps</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>50-125</td>
<td>1 caps</td>
<td>–</td>
</tr>
<tr>
<td>26</td>
<td>52-130</td>
<td>1 caps</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>54-135</td>
<td>1 caps</td>
<td>–</td>
</tr>
<tr>
<td>28</td>
<td>56-140</td>
<td>1 caps</td>
<td>–</td>
</tr>
<tr>
<td>29</td>
<td>58-145</td>
<td>1 caps</td>
<td>–</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with history of allergy to clofazimine.
- Administer with caution to patients with severe hepatic impairment.
- May cause:
  - orange-brown discolouration of skin and body fluids;
  - strong QT prolongation;
  - gastrointestinal intolerance (nausea, vomiting, abdominal pain);
  - severe abdomen pain, bowel obstruction, intestinal bleeding;
  - eye and skin dryness and irritation, hypersensitivity reactions, photosensitivity.
- For the management of adverse effects see Appendix 17.
- Avoid or use with caution and under close monitoring in patients taking other QT prolonging drugs (Appendix 19).
- Pregnancy: use only if benefits outweigh the risks (safety is not established).
- Breast-feeding: avoid breastfeeding during treatment (safety not established). If used, may cause breast milk discolouration and reversible skin discolouration in breastfed infants.

Monitoring

- Symptomatic monitoring.
- ECG.

Patient instructions

- Take with food to improve gastrointestinal tolerance.
- Protect your skin from sun.
- Harmless orange-brown discoloration of the skin and body fluids (urine, sweat, saliva, sputum, tears, breast milk, etc.). It is reversible but may take months to disappear after stopping treatment.

Storage

- Below 25 °C

Cycloserine (Cs) or terizidone (Trd)

Update: January 2022

Forms and strengths

<table>
<thead>
<tr>
<th>Age group</th>
<th>Dose</th>
<th>Form</th>
<th>Monitoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>30-35</td>
<td>100</td>
<td>1 caps</td>
<td></td>
</tr>
<tr>
<td>36-45</td>
<td>100</td>
<td>1 caps</td>
<td></td>
</tr>
<tr>
<td>46-55</td>
<td>100</td>
<td>1 caps</td>
<td></td>
</tr>
<tr>
<td>56-70</td>
<td>100</td>
<td>1 caps</td>
<td></td>
</tr>
<tr>
<td>&gt; 70</td>
<td>100</td>
<td>1 caps</td>
<td></td>
</tr>
</tbody>
</table>

(a) Capsule or tablet
• 250 mg and 125 mg capsules

**Dosage**

• Child under 30 kg: 7.5 to 10 mg/kg 2 times daily (or 15 to 20 mg/kg once daily if tolerated)
• Child 30 kg and over and adult: 5 to 7.5 mg/kg 2 times daily (or 10 to 15 mg/kg once daily if tolerated)
• Maximum dose: 1000 mg daily
• Renal insufficiency: 250 mg once daily or 500 mg 3 times a week
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>250 mg capsule</th>
<th>125 mg capsule</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75-100</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>6</td>
<td>90-120</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>7</td>
<td>105-140</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>8</td>
<td>120-160</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>9</td>
<td>135-180</td>
<td>–</td>
<td>1 caps</td>
</tr>
<tr>
<td>10</td>
<td>150-200</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>11</td>
<td>165-220</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>12</td>
<td>180-240</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>13</td>
<td>195-260</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>14</td>
<td>210-280</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>15</td>
<td>225-300</td>
<td>–</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>16</td>
<td>240-320</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>17</td>
<td>255-340</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>18</td>
<td>270-360</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>19</td>
<td>285-380</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>20</td>
<td>300-400</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>21</td>
<td>315-420</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>22</td>
<td>330-440</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>23</td>
<td>345-460</td>
<td>–</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>24</td>
<td>360-480</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>375-500</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
<tr>
<td>26</td>
<td>390-520</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>405-540</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
<tr>
<td>28</td>
<td>420-560</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
<tr>
<td>29</td>
<td>435-580</td>
<td>1 caps x 2</td>
<td>–</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Avoid in patients with epilepsy, depression, psychosis, severe anxiety, history of neurological or psychiatric disorders, chronic alcohol use. However, if essential to the regimen, it can be administered under close monitoring.
- May cause:
  - neurotoxicity: seizure, headache, lethargy, confusion, mood change, drowsiness, anxiety, psychosis, depression, suicidal ideation, peripheral neuropathy; rarely, vestibular toxicity;
  - hypersensitivity reactions.
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with isoniazid and thionamides (increased risk of neurotoxicity).
- Administer concomitantly pyridoxine (vitamin B₆); child: 1 to 2 mg/kg (usual range: 10 to 50 mg) once daily; adult: 100 mg once daily.
- Pregnancy: use if the benefits outweigh the risks. Administer pyridoxine to the mother (as above).
- Breastfeeding: no contra-indication. Administer pyridoxine to the mother (as above) and the breast-fed neonate or infant (1 to 2 mg/kg once daily).

Monitoring

- Symptomatic monitoring.

Patient instructions

- Take capsules with water before or after meals.
- Avoid alcohol during treatment.

Remarks

- To increase tolerance, start with a low dose (e.g. 250 mg daily in adults), then increase over 1 to 2 weeks to achieve the requested dose.

Storage

- Below 25 °C

Delamanid (Dlm)

Update: September 2022

Forms and strengths

- 50 mg tablet
- 25 mg dispersible tablet

Forms and strengths

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>30-35</td>
<td>500</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>36-45</td>
<td>500</td>
<td>1 caps x 2</td>
</tr>
<tr>
<td>46-55</td>
<td>750</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>56-70</td>
<td>750</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>750</td>
<td>1 caps (morning) + 2 caps (evening)</td>
</tr>
</tbody>
</table>
Dosage

- Child under 10 kg: according to weight and age
- Child 10 to 15 kg: 25 mg 2 times daily
- Child 16 to 29 kg: 50 mg morning and 25 mg evening
- Child 30 to 45 kg and under 15 years: 50 mg 2 times daily
- Child 46 kg and over and adult: 100 mg 2 times daily

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>50 mg tablet</th>
<th>25 mg dispersible tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-9</td>
<td>25-50</td>
<td>–</td>
<td>&lt; 3 months: 1 tab</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>≥ 3 months: 1 tab x 2</td>
</tr>
<tr>
<td>10-15</td>
<td>50</td>
<td>–</td>
<td>1 tab x 2</td>
</tr>
<tr>
<td>16-29</td>
<td>75</td>
<td>–</td>
<td>2 tab (morning) + 1 tab (evening)</td>
</tr>
<tr>
<td>30-45</td>
<td>100-200</td>
<td>&lt; 15 years: 1 tab x 2</td>
<td>≥ 15 years: 2 tab x 2</td>
</tr>
</tbody>
</table>

- If 25 mg dispersible tablets are not available, 50 mg tablets can be crushed and suspended in 10 ml of water or fruit juice to obtain a solution of 5 mg of delamanid per ml, administered as follows:

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>50 mg tablet in 10 ml (5 mg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-9</td>
<td>25-50</td>
<td>&lt; 3 months: 5 ml</td>
</tr>
<tr>
<td></td>
<td></td>
<td>≥ 3 months: 5 ml x 2</td>
</tr>
<tr>
<td>10-15</td>
<td>50</td>
<td>5 ml x 2</td>
</tr>
<tr>
<td>16-29</td>
<td>75</td>
<td>10 ml (morning) + 5 ml (evening)</td>
</tr>
</tbody>
</table>

Contra-indications, adverse effects, precautions

- Do not administer (or discontinue) to patients with QTcF > 500 ms or albumin level < 2.8 g/dl.
- Avoid or use with caution and under close monitoring in patients with:
  - history of syncopal episodes or torsades de pointes, congenital QT prolongation, cardiac disease;
  - electrolyte disturbances (correct first K, Ca, Mg);
  - severe renal or hepatic impairment.
- Use with caution and under close monitoring in patients taking QT-prolonging drugs (Appendix 19).
- May cause: nausea, vomiting, dizziness, insomnia, mild QT prolongation.
- For the management of adverse effects see Appendix 17.

Pregnancy: use if benefits outweigh the risks (safety not established).

Breastfeeding: avoid breastfeeding during treatment (safety not established).

Monitoring

- Symptomatic monitoring.
- ECG.

Patient instructions
• Take with food.
• 50 mg tablets should be swallowed whole if possible.
• 25 mg tablets should be dispersed in water or fruit juice.

Storage

❄️ – Below 25 °C

Ethambutol (E)

Update: January 2022

Forms and strengths

• 100 mg and 400 mg tablets
• 100 mg dispersible tablet, to be dispersed in 10 ml water

Dosage

• Child and adult: 15 to 25 mg/kg once daily
• Maximum dose: 1200 mg daily
• Renal insufficiency: 15 to 25 mg/kg 3 times a week
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>400 mg tablet</th>
<th>100 mg tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75-125</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>6</td>
<td>90-150</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>7</td>
<td>105-175</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>8</td>
<td>120-200</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>9</td>
<td>135-225</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>10</td>
<td>150-250</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>11</td>
<td>165-275</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>12</td>
<td>180-300</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>13</td>
<td>195-325</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>14</td>
<td>210-350</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>15</td>
<td>225-375</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>16</td>
<td>240-400</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>17</td>
<td>255-425</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>18</td>
<td>270-450</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>19</td>
<td>285-475</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>20</td>
<td>300-500</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>21</td>
<td>315-525</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>22</td>
<td>330-550</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>23</td>
<td>345-575</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>24</td>
<td>360-600</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>375-625</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>26</td>
<td>390-650</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>405-675</td>
<td>1½ tab</td>
<td>–</td>
</tr>
<tr>
<td>28</td>
<td>420-700</td>
<td>1½ tab</td>
<td>–</td>
</tr>
<tr>
<td>29</td>
<td>435-725</td>
<td>1½ tab</td>
<td>–</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with severe renal impairment or pre-existing optic neuritis (e.g. diabetic retinopathy).
- May cause: dose-related retrobulbar optic neuritis, exacerbated in renal impairment.
- The dosage must be carefully adjusted to the weight, especially for children under 5 years, as it is more difficult to detect visual changes at this age.
- For the management of adverse effects see Appendix 17.
- Pregnancy: no contra-indication
- Breastfeeding: no contra-indication

Monitoring

- Symptomatic monitoring.

Patient instructions

- Take with or without food.
- 100 mg dispersible tablets should be dispersed in 10 ml water.

Remarks

- For adults on drug-susceptible TB treatment, ethambutol is given as part of a fixed-dose combination.
- Ethambutol is also used in the treatment of drug-resistant TB treatment for longer duration. For treatment > 2 months, daily doses should be closer to 15 mg/kg and visual acuity and colour discrimination should be monitored.

Storage

- Below 25 °C

Ethionamide (Eto) or prothionamide (Pto)

Update: January 2022

Forms and strengths

- 250 mg tablet (ethionamide or prothionamide)
- 125 mg dispersible tablet (ethionamide), to be dispersed in 10 ml water

Dosage

- Child and adult: 15 to 20 mg/kg once daily
- Maximum dose: 1000 mg daily

<table>
<thead>
<tr>
<th>Weight Range</th>
<th>Dosage</th>
<th>Tablets</th>
</tr>
</thead>
<tbody>
<tr>
<td>30-35</td>
<td>800</td>
<td>2 tab</td>
</tr>
<tr>
<td>36-45</td>
<td>800</td>
<td>2 tab</td>
</tr>
<tr>
<td>46-55</td>
<td>1200</td>
<td>3 tab</td>
</tr>
<tr>
<td>56-70</td>
<td>1200</td>
<td>3 tab</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>1200</td>
<td>3 tab</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>Daily dose (mg)</td>
<td>250 mg tablet</td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>---------------</td>
</tr>
<tr>
<td>5</td>
<td>75-100</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>90-120</td>
<td>–</td>
</tr>
<tr>
<td>7</td>
<td>105-140</td>
<td>–</td>
</tr>
<tr>
<td>8</td>
<td>120-160</td>
<td>–</td>
</tr>
<tr>
<td>9</td>
<td>135-180</td>
<td>–</td>
</tr>
<tr>
<td>10</td>
<td>150-200</td>
<td>–</td>
</tr>
<tr>
<td>11</td>
<td>165-220</td>
<td>–</td>
</tr>
<tr>
<td>12</td>
<td>180-240</td>
<td>–</td>
</tr>
<tr>
<td>13</td>
<td>195-260</td>
<td>–</td>
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<tr>
<td>14</td>
<td>210-280</td>
<td>–</td>
</tr>
<tr>
<td>15</td>
<td>225-300</td>
<td>–</td>
</tr>
<tr>
<td>16</td>
<td>240-320</td>
<td>–</td>
</tr>
<tr>
<td>17</td>
<td>255-340</td>
<td>–</td>
</tr>
<tr>
<td>18</td>
<td>270-360</td>
<td>–</td>
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<tr>
<td>19</td>
<td>285-380</td>
<td>–</td>
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<tr>
<td>20</td>
<td>300-400</td>
<td>–</td>
</tr>
<tr>
<td>21</td>
<td>315-420</td>
<td>–</td>
</tr>
<tr>
<td>22</td>
<td>330-440</td>
<td>–</td>
</tr>
<tr>
<td>23</td>
<td>345-460</td>
<td>–</td>
</tr>
<tr>
<td>24</td>
<td>360-480</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>375-500</td>
<td>2 tab</td>
</tr>
<tr>
<td>26</td>
<td>390-520</td>
<td>2 tab</td>
</tr>
<tr>
<td>27</td>
<td>405-540</td>
<td>2 tab</td>
</tr>
<tr>
<td>28</td>
<td>420-560</td>
<td>2 tab</td>
</tr>
<tr>
<td>29</td>
<td>435-580</td>
<td>2 tab</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with severe hepatic impairment.
- Administer with caution to patients with hepatic disease, diabetes or depression.
- May cause:
  - frequently: gastrointestinal disturbances (abdominal or epigastric pain, diarrhoea, metallic taste, nausea and vomiting, stomatitis, etc.);
  - occasionally: endocrine disorders (gynecomastia, hypothyroidism), alopecia, depression, anxiety, psychosis, hypoglycaemia, vestibular disorders, hepatotoxicity, peripheral neuropathy, optic neuritis, hypersensitivity reactions, seizures.
- For the management of adverse effects see Appendix 17.
- Monitor combination with: cycloserine or terizidone (increased risk of seizures) and para-aminosalicylic acid (increased risk of gastrointestinal disturbances and hypothyroidism).
- Administer concomitantly pyridoxine (vitamin B₆); child: 1 to 2 mg/kg (usual range: 10 to 50 mg) once daily; adult: 100 mg once daily.
- Pregnancy: CONTRA-INDICATED
- Breastfeeding: administer pyridoxine to the mother (as above). Observe the breast-fed neonate or infant for adverse effects and supplement it with pyridoxine (1 to 2 mg/kg once daily).

Monitoring

- Symptomatic monitoring.
- Liver function and thyroid function.

Patient instructions

- Take with food and/or at bedtime to limit gastrointestinal disturbances.
- 125 mg tablets should be dispersed in 10 ml water.
- Avoid alcohol during treatment.

Remarks

- To improve tolerance, start with a low dose (e.g. 250 mg daily in adults), then increase over 1 to 2 weeks to achieve the requested dose.
- For the 6HRZETo regimen for drug-susceptible TB meningitis, the dose is 20 mg/kg once daily (max. 750 mg daily).

Storage

- Below 25 °C

Imipenem/cilastatin (Ipm/Cln)

Update: January 2022
**Forms, strengths and route of administration**

- Powder for injection, in vial of 500 mg imipenem monohydrate/500 mg cilastatin sodium, to be reconstituted with 20 ml of 0.9% sodium chloride (25 mg imipenem/ml).
- Each dose is to be diluted in 100 ml of 0.9% sodium chloride and to be administered by IV infusion:
  - over 30 minutes for doses ≤ 500 mg/500 mg
  - over 60 minutes for doses > 500 mg/500 mg
- Use a deep line, preferably an implantable venous access device (Port-a-Cath).

**Dosage (expressed in imipenem)**

- Adolescent 15 years and over (and ≥ 30 kg) and adult: 1000 mg (2 vials) 2 times daily with 10 hours minimum between infusions
- Maximum dose: 2000 mg daily
- Renal insufficiency: 750 mg every 12 hours for CrCl 20-40 ml/minute; 500 mg every 12 hours for CrCl < 20 ml/minute

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>Daily dose (ml) - IV infusion (500 mg/500 mg per vial)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-29</td>
<td></td>
<td>Do not used in patients &lt; 15 years and &lt; 30 kg</td>
</tr>
<tr>
<td>30-33</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
<tr>
<td>34-40</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
<tr>
<td>41-45</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
<tr>
<td>46-50</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
<tr>
<td>51-70</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>2000</td>
<td>2 vials (40 ml) in 100 ml of 0.9% NaCl x 2</td>
</tr>
</tbody>
</table>

**Contra-indications, adverse effects, precautions**

- Do not administer to patients with hypersensitivity to carbapenems.
- Administer with caution to patients allergic to other beta-lactams (cross-hypersensitivity may occur).
- May cause:
  - nausea, vomiting (the infusion rate may be slowed down in case of nausea), diarrhoea;
  - neurotoxicity: confusional state, seizures (most frequently in patients with history of seizures or renal impairment);
  - hypersensitivity reactions;
  - local reactions (phlebitis/thrombophlebitis).
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with: valproic acid (decreased plasma concentration of valproic acid and risk of seizures), oral or injectable ganciclovir (risk of seizures).
- **Pregnancy and breastfeeding**: avoid unless the benefits outweigh the risks

**Monitoring**

- Symptomatic monitoring.
Remarks

- Administer clavulanic acid 60 minutes before each dose of imipenem/cilastatin.
- Do not mix with Ringer lactate (incompatibility) but may be administered via Y-site.
- Do not mix with other drugs in the infusion bag.

Storage

- Below 25 °C
- Once reconstituted, solution:
  - remains stable 4 hours at room temperature or 24 hours between 2 to 8 °C;
  - may darken from colourless to yellow (this does not indicate a loss of potency);
  - should be discarded if it becomes brown.

Isoniazid - Standard dose (H)

Update: January 2022

Forms and strengths

- 300 mg and 100 mg tablets
- 100 mg and 50 mg dispersible tablets, to be dispersed in 10 ml water

Dosage

- Child under 30 kg: 10 mg/kg (7 to 15 mg/kg) once daily
- Child 30 kg and over and adult: 5 mg/kg (4 to 6 mg/kg) once daily
- Maximum dose: 300 mg daily
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>300 mg tablet</th>
<th>100 mg tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>35-75</td>
<td>–</td>
<td>½ tab</td>
</tr>
<tr>
<td>6</td>
<td>42-90</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>7</td>
<td>49-105</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>8</td>
<td>56-120</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>9</td>
<td>63-135</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>10</td>
<td>70-150</td>
<td>–</td>
<td>1½ tab</td>
</tr>
<tr>
<td>11</td>
<td>77-165</td>
<td>–</td>
<td>1½ tab</td>
</tr>
<tr>
<td>12</td>
<td>84-180</td>
<td>–</td>
<td>1½ tab</td>
</tr>
<tr>
<td>13</td>
<td>91-195</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>14</td>
<td>98-210</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>15</td>
<td>105-225</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>16</td>
<td>112-240</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>17</td>
<td>119-255</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>18</td>
<td>126-270</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>19</td>
<td>133-285</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>20</td>
<td>140-300</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>21</td>
<td>147-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>22</td>
<td>154-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>23</td>
<td>161-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>24</td>
<td>168-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>175-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>26</td>
<td>182-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>189-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>28</td>
<td>196-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>29</td>
<td>203-300</td>
<td>1 tab</td>
<td>–</td>
</tr>
</tbody>
</table>
Alternatively, 50 mg dispersible tablets may be used instead of ½ tablets of 100 mg.

### Contra-indications, adverse effects, precautions

- Do not administer to patients with severe hepatic impairment.
- May cause:
  - peripheral neuropathy;
  - hepatotoxicity;
  - hypersensitivity reactions, arthragias, optic neuritis, psychotic reactions, seizures and depression.
- Monitor closely:
  - pregnant and breastfeeding women, patients with renal impairment or diabetes; malnourished or HIV-infected patients (increased risk of neuropathy);
  - alcoholic patients (increased risk of neuropathy and hepatotoxicity);
  - patients with chronic hepatic disease or taking rifampicin or ≥ 35 years (increased risk of hepatotoxicity);
  - patients taking anticonvulsants, benzodiazepines (risk of toxicity), warfarin (risk of bleeding). Dose adjustment may be required.
- For the management of adverse effects see Appendix 17.
- Administer concomitantly pyridoxine (vitamin B₆) to patients at risk of peripheral neuropathy (child: 5 to 10 mg once daily; adult: 10 mg once daily).
- **Pregnancy and breastfeeding**: no contra-indication. Administer pyridoxine to the mother (as above) and the breast-fed neonate or infant (5 mg once daily).

### Monitoring

- Symptomatic monitoring.
- Liver function in patients with hepatic disease.

### Patient instructions

- Take without food.
- 100 mg dispersible tablet should be dispersed in 10 ml water.
- Avoid alcohol during treatment.

### Remarks

- For patients on drug-susceptible TB treatment, isoniazid is given as part of a fixed-dose combination.
- For the 6HRZ-Eto regimen for drug-susceptible TB meningitis, the dose of isoniazid is 20 mg/kg once daily (max. 400 mg daily).
- Isoniazid is also used in the treatment of latent TB infection and multidrug-resistant TB treatment (at high dose - H₆)

### Storage

- Below 25 °C

---

**Levofloxacin (Lfx)**
Forms and strengths

- 250 mg and 500 mg tablets
- 100 mg dispersible tablet, to be dispersed in 10 ml water

Dosage

- Child under 30 kg: 15 to 20 mg/kg once daily
- Child 30 kg and over and adult: 750 to 1000 mg once daily
- Maximum dose: 1500 mg daily
- Renal insufficiency: 750 to 1000 mg 3 times a week
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>500 mg tablet</th>
<th>250 mg tablet</th>
<th>100 mg dispersible tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75-100</td>
<td>-</td>
<td>-</td>
<td>1 tab</td>
</tr>
<tr>
<td>6</td>
<td>90-120</td>
<td>-</td>
<td>-</td>
<td>1 tab</td>
</tr>
<tr>
<td>7</td>
<td>105-140</td>
<td>-</td>
<td>-</td>
<td>1½ tab</td>
</tr>
<tr>
<td>8</td>
<td>120-160</td>
<td>-</td>
<td>-</td>
<td>1½ tab</td>
</tr>
<tr>
<td>9</td>
<td>135-180</td>
<td>-</td>
<td>-</td>
<td>1½ tab</td>
</tr>
<tr>
<td>10</td>
<td>150-200</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>11</td>
<td>165-220</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>12</td>
<td>180-240</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>13</td>
<td>195-260</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>14</td>
<td>210-280</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>15</td>
<td>225-300</td>
<td>-</td>
<td>-</td>
<td>2 tab</td>
</tr>
<tr>
<td>16</td>
<td>240-320</td>
<td>-</td>
<td>-</td>
<td>3 tab</td>
</tr>
<tr>
<td>17</td>
<td>255-340</td>
<td>-</td>
<td>-</td>
<td>3 tab</td>
</tr>
<tr>
<td>18</td>
<td>270-360</td>
<td>-</td>
<td>-</td>
<td>3 tab</td>
</tr>
<tr>
<td>19</td>
<td>285-380</td>
<td>-</td>
<td>-</td>
<td>3 tab</td>
</tr>
<tr>
<td>20</td>
<td>300-400</td>
<td>-</td>
<td>-</td>
<td>3 tab</td>
</tr>
<tr>
<td>21</td>
<td>315-420</td>
<td>-</td>
<td>-</td>
<td>4 tab</td>
</tr>
<tr>
<td>22</td>
<td>330-440</td>
<td>-</td>
<td>-</td>
<td>4 tab</td>
</tr>
<tr>
<td>23</td>
<td>345-460</td>
<td>-</td>
<td>-</td>
<td>4 tab</td>
</tr>
<tr>
<td>24</td>
<td>360-480</td>
<td>-</td>
<td>-</td>
<td>4 tab</td>
</tr>
<tr>
<td>25</td>
<td>375-500</td>
<td>-</td>
<td>2 tab</td>
<td>-</td>
</tr>
<tr>
<td>26</td>
<td>390-520</td>
<td>-</td>
<td>2 tab</td>
<td>-</td>
</tr>
<tr>
<td>27</td>
<td>405-540</td>
<td>-</td>
<td>2 tab</td>
<td>-</td>
</tr>
<tr>
<td>28</td>
<td>420-560</td>
<td>-</td>
<td>2 tab</td>
<td>-</td>
</tr>
<tr>
<td>29</td>
<td>435-580</td>
<td>-</td>
<td>2 tab</td>
<td>-</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity or tendon damage during a previous treatment with a fluoroquinolone.
- Administer with caution to patients:
  - over 60 years or on corticosteroid treatment (increased risk of tendon damage);
  - with diabetes or history of mental disorders or seizures.
- May cause:
  - tendinitis, tendon rupture, mild QT prolongation;
  - gastrointestinal disturbances (abdominal or epigastric pain, diarrhoea);
  - neurological disorders (headache, psychosis, seizures, etc.);
  - photosensitivity;
  - hypersensitivity reactions, hypo/hyperglycaemia;
  - rarely: crystalluria, peripheral neuropathy, ototoxicity.
- For the management of adverse effects see Appendix 17.
- Avoid or use with caution and under close monitoring in patients taking other QT prolonging drugs (Appendix 19) or warfarin.
- Do not administer simultaneously with: antacids containing magnesium/aluminium, calcium, iron and zinc salts (administer 2 hours apart).

Pregnancy: use if benefits outweigh the risks (safety not established).
Breastfeeding: avoid breastfeeding during treatment (no absolute contra-indication).

Monitoring
- Symptomatic monitoring.

Patient instructions
- Take 2 hours apart from milk-based product, antacids, calcium, iron and zinc salts.
- 100 mg tablets should be dispersed in 10 ml water.
- Maintain a good fluid intake.
- Protect your skin from sun.

Storage

°C – Below 25 °C

Linezolid (Lzd)

Update: October 2022

Forms and strengths
- 600 mg tablet (breakable and non-breakable)
• 150 mg dispersible tablet
• 100 mg/5 ml, granules for oral suspension

**Dosage**

- Child under 15 kg: 15 mg/kg once daily
- Child 15 to 45 kg: 10 to 12 mg/kg once daily
- Patient 46 kg and over: 600 mg once daily
- Maximum dose: 600 mg daily

**Note:** in the BPaLM therapeutic regimen for patients 15 years and over, the dose of linezolid is 600 mg once daily for 16 weeks then 300 mg once daily up to the end of treatment.

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>600 mg tablet</th>
<th>150 mg dispersible tablet</th>
<th>100 mg per 5 ml oral suspension</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75</td>
<td>–</td>
<td>–</td>
<td>3 ml</td>
</tr>
<tr>
<td>6</td>
<td>90</td>
<td>–</td>
<td>–</td>
<td>4 ml</td>
</tr>
<tr>
<td>7</td>
<td>105</td>
<td>–</td>
<td>–</td>
<td>5 ml</td>
</tr>
<tr>
<td>8-9</td>
<td>120-135</td>
<td>–</td>
<td>–</td>
<td>6 ml</td>
</tr>
<tr>
<td>10-15</td>
<td>150-180</td>
<td>–</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>16-23</td>
<td>160-276</td>
<td>–</td>
<td>1½ tab</td>
<td>–</td>
</tr>
<tr>
<td>24-29</td>
<td>240-348</td>
<td>–</td>
<td>2 tab</td>
<td>–</td>
</tr>
<tr>
<td>30-35</td>
<td>300</td>
<td>–</td>
<td>2 tab</td>
<td>–</td>
</tr>
<tr>
<td>36-45</td>
<td>450</td>
<td>–</td>
<td>3 tab</td>
<td>–</td>
</tr>
<tr>
<td>46-55</td>
<td>600</td>
<td>1 tab</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>56-70</td>
<td>600</td>
<td>1 tab</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>600</td>
<td>1 tab</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

• Alternatively, for children 5 to 6 kg, if oral suspension is not available: one half of a 150 mg dispersible tablet (75 mg) once daily.
• If 150 mg dispersible tablets are not available, 600 mg tablets can be crushed and suspended in 10 ml of water or fruit juice to obtain a solution of 60 mg of linezolid per ml, administered as follows:
### Contra-indications, adverse effects, precautions

- Administer with caution to patients with haematological disorders or hypertension.
- May cause:
  - anaemia, neutropenia and/or thrombocytopenia;
  - lactic acidosis;
  - peripheral neuropathy (can be irreversible); rarely, optic neuritis;
  - abdominal pain, diarrhoea, nausea.
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with serotonergic drugs such as tricyclic antidepressants (e.g. amitriptyline) or selective serotonin reuptake inhibitors (e.g. fluoxetine, paroxetine): risk of serotonin syndrome.
- Administer concomitantly pyridoxine (vitamin B₆); child: 1 to 2 mg/kg (usual range: 10 to 50 mg) once daily; adult: 100 mg once daily.
- **Pregnancy**: use if the benefits outweigh the risks. Administer pyridoxine to the mother (as above).
- **Breastfeeding**: avoid breastfeeding during treatment (safety not established).

### Monitoring

- Symptomatic monitoring.
- Full blood count.
- Visual acuity and colour discrimination.

### Patient instructions

- Take with or without food.

### Storage

- Below 25 °C

Once reconstituted, the oral suspension may be kept at room temperature for 21 days, protected from light.

### Meropenem (Mpm)

**Update:** January 2022

### Forms, strengths and route of administration

- Powder for injection, in 500 mg vial, to be reconstituted with 10 ml of water for injection (50 mg meropenem/ml).
- Each dose is to be diluted in 5 ml/kg of 0.9% sodium chloride in children under 20 kg and in 100 ml of 0.9% sodium chloride in children 20 kg and over and adults and to be administered by IV infusion over 15 to 30 minutes.
- Use a deep line, preferably an implantable venous access device (Port-a-Cath).

<table>
<thead>
<tr>
<th>Weight (mg)</th>
<th>Daily dose (mg)</th>
<th>600 mg tablet in 10 ml (60 mg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>75</td>
<td>1.25 ml</td>
</tr>
<tr>
<td>6</td>
<td>90</td>
<td>1.5 ml</td>
</tr>
<tr>
<td>7-9</td>
<td>105-135</td>
<td>2 ml</td>
</tr>
<tr>
<td>10-15</td>
<td>150-180</td>
<td>2.5 ml</td>
</tr>
</tbody>
</table>
Dosage

- Child under 30 kg: 20 to 40 mg/kg every 8 hours
- Child 30 kg and over and adult: 1500 to 2000 mg 2 times daily with 10 hours minimum between infusions
- Maximum dose: 6000 mg daily
- Renal insufficiency: 750 mg every 12 hours for CrCl 20-40 ml/minute; 500 mg every 12 hours for CrCl < 20 ml/minute
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>Daily dose (ml) – IV infusion (500 mg per vial)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>300</td>
<td>2 ml in 25 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>6</td>
<td>300</td>
<td>2 ml in 30 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>7</td>
<td>600</td>
<td>4 ml in 35 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>8</td>
<td>600</td>
<td>4 ml in 40 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>9</td>
<td>600</td>
<td>4 ml in 45 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>10</td>
<td>900</td>
<td>6 ml in 50 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>11</td>
<td>900</td>
<td>6 ml in 55 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>12</td>
<td>900</td>
<td>6 ml in 60 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>13</td>
<td>900</td>
<td>6 ml in 65 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>14</td>
<td>900</td>
<td>6 ml in 70 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>15</td>
<td>900</td>
<td>6 ml in 75 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>16</td>
<td>1200</td>
<td>8 ml in 80 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>17</td>
<td>1200</td>
<td>8 ml in 85 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>18</td>
<td>1200</td>
<td>8 ml in 90 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>19</td>
<td>1200</td>
<td>8 ml in 95 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>20</td>
<td>1200</td>
<td>8 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>21</td>
<td>1200</td>
<td>8 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>22</td>
<td>1200</td>
<td>8 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>23</td>
<td>1200</td>
<td>8 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>24</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>25</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>26</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>27</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>28</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
<tr>
<td>29</td>
<td>1650</td>
<td>11 ml in 100 ml of 0.9% NaCl x 3</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to carbapenems.
- Administer with caution to patients allergic to cephalosporins (cross-hypersensitivity may occur).
- May cause:
  - nausea, vomiting (the infusion rate may be slowed down in case of nausea), diarrhoea;
  - neurotoxicity: confusional state, seizures (rarely compared to imipenem/cilastatin, most frequently in patients with history of seizures or renal impairment);
  - hypersensitivity reactions;
  - local reactions (phlebitis/thrombophlebitis).
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with valproic acid (decreased concentration of valproic acid and risk of seizures).
- **Pregnancy and breastfeeding**: avoid unless the benefits outweigh the risks.

Monitoring

- Symptomatic monitoring.

Remarks

- Administer clavulanic acid 60 minutes before each dose of meropenem.
- Do not mix with other drugs in the infusion bag.

Storage

- Below 25 °C
- Once reconstituted, solution should be used immediately (within 1 hour of preparation)

**Moxifloxacin (Mfx)**

Update: September 2022

Forms and strengths

- 400 mg tablet
- 100 mg dispersible tablet, to be dispersed in 10 ml water

Dosage (standard dose)

- Child under 30 kg: 10 to 15 mg/kg once daily
- Child 30 kg and over and adult: 400 mg once daily
- Maximum dose: 400 mg daily
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>400 mg tablet</th>
<th>100 mg dispersible tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>50-75</td>
<td>–</td>
<td>7 ml</td>
</tr>
<tr>
<td>6</td>
<td>60-90</td>
<td>–</td>
<td>7 ml</td>
</tr>
<tr>
<td>7</td>
<td>70-105</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>8</td>
<td>80-120</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>9</td>
<td>90-135</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>10</td>
<td>100-150</td>
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<td>2 tab</td>
</tr>
<tr>
<td>11</td>
<td>110-165</td>
<td>–</td>
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<td>12</td>
<td>120-180</td>
<td>–</td>
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<td>13</td>
<td>130-195</td>
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<td>2 tab</td>
</tr>
<tr>
<td>14</td>
<td>140-210</td>
<td>–</td>
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</tr>
<tr>
<td>15</td>
<td>150-225</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>16</td>
<td>160-240</td>
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<td>17</td>
<td>170-255</td>
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<td>18</td>
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<td>19</td>
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<tr>
<td>22</td>
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<tr>
<td>23</td>
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<td>24</td>
<td>240-360</td>
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<tr>
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<td>250-375</td>
<td>–</td>
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</tr>
<tr>
<td>26</td>
<td>260-390</td>
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</tr>
<tr>
<td>29</td>
<td>290-435</td>
<td>–</td>
<td>4 tab</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity or tendon damage during a previous treatment with a fluoroquinolone.
- Administer with caution to patients:
  - over 60 years or on corticosteroid treatment (increased risk of tendon damage);
  - with diabetes or history of mental disorders or seizures.
- May cause:
  - tendinitis, tendon rupture, moderate QT prolongation;
  - gastrointestinal disturbances (abdominal or epigastric pain, diarrhoea);
  - neurological disorders (headache, psychosis, seizures, etc.);
  - photosensitivity;
  - hypersensitivity reactions, hypo/hyperglycaemia;
  - rarely: crystalluria, peripheral neuropathy, ototoxicity.
- For the management of adverse effects see Appendix 17.
- Avoid or use with caution and under close monitoring in patients taking other QT prolonging drugs (Appendix 19) or warfarin.
- Do not administer simultaneously with: antacids containing magnesium/aluminium, calcium, iron and zinc salts (administer 2 hours apart).
- Pregnancy:
  - DR-TB: use if benefits outweigh the risks (safety not established).
  - DS-TB: do not use.
- Breastfeeding:
  - DR-TB: avoid breastfeeding during treatment (no absolute contra-indication).

Monitoring

- Symptomatic monitoring

Patient instructions

- Take 2 hours apart from milk-based product, antacids, calcium, iron and zinc salts.
- 100 mg tablets should be dispersed in 10 ml water.
- Maintain a good fluid intake.
- Protect your skin from sun.

Remarks

- Higher dose moxifloxacin (Mfx⁴), i.e. 600 to 800 mg once daily in patients over 30 kg may be used in the presence of certain mutations conferring low level fluoroquinolone resistance. Mfx⁴ may cause strong QT prolongation.

Storage

☀️ – 🌂 – Below 25 °C
Para-aminosalicylate sodium (PAS)

Update: October 2022

Forms and strengths

- Powder for oral solution, 5.52 g sachet of para-aminosalicylate sodium (equivalent to 4 g PAS acid), to be dissolved in 100 ml water

Dosage (expressed in PAS acid)

- Child under 30 kg: 100 to 150 mg/kg 2 times daily
- Child 30 kg and over and adult: 4 g 2 times daily (max. 12 g daily)
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>Oral solution or sachet PAS sodium</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>1000-1500</td>
<td>19 ml x 2</td>
</tr>
<tr>
<td>6</td>
<td>1200-1800</td>
<td>19 ml x 2</td>
</tr>
<tr>
<td>7</td>
<td>1400-2100</td>
<td>25 ml x 2</td>
</tr>
<tr>
<td>8</td>
<td>1600-2400</td>
<td>25 ml x 2</td>
</tr>
<tr>
<td>9</td>
<td>1800-2700</td>
<td>25 ml x 2</td>
</tr>
<tr>
<td>10</td>
<td>2000-3000</td>
<td>50 ml x 2</td>
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<tr>
<td>11</td>
<td>2200-3300</td>
<td>50 ml x 2</td>
</tr>
<tr>
<td>12</td>
<td>2400-3600</td>
<td>50 ml x 2</td>
</tr>
<tr>
<td>13</td>
<td>2600-3900</td>
<td>50 ml x 2</td>
</tr>
<tr>
<td>14</td>
<td>2800-4200</td>
<td>50 ml x 2</td>
</tr>
<tr>
<td>15</td>
<td>3000-4500</td>
<td>50 ml x 2</td>
</tr>
<tr>
<td>16</td>
<td>3200-4800</td>
<td>75 ml x 2</td>
</tr>
<tr>
<td>17</td>
<td>3400-5100</td>
<td>75 ml x 2</td>
</tr>
<tr>
<td>18</td>
<td>3600-5400</td>
<td>75 ml x 2</td>
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<tr>
<td>19</td>
<td>3800-5700</td>
<td>75 ml x 2</td>
</tr>
<tr>
<td>20</td>
<td>4000-6000</td>
<td>75 ml x 2</td>
</tr>
<tr>
<td>21</td>
<td>4200-6300</td>
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<tr>
<td>22</td>
<td>4400-6600</td>
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<td>23</td>
<td>4600-6900</td>
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</tr>
<tr>
<td>24</td>
<td>4800-7200</td>
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<tr>
<td>25</td>
<td>5000-7500</td>
<td>80 ml x 2</td>
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<tr>
<td>26</td>
<td>5200-7800</td>
<td>80 ml x 2</td>
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<tr>
<td>27</td>
<td>5400-8000</td>
<td>80 ml x 2</td>
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<tr>
<td>28</td>
<td>5600-8000</td>
<td>80 ml x 2</td>
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<tr>
<td>29</td>
<td>5800-8000</td>
<td>80 ml x 2</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Avoid in patients with severe renal disease.
- Avoid or use with caution in patients with hepatic impairment or gastric ulcer.
- May cause:
  - frequent gastrointestinal disturbances (nausea, vomiting, gastritis, diarrhoea);
  - hypothyroidism, hepatotoxicity, hypersensitivity reactions.
- Monitor combination with ethionamide/prothionamide (increased risk of gastrointestinal disturbances and hypothyroidism).
- For the management of adverse effects see Appendix 17.
- Pregnancy: use only if benefits outweigh the risks (safety not established).
- Breastfeeding: avoid breastfeeding during treatment (safety not established).

Monitoring

- Symptomatic monitoring.
- Liver and thyroid function.

Patient instructions

- Mix the powder with 100 ml water.
- Take with food to limit gastrointestinal disturbances.

Remarks

- To increase gastrointestinal tolerance, start with a low dose, e.g. for an adult: 2 g twice daily for 1 to 2 weeks, then 4 g twice daily.

Storage

- Below 25 °C

Pretomanid (Pa)

Update: October 2022

Forms and strengths

- 200 mg tablet

Dosage

- Adolescent 15 years and over and adult: 200 mg once daily, in combination with:
  - bedaquiline, linezolid and moxifloxacin (BPaLM)
  - bedaquiline, linezolid and clofazimine (BPaLC)
  - bedaquiline and linezolid (BPaL)
- Maximum dose: 200 mg daily

<table>
<thead>
<tr>
<th>30-70</th>
<th>8 g</th>
<th>1 sachet x 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; 70</td>
<td>8-12 g</td>
<td>1 to 1½ sachet x 2</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer if one of the drugs included in the regimen is contraindicated.
- The contribution of pretomanid to the adverse effects of pretomanid-containing regimens is not determined.
- For adverse effects of companion drugs see individual drug information sheets.
- Pregnancy: use if benefits outweigh the risks (safety not established).
- Breastfeeding: avoid breastfeeding during treatment (safety not established).

Monitoring

- Symptomatic monitoring.
- For monitoring of companion drugs see individual drug information sheets.

Patient instructions

- Take with food.

Storage

🌡 – 🏨 – Below 25 °C

Pyrazinamide (Z)

Update: January 2022

Forms and strengths

- 400 mg tablet
- 150 mg dispersible tablet, to be dispersed in 10 ml water

Dosage

- Child under 30 kg: 35 mg/kg (30 to 40 mg/kg) once daily
- Child 30 kg and over and adult: 25 mg/kg (20 to 30 mg/kg) once daily
- Maximum dose: 2000 mg daily
- Renal insufficiency: 25 mg/kg 3 times a week
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>400 mg tablet</th>
<th>150 mg dispersible tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>150-200</td>
<td>–</td>
<td>1 tab</td>
</tr>
<tr>
<td>6</td>
<td>180-240</td>
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<td>1 tab</td>
</tr>
<tr>
<td>7</td>
<td>210-280</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>8</td>
<td>240-320</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>9</td>
<td>270-360</td>
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<td>3 tab</td>
</tr>
<tr>
<td>13</td>
<td>390-520</td>
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</tr>
<tr>
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</tr>
<tr>
<td>15</td>
<td>450-600</td>
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</tr>
<tr>
<td>16</td>
<td>480-640</td>
<td>–</td>
<td>4 tab</td>
</tr>
<tr>
<td>17</td>
<td>510-680</td>
<td>–</td>
<td>4 tab</td>
</tr>
<tr>
<td>18</td>
<td>540-720</td>
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<td>4 tab</td>
</tr>
<tr>
<td>19</td>
<td>570-760</td>
<td>–</td>
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</tr>
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<td>20</td>
<td>600-800</td>
<td>–</td>
<td>5 tab</td>
</tr>
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</tr>
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<td>22</td>
<td>660-880</td>
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<td>24</td>
<td>720-960</td>
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</tr>
<tr>
<td>25</td>
<td>750-1000</td>
<td>2½ tab</td>
<td>–</td>
</tr>
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<td>26</td>
<td>780-1040</td>
<td>2½ tab</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>810-1080</td>
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<td>28</td>
<td>840-1120</td>
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<td>–</td>
</tr>
<tr>
<td>29</td>
<td>870-1160</td>
<td>2½ tab</td>
<td>–</td>
</tr>
</tbody>
</table>
### Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to pyrazinamide, severe hepatic impairment or severe gout.
- May cause: gout and arthralgias, hepatotoxicity, gastrointestinal disturbances (epigastric pain, nausea and vomiting), hypersensitivity reactions; rarely, photosensitivity.
- For the management of adverse effects see Appendix 17.
- **Pregnancy:** no contra-indication
- **Breastfeeding:** no contra-indication

### Monitoring

- Symptomatic monitoring.
- Liver function in patients with hepatic impairment or under drug-resistant TB treatment.

### Patient instructions

- Take with or without food.
- 150 mg tablets should be dispersed in 10 ml water.
- Protect your skin from sun.

### Remarks

- For patients on drug-susceptible TB treatment, pyrazinamide is given as part of a fixed-dose combination.
- For the 6HRZ-Eto regimen for drug-susceptible TB meningitis, the dose of pyrazinamide is 40 mg/kg once daily (max. 2000 mg daily).

### Storage

≥ – ≤ – Below 25 °C

### Rifabutin (Rfb)

**Update:** January 2022

### Forms and strengths

- 150 mg capsule

### Dosage

- Child and adult: 5 to 10 mg/kg once daily
- Maximum dose: 300 mg daily
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>150 mg capsule</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
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<tr>
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<td>1 caps</td>
</tr>
<tr>
<td>19</td>
<td>95-190</td>
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</tr>
<tr>
<td>20</td>
<td>100-200</td>
<td>1 caps</td>
</tr>
<tr>
<td>21</td>
<td>105-210</td>
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</tr>
<tr>
<td>22</td>
<td>110-220</td>
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</tr>
<tr>
<td>23</td>
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</tr>
<tr>
<td>25</td>
<td>125-250</td>
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<tr>
<td>26</td>
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<td>28</td>
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<td>1 caps</td>
</tr>
<tr>
<td>29</td>
<td>145-290</td>
<td>1 caps</td>
</tr>
</tbody>
</table>
Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to rifamycins or history of severe haematological disorders (thrombocytopenia, purpura) during a previous treatment with a rifamycin.
- Administer with caution to patients with severe renal impairment or hepatic or haematological disorders.
- May cause:
  - gastrointestinal disturbances, hepatotoxicity;
  - haematological disorders (leukopenia, anaemia, thrombocytopenia), hypersensitivity reactions;
  - reversible uveitis.
- For the management of adverse effects see Appendix 17.
- Reduce the dose of rifabutin:
  - in patients taking boosted protease inhibitors (Appendix 19);
  - if rifabutin toxicity is suspected in patients taking clarithromycin, fluconazole or itraconazole.
- Rifabutin reduces the effect of many drugs (macrolides, some antiretrovirals, some hormones, warfarin, etc.):
  - in patients taking antiretrovirals see Appendix 19;
  - in women using contraception, use injectable medroxyprogesterone or an intrauterine device;
  - for the other drugs, adjust dosage if necessary.
- Pregnancy and breastfeeding: avoid (safety not established). If used in late pregnancy, administer phytomenadione (vitamin K₁) to the mother and the neonate.

Monitoring

- Symptomatic monitoring.
- Liver function in patients with hepatic disease.
- Full blood count.

Patient instructions

- Take with or without food.
- Harmless orange-red discoloration of the urine, faeces, sweat, saliva, sputum, tears and other body fluids.

Storage

- Below 25 °C

Rifampicin (R)

Update: January 2022

Forms and strengths

<table>
<thead>
<tr>
<th>30-35</th>
<th>300</th>
<th>2 caps</th>
</tr>
</thead>
<tbody>
<tr>
<td>36-45</td>
<td>300</td>
<td>2 caps</td>
</tr>
<tr>
<td>46-55</td>
<td>300</td>
<td>2 caps</td>
</tr>
<tr>
<td>56-70</td>
<td>300</td>
<td>2 caps</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>300</td>
<td>2 caps</td>
</tr>
</tbody>
</table>
Dosage

- 300 mg capsule and 150 mg tablet

- Child under 30 kg: 15 mg/kg (10 to 20 mg/kg) once daily
- Child 30 kg and over and adult: 10 mg/kg (8 to 12 mg/kg) once daily
- Maximum dose: 600 mg daily
- Hepatic impairment: 8 mg/kg once daily max.
<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>300 mg capsule</th>
<th>150 mg tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>50-100</td>
<td>–</td>
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<td>7</td>
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<td>–</td>
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<td>8</td>
<td>80-160</td>
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<td>14</td>
<td>140-280</td>
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</tr>
<tr>
<td>15</td>
<td>150-300</td>
<td>–</td>
<td>1½ tab</td>
</tr>
<tr>
<td>16</td>
<td>160-320</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>17</td>
<td>170-340</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>18</td>
<td>180-360</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>19</td>
<td>190-380</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>20</td>
<td>200-400</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>21</td>
<td>210-420</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>22</td>
<td>220-440</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>23</td>
<td>230-460</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>24</td>
<td>240-480</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>25</td>
<td>250-500</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>26</td>
<td>260-520</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>27</td>
<td>270-540</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>28</td>
<td>280-560</td>
<td>1 tab</td>
<td>–</td>
</tr>
<tr>
<td>29</td>
<td>290-580</td>
<td>1 tab</td>
<td>–</td>
</tr>
</tbody>
</table>
### Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to rifamycins or history of severe haematological disorders (thrombocytopenia, purpura) during a previous treatment with a rifamycin.
- Avoid or administer with caution to patients with hepatic disorders.
- May cause:
  - hepatotoxicity;
  - influenza-like symptoms, thrombocytopenia, hypersensitivity reactions.
- For the management of adverse effects see Appendix 17.
- Rifampicin reduces the effect of many drugs (antimicrobials, some antiretrovirals, some hormones, antidiabetics, corticosteroids, phenytoin, direct-acting antivirals for chronic hepatitis C, warfarin, etc.):
  - in patients taking antiretrovirals see Appendix 19;
  - in women using contraception, use injectable medroxyprogesterone or an intrauterine device;
  - in the event of concomitant fluconazole administration, administer each drug 12 hours apart (rifampicin in the morning, fluconazole in the evening);
  - for the other drugs, adjust dosage if necessary.
- **Pregnancy and breastfeeding:** no contra-indication. If used in late pregnancy, administer phytomenadione (vitamin K₁) to the mother and the neonate.

### Monitoring

- Symptomatic monitoring.
- Liver function in patients with hepatic disease.

### Patient instructions

- Take without food (or with a small amount of food to increase gastrointestinal tolerance).
- Harmless orange-red discoloration of the urine, faeces, sweat, saliva, sputum, tears and other body fluids.

### Remarks

- For patients on drug-susceptible TB treatment, rifampicin is given as part of a fixed-dose combination.
- For the 6HRZ-Eto regimen for drug-susceptible TB meningitis, the dose of rifampicin is 20 mg/kg once daily (max. 600 mg daily).
- Rifampicin is also used in the treatment of latent TB infection.

### Storage

- Below 25 °C

---

**Rifapentine (P)**

Update: October 2022
Forms and strengths

- 300 mg and 150 mg coated tablets

Dosage

- Child 12 years and over and adult 40 kg and over: 1200 mg once daily

<table>
<thead>
<tr>
<th>Age</th>
<th>Daily dose (mg)</th>
<th>300 mg tablet</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 12 years</td>
<td>Do not administer</td>
<td>–</td>
</tr>
<tr>
<td>≥ 12 years</td>
<td>1200</td>
<td>4</td>
</tr>
</tbody>
</table>

Contra-indications, adverse effects, precautions

- Do not administer to patients with hypersensitivity to rifamycins or history of severe haematological disorders (thrombocytopenia, purpura) during a previous treatment with rifamycins.
- Avoid or administer with caution to patients with hepatic disorders.
- May cause:
  - hepatotoxicity;
  - influenza-like symptoms, thrombocytopenia, hypersensitivity reactions.
- For the management of adverse effects see Appendix 17.
- Rifapentine reduces the effect of many drugs (antimicrobials, some antiretrovirals, some hormones, antidiabetics, corticosteroids, phenytoin, direct-acting antivirals for chronic hepatitis C, warfarin, etc.):
  - in patients taking antiretrovirals see Appendix 19.
  - in women using contraception, use injectable medroxyprogesterone or an intrauterine device;
  - in the event of concomitant fluconazole administration, administer each drug 12 hours apart (rifampicin in the morning, fluconazole in the evening);
  - for the other drugs, adjust dosage if necessary.
- Pregnancy and breastfeeding: not recommended (safety not established).

Monitoring

- Symptomatic monitoring.
- Liver function in patients with hepatic disease.

Patient instructions

- Take with food.
- Harmless orange-red discoloration of the urine, faeces, sweat, saliva, sputum, tears and other body fluids.

Remarks

- While rifampicin should be taken on an empty stomach, rifapentine is better absorbed if taken with food.
- Also comes in fixed dose combination containing 300 mg of rifapentine/300 mg of isoniazid which can be used in the treatment regimen 2HPZ-Mfx/2HP-Mfx for drug-susceptible TB.
- Rifapentine is also used in the treatment of latent TB infection in children, adolescents, and adults.

Storage

🌞 – ❄ – Below 25 °C
Streptomycin (S)
Update: January 2022

Forms, strengths and route of administration

- Powder for injection, in vial of 1 g streptomycin base, to be dissolved in 4 ml of water for injection, for IM injection
  DO NOT ADMINISTER BY IV INJECTION.

Dosage

- Adolescent 30 kg and over and adult: 12 to 18 mg/kg once daily
- Adult 60 years and over: 15 mg/kg 3 times a week
- Maximum dose: 1000 mg daily
- Renal insufficiency: 12 to 15 mg/kg 2 or 3 times a week

The daily doses take into account the displacement volume (see note below).

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Daily dose (mg)</th>
<th>Daily dose (ml) - IM injection (1 g in 4 ml of water for injection; final volume 4.83 ml; 207 mg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-29</td>
<td>Not used in patients &lt; 30 kg</td>
<td></td>
</tr>
<tr>
<td>30-33</td>
<td>500</td>
<td>2.4 ml</td>
</tr>
<tr>
<td>34-40</td>
<td>600</td>
<td>2.8 ml</td>
</tr>
<tr>
<td>41-45</td>
<td>700</td>
<td>3.4 ml</td>
</tr>
<tr>
<td>46-50</td>
<td>800</td>
<td>4 ml</td>
</tr>
<tr>
<td>51-70</td>
<td>900</td>
<td>4.4 ml</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>1000</td>
<td>Entire volume</td>
</tr>
</tbody>
</table>

Note: displacement volume

Powders for injection are usually formulated such that after reconstitution the final content of the vial corresponds to an adult dose. Errors may occur when only part of the reconstituted solution is to be administered and no allowance is made for the displacement volume. The risk of error increases the greater the weight of the powder and the smaller the volume of solvent used.

Contra-indications, adverse effects, precautions

- Do not administer to children or adolescents under 30 kg and patients with allergy to aminoglycosides.
- Administer with caution to patients 60 years and over or patients with pre-existing renal, vestibular, auditory or severe hepatic impairment.
- May cause:
  - ototoxicity, nephrotoxicity, electrolyte disturbances; rarely, hypersensitivity reactions;
  - local pain after injection.
- For the management of adverse effects see Appendix 17.
- Avoid or monitor combination with other ototoxic and/or nephrotoxic drugs (furosemide, amphotericin B, tenofovir, etc.)
Monitoring

- Symptomatic monitoring.
- Audiometry, serum creatinine and electrolytes (K, Ca, Mg).

Patient instructions

- Maintain a good fluid intake to limit renal problems.

Remarks

- Use a different site for each injection (absorption may be delayed if the same site is used repeatedly).

Storage

- Below 25 °C

Patient instructions

Update: January 2022

- Patients on drug-susceptible TB treatment
- Patients on drug-resistant TB treatment

Patients on drug-susceptible TB treatment

TB drugs are usually well tolerated. However, inform patients that they should immediately seek medical attention in the event of:

- Skin rash
- Yellowing of the skin or eyes or dark urine
- Numbness or tingling of fingers or toes
- Decreased urination
- Palpitations
- Blurred vision, reduced visual acuity, blind spot, green-red colour blindness, eye pain, sensitivity to light
- Pain, burning, swelling of a tendon or muscle
- Pain or swelling in the joints

Patients on drug-resistant TB treatment

Inform patients that they should immediately seek medical attention in the event of:

- Skin rash
- Yellowing of the skin or eyes or dark urine
- Numbness or tingling of fingers or toes
- Decreased urination
- Palpitations
Appendix 11. Use of tuberculosis drugs in pregnant or breastfeeding women

Update: October 2022
<table>
<thead>
<tr>
<th>TB drugs</th>
<th>Evidence and recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FQs</strong></td>
<td>For DR-TB: commonly used in pregnant women despite limited data. Associated with low birth weight in one observational study[1]. As FQs reduce mortality from DR-TB, the benefits often outweigh the risks. Avoid breastfeeding if possible[2] (no absolute contra-indication). For DS-TB: do not use the regimen 2HPZ-Mfx/2HP-Mfx in pregnant or breastfeeding women.</td>
</tr>
<tr>
<td><strong>Bdq</strong></td>
<td>No evidence of fetal harm in animal studies. Associated with low birth weight in one observational study[1]. As Bdq reduces mortality from DR-TB, the benefits often outweigh the risks. Avoid breastfeeding if possible (high concentrations in human and animal breast milk)[3][4].</td>
</tr>
<tr>
<td><strong>Lzd</strong></td>
<td>Few reported cases of use in pregnant women. Fetal harm in animal studies. As Lzd reduces mortality from DR-TB, the benefits often outweigh the risks. Avoid breastfeeding if possible (no data).</td>
</tr>
<tr>
<td><strong>Cfz</strong></td>
<td>Despite common use for leprosy and MDR-TB in pregnant women, few data on pregnancy outcomes. Fetal harm in animal studies. Use during pregnancy only if the benefits outweigh the risks. Avoid breastfeeding if possible (no data). If used, inform mother of possible (and reversible) skin discolouration of the breastfed infant.</td>
</tr>
<tr>
<td><strong>Cs, Trd</strong></td>
<td>Use during pregnancy only if the benefits outweigh the risks (no data). No contra-indication during breastfeeding.</td>
</tr>
<tr>
<td><strong>Dlm</strong></td>
<td>Use during pregnancy only if benefits outweigh the risks (limited human data, fetal harm in animal studies). Avoid breastfeeding if possible (high concentrations in animal breast milk).</td>
</tr>
<tr>
<td><strong>Ipm/Cln, Mpm</strong></td>
<td>Use during pregnancy and breastfeeding only if the benefits outweigh the risks (no data).</td>
</tr>
<tr>
<td><strong>Am, S</strong></td>
<td>Contra-indicated in pregnancy. No contra-indication during breastfeeding[6].</td>
</tr>
<tr>
<td><strong>Eto, Pto</strong></td>
<td>For DR-TB: contra-indicated in pregnancy (fetal harm in animal studies[5]). In breastfeeding women, use only if the benefits outweigh the risks (limited data). For DS-TB: do not use the regimen 6HRZ-Eto in pregnant or breastfeeding women.</td>
</tr>
<tr>
<td><strong>PAS</strong></td>
<td>Use in pregnancy only if the benefits outweigh the risks (limited human data, no fetal harm in animal studies). Avoid breastfeeding if possible (no data).</td>
</tr>
<tr>
<td><strong>Pa</strong></td>
<td>Use during pregnancy and breastfeeding only if the benefits outweigh the risks (no human data, no fetal harm in animal studies[6]).</td>
</tr>
<tr>
<td><strong>P, Rfb</strong></td>
<td>Not recommended during pregnancy and breastfeeding.</td>
</tr>
</tbody>
</table>

For more specific recommendations for pregnant and breastfeeding women see [Chapter 9](#), [Chapter 10](#), and [Appendix 10](#).
Appendix 12. Dose adjustments in renal insufficiency

Update: January 2022

12.1 Normal values for creatinine clearance (CrCl)

Women: 88 to 128 ml/minute
Men: 97 to 137 ml/minute

12.2 Estimation of CrCl (Cockcroft–Gault method)

12.2.1 If serum creatinine is in µmol/litre

\[
\text{Weight (kg)} \times (140 - \text{age}) \times (\text{constant})
\]

\[
\frac{\text{Serum creatinine (µmol/litre)}}{}
\]

The constant = 1.04 for women and 1.23 for men

12.2.2 If serum creatinine is in mg/dl

\[
\frac{\text{Weight (kg)} \times (140 - \text{age})}{72 \times \text{serum creatinine (mg/dl)}}
\]

For women, the result must be multiplied by 0.85.

Example (calculation with serum creatinine in µmol/litre): A woman on cycloserine (Cs), 50 kg, 46 years, serum creatinine = 212 µmol/litre

- **Step 1** - Calculate the CrCl:
  \[
  50 \times (140 - 46) \times 1.04 = 4,888
  \]
  \[
  4,888 \div 212 = 23.1
  \]
  For this patient, the CrCl is 23.1 ml/minute

- **Step 2** - CrCl is < 30 ml/minute, administer 250 mg of Cs once daily or 500 mg 3 times a week.

- **Step 3** - Adjust each drug as required according to the table below.
12.2.3 Overweight and obese patients

For overweight (BMI > 25) or obese (BMI > 30) patients, use the ideal body weight (IBW) rather than the actual body weight to avoid overestimation of the CrCl.

The IBW is calculated using the patient’s height:\nIBW women (kg) = 45.4 + 0.89 (height in cm – 152.4)\nIBW men (kg) = 49.9 + 0.89 (height in cm – 152.4)

Example:\nA woman, weight 70 kg, height 160 cm (BMI = 27.3, i.e. overweight)\n45.4 + 0.89 (160 – 152.4) = 45.4 + 0.89 (7.6) = 45.4 + 6.76 = 52.2
For this patient, the IBW is 52 kg.

12.3 Dosing of TB drugs in renal insufficiency
<table>
<thead>
<tr>
<th>Drugs</th>
<th>Dose and frequency if Clcr &lt; 30 ml/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>No change</td>
</tr>
<tr>
<td>R</td>
<td>No change</td>
</tr>
<tr>
<td>Z</td>
<td>25 mg/kg 3 times a week (not daily)</td>
</tr>
<tr>
<td>E</td>
<td>15-25 mg/kg 3 times a week (not daily)</td>
</tr>
<tr>
<td>Rfb</td>
<td>No change</td>
</tr>
<tr>
<td>Mfx</td>
<td>No change</td>
</tr>
<tr>
<td>Lfx</td>
<td>750-1000 mg 3 times a week (not daily)</td>
</tr>
<tr>
<td>Bdq(^{(a)})</td>
<td>No change</td>
</tr>
<tr>
<td>Lzd</td>
<td>No change</td>
</tr>
<tr>
<td>Cfz</td>
<td>No change</td>
</tr>
<tr>
<td>Cs(^{(b)})</td>
<td>250 mg once daily or 500 mg 3 times a week</td>
</tr>
<tr>
<td>Dlm(^{(a)})</td>
<td>No change</td>
</tr>
</tbody>
</table>
| Ipm/Cin| 750 mg every 12 hours for CrCl 20-40 ml/min  
500 mg every 12 hours for CrCl < 20 ml/min |
| Mpm    | 750 mg every 12 hours for CrCl 20-40 ml/min  
500 mg every 12 hours for CrCl < 20 ml/min |
| Am\(^{(c)}\) | 12-15 mg/kg 2 or 3 times a week (not daily) |
| S\(^{(c)}\) | 12-15 mg/kg 2 or 3 times a week (not daily) |
| Eto/Pto| No change                            |
| PAS\(^{(d)}\) | 4 g 2 times daily                    |
| H\(^{h}\) | No information                      |
| Amx/Clv\(^{(d)}\) | No change                  |
| P      | No change                            |
| Pa     | No information                      |

(a) Use with caution in case of severe renal insufficiency or dialysis (limited data).
(b) Monitor carefully for signs of neurotoxicity.
(c) Use with caution in case of severe renal insufficiency or dialysis (increased risk of nephrotoxicity and ototoxicity).
(d) Avoid sodium salt formulations of PAS in patients with severe renal disease (risk of excessive sodium load).
(e) On a case-by-case basis, consider once daily dosing (e.g. 500/125 mg every 24 hours) for patients with CrCl < 10 ml/minute.

**Notas**
(a) If possible use a calculator to avoid errors, e.g.: [https://www.mdcalc.com/creatinine-clearance-cockcroft-gault-equation](https://www.mdcalc.com/creatinine-clearance-cockcroft-gault-equation)

(b) If possible use a calculator to avoid errors, e.g.: [https://www.mdcalc.com/ideal-body-weight-adjusted-body-weight](https://www.mdcalc.com/ideal-body-weight-adjusted-body-weight)

**Appendix 13. Daily dose of tuberculosis drugs using fixed-dose combinations**

Update: October 2022

**13.1 Conventional regimens for drug-susceptible tuberculosis**

**Intensive phase**

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Paediatric formulations</th>
<th>Adult formulations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HZR 50/150/75 mg</td>
<td>E 100 mg</td>
</tr>
<tr>
<td>4-7</td>
<td>1 tab</td>
<td>1 tab</td>
</tr>
<tr>
<td>8-11</td>
<td>2 tab</td>
<td>2 tab</td>
</tr>
<tr>
<td>12-13</td>
<td>3 tab</td>
<td>2 tab</td>
</tr>
<tr>
<td>14-15</td>
<td>3 tab</td>
<td>3 tab</td>
</tr>
<tr>
<td>16-17</td>
<td>4 tab</td>
<td>3 tab</td>
</tr>
<tr>
<td>18-22</td>
<td>4 tab</td>
<td>–</td>
</tr>
<tr>
<td>23-29</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>30-34</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>35-39</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>40-54</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>55-70</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>
For example:
- A child weighing 9 kg takes 2 tablets of HZR (50 mg/150 mg/75 mg) + 2 tablets of E (100 mg) once daily.
- A child weighing 20 kg takes 4 tablets of HZR (50 mg/150 mg/75 mg) + 1 tablet of E (400 mg) once daily.

**Note:** ethambutol is not routinely given to all children: see Chapter 9.

### Continuation phase

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>Paediatric formulation HR 50/75 mg</th>
<th>Adult formulation HR 75/150 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-7</td>
<td>1 tab</td>
<td></td>
</tr>
<tr>
<td>8-11</td>
<td>2 tab</td>
<td></td>
</tr>
<tr>
<td>12-14</td>
<td>3 tab</td>
<td></td>
</tr>
<tr>
<td>15-21</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>22-29</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>30-34</td>
<td>–</td>
<td>2 tab</td>
</tr>
<tr>
<td>35-39</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>40-54</td>
<td>–</td>
<td>3 tab</td>
</tr>
<tr>
<td>55-70</td>
<td>–</td>
<td>4 tab</td>
</tr>
<tr>
<td>&gt; 70</td>
<td>–</td>
<td>4 tab</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TB drugs</th>
<th>Daily dosing in patients &lt; 30 kg</th>
<th>Daily dosing in patients ≥ 30 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>E</td>
<td>15 to 25 mg/kg</td>
<td>15 to 25 mg/kg</td>
</tr>
<tr>
<td>H</td>
<td>7 to 15 mg/kg</td>
<td>4 to 6 mg/kg</td>
</tr>
<tr>
<td>Z</td>
<td>30 to 40 mg/kg</td>
<td>20 to 30 mg/kg</td>
</tr>
<tr>
<td>R</td>
<td>10 to 20 mg/kg</td>
<td>8 to 12 mg/kg</td>
</tr>
</tbody>
</table>

### 13.2 2HPZ-Mfx/2HP-Mfx regimen for drug-susceptible tuberculosis
Appendix 14. Monitoring of patients on drug-susceptible tuberculosis treatment

Update: October 2022

A cross "X" with no brackets indicates that the exam should be performed in all patients.
A cross between brackets "(X)" indicates that the exam should only be performed in certain patients.
<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>Treatment</th>
<th></th>
<th>End of treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>W2</td>
<td>M1</td>
<td>M2</td>
<td>M3</td>
</tr>
<tr>
<td>Clinical visits</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adverse events</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Bacteriological tests</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rapid molecular tests&lt;sup&gt;(b)&lt;/sup&gt;</td>
<td>X</td>
<td></td>
<td>(X)</td>
<td></td>
</tr>
<tr>
<td>Smear microscopy</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Culture and pDST&lt;sup&gt;(c)&lt;/sup&gt;</td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
<td></td>
</tr>
<tr>
<td>Other investigations</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Radiography&lt;sup&gt;(d)&lt;/sup&gt;</td>
<td>(X)</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
</tr>
<tr>
<td>Full blood count&lt;sup&gt;(e)&lt;/sup&gt;</td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
<td></td>
</tr>
<tr>
<td>Liver function&lt;sup&gt;(f)&lt;/sup&gt;</td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
</tr>
<tr>
<td>Serum creatinine&lt;sup&gt;(g)&lt;/sup&gt;</td>
<td>(X)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c, blood glucose&lt;sup&gt;(h)&lt;/sup&gt;</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HIV, HBV, HCV&lt;sup&gt;(i)&lt;/sup&gt;</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CD4 and viral load&lt;sup&gt;(j)&lt;/sup&gt;</td>
<td>(X)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(a) For treatments longer than 6 months.
(b) Rapid molecular tests:
   - Xpert MTB/RIF (or Ultra) and Xpert MTB/XDR (or GenoType MTBDRsl if Xpert MTB/XDR not available).
   - Repeat RMTs if microscopy or culture is positive at Month 2 or later.
(c) Culture and pDST to first- and second-line drugs:
   - At baseline if RMTs are not available, to detect rifampicin and isoniazid resistance or rifampicin resistance mutations not detected by RMTs.
   - At Month 2 or later, if RMTs show a new resistance.
   - At Month 4, if microscopy is still positive.
(d) Radiography:
   - Chest: at baseline for children with presumptive PTB, patients with non-bacteriologically confirmed PTB, suspicion of other intra-thoracic TB, then if indicated (e.g. worsening respiratory symptoms, non-response to TB treatment).
   - Bone: at baseline then every 6 months for patients with bone and joint TB.
Appendix 15. Monitoring of patients on drug-resistant tuberculosis treatment

Update: October 2022

A cross "X" with no brackets indicates that the exam should be performed in all patients.
A cross between brackets "(X)" indicates that the exam should only be performed in certain patients.

(e) For patients on AZT or rifabutin.
(f) For patients with pre-existing hepatic disease: AST and ALT (and bilirubin if AST or ALT are elevated).
(g) For patients with renal insufficiency.
(h) For all patients to detect diabetes. If diabetes is detected, monitor according to standard protocols.
(i) For all patients, unless documented HIV, hepatitis B and C status; HIV test every 6 months in high HIV prevalence areas.
(j) For HIV-infected patients.
<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>W1</th>
<th>W2</th>
<th>W3</th>
<th>W4</th>
<th>W5</th>
<th>W6</th>
<th>W7</th>
<th>M2</th>
<th>M3</th>
<th>Until end of treatment</th>
<th>End of treatment</th>
<th>Post end of treatment</th>
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</thead>
<tbody>
<tr>
<td><strong>Clinical visits</strong></td>
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<tr>
<td>Vital signs, weight, etc.</td>
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<td>X</td>
<td>X</td>
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<td>X</td>
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<td>X</td>
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<td>At each visit</td>
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<tr>
<td>Adverse events</td>
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<td>At each visit</td>
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<tr>
<td>BPNS&lt;sup&gt;(a)&lt;/sup&gt;</td>
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<tr>
<td>Visual function tests&lt;sup&gt;(b)&lt;/sup&gt;</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
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<tr>
<td>Audiometry&lt;sup&gt;(c)&lt;/sup&gt;</td>
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<td>(X)</td>
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<td>(Monthly)</td>
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<tr>
<td>ECG&lt;sup&gt;(d)&lt;/sup&gt;</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
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<tr>
<td><strong>Bacteriological tests</strong></td>
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<tr>
<td>Smear microscopy</td>
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<td>Monthly</td>
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<td>Culture</td>
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<td></td>
<td>Monthly</td>
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<tr>
<td>Rapid molecular tests&lt;sup&gt;(e)&lt;/sup&gt;</td>
<td></td>
<td>X</td>
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<td></td>
<td></td>
<td>If culture or microscopy positive at M4 or later</td>
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<td></td>
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<tr>
<td>Full pDST&lt;sup&gt;(f)&lt;/sup&gt;</td>
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<td>X</td>
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<td></td>
<td></td>
<td>If culture positive at M4 or later</td>
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<tr>
<td><strong>Other investigations</strong></td>
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<tr>
<td>Radiography&lt;sup&gt;(g)&lt;/sup&gt;</td>
<td></td>
<td>X</td>
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<td></td>
<td>X</td>
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<tr>
<td>Full blood count&lt;sup&gt;(h)&lt;/sup&gt;</td>
<td></td>
<td>X</td>
<td>(X)</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
<td>(X)</td>
<td></td>
<td></td>
<td>(Monthly)</td>
<td></td>
<td>(X)</td>
</tr>
<tr>
<td>Liver function&lt;sup&gt;(i)&lt;/sup&gt;</td>
<td></td>
<td>X</td>
<td>(X)</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
<td></td>
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<td>(Monthly)</td>
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<tr>
<td>Serum creatinine and potassium&lt;sup&gt;(j)&lt;/sup&gt;</td>
<td></td>
<td>X</td>
<td>(X)</td>
<td></td>
<td>(X)</td>
<td>(X)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(Monthly)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HbA1c, blood</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
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<td></td>
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</tbody>
</table>
Appendix 16. Additional investigations in drug-resistant tuberculosis

Update: October 2022

16.1 Electrocardiogram (ECG)

The QT interval is measured in milliseconds (ms) from the start of the QRS complex to the end of the T wave of the ECG. Its value varies depending on the heart rate and should be corrected accordingly (QTc).

To calculate the QTc interval it is recommended to use the Fridericia formula (QTcF):

$$\text{QTcF} = \frac{\text{QT interval}}{\sqrt[3]{\text{interval between two waves R}}}$$

<table>
<thead>
<tr>
<th>Investigation</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose (k)</td>
<td>If indicated</td>
</tr>
<tr>
<td>HIV, HBV, HCV (l)</td>
<td>If indicated</td>
</tr>
<tr>
<td>CD4 and viral load (m)</td>
<td>(Every 6 months)</td>
</tr>
<tr>
<td>TSH (n)</td>
<td>(Every 3 months)</td>
</tr>
<tr>
<td>Pregnancy test (o)</td>
<td>If indicated</td>
</tr>
</tbody>
</table>

(a) For patients on Lzd.
(b) For patients on E, Lzd or thionamides: visual acuity and colour vision deficiency.
(c) For patients on Am or S.
(d) Electrocardiogram, for patients taking:
   - < 2 moderate or severe QT-prolonging TB drugs or < 3 QT-prolonging drugs (TB and non-TB): at baseline then monthly.
   - ≥ 2 moderate or severe QT-prolonging TB drugs or ≥ 3 QT-prolonging drugs (TB and non-TB) or with other risk factors for QT prolongation or TdP: once a week for the first month, then once a month.
(e) Rapid molecular tests:
   - Xpert MTB/RIF (or Ultra) and Xpert MTB/XDR (or GenoType MTBDRsl if Xpert MTB/XDR not available).
   - Repeat Xpert MTB/XDR (or GenoType MTBDRsl) if culture or microscopy is positive at Month 4 or later.
(f) For first- and second-line drugs. Repeat if culture is positive at Month 4 or later.
(g) For all patients at baseline, then every 6 months.
(h) For all patients at baseline, then:
   - Patients on Lzd: every 2 weeks for the first 2 months, then once a month.
   - Patients on AZT: once a month for the first 2 months, then if indicated.
(i) For all patients: AST and ALT (and bilirubin if AST or ALT are elevated).
(j) For all patients at baseline. Repeat if indicated. For patients on Am or S: once a month or more frequently if indicated.
(k) For all patients to detect diabetes. If diabetes is detected, monitor according to standard protocols.
(l) For all patients, unless documented HIV, hepatitis B and C status; HIV test every 6 months in high HIV prevalence areas.
(m) For HIV-infected patients.
(n) For patients on thionamides or PAS.
(o) For adolescents and women of childbearing age. Repeat if indicated.
Normal QTc values:
< 470 ms in women
< 450 ms in men

16.2 Brief peripheral neuropathy screen (BPNS)

Adapted from AIDS Clinical Trial Group (ACTG)¹².

Step 1. Grade subjective symptoms

- Ask the patient to rate the severity of symptoms on a scale from 0 (no symptoms) to 10 (most severe symptoms) for right (R) and left (L) feet and legs.
- Enter the score for each symptom in the corresponding column.

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>R</th>
<th>L</th>
</tr>
</thead>
<tbody>
<tr>
<td>a. Pain or burning sensation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b. Pins and needles sensation (tingling sensation)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c. Numbness (lack of feeling)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Symptoms may be unilateral or bilateral and of different intensity. Use the highest subjective sensory neuropathy score to obtain the severity grade.

<table>
<thead>
<tr>
<th>Subjective sensory neuropathy score</th>
<th>Severity grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1-3</td>
<td>1</td>
</tr>
<tr>
<td>4-6</td>
<td>2</td>
</tr>
<tr>
<td>7-10</td>
<td>3</td>
</tr>
</tbody>
</table>
**Step 2. Evaluate vibration perception**

- Place the vibrating 128 Hz tuning fork on the top of the distal joint of the right and left big toes and begin counting the seconds.
- Ask the patient to say when they no longer feel the vibration.

There is a decrease in vibration perception if the patient feels the vibration for 10 seconds or less on both sides.

<table>
<thead>
<tr>
<th>Vibration perception</th>
<th>Result</th>
<th>Grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Felt &gt; 10 seconds</td>
<td>Normal</td>
<td>0</td>
</tr>
<tr>
<td>Felt 6-10 seconds</td>
<td>Mild loss</td>
<td>1</td>
</tr>
<tr>
<td>Felt &lt; 5 seconds</td>
<td>Moderate loss</td>
<td>2</td>
</tr>
<tr>
<td>Not felt</td>
<td>Severe loss</td>
<td>3</td>
</tr>
</tbody>
</table>

**Step 3. Evaluate tendon reflexes**

Using a reflex hammer, tap the Achilles tendon on each ankle.

**Step 4. Make a diagnosis**

Diagnosis of peripheral neuropathy is based on the combination of:
- subjective symptoms of grade 1, 2 or 3, and
- at least one bilateral objective finding:
  - reduced vibration perception (grade 1, 2 or 3), or
  - decreased reflexes (absent or hypoactive reflexes)

**16.3 Ishihara test**

The patient is asked to look at a set of plates with circles made of dots of different sizes and colours.

Some circles contain dots that form a number or a shape clearly visible to patients with normal colour vision. Patients who cannot see or have difficulty distinguishing numbers or shapes have a red-green colour vision defect.

Some circles contain dots that form a number or a shape visible to patients with red-green colour vision defect, but invisible to patients with normal colour vision.

The test should be performed as per the manufacturer’s instructions.

**Notas**

(a) When possible, use a calculator to avoid errors, e.g. [https://www.mdcalc.com/corrected-qt-interval-qtc](https://www.mdcalc.com/corrected-qt-interval-qtc)

**Referencias**

   [https://n.neurology.org/content/65/11/1778.long](https://n.neurology.org/content/65/11/1778.long)

Appendix 16. Basic TB infection control risk assessment tool

Appendix 17. Management of adverse effects

Update: January 2022

- Gastrointestinal disorders
  - Abdominal pain
  - Diarrhoea
  - Epigastric pain
  - Hepatotoxicity
  - Metallic taste
  - Nausea and vomiting
- Neurotoxicity
  - Depression
  - Headache
  - Optic neuritis
  - Ototoxicity
  - Peripheral neuropathy
  - Psychosis
  - Seizures
- Endocrine disorders
  - Gynecomastia
  - Hypothyroidism
- Dermatological disorders
  - Alopecia
  - Fungal infection
  - Photosensitivity
  - Skin reactions
- Musculoskeletal disorders
  - Arthralgias
  - Tendinitis/tendon rupture
- Miscellaneous
  - Electrolyte disorders
  - Haematologic disorders
  - Lactic acidosis
  - Nephrotoxicity
  - QT prolongation

Gastrointestinal disorders
Abdominal pain

Eto or Pto, PAS, Cfz, Lzd, FQs, H, Z

Abdominal pain is common with MDR/RR-TB treatment. It can be the early sign of severe adverse effects such as hepatitis, pancreatitis, or lactic acidosis.

Deposition of Cfz crystals may cause severe abdominal pain (presentation of acute abdomen). In this case, stop Cfz until symptoms resolve.

Diarrhoea

PAS, FQs, Eto or Pto, Amx/Clv, Ipm/Cln or Mpm

Diarrhoea, along with cramping, can cause significant difficulty and lead to discontinuation of treatment.

PAS often causes diarrhoea at treatment initiation. It usually resolves or improves substantially after some weeks.

For diarrhoea with no blood in stools and no fever, loperamide PO (adult: 4 mg followed by 2 mg after each loose stool to a maximum of 10 mg daily) may be used intermittently, especially when the patient needs to attend social functions or return to work, but not on a daily basis.

Encourage the patient to tolerate some degree of diarrhoea. Prevent (encourage fluid intake including oral rehydration solution) or treat dehydration.

In the event of severe diarrhoea, particularly if associated with blood in stools, severe abdominal pain, or fever > 38.5 °C, consider other causes such as acute bacterial enteritis, or pseudo-membranous colitis (C. difficile) due to FQs. Do not use loperamide in bloody diarrhoea or diarrhoea associated with fever.

Monitor serum electrolytes in patients with severe diarrhoea on QT prolonging drugs.

Epigastric pain

PAS, Eto or Pto, FQs, E, Z

Gastritis (epigastric burning or cramp relieved by eating) or dyspepsia (epigastric pain or discomfort following meals, often accompanied by bloating, sensation of fullness and nausea) are frequent with PAS, Eto or Pto.
Haematemesis (vomiting of blood) and melena (black stools) are symptoms of a bleeding gastric ulcer and require urgent intervention.

Hepatotoxicity

Z, H, R, P, Eto or Pto, PAS, Bdq, Amx/Clv

All TB drugs may cause hepatotoxicity. However, certain drugs are likely more responsible than others for this adverse effect.

The liver function tests (LFTs) used for the diagnosis and monitoring of hepatotoxicity are serum levels of alanine aminotransferase (ALT), aspartate aminotransferase (AST) and bilirubin.

A mild, transient elevation of ALT and AST may be observed during treatment and usually remains asymptomatic. Significant hepatotoxicity is usually symptomatic.

Clinical features resemble that of viral hepatitis. Early symptoms include malaise, fatigue, loss of appetite, muscle and joint pain. Nausea, vomiting and abdominal pain are common in severe toxicity. Jaundice, scleral icterus, dark (tea-coloured) urine and discoloured stool are signs of clinical worsening.

Differential diagnosis includes infections (e.g. viral hepatitis, cytomegalovirus, leptospirosis, yellow fever, rubella), chronic alcohol use and hepatotoxicity due to other drugs (e.g. anti-epileptics, paracetamol, sulfa drugs, erythromycin).

Clinical hepatitis can be fatal and action should be taken immediately.

1) General management

- Patient with symptoms of hepatitis:
  Stop all TB drugs and perform LFTs:
  a) AST or ALT or bilirubin ≥ 3 times upper limit of normal (ULN): wait for resolution of symptoms, perform LFTs weekly and restart TB treatment when LFTs are < 3 times ULN.
  b) AST, ALT and bilirubin < 3 times ULN and mild symptoms (no jaundice): restart TB treatment, closely monitor the patient and perform LFTs weekly. Continue TB treatment as long as LFTs levels remain < 3 ULN and there are no signs of worsening hepatitis.

- Patient without symptoms of hepatitis, but elevated LFTs:
  a) AST or ALT ≥ 5 times ULN or bilirubin ≥ 3 ULN: stop all TB drugs and perform LFTs weekly. Restart TB treatment when LFTs return < 3 times ULN.
  b) AST and ALT < 5 times ULN and bilirubin < 3 ULN: continue TB treatment and perform LFTs weekly.

If LFTs continue to increase after stopping TB treatment, then ongoing progressive drug-induced hepatitis or an unrelated cause of hepatitis should be suspected.

2) Patient on DS-TB treatment

In most cases, the same treatment can be resumed without incident. The objective is to resume the initial regimen or an alternative regimen as rapidly as possible.

If symptoms reappear or LFTs re-increase, try to reintroduce the TB drugs one by one. Start with E and R and reintroduce H three to 7 days later. If E, R and H have been introduced and the LFT abnormalities have not recurred, do not introduce Z as it is most likely the causative agent.

The alternative regimen depends on the drug causing hepatotoxicity:

- **For gastritis:**
  omeprazole PO: 20 mg once daily in the morning for 7 to 10 days. In severe or recurrent cases, dose may be increased to 40 mg once daily and the treatment may be prolonged for up to 8 weeks.
  Histamine H2-antagonists (e.g., ranitidine) may be an alternative.

- **For dyspepsia:**
  omeprazole PO: 10 mg once daily in the morning for 4 weeks

Haematemesis (vomiting of blood) and melena (black stools) are symptoms of a bleeding gastric ulcer and require urgent intervention.
3) Patient on DR-TB treatment

When restarting TB treatment, start with the drugs least hepatotoxic (E, Lfx or Mfx, Cs or Trd, Dlm, Am or S, Ipm/Cln or Mpm), then drugs moderately hepatotoxic (Bdq, Cfz, Amx/Clav), then give the most hepatotoxic (Z, H, R, Eto or Pto, PAS). Add drugs one at a time every 5 to 7 days, and check LFTs.

The causative agent can generally be identified in this manner. It can be discontinued if not essential and replaced with another less hepatotoxic TB drug.

Note: hepatotoxicity may occur in patients receiving regimens containing pretomanid (Pa-Mfx-Z and Bdq-Pa-Lzd). However, the responsible drug has not been determined.

Metallic taste

Eto or Pto, FQs

Encourage the patient to tolerate this adverse effect. Normal taste returns when TB treatment is stopped.

Nausea and vomiting

Eto or Pto, PAS, Z, Amx/Clv, Cfz, Lzd, Ipm/Cln or Mpm, Bdq

Nausea and vomiting are frequent, especially with Eto or Pto and PAS during the first few weeks of treatment. To avoid nausea and vomiting, these drugs can be initiated at low dose with gradual increase over one to 2 weeks.

- Always look for:
  - Signs of dehydration (thirst, dry mouth, sunken eyes)
  - Serum electrolytes disorders if vomiting
  - Signs of hepatitis
  - Haematemesis and melena
- Dehydration and electrolyte disorders should be corrected as necessary.
- Treat nausea and vomiting aggressively, using a stepwise approach:

  First phase - Adjust administration of the responsible drug
  - Try to identify the drug(s) causing nausea and vomiting. Stop it/them for 2 or 3 days, then gradually reintroduce.
  - Administer the drug(s) causing nausea at bedtime.
  - Patient on PAS:
    - Take one hour after taking other TB drugs.
    - If PAS is taken once daily, take in 2 divided doses.
  - Encourage the patient: nausea and vomiting often improve over the first weeks and may resolve entirely with time.

  Second phase - Administer an antiemetic
**ondansetron** PO 30 minutes before TB drugs:
Child 6 months to < 2 years: 2 mg once daily
Child 2 to < 4 years: 2 mg 2 times daily
Child 4 to < 12 years: 4 mg 2 times daily
Child ≥ 12 years and adult: 4 to 8 mg 2 times daily

Ondansetron is a QT prolonging drug and should be avoided in patients on Cfz, Bdq, Mfx, Dlm, Lfx.

In adults, when ondansetron is not available or is to be avoided:
**metoclopramide** PO:
Adult < 60 kg: 5 mg 3 times daily
Adult ≥ 60 kg: 10 mg 3 times daily
The interval between each dose should be at least 6 hours (even in the event of vomiting). Do not use metoclopramide if neurological problems develop.

or
**promethazine** PO 30 minutes before TB drugs:
Adult: 25 mg

**Third phase - Reduce the dose or temporarily stop the responsible drug**

- Patient on Eto or Pto: consider reducing dose by one weight class (e.g. if taking 1000 mg daily, reduce to 750 mg). Avoid giving an adult weighing more than 33 kg less than 500 mg daily of Eto or Pto.
- Patient on Cfz: reduce the dose by half.
- In the event of intractable nausea and vomiting, stop all TB drugs until symptoms resolve.

**Note:** if there is excessive anxiety over the nausea caused by TB drugs, consider adding **diazepam** PO (adult: 5 mg 30 minutes before TB drugs). This can help to avoid “anticipation nausea”. The treatment must be short as benzodiazepines may cause dependence and tolerance. Do not exceed 10 days of treatment.

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**Neurotoxicity**

- Depression
- Headache
- Optic neuritis
- Ototoxicity
- Peripheral neuropathy
- Psychosis
- Seizures

---

**Depression**

**Cs or Trd, Eto or Pto**

The treatment of MDR/RR-TB may contribute to depression. Depressive symptoms may fluctuate during TB treatment. History of depression may increase the risk of developing depression during treatment, but is not a contra-indication to use of any of the above TB drugs.

Consider lowering the dose or discontinuing a suspected TB drug, provided this does not compromise the effectiveness of TB treatment.
Other interventions include psychological support to patient (and family if needed) and, when necessary antidepressant treatment. Avoid selective serotonin reuptake inhibitors and tricyclic antidepressants with Lzd (risk of serotonin syndrome).

Suicidal ideation is more commonly associated with Cs or Trd. Evidence of suicidal ideation should prompt immediate action:
- Keep the patient in the hospital for surveillance.
- Stop Cs or Trd.
- Lower the dose of Eto or Pto to 500 mg daily until the patient is stable.
- Refer to mental health consultation.

### Headache

**Cs or Trd, Bdq, Dlm, FQs**

Headache is common during the first months of treatment. It can be treated with analgesics. Headache due to Cs or Trd can be prevented by starting at low dose (250 to 500 mg daily), with gradual increase over 1 to 2 weeks.

### Optic neuritis

**Lzd, E; rarely H, Eto or Pto**

This adverse effect is typically due to Lzd and E.

Symptoms include loss of red-green colour distinction, reduced visual acuity and central scotoma. Loss of red-green colour distinction is the first sign. In this case, stop the suspect drug immediately and permanently.

Symptoms are usually reversible after discontinuation of the drug, but optic neuritis due to Lzd may be irreversible.

### Ototoxicity

**Aminoglycosides; rarely: Cs or Trd, FQs, Eto or Pto, Lzd**

Hearing loss, tinnitus and/or vestibular disorders (vertigo, dizziness, imbalance) are signs of ototoxicity.

Ototoxicity is most commonly observed in patients receiving large cumulative doses of aminoglycosides. Concomitant use of loop diuretics (furosemide), particularly in patients with renal insufficiency, may exacerbate ototoxicity.

Baseline and follow-up audiometry is required to detect early hearing loss. Hearing loss in high frequencies (> 4000 Hz) is often the first sign of auditory toxicity due to aminoglycosides and can be unnoticed by the patient.

In case of hearing loss, tinnitus or vestibular disorders, discontinue the suspected drug if this does not compromise the effectiveness of TB treatment.

If no alternative is available, reduce the dose of aminoglycoside (3 times weekly rather than daily, e.g. on Monday, Wednesday and Friday). Continuation of aminoglycoside therapy despite hearing loss almost always results in deafness.
Tinnitus and vestibular disorders can rarely be due to the following drugs: Cs or Trd, FQs, Eto or Pto and Lzd. If stopping the aminoglycoside does not improve symptoms, other drugs can be discontinued to see if the symptoms improve, then reintroduced one by one to see if symptoms return. Drug-induced tinnitus and vestibular disorders can be irreversible.

**Peripheral neuropathy**

**Lzd, Cs or Trd, H, Eto or Pto; rarely E, FQs**

Peripheral neuropathy refers to damage to the nerves located outside of the central nervous system. This adverse effect is associated to several TB drugs but is commonly due to Lzd, Cs or Trd and H.

Peripheral neuropathy occurs most commonly in the lower extremities. Signs and symptoms include sensory disturbances (e.g. numbness, tingling, burning, pain, loss of temperature sensation), difficulty walking, weakness and decreased or absent deep tendon reflexes. At times, sensory changes may occur in upper extremities.

Linezolid-induced neuropathy is extremely painful and may be non-reversible.

1) **Patient on DS-TB treatment**

- To prevent isoniazid-induced peripheral neuropathy:
  
  Administer **pyridoxine** PO to patients at risk (pregnant and breastfeeding women, neonates and breastfed infants, and patients with HIV infection, alcohol dependency, malnutrition, diabetes, chronic hepatic disease, and renal impairment) along with their TB treatment:
  
  Neonate, infant and child < 5 kg: 5 mg once daily  
  Child ≥ 5 kg and adult: 10 mg once daily

- If peripheral neuropathy develops:
  
  Administer **pyridoxine** PO
  
  Child < 12 years: 10 to 20 mg 2 times daily  
  Child ≥ 12 years: 50 mg 2 times daily  
  Adult: 50 mg 3 times daily
  
  For pain management: ibuprofen or paracetamol.

2) **Patient on DR-TB treatment**

- To prevent peripheral neuropathy:
  
  Administer **pyridoxine** PO:
  
  - Patient on H: all patients at risk, as for DS-TB.
  - Patient on Cs or Trd, Lzd, H11 and Eto or Pto:
    
    Neonate, infant, child: 1 to 2 mg/kg (usual range in child: 10 to 50 mg) once daily  
    Adult: 100 mg once daily

- If peripheral neuropathy develops:
  
  - Patient on Lzd: stop Lzd immediately. For mild symptoms not requiring analgesics, Lzd can be restarted at a lower dose once symptoms subside. For moderate or severe symptoms, stop Lzd permanently. Consider additional TB drugs to reinforce the therapeutic regimen.
  
  - Patient on Cs or Trd or H11: stop these drugs. If they are essential to the regimen, they may be re-introduced once symptoms subside.

Other contributing causes should be addressed (e.g., diabetes or malnutrition).

Administer **pyridoxine** PO: 100 mg daily in adults until symptoms resolve.

For pain management: ibuprofen or paracetamol.

Physiotherapy may be of benefit.

If these measures are insufficient, treat as chronic neuropathic pain, but avoid tricyclic antidepressants in patients on Lzd (risk of serotonin syndrome).
Psychosis

Cs or Trd, FQs, H, Eto or Pto

Visual or auditory hallucinations, delusions, paranoia and bizarre behaviour are hallmarks of psychosis. Health personnel should be familiar with these symptoms to allow early detection.

The most likely TB drug involved is Cs or Trd, but psychotic symptoms may occur with FQs, H, Eto or Pto.

History of psychosis is not a contra-indication to the use of the above-mentioned drugs, though psychiatric symptoms are more likely to occur in such circumstances.

Some patients may need antipsychotic treatment throughout the duration of TB treatment.

Psychosis is generally reversible upon discontinuation of TB treatment.

For acute psychosis:
- If patients are at risk of harming themselves or others: urgent hospitalisation.
- Stop Cs or Trd.
- Treat the acute psychosis.

Once psychotic symptoms have resolved, antipsychotic treatment can be tapered most of the time. Cs or Trd can be resumed, generally at lower dose.

Antipsychotic treatment should be continued until the end of Cs or Trd treatment and then can usually be stopped gradually (do not stop it abruptly).

If the patient does not tolerate the reintroduction of Cs or Trd, another TB drug should be considered.

Whenever psychosis occurs in a patient on Cs or Trd, check the serum creatinine. Cs or Trd is 100% renally excreted and a decrease in renal function can result in toxic levels of Cs or Trd. In this case, a temporary suspension of Cs or Trd and re-introduction at an adjusted dose may be needed (Appendix 12).

Seizures

Cs or Trd, H, FQs, Eto or Pto, Ipm/Cln or Mpm

All the above-mentioned drugs may cause seizures. However, rule out or treat other possible causes (e.g., epilepsy, meningitis, encephalitis, alcohol withdrawal, hypoglycaemia, stroke, cancer, or toxoplasmosis in HIV-infected patients).

In the event of seizures, measure blood glucose level and blood electrolytes. Measure also serum creatinine. With impaired renal function, TB drugs can reach toxic levels, causing seizures. Dosage adjustment may be necessary (Appendix 12).

A history of seizures is not an absolute contra-indication to the use of the above-mentioned drugs. However, do not use Cs or Trd if there is an alternative. In patients with epilepsy, seizures should be controlled with anti-epileptic therapy before starting TB treatment.

The use of TB drugs (especially H and R) in patients on antiepileptics may lead to decreased blood levels of antiepileptics and seizures.

In patients without history of seizures, a first episode of seizures on TB treatment is likely due to the TB drugs. However, none of the above drugs leave permanent damage.

If a patient has a seizure for the first time:
- Stop suspected TB drugs for a short period.
• Start antiepileptic treatment, especially in the event of repeated seizures after stopping suspected drugs. Do not use carbamazepine or phenytoin in patients receiving Bdq or Dlm (strong CYP450 inducers).
• Reintroduce TB drugs that are essential to TB treatment. Usually, they can be resumed at a lower dose, but the effective dose should be reached as soon as possible.

Antiepileptic treatment may be necessary until the end of the TB treatment.

**Endocrine disorders**

- [Gynecomastia](#)
- [Hypothyroidism](#)

**Gynecomastia**

**Eto or Pto**

Eto or Pto may cause breast enlargement in men and women. Galactorrhoea has been reported. Encourage the patient to tolerate this adverse effect. Symptoms resolve when Eto or Pto is stopped.

**Hypothyroidism**

**Eto or Pto, PAS**

Symptoms appear slowly, are nonspecific and may include fatigue, muscle weakness, daytime sleepiness, excessive sensitivity to cold, dry skin, coarse hair, constipation, facial puffiness, and depression. Thyroid enlargement and delayed deep tendon reflexes may be seen on examination.

The diagnosis is confirmed by a serum level of thyroid-stimulating hormone (TSH) ≥ 10 mIU/litre.

Eto or Pto and PAS may cause hypothyroidism, even more frequently when used together. If possible the responsible TB drugs should be replaced but may be continued if there is no alternative.

In both cases, replacement hormone therapy is required:

**Levotiroxine** PO

Adult < 60 years: initially 75 to 100 micrograms once daily then, adjust in 25 microgram increments every 4 to 12 weeks according to response. Usual maintenance dose is 100 to 200 micrograms daily.

Adult ≥ 60 years and/or with significant cardiovascular disease: initially 25 micrograms once daily then, adjust in 25 microgram increments every 4 to 12 weeks according to response. Usual maintenance dose is 100 to 125 micrograms daily.

The daily dose should be taken at the same time each day, 30 to 60 minutes before a meal or a caffeine-containing drink (e.g. coffee, tea) or other drugs to improve absorption.

Monitor TSH until it normalizes below 5 mlU/litre.

Thyroid dysfunction resolves upon discontinuation of TB treatment. Hormone replacement may be discontinued several months after TB treatment completion.
Dermatological disorders

- **Alopecia**
- **Fungal infection**
- **Photosensitivity**
- **Skin reactions**

### Alopecia

**H, Eto or Pto**

Temporary and mild hair loss may (rarely) occur in the first months of treatment. Encourage the patient to tolerate this adverse effect. Symptoms resolve when TB treatment is stopped.

### Fungal infection

**FQs**

Vaginal, penile, skin fold and oral candidiasis may occur in patients taking FQs. Topical antifungals or short-course oral antifungals are usually effective.

### Photosensitivity

**Cfz, FQs; rarely Z**

Advise patient to avoid direct exposure to the sun, wear protecting clothes (e.g. long sleeves) and use sunscreen.

### Skin reactions

**All TB drugs**

Skin reactions such as itch and skin rash may be hypersensitivity reactions due to any TB drug. General signs of hypersensitivity such as fever, dizziness, vomiting and headache may also occur. Skin reactions usually appear early during treatment, often in the first month, but rarely during the first week. Most skin reactions are mild or moderate. Severe – even lethal – exfoliative dermatitis (Stevens Johnson's syndrome) may occasionally occur, particularly if administration of the TB drug continues after first signs of hypersensitivity appear.
**Minor skin reactions**

- Simple itching: symptomatic treatment (e.g. antihistamine) without interrupting or modifying the TB treatment.
- Localised, mild skin rash, with or without itching:
  - Rule out other possible causes unrelated to TB drugs (i.e. scabies, contact dermatitis).
  - If no obvious other cause, stop all TB drugs.
  - Give symptomatic treatment (an antihistamine, no corticosteroids except in emergencies) and wait for disappearance of symptoms.
  - Once the reaction has resolved, try to determine which drug caused the reaction (see rechallenge of TB drugs below).

**Major skin reactions**

- Stop all TB drugs.
- In the event of anaphylaxis, manage according to standard emergency protocol (epinephrine, etc.).
- For severe generalised rash, a parenteral corticosteroid may be needed.
- Once the reaction has resolved, try to determine which TB drug caused the reaction (see rechallenge of TB drugs below).
- Never re-introduce any drug resulting in Stevens-Johnson syndrome or anaphylaxis.

**Rechallenge of TB drugs**

Each TB drug can be reinstated as a “challenge” (a test-dose). Introduce one drug at a time, starting with the drugs least likely to have caused the reaction.

Give the drugs in a setting where a health care provider can respond to any severe allergic reaction.

If a test-dose of any drug causes a reaction, discontinue this drug, unless it is deemed essential to the regimen (in this case, desensitisation can be considered).

- **First-line TB drugs**
  
  Start with isoniazid over 3 days then add rifampicin over 3 days, etc.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Likelihood</th>
<th>Trial dose 1</th>
<th>Trial dose 2</th>
<th>Trial dose 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>Least likely</td>
<td>50 mg</td>
<td>Full dose</td>
<td>Full dose</td>
</tr>
<tr>
<td>R</td>
<td>Least likely</td>
<td>75 mg</td>
<td>300 mg</td>
<td>Full dose</td>
</tr>
<tr>
<td>Z</td>
<td>Likely</td>
<td>250 mg</td>
<td>1000 mg</td>
<td>Full dose</td>
</tr>
<tr>
<td>E</td>
<td>Likely</td>
<td>100 mg</td>
<td>500 mg</td>
<td>Full dose</td>
</tr>
</tbody>
</table>

**Note:** if the initial reaction to treatment is severe, a weaker trial dose should be used (approximately 1/10th of the dose indicated for trial dose 1).

- **Second-line TB drugs**
  
  Start with the most important drug in a regimen unless there is suspicion that it is the cause of the reaction. Restart each TB drug one after the other, starting at about 1/10 of the dose on Day 1, half-dose on Day 2 and full dose on Day 3.

**Musculoskeletal disorders**

- **Arthralgias**
- **Tendinitis/tendon rupture**
**Arthralgias**

*Z, Rfb, H, Bdq, FQs*

Arthralgias generally diminish over time. Serum uric acid levels are frequently elevated, but this is of little clinical relevance. Anti-hyperuricaemic therapy is of no proven benefit in these patients.

Begin therapy with an anti-inflammatory agent, e.g. **ibuprofen** PO (adult: 400 to 800 mg 3 times daily). **Paracetamol** PO (adult: 500 to 1000 mg 3 times daily) may also help bring relief when given together with an anti-inflammatory drug.

If symptoms fail to resolve, consider lowering the dose of the suspected agent (most often Z), if this does not compromise the effectiveness of TB treatment.

**Tendinitis/tendon rupture**

*FQs*

In the acute phase, the main symptom of tendinitis is pain when moving the affected joint or palpating the tendon. In later phase, continuous pain and tendon thickening or nodularity may be present.

The Achilles tendon is involved in most cases, but other joints may be affected (shoulder, hand, etc.).

New and intense physical activities are not recommended during a treatment with a FQ.

Tendinitis is more common in older patients, patients with renal insufficiency or under corticosteroids.

Tendon rupture is a complication of tendinitis. Signs and symptoms include a snap or pop sound at the time of rupture, bruising, inability to move the joint and a lack of continuity of the tendon on palpation.

Early detection of tendinitis, symptomatic treatment, and discontinuation of FQ can prevent tendon rupture. If the TB treatment is likely to fail without the FQ, try to continue the FQ. Inform the patient that tendon rupture may occur, but that FQ is essential to prevent TB treatment failure.

Symptomatic treatment:
- Rest the joint involved.
- Pain management: application of ice, and **ibuprofen** PO:
  - Adult: 400 to 600 mg every 4 to 6 hours when required, maximum dose: 2400 mg daily.

**Miscellaneous**

- Electrolyte disorders
- Haematologic disorders
- Lactic acidosis
- Nephrotoxicity
- QT prolongation

**Electrolyte disorders**
Aminoglycosides

Electrolyte disorders can occur with the aminoglycosides and are typically reversible with discontinuation of therapy.

Other potential causes (vomiting and diarrhoea) should be treated if present.

If clinical signs of mild to moderate hypokalaemia develop (i.e. muscle cramps, spasms or weakness) or if serum potassium level is between 2.5-3.4 mmol/litre, potassium replacement is required:

**potassium chloride** PO:
Child under 45 kg: 2 mmol/kg (2 ml/kg) daily in divided doses
Child 45 kg and over and adult: 30 mmol (30 ml) 3 times daily

If clinical signs of severe hypokalaemia develop (i.e. marked muscle weakness, cardiac arrhythmias) or if serum potassium level is < 2.5 mmol/litre, hospitalise and urgently administer potassium chloride by slow IV infusion.

For a patient with hypokalaemia:
- Monitor serum potassium levels and QT interval until they return to normal.
- Consider magnesium PO if serum magnesium cannot be measured. Untreated hypomagnesaemia may lead to "resistance" to correction of hypokalaemia. Magnesium should be taken at least 2 hours before or 4 to 6 hours after the FQs.

**Haematologic disorders**

Lzd, R, P, Rfb, E

Most TB drugs can cause hematological disorders that may involve any blood cells (red cells, white cells, platelets). However, the TB drugs most involved are Lzd and rifamycins.

<table>
<thead>
<tr>
<th>Severity grade in adults (a)</th>
<th>Anaemia</th>
<th>Neutropenia</th>
<th>Thrombocytopenia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild</td>
<td>10.5 - 9.5 g/dl</td>
<td>1500 - 1000/mm³</td>
<td>100,000 - 75,000/mm³</td>
</tr>
<tr>
<td>Moderate</td>
<td>&lt; 9.5 - 8.0 g/dl</td>
<td>&lt; 1000 - 750/mm³</td>
<td>&lt; 75,000 - 50,000/mm³</td>
</tr>
<tr>
<td>Severe</td>
<td>&lt; 8.0 - 6.5 g/dl</td>
<td>&lt; 750 - 500/mm³</td>
<td>&lt; 50,000 - 20,000/mm³</td>
</tr>
<tr>
<td>Life-threatening</td>
<td>&lt; 6.5 g/dl</td>
<td>&lt; 500/mm³</td>
<td>&lt; 20,000/mm³</td>
</tr>
</tbody>
</table>

(a) Adapted from NIAID Division of Microbiology and Infectious Diseases, severity scale, Nov-2007.

1) Patient on DS-TB treatment

Rifamycins can cause potentially life-threatening thrombocytopenia. This is more common when used intermittently. Clinical features may include minor haemorrhage (e.g. epistaxis) or severe haemorrhage and thrombocytopenic purpura.

Measure platelets when thrombocytopenia is suspected:
- Moderate thrombocytopenia: stop the rifamycin and monitor platelets weekly until > 75,000/mm³.
- Severe thrombocytopenia: stop all TB drugs. Hospitalise. Treat shock or severe haemorrhage.

In any event rifamycins should not be reintroduced.

2) Patient on DR-TB treatment
Lzd may cause anemia, neutropenia and/or thrombocytopenia.

<table>
<thead>
<tr>
<th>Toxicity</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild to moderate</td>
<td>- In all cases:</td>
</tr>
<tr>
<td></td>
<td>▪ Monitor carefully.</td>
</tr>
<tr>
<td></td>
<td>▪ Consider reduction of dose of Lzd (e.g. 300 mg once daily or 600 mg 3 times weekly in adults).</td>
</tr>
<tr>
<td></td>
<td>▪ For moderate anemia: consider adding erythropoietin (EPO).</td>
</tr>
<tr>
<td></td>
<td>▪ For moderate neutropenia:</td>
</tr>
<tr>
<td></td>
<td>▪ Stop Lzd.</td>
</tr>
<tr>
<td></td>
<td>▪ Restart at reduced dose once toxicity has decreased to ‘mild’.</td>
</tr>
<tr>
<td>Severe</td>
<td>- In all cases:</td>
</tr>
<tr>
<td></td>
<td>▪ Stop Lzd and monitor carefully.</td>
</tr>
<tr>
<td></td>
<td>▪ If Lzd is essential to the regimen, restart at reduced dose once toxicity has decreased to ‘mild’.</td>
</tr>
<tr>
<td></td>
<td>▪ For severe anemia: consider adding EPO.</td>
</tr>
<tr>
<td>Life-threatening</td>
<td>- Stop Lzd and monitor carefully.</td>
</tr>
<tr>
<td></td>
<td>- Hospitalise.</td>
</tr>
<tr>
<td></td>
<td>- Perform blood transfusion</td>
</tr>
<tr>
<td></td>
<td>- If Lzd is essential to the regimen consider restarting at reduced dose once toxicity has decreased to ‘mild’.</td>
</tr>
</tbody>
</table>

### Lactic acidosis

**Lzd**

Lactic acidosis is a rare but potentially life-threatening increase of lactic acid in the bloodstream, that can be due to mitochondrial toxicity of certain TB drugs, usually Lzd. Signs and symptoms include nausea and vomiting, abdominal pain, extreme fatigue, muscle cramps and increased respiratory rate.

If lactic acidosis is suspected, measure blood lactate and pH. Blood lactate ≥ 4 mmol/litre and pH < 7.35 confirm the diagnosis. Stop Lzd and hospitalise for adequate management.

Note that lactic acidosis may also be due to ART (NRTIs).

### Nephrotoxicity

#### Aminoglycosides

Nephrotoxicity is diagnosed by a rise in serum creatinine above baseline. In its early form it is usually asymptomatic, which means it is very important to monitor serum creatinine while on aminoglycosides.

Symptomatic cases may present with decreased urine output, evidence of volume overload (edema, anasarca or shortness of breath) or uremic symptoms such as mental status changes (confusion, somnolence).
Comorbidities such as diabetes or chronic renal failure are not a contra-indication to treatment with aminoglycosides, though caution must be exercised in such circumstances.

- If renal failure occurs:
  - Stop the aminoglycoside.
  - Rule out other causes of renal failure (e.g. diabetes, dehydration, other drugs, congestive heart failure, urinary obstruction, urinary tract infection, prostate hypertrophy).
  - Adjust doses of other TB drugs to creatinine clearance (Appendix 12).
  - Monitor serum creatinine and electrolytes every 1 to 2 weeks until stable.
- If renal function stabilises or improves and if the drug is essential, resume the aminoglycoside adjusted to creatinine clearance (Appendix 12).

**QT prolongation**

*Cfz, Mfx*, Bdq, Mfx, Dlm, Lfx

Some TB drugs may cause QT prolongation and predispose to torsades de pointes, arrhythmias, and sudden death. ECG should be performed before starting TB treatment then monitored throughout the course of treatment in patients taking these drugs. Possible other causes include other QT prolonging drugs (Appendix 19), hypothyroidism and genetic causes such as long QT syndrome.

Mild or moderate QT prolongation (QTcF > 470 in women and > 450 ms in men and ≤ 500 ms) is common. Severe QT prolongation (QTcF > 500 ms or increase > 60 ms from baseline) is relatively rare.

- In all cases:
  - Measure serum electrolytes and correct electrolyte disorders if necessary.
  - Measure thyroid stimulating hormone (TSH) and, if necessary, treat hypothyroidism.
- For mild and moderate QT prolongation: monitor ECG at least weekly.
- For severe QT prolongation: stop QT prolonging drugs, hospitalise, perform continuous ECG monitoring until QT returns to normal. Once the patient is stable (normal QTcF and no electrolyte disorders), critical QT prolonging TB drugs can be reintroduced:
  - Patient on Bdq: consider resuming while suspending all other QT prolonging drugs.
  - Patient on Mfx: use Lfx instead.
  - Patient on Cfz or Dlm: consider stopping if alternatives are available.
  - Patient on QT prolonging non-TB drug: consider stopping it.

**Appendix 17. Air change per hour (ACH) measurement recommendations**

The ACH in a mechanically ventilated room should remain more or less constant, whereas natural ventilation will vary according to:

- Whether the doors/windows/vents in that room are open or not;
- Wind speed and direction;
- Temperature and humidity differential between inside and outside.

The ACH rate is one tool among others to assess if:

- The efficiency of the system in delivering the outdoor air and in removing the pollutants to each location in the room;
- The overall airflow direction is from clean to dirty zones.
To calculate the ACH in a given room:

- Start by drawing a sketch of the room;
- Measure the dimensions of the room and calculate the volume (in m³);
- Measure the surface (in m²) of all the openings/vents in the room and air direction across the openings/vents;
- Measure the air speed (in meters per second) using an anemometer.

\[
ACH = 0.65 \times \text{air speed (m/s)} \times \text{opening area (m}^2\text{)} \times 3600
\]

\[
\text{Room volume (m}^3\text{)}
\]

Summary of proposed specifications:

<table>
<thead>
<tr>
<th></th>
<th>Surface (m²)</th>
<th>Height (m)</th>
<th>ACH</th>
<th>Opening window surface area (m²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single rooms</td>
<td>&gt; 7.5 (2.5 x 3)</td>
<td>&gt; 3</td>
<td>&gt; 12</td>
<td>&gt; 25%</td>
</tr>
<tr>
<td>Wards</td>
<td>4.5 m²/patient</td>
<td>&gt; 3.5</td>
<td>&gt; 12</td>
<td>&gt; 15%</td>
</tr>
<tr>
<td>Waiting rooms (preferably outside)</td>
<td>3 m²/patient</td>
<td>&gt; 3.5</td>
<td>&gt; 12</td>
<td>&gt; 15%</td>
</tr>
<tr>
<td>Sputum collection areas (preferably outside)</td>
<td>&gt; 1.5</td>
<td>&gt; 2.5</td>
<td>&gt; 20</td>
<td>&gt; 50%</td>
</tr>
<tr>
<td>Toilets</td>
<td>&gt; 1.2</td>
<td>&gt; 2.5</td>
<td>&gt; 12</td>
<td>&gt; 25%</td>
</tr>
<tr>
<td>Consultation rooms</td>
<td>&gt; 7.5 (2.5 x 3)</td>
<td>&gt; 3</td>
<td>&gt; 12</td>
<td>&gt; 25%</td>
</tr>
<tr>
<td>Central corridors (avoid in new buildings)</td>
<td>&gt; 2</td>
<td>&gt; 3</td>
<td>&gt; 12</td>
<td>&gt; 25%</td>
</tr>
</tbody>
</table>

There are two main techniques to measure the ventilation. The most commonly used is the anemometer that measures the velocity (speed) of air (see manufacturer’s recommendations for various types of anemometers). The technique using the gas analyser is difficult and should only be used by trained staff.

**Appendix 18. Compassionate use**

**11.1 Definitions**

The term “compassionate use” refers to the use of potentially life-saving experimental treatments to patients suffering from a disease for which no satisfactory authorised therapy exists and/or who cannot enter a clinical trial. For many patients, these treatments represent their last hope.

Experimental treatment is below referred to as investigational new drug (IND).

**11.2 Indications**

Both MDR-TB and XDR-TB can be life-threatening diseases for which approved drugs alone may be ineffective. In some cases, experimental TB drugs, used in combination with approved drugs, could potentially be effective or life-saving.

Compassionate use may be considered for patients presenting with a life-threatening condition (e.g. deteriorating clinical condition due to TB and/or severe immune depression) when:

- Available treatments have failed or are very likely to fail (e.g. regimen comprises less than 3 highly likely effective drugs and/or clinical evolution shows that the treatment is not effective).
Compassionate use might be considered for a single patient or a group of patients presenting similar characteristics.

The use of two INDs would basically follow the same indications and conditions. Possible interactions and overlapping toxicity between the INDs have to be taken into consideration.

11.3 Minimal requirements

Compassionate use should only be considered if conditions for an adequate management of DR-TB patients are in place: optimal treatment regimen; clinical, biological and bacteriological monitoring; adherence support and follow-up. Results of DST by a validated laboratory are critical to decision making.

In addition to the basic components of regular DR-TB case management a specific monitoring might be required for the use of an IND.

It is essential that a reporting system is in place in order to diligently report any adverse events.

11.4 National regulations

In most countries, only drugs for which a marketing authorization has been granted by the national regulatory agency can be used in humans. Some national regulatory agencies have developed mechanisms to facilitate the access to new drugs at different stages of development, but before market approval. In this case, a party can apply for approval of an IND and then seek the proper permission to import the drug to a country. The use of an IND requires permission from the proper national regulatory authorities and/or country ethic boards.

Appendix 18. Advantages and disadvantages of ventilation techniques
Appendix 19. Drug interactions and overlapping toxicities

Update: October 2022

19.1 Interactions between cytochrome P450 inducers/inhibitors and bedaquiline

Drugs interfering with the cytochrome P450 (CYP450) enzyme system should be avoided with bedaquiline.
This list is not exhaustive. Clinicians should be informed of any cytochrome P450 inducers and inhibitors their patients may be taking.

### 19.2 Overlapping toxicity of QT-prolonging drugs

**TB drugs** (mean QT interval prolongation)
- Mild QT prolongation: delamanid (8.6 ms), levofloxacin (4.6 ms),
- Moderate QT prolongation: bedaquiline (12.3 ms), moxifloxacin (12.3 ms),
- Strong QT prolongation: clofazimine (28.5 ms), moxifloxacin high dose (23.14 ms).

**Non-TB drugs**
- Antimalarials: artemisinine derivatives (high risk), quinine
- Antipsychotics: haloperidol (high risk), chlorpromazine, fluphenazine, olanzapine, risperidone
- Cardiac drugs: amiodarone (high risk), beta-blockers, digoxin
- Oral azole antifungals: fluconazole, itraconazole
- Macrolides: azithromycin, clarithromycin, erythromycin
- Anti-nausea drugs: ondansetron
- Antiretrovirals: boosted protease inhibitors, efavirenz

This list is not exhaustive. Clinicians should be informed of any QT-prolonging drugs their patients may be taking.

### 19.3 Interactions between tuberculosis and antiretroviral drugs

AZT: zidovudine; ATV: atazanavir; 3TC: lamivudine; RAL: raltegravir; ABC: abacavir; DTG: dolutegravir; FTC: emtricitabine; TDF: tenofovir disoproxil fumarate; LPV/r: lopinavir/ritonavir; EFV: efavirenz; RTV or r: ritonavir.
R: rifampicin; Rfb: rifabutin; P: rifapentine; Bdq: bedaquiline.

<table>
<thead>
<tr>
<th>Strong CYP450 inducers</th>
<th>Moderate CYP450 inducers</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rifampicin</td>
<td>Efavirenz</td>
<td>Decrease bedaquiline plasma concentrations</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Rifapentine</td>
<td></td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>Rifabutin</td>
<td></td>
</tr>
<tr>
<td>Phenobarbital</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Strong CYP450 inhibitors</th>
<th>Moderate CYP450 inhibitors</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atazanavir</td>
<td>Erythromycin</td>
<td>Increase bedaquiline plasma concentrations</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>Fluconazole</td>
<td></td>
</tr>
<tr>
<td>Clarithromycin</td>
<td>Verapamil</td>
<td></td>
</tr>
<tr>
<td>Lopinavir</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nelfinavir</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ritonavir</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Drugs metabolized by CYP</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emtricitabine</td>
<td>Can increase bedaquiline plasma concentrations</td>
</tr>
</tbody>
</table>
### 19.4 Overlapping toxicities of antiretrovirals and TB drugs

Drugs strongly associated with the listed toxicities appear in bold lettering.

<table>
<thead>
<tr>
<th>TB drugs</th>
<th>NRTI (ABC, 3TC, TDF, AZT)</th>
<th>INI (DTG, RAL)</th>
<th>NNRTI (NVP, EFV)</th>
<th>Boosted PI (LPV/r, ATV/r, DRV/r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R[^8][^7]</td>
<td>All NRTI</td>
<td>DTG</td>
<td>NVP</td>
<td>ATV/r or DRV/r</td>
</tr>
<tr>
<td></td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
<td>• Do not combine.</td>
<td>• Do not combine.</td>
</tr>
<tr>
<td></td>
<td>• No dose adjustment.</td>
<td></td>
<td>• Replace NVP with DTG or EFV.</td>
<td>• Replace R with Rfb.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• If not possible, replace R with Rfb.</td>
<td>• LPV/r</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>EFV</td>
<td>If not possible and a PI is essential, dose adjustment is required[^6].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Can be combined.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• No dose adjustment.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rfb[^7]</td>
<td>All NRTI</td>
<td>All INI</td>
<td>NVP</td>
<td>All boosted PI</td>
</tr>
<tr>
<td></td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
</tr>
<tr>
<td></td>
<td>• No dose adjustment.</td>
<td>• No dose adjustment.</td>
<td>• No dose adjustment.</td>
<td>• No dose adjustment.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Monitor Rfb toxicity.</td>
<td>• Monitor Rfb toxicity.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>EFV</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Do not combine.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P[^6][^7]</td>
<td>All NRTI</td>
<td>All INI</td>
<td>NVP</td>
<td>All boosted PI</td>
</tr>
<tr>
<td></td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
<td>• Do not combine.</td>
<td>• Do not combine.</td>
</tr>
<tr>
<td></td>
<td>• No dose adjustment.</td>
<td>• No dose adjustment.</td>
<td>• Replace NVP with DTG or EFV.</td>
<td>• Reduce the dose of Rfb by half</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>EFV</td>
<td>• Monitor Rfb toxicity.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Can be combined.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• No dose adjustment.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bdq[^7]</td>
<td>[^8]</td>
<td>All INI</td>
<td>NVP</td>
<td>All boosted PI</td>
</tr>
<tr>
<td></td>
<td>• Can be combined.</td>
<td>• Can be combined.</td>
<td>• Do not combine.</td>
<td>• Do not combine.</td>
</tr>
<tr>
<td></td>
<td>• No dose adjustment.</td>
<td>• No dose adjustment.</td>
<td>• Replace NVP with DTG or EFV.</td>
<td>• Replace boosted PI with DTG.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>EFV</td>
<td>• If no alternative, closely monitor ECG.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Do not combine.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Replace EFV with DTG or NVP.</td>
<td></td>
</tr>
</tbody>
</table>

(a) DTG: administer 50 mg 2 times daily, rather than the usual dose of 50 mg once daily.
(b) RAL: e.g. administer 12 mg/kg 2 times daily, rather than the usual dose of 6 mg/kg 2 times daily.
(c) LPV/r:
- Child: increase the dose of RTV to obtain a one-to-one (1:1) LPV/r ratio
- Adult: double the dose (e.g. 800/200 mg 2 times daily, rather than the usual dose of 400/100 mg 2 times daily)

For more information, see University of Liverpool HIV Drug Interaction Checker: [https://www.hiv-druginteractions.org/checker](https://www.hiv-druginteractions.org/checker).
<table>
<thead>
<tr>
<th>Toxicity</th>
<th>ARVs</th>
<th>TB drugs</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal pain</td>
<td>All ARVs</td>
<td>Eto or Pto, PAS, Cfz, Lzd, FQs, H, Z</td>
<td>Common. Often benign, but can be an early symptom of severe adverse effects (<a href="#">Appendix 17</a>).</td>
</tr>
</tbody>
</table>
| Depression               | EFV, DTG                  | Cs or Trd, FQs, Eto or Pto, H                 | • EFV: consider replacing EFV in the event of severe depression.  
• DTG: can cause depression, but less frequently [9].                                                                                                                                                  |
| Diarrhoea                | All PI, DTG               | Eto or Pto, PAS, FQs, Amx/Clv, Ipm/Cln         | Common. Also consider opportunistic infections as a cause of diarrhoea or *Clostridium difficile* infection (pseudomembranous colitis).                                                                 |
| Electrolyte disorders    | TDF (rare)                | Am, S                                         | See Nephrotoxicity.                                                                                                                                                                                      |
| Haematological disorders | AZT                       | Lzd                                           | • Monitor blood count.  
• Replace AZT in the event of bone marrow suppression.  
For Lzd, see Appendix 17.  
• If the patient takes CMX, also consider CMX as a cause of haematological disorders.                                                                                                              |
| Headache                 | AZT, EFV, DTG             | Cs or Trd, Bdq, Dlm                          | Rule out bacterial or cryptococcal meningitis, toxoplasmosis, etc. Headache secondary to AZT, EFV, DTG and Cs or Trd are usually transient.                                                                 |
| Hepatotoxicity           | NVP, EFV, boosted PIs, DTG| Z, H, R, E, PAS, Eto or Pto, Bdq, Amx/Clav    | • If severe, stop ART and TB drugs. When treatment is resumed, start the TB drugs first ([Appendix 17](#)).  
• If the patient takes CMX, also consider CMX as a cause of hepatotoxicity.                                                                                                                          |
| Nausea and vomiting      | RTV, NVP, and most other ARVs | Eto or Pto, PAS, Z, Amx/Clv, Cfz, Lzd, Ipm/Cln, Bdq | Persistent vomiting can be a result of more severe conditions, such as lactic acidosis and/or drug-induced hepatitis.                                                                                     |
| Nephrotoxicity           | TDF                       | Am, S                                         | • Avoid TDF in patients on aminoglycosides.  
• If an aminoglycoside is essential:  
  ▪ For patients already on ART, replace TDF with ABC  
  ▪ For new patients, start with AZT or ABC.  
• If TDF and aminoglycoside cannot be avoided, monitor serum creatinine, creatinine clearance and electrolytes at least every 2 weeks.                                                          |
| Neurotoxicity            | EFV, DTG                  | Cs or Trd, H, Eto or Pto, FQs                 | • EFV: numerous transient effects on the central nervous system during first 2-3 weeks of treatment. If they do not resolve, consider replacing EFV. There is limited data on the use of EFV with Cs or Trd; concomitant use is accepted practice provided the patient is closely monitored for neurotoxicity. |
Appendix 19. Upper room ultraviolet germicidal irradiation (UVGI) system

The use of UVGI in the upper part of rooms may be effective in killing or inactivating *M. tuberculosis* generated by infected persons.

**Referencias**


**Referencias**

1. DTG: can cause insomnia and dizziness. Administer in the morning or consider replacing with EFV, a boosted PI or RAL.

<table>
<thead>
<tr>
<th>QT prolongation</th>
<th>Boosted PIs, EFV</th>
<th>Cfx, Mfx&lt;sup&gt;h&lt;/sup&gt;, Bdq, Mfx, Dlm, Lfx</th>
<th>For monitoring see Appendix 15.</th>
</tr>
</thead>
</table>

| Skin reactions | ABC, NVP, EFV, and all other ARVs | All TB drugs | Do not re-introduce ABC (risk of anaphylaxis).  
Do not re-introduce any agent that may have caused Stevens-Johnson syndrome.  
If the patient takes CMX, also consider CMX as a cause of skin reactions. |
19.1 Mechanism of action

UV lamps are installed into fixtures suspended from a ceiling or mounted on a wall. Fixtures are shielded with louvers or bafflers in order to block radiation below the horizontal plane of the fixtures. UV lights create in the upper portion of the room a germicidal zone where the bacilli are killed (Figure 1). Patients in the lower portion of the room are not exposed to UVGI lights. Good air mixing is needed to transport the air (and thereby the bacilli) to the upper portion of the room. Disinfection is achieved through the rapid dilution of contaminated lower room air with clean irradiated upper room air.

Figure 1

![UV measurements given in microwatts per square centimeter (µW/cm²)](image)

Distance from the floor (feet)

Distance from the back of fixture (feet)

*1 foot = 0.3048 m

From the WHO, Implementing the WHO Policy on TB Infection Control in Health-Care Facilities, Congregate Settings and Households

The lamps should irradiate the entire surface of the upper part of the room (Figure 2), in order to disinfect the largest possible volume of air mixed at a low speed between the upper and lower part of the room.

UVGI Upper-room Irradiation

Figure 2

![UVGI fixture](image)

UVGI FIXTURE

AIR FLOW (NATURAL CONVECTION)

From Guidelines for the Utilization of Ultraviolet Germicidal Irradiation technology in controlling transmission of tuberculosis in health care facilities in South Africa

Several factors influence the efficiency of UVGI systems:

- Ventilation rate: in controlled environment, at rates up to 6 air change per hour (ACH), UVGI systems increase the effect of air cleaning to > 12 ACH. But when ventilation rates are increased above 6 ACH, UVGI system effectiveness could be reduced because the time for bacteria irradiation is shorter.

- Effective mixing within the room may be provided by natural convection currents or fans, preferably, ceiling ones. Low velocity ceiling fans boosted UVGI system's effectiveness up to 33% when ACH was below 6.
• Relative humidity: studies\(^8\)\(^9\)\(^10\) have reported rapidly decreasing air cleaning effectiveness in UVGI systems when the relative humidity goes above 70%.

• Installation: the height of the room should be minimum 2.5 m and UVGI fixtures should be installed at the minimum height of 2.1 m. As a thumb rule, a 30W lamp should be sufficient for 18 m\(^2\) of surface\(^11\)\(^12\), but room shape and type of fixture should be taken into consideration when calculating the needs. For instance, wall-mounted lamps would have a smaller germicidal area than ceiling-mounted ones. Lamps should be on whenever there is a risk of TB transmission. For example, in rooms with hospitalized patients, the lamps should be turned on 24 hours a day.

• Maintenance: see below.

### 19.2 Maintenance

Dust-covered and/or old UVGI lamps are less effective, hence the need for a careful maintenance, including regular cleaning:

- Lamps and fixture surfaces should be wiped at least monthly (more often if necessary) with a cloth dampened with 70% alcohol. Do not use water and soap or any detergent. The cleaning should be performed when lamps and fixtures are cool.

- Measurement of UVGI level must be done at installation and at least once a year. A UV light meter programmed to detect UV light on a wavelength of 254 nm is needed. Measurements should be performed at eye level in the occupied zone (~ 1.6 m) and in upper irradiated portion of the room, at a distance of 1.2 m from the fixture in all possible directions (imitating a circle with measurements done while moving in circumference spaced of 1 m). Ideally, all upper room measurements should be around 30 μW/cm\(^2\) to 50 μW/cm\(^2\). Persons doing these measurements should wear protective equipment (UV protective glasses, clothing made of tightly woven fabric, soft cotton gloves) and cover exposed skin with opaque creams with solar-protection factors > 15.

- UV lamps last between 5 000 and 10 000 hours of continue use (7 to 14 months). Check manufacturer's information. After this period, UV lamps rapidly lose effectiveness and need to be changed.

### 19.3 Disposal

UV lamps contain mercury and quartz and are considered as hazardous waste. Disposal is extremely difficult in many countries; this should be considered before implementing them. If adequate disposal of the lamps by specialized enterprises is not possible in the country, neither their repatriation; UV lamps should be disposed of by encapsulation (sealed in a metal 200 litre drum filled with concrete and then buried away from water sources).

**Safety considerations**

Reflecting surfaces in the irradiation area of UV lamps must be avoided (i.e. oil painted ceilings, etc.).

At certain wavelengths (including UV-C) UV exposure may be harmful. Skin exposure can produce sunburn (erythema). Exposure of the eyes can produce conjunctivitis (feeling of sand in the eyes, tearing) and/or keratitis (intense pain, sensitivity to light). These symptoms typically commence 6 to 12 hours after exposure.

Despite the fact that these are reversible conditions, health care workers should immediately report them to the IC officer. This could mean that UV irradiation is higher than previously thought at lower room level (lamp poorly positioned? Reflecting surface?).

The USA National Institute for Occupational Safety and Health (NIOSH) states that safe exposure limits are set below those found to initiate eye irritation, the body surface most susceptible to UV. Next table shows the permissible exposure times for given effective irradiances at 254 nm wavelength.
Exposures exceeding this limit would require the use of personal protection equipment to protect the skin and eyes.

In order to avoid overexposure of UVGI, education of health care workers should include basic information on UVGI systems and their potential harmful effects if overexposure occurs.

<table>
<thead>
<tr>
<th>Permissible exposure time&lt;sup&gt;(a)&lt;/sup&gt;</th>
<th>Effective irradiance&lt;sup&gt;(µW/cm²)&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Units given)</td>
<td>(Seconds)</td>
</tr>
<tr>
<td>8 h</td>
<td>28,800</td>
</tr>
<tr>
<td>4 h</td>
<td>14,400</td>
</tr>
<tr>
<td>2 h</td>
<td>7,200</td>
</tr>
<tr>
<td>1 h</td>
<td>3,600</td>
</tr>
<tr>
<td>30 min</td>
<td>1,800</td>
</tr>
<tr>
<td>15 min</td>
<td>900</td>
</tr>
<tr>
<td>10 min</td>
<td>600</td>
</tr>
<tr>
<td>5 min</td>
<td>300</td>
</tr>
<tr>
<td>1 min</td>
<td>60</td>
</tr>
<tr>
<td>30 s</td>
<td>30</td>
</tr>
<tr>
<td>10 s</td>
<td>10</td>
</tr>
<tr>
<td>1 s</td>
<td>1</td>
</tr>
<tr>
<td>0.5 s</td>
<td>0.5</td>
</tr>
<tr>
<td>0.1 s</td>
<td>0.1</td>
</tr>
</tbody>
</table>

(a) The occupational exposure limit for UV-C at 254 nm is 6,000 µJ/cm². This can be also calculated with the following formula: Dose (in µJ/cm²) = Time (in seconds) * Irradiance (in µW/cm²).

Exposures exceeding this limit would require the use of personal protection equipment to protect the skin and eyes.

In order to avoid overexposure of UVGI, education of health care workers should include basic information on UVGI systems and their potential harmful effects if overexposure occurs.

**Referencias**


Appendix 20. Treatment supporters

Update: January 2022

Treatment supporters need specific training to know and understand their role in order to provide the patient with adequate treatment education and support. They should be compensated for their time and services and reimbursed for expenses incurred.

20.1 Selecting a treatment supporter

The treatment supporter:\n- Is someone from the patient’s community;
- Is preferably a community health worker or a person with a background in health (e.g. pharmacist), but can also be a non health worker (co-worker or neighbour);
- Is chosen by, or is acceptable to, the patient and their family (e.g. supporter and patient of the same sex);
- Is able to observe the patient’s confidentiality;
- Has a stable living situation;
- Has basic literacy skills (can read and write and has basic numeracy skills);
- Is motivated to care for TB patients and committed to supporting them for the full duration of treatment;
- Lives near enough to the patient to be able to make regular visits (daily or weekly) and go to their home immediately in the event of an emergency;
- Is in good physical condition and not immunedepressed.\n
It is usually not recommended to have family members as treatment supporters. The family relationship may interfere with the ability to administer TB treatment, especially if the patient is a child.

20.2 Roles and responsibilities

Role and responsibilities of a treatment supporter may include:
- Supervision of all drug intakes and keeping records on TB treatment card.
- Detection of adverse effects and, when necessary, prompt referral of the patient to a health facility.
- Accompanying the patient to medical consultations.
- Collection and transport of sputum specimens for smear and culture.
- Provision of health education to family members, including the risk of transmission and implementation of infection control measures in the home.
Appendix 21. Informing the patient

21.1 At the start of treatment

Arrange two interviews (allow about 20 minutes for each): one to supply the patients with the information they need to follow the treatment, the second to make sure they have assimilated the information. These interviews should coincide with the first two clinical visits. The first interview should occur before the treatment begins. Depending on how the clinic is organized, the interviews are done either by the prescribing clinician alone at the time of the clinical visit, or with the help of a specially-trained staff member at an interview just for this purpose. Patients may bring someone with them, if they wish.

Outpatients

First interview

- Explain:
  - The disease and how it spreads
    For example: this is a serious, but generally curable, infection that affects the lungs and can be spread if not treated (tailor the information according to the focus of the infection, the resistance pattern).
  - The treatment process:
    Length, intensive/continuation phases, clinical and bacteriological monitoring, visit schedule (tailor the information according to the regimen); how DOT will work and why it is important when relevant.
  - The medications:
    - Management:
      Where, when, and from whom to get medications;
      Number of tablets per day; number of doses per day, etc.;
      Keep tablets in their blister pack until taken, no removing them from their package ahead of time.
    - Main adverse effects and what to do if they occur:
      For example: for rifampicin, point out that it turns the urine, stools, tear, etc. reddish-orange, that this is normal and not a cause for concern. For ethambutol, advise the patient to consult the doctor immediately if s/he notices a decrease in his/her vision or ability to correctly distinguish colours, etc.
    - Special precautions (depending on concomitant treatment):
      For example: take rifampicin in the morning, and fluconazole at night.
  - Any incentives or enablers the patient may qualify for and how the patient can access them.
- Stress the importance of adherence, anticipate problems, and think about possible solutions.
- Answer any questions.
- Give the date of the second interview (one week later).

Second interview (one week later)

- Check to make sure that information has been assimilated; ask open-ended questions, give the patient time to answer. Give more information, if necessary.
- Answer any questions.
- Remind the patient of the date of the next visit.

Notas


(b) The most common cause of immunosuppression is HIV infection, but chronic illnesses such as diabetes also alter the immune system and are a risk factor for TB infection and active TB.
Hospitalized patients

First interview

Same as above, plus explain:

- Hospital infection control measures:
- Isolation and why it is indicated; the importance of covering the mouth when coughing or sneezing, the use of sputum containers, visits outside the building, face masks/respirators (who, when, why), airing out the room, etc.
- Timetable for injections and distribution of drugs.

Second interview (when patient is ready for discharge)

- Explain:
  - Where and when to get medications, the visit schedule;
  - DOT and other treatment support as relevant.
- Make sure that the information the patient needs to continue treatment as an outpatient has been assimilated (treatment process, medications, adverse effects and what to do, etc.).
- Stress the importance of adherence, anticipate problems, and think about possible solutions.
- Answer any questions.

21.2 In the course of treatment

Adherence interviews should take place at least monthly (more frequently if needed) throughout the entire course of treatment. Their purpose is to identify/resolve any problems resulting in poor adherence. Assessment is done either by the clinician at the monthly clinical visit, or by the nurse responsible for individual distribution of drugs.
Adherence interviews should be quick (about 5 minutes); on the other hand, devote as much time as necessary to resolving any problems.
The interview at the end of the intensive phase is more specifically devoted to informing the patient, because of the changes in drug regimen for the continuation phase.

Appendix 23. Treatment card for patients on first-line anti-TB therapy

[Download PDF]

Appendix 24. Tuberculosis register for patients on first-line anti-TB therapy

[Download PDF]
Appendix 27. Respirators

Update: January 2022

27.1 Introduction

Respirators are masks designed to protect the wearer from inhaling bacilli.

They must be worn by all staff in areas where the risk of TB transmission is high:
- Inpatient department with smear positive or drug-resistant TB patients
- Consultation room for TB diagnosis
- Laboratory (sputum smear preparation and culture/drug susceptibility testing)
- Sputum collection area
- Radiology department
- Waiting areas in high TB prevalence settings

Visitors and attendants must wear a respirator when entering a contagious TB patient’s room.

Recommended respirators include:
- The CE-certified filtering facepiece EN 149 FFP2, filtering efficiency 94% if challenged with 0.4μm particles;
  or
- The United States Centre for Disease Control and Prevention/National Institute for Occupational Safety and Health
  (NIOSH) certified N95, filtering efficiency > 95% if challenged with 0.3μm particles.

27.2 Instructions for use

Respirators are for personal use. The same respirator cannot be shared between staff members or between caregivers.

The respirator should be put on before entering the room and removed after exiting the room.

Respirators must be worn covering the nose, mouth and chin and provide a tight seal around the edge. Every time that a respirator is put on, a seal check has to be performed:
- Fully open the respirator and slightly bend the nose wire to form a curve.
Different factors may not allow proper sealing of respirators to the face: respirator size and/or model; respirator wearer's facial features, including beard and facial hair; headscarves, etc.

There is limited evidence on the acceptable length of time a respirator can be worn with maintained efficiency. The filter materials remain functional for weeks or months, but with frequent wearing the respirator will become less adjusted.

An extensively used respirator should be discarded after 7 days. However, if for example, it is only used a few hours 2 to 3 times a week, it can be reused for several weeks. During this period, staff can reuse their respirator provided it is not wet or damaged and its straps are not loosened. Each staff member should keep their respirator in the pocket of their personal gown without creasing it. If the filter material is damaged or the mask has loose straps, the respirator should be discarded immediately.

**Note:** TB bacillus is trapped in the filter of a mask and will not be released with shaking or other physical movements of the mask.

### 27.3 Storage

Store in a dry, well ventilated place. Respirators should not be crushed during storage.

### 27.4 Disposal

Respirators are disposed of as “soft waste” and do not need to be disinfected before being discarded.

### 27.5 Fit testing

Proper fit of a respirator is critical to ensure respiratory protection. Therefore, all staff members who could be exposed to *M. tuberculosis* should before being required to wear a respirator perform a “fit testing” to determine if the respirators being used fit them properly.

At least two models of respirators should be available. If a worker cannot be fitted with one model, the other one should be used.

Testing is performed using a fit testing kit. The kit contains all the supplies and instructions needed to perform the test.

**Fit testing kit**
Appendix 27. Request form for microscopy and Xpert MTB/RIF

Appendix 28. Surgical masks

Update: January 2022

28.1 Introduction

The purpose of surgical masks is to catch droplet nuclei that patients expel while talking, breathing or coughing. Surgical masks should be worn by contagious or potentially contagious patients (confirmed or presumed cases) when they leave their rooms to go to another department or any other enclosed area. Wearing a surgical mask is not necessary when patients are alone in their room or outdoors.

The terms “surgical”, “medical” or “procedure” are sometimes used interchangeably to qualify masks. Only masks that conform to the norms EN 14683 or ASTM F2100 should be used.

28.2 Instructions for use

Surgical masks are for personal use. The same mask cannot be shared.

- Open the mask.
- Bend the nasal bar (if included).
- Put the chin into the mask.
- Attach the two straps behind the head or over the ears.

Surgical masks must be replaced at least once a day and when they become wet or damaged.

It is not recommended to wear masks for large portions of the day or while sleeping, as they restrict air movement and are not comfortable.

28.3 Storage

Store in a dry, well ventilated place.

28.4 Disposal

Masks are disposed of as “soft waste” and do not need to be disinfected before being discarded.
Appendix 29. BCG vaccine

Update: January 2022

Composition, forms and route of administration

- Live attenuated bacterial vaccine
- Powder for injection, to be dissolved with the entire vial of the specific solvent supplied by the manufacturer, in multidose vial, for intradermal injection

Dosage and vaccination schedule

Refer to national recommendations. In countries with a high incidence of TB (> 40 cases per 100,000), WHO recommends[1]:
- Child under 12 months: 0.05 ml single dose as soon as possible after birth
- Child 12 months and over and adult: 0.1 ml single dose

Technique and site of administration

- Clean the injection site with clean water. Do not use antiseptics as risk of inactivation of vaccine). Allow to dry.
- Administer intradermally. If the injection is correctly performed, an “orange-skin” papule measuring 5-8 mm in diameter should appear at the injection site.
- The vaccine is administered in the deltoid region of the arm, about one-third down the upper arm over the insertion of the deltoid muscle.
- The vaccine should be injected in the same place for each child so that the BCG scar is easier to locate.

Contra-indications

- Do not administer to patients with congenital or acquired immunodeficiency (e.g. HIV infection or serologic status unknown, but symptoms consistent with HIV infection, immunosuppressive therapy, malignant haemopathy).
- Postpone vaccination until recovery in the event of acute extensive dermatosis, acute complicated malnutrition or severe acute febrile illness (minor infections are not contra-indications).

Adverse effects

- Local reaction 2-4 weeks after injection: papule that ends up as an ulcer and usually heals spontaneously (dry dressing only) after 2 to 5 months, leaving a permanent
- Complications requiring no specific treatment and which almost always evolve favourably:
  - persistent ulcer with serous discharge for over 4 months after injection;
  - non-suppurated adenitis, most often axillary, sometimes cervical;
  - abscess at the injection site due to infection (red, hot and painful abscess) or inadvertent intradermal injection (cold and painless abscess).
- Uncommon complications:
- suppurative lymphadenitis, mostly observed in neonates, usually due to inadvertent intradermal injection. The lymph node, which can have a diameter of over 3 cm, evolves toward softening and fistulisation with chronic osteomyelitis in exceptional cases.
- disseminated BCG disease\(^b\), most commonly in immunocompromised children under 2 years old (mortality rate > 70\%)\(^2\).

### Precautions

- If administered simultaneously with other vaccines, use different syringes and injection sites. Do not mix with other vaccines in the same syringe.
- **Pregnancy:** CONTRA-INDICATED
- **Breastfeeding:** no contra-indication

### Storage

- Reconstituted vaccine: between 2 °C and 8 °C for 6 hours max.
- Powder: between 2 °C and 8 °C.
- Solvent: a cold chain is not required for However, at least 24 hours before reconstitution of the vaccine, the solvent must be refrigerated between 2 °C and 8 °C so that the solvent and lyophilised powder are at the same temperature: a temperature difference during reconstitution may reduce vaccine efficacy. Do not freeze.

### Notas

(a) BCG vaccine provides high protection for neonates, but only moderate for school age TST negative children.

(b) If disseminated BCG disease is diagnosed, a 6-month TB treatment should be administered.

### Referencias


### Appendix 29. Sputum smear microscopy register

[Appendix 29. Sputum smear microscopy register](#)

### Appendix 30. Xpert MTB/RIF register

[Appendix 30. Xpert MTB/RIF register](#)
Appendix 31. Drug-o-gram

Appendix 32. Quaterly report

Appendix 33. Report on detection and enrolment of TB cases with rifampicin and multidrug-resistance

Appendix 34. Report of final outcomes of drug-resistant tuberculosis

Appendix 35. Check-list for the evaluation of a TB service