Multidrug and extensively drug-resistant tuberculosis from a general practice perspective

BM Yashodhara 1
Choo Beng Huat 1
Lakshmi Nagappa Naik 1
Shashikiran Umakanth 2
Manjunatha Hande 3
Joseph M Pappachan 3

1Department of Medicine, Melaka Manipal Medical College, Melaka, India; 2Department of Medicine, Kasturaba Medical College, Manipal; 3Department of Medicine, Kottayam Medical College, Kerala, India

Abstract: Despite intensive efforts to eradicate the disease, tuberculosis continues to be a major threat to Indian society, with an estimated prevalence of 3.45 million cases in 2006. Emergence of multidrug-resistant tuberculosis has complicated eradication attempts in recent years. Incomplete and inadequate treatment are the main causes for development of drug resistance. Directly observed therapy, short-course (DOTS) is the World Health Organization (WHO) strategy for worldwide eradication of tuberculosis, and our country achieved 100% coverage for DOTS through the Revised National Tuberculosis Control Program in 2006. For patients with multidrug-resistant tuberculosis, the WHO recommends a DOTS-Plus treatment strategy. Early detection and prompt treatment of multidrug-resistant tuberculosis is crucial to avoid spread of the disease and also because of the chances of development of potentially incurable extensively drug-resistant tuberculosis in these cases. This review discusses the epidemiologic, diagnostic, and therapeutic aspects of multidrug-resistant tuberculosis, and also outlines the role of primary care doctors in the management of this dangerous disease.

Keywords: multidrug-resistant, extensively drug-resistant, tuberculosis, general practice

Introduction

Tuberculosis is among the top three causes of death from a single infectious agent, along with malaria and human immunodeficiency virus (HIV). One-third of the world’s population is infected with Mycobacterium tuberculosis, the etiologic agent of tuberculosis. Of the 9.2 million tuberculosis cases reported worldwide in 2006, 0.7 million were HIV-positive and 1.7 million cases were fatal, of which 0.2 million occurred in HIV-positive people. India continues to occupy a unique position with respect to the global tuberculosis burden. In 2006, of the 9.2 million new cases of tuberculosis reported globally, 1.93 million were reported in India, which constitutes approximately one-fifth of the total global burden. This included 0.87 million smear-positive cases of the total estimated prevalence and incidence of tuberculosis of 3.45 million and 1.93 million, respectively, in India for 2006.

Pulmonary tuberculosis is a highly contagious and life-threatening infection, and the mode of spread is droplet infection. The usual source of infection is a human case whose sputum is positive for tuberculous bacilli and who has received no treatment or has not been treated fully. The World Health Organization (WHO) classifies tuberculosis infection into four categories for the purpose of treating cases effectively and in a uniform manner worldwide. In recent years, emergence of multidrug-resistant tuberculosis has become a major issue in the successful management and eradication of the disease.
Multidrug-resistant tuberculosis has now reached an approximate prevalence rate of 5% of the total global tuberculosis burden,1 amounting to half a million cases, of which most cases are from Indian, China, and Russia.1,5 Although the measures adopted by various countries have resulted in a 51% case detection rate for tuberculosis and a 70% successful treatment rate worldwide, there is still a long way to go to achieve a 70% case detection rate and 85% successful treatment rate for nonresistant tuberculosis, as suggested by the WHO.5 Also, the WHO has suggested 70% case detection and treatment rates for multidrug-resistant tuberculosis to curb the development of extensively drug-resistant tuberculosis.4 If an adequate drug regimen is followed, tuberculosis is considered to be 100% curable, as proven to some extent by reduction in the case fatality rate from 29% to 4%, under the Revised National Tuberculosis Control Program (RNTCP) in India which adopted directly observed therapy, short-course (DOTS) in 1993,7 but tuberculosis still has a high mortality rate, as evidenced by nearly two million deaths every year worldwide.8 To reduce this burden further, India is planning to implement DOTS-Plus by 2010.7 The incidence of extensively drug-resistant tuberculosis is about 6%–7.3% in patients treated for multidrug-resistant tuberculosis worldwide.9,10 Table 1 shows the WHO categories of tuberculosis treatment.

Definitions and epidemiology
Multidrug resistance is defined as resistance of *M. tuberculosis* to two or more first-line antituberculous drugs that include isoniazid and rifampicin.11 Extensively drug-resistant tuberculosis refers to cases of multidrug-resistant tuberculosis which show resistance to second-line antituberculous drugs including any one of the fluoroquinolones and one of the three injectable drugs (aminoglycosid, kanamycin, or capreomycin).12 The most recent estimates suggest that globally there were about 489,000 cases of multidrug-resistant tuberculosis in 2006, of which 110,132 cases were in India.2 This is about 4.9% of the total number of tuberculosis cases in the country. About 0.9%–1.5% of multidrug-resistant cases in India may fall into the category of extensively drug-resistant tuberculosis.12

Multidrug resistance can be primary or acquired. Primary resistance is defined as resistance in patients without a history or other evidence of previous antituberculous drug treatment. Acquired drug resistance occurs in those who have previously received antituberculous treatment for at least one month and in those with treatment failures and relapses. The prevalence of primary multidrug resistance is 3% and that of acquired multidrug resistance is 12% in India. Although drug resistance started appearing in patients with *M. tuberculosis* infection soon after the introduction of effective antituberculous drugs, multidrug-resistant tuberculosis was not a major public health problem until the early 1990s13 when HIV infection became a global epidemic.

Causes of multidrug-resistant tuberculosis
A history of tuberculosis and previous antituberculous treatment are the most important factors identified in the causation of multidrug-resistant tuberculosis.14 Factors which predispose a patient to development of multidrug-resistant tuberculosis include:

- Incomplete treatment
- Inadequate treatment
- Errors in tuberculosis management, such as use of a single antituberculous drug
- Addition of a single drug to a failing regimen
- Failure to identify pre-existing resistance
- Initiation of an inadequate primary regimen
- Failure to identify and address nonadherence to treatment
- Noncompliance
- Inappropriate isoniazid preventive therapy
- Variations in bioavailability of antituberculous drugs.15

There are various other factors favoring the spread of tuberculosis, which include increasing high-risk groups (eg, the elderly, prisoners, immigrants, drug addicts6), recurrent defaulters, multidrug-resistant and extensively drug-resistant treatment failures,1 increasing age of the general population, increasing number of patients with immunosuppression due to bone marrow and solid organ transplantation,16 and lack of optimal recommended duration of treatment for multidrug-resistant and extensively drug-resistant tuberculosis in controlled clinical trials.3 In addition, certain host genetic factors and coexistent HIV infection may also predispose to the development of multidrug-resistant tuberculosis.11,17

<table>
<thead>
<tr>
<th>Category of treatment</th>
<th>Patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category I</td>
<td>New sputum smear-positive</td>
</tr>
<tr>
<td></td>
<td>Seriously ill, sputum smear-negative</td>
</tr>
<tr>
<td></td>
<td>Seriously ill, extrapulmonary</td>
</tr>
<tr>
<td>Category II</td>
<td>Sputum smear-positive, relapse</td>
</tr>
<tr>
<td></td>
<td>Sputum smear-positive, failure</td>
</tr>
<tr>
<td></td>
<td>Sputum smear-positive treatment after default</td>
</tr>
<tr>
<td>Category III</td>
<td>Sputum smear-negative, not seriously ill</td>
</tr>
<tr>
<td></td>
<td>Extrapulmonary, not seriously ill</td>
</tr>
<tr>
<td>Category IV</td>
<td>Multidrug-resistant tuberculosis</td>
</tr>
</tbody>
</table>
Diagnosis of drug resistance

Drug-resistant tuberculosis needs to be suspected based on history of prior treatment (smear-positive cases after repeated treatment courses and category II treatment failures), and such cases should undergo drug susceptibility testing. Lowenstein-Jensen culture is the method traditionally used for drug sensitivity testing in tuberculosis. A duration of 6–8 weeks is required before sensitivity results are known with this method. Several new methods are now available for testing drug sensitivity patterns, but they are costlier and are not widely available. The methods for diagnosis of multidrug-resistant tuberculosis include:

- Conventional (Lowenstein-Jensen) culture, ie, proportional, resistance ratio, absolute concentration
- BACTEC-460
- Mycobacterial growth indicator tube
- E test
- Mycobacteriophage method
- Molecular methods, ie, polymerase chain reaction, ligase chain reaction, restriction fragment length polymorphism.

Identification and proper management

Drug marketing surveys done in India for 2006 have shown that 75% of the total cost for antituberculous drugs was spent outside of the RNTCP, mainly in the private sector. The vast majority of multidrug-resistant tuberculosis cases might have been managed in the private sector in recent years. Improper supervision of treatment in the private sector might lead to treatment defaulting and hence incomplete management of cases. This may result in development of extensively drug-resistant tuberculosis, which is a major public health challenge.

Multidrug-resistant tuberculosis cases are difficult to manage in view of the need for prolonged treatment (up to 24 months), high rates of treatment failures, higher costs and potential toxicities of medications, need for surgery in locally confined untreatable cases, and risk of transmission to unexposed individuals, including health care workers. Cases of extensively drug-resistant tuberculosis may be incurable, making the situation worse. A comparison of multidrug-resistant and extensively drug-resistant tuberculosis is made in Table 2.

Treatment

Tuberculosis is a disease with high cure rates if managed properly, and an effective national control program can reduce the risk of transmission to the unexposed public.

The Indian tuberculosis control program is now one of the largest public health programs in the world. The RNTCP was established under WHO guidance in 1993. The strategy of the RNTCP for control of tuberculosis is DOTS, which is familiar to most clinicians now. With extensive efforts on the part of government and health departments, India achieved a DOTS coverage of 100% in 2006. The reported cure rate with DOTS treatment was 86% in 2005.

Management of multidrug-resistant tuberculosis is a challenge for clinicians. Adherence to appropriate and prolonged drug therapy is the cornerstone of treatment success in multidrug-resistant tuberculosis. Hence, the standard DOTS regimen may not be helpful for treating these cases.

In low- and middle-income countries that have adopted the DOTS strategy, the WHO and its international partners have emphasized DOTS-Plus for multidrug resistant tuberculosis programs since 1998, in order to promote effective treatment of multidrug-resistant tuberculosis.

A comparison between DOTS and DOTS-Plus is shown in Table 3. The major differences between DOTS and DOTS-Plus are that DOTS employs sputum microscopy for diagnosis of tuberculosis cases and multidrug-resistant tuberculosis cases, and uses first-line antituberculosis drugs over a short period of treatment, while DOTS-Plus uses sputum culture and drug sensitivity testing for diagnosis of multidrug-resistant tuberculosis cases and uses both first- and second-line antituberculosis drugs over a longer period of time, ie, up to 24 months.

Only experienced clinicians at centers equipped with reliable laboratory services for mycobacterial culture and in vitro sensitivity testing should treat multidrug-resistant
Table 3 Comparison of DOTS and DOTS-Plus\textsuperscript{11}

<table>
<thead>
<tr>
<th></th>
<th>DOTS</th>
<th>DOTS-Plus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Political and administrative</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>commitment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diagnosis by Sputum microscopy</td>
<td>Sputum</td>
<td>Sputum</td>
</tr>
<tr>
<td></td>
<td>microscopy</td>
<td>microscopy</td>
</tr>
<tr>
<td></td>
<td>Sputum</td>
<td>culture</td>
</tr>
<tr>
<td></td>
<td>and drug</td>
<td>susceptibility</td>
</tr>
<tr>
<td></td>
<td>susceptibility</td>
<td></td>
</tr>
<tr>
<td>Drugs used</td>
<td>First-line</td>
<td>First-line</td>
</tr>
<tr>
<td></td>
<td>antituberculous</td>
<td>and second-line</td>
</tr>
<tr>
<td>Duration of intensive phase</td>
<td>Short course</td>
<td>18–24 months</td>
</tr>
<tr>
<td>of treatment</td>
<td>2–3 months</td>
<td>Six months</td>
</tr>
<tr>
<td>Individualized treatment</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Directly observed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Purpose of therapy</td>
<td>Cure tuberculosis</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>and prevent</td>
<td>Treat multidrug</td>
</tr>
<tr>
<td></td>
<td>multidrug</td>
<td>resistance</td>
</tr>
<tr>
<td>Monitoring of treatment</td>
<td>Monitoring</td>
<td>Monitoring,</td>
</tr>
<tr>
<td></td>
<td>and evaluation</td>
<td>reporting,</td>
</tr>
<tr>
<td></td>
<td>of treatment</td>
<td>evaluation</td>
</tr>
<tr>
<td></td>
<td>outcome</td>
<td>of treatment</td>
</tr>
</tbody>
</table>

Two newer classes of drugs targeting mycobacterial cell wall synthesis have been discovered recently, ie, benzothiazinones and dinitrobenzamide, which are highly active against \textit{M. tuberculosis}, including extensively drug-resistant tuberculosis strains.\textsuperscript{31}

Future inventions include drugs with activity on different bacterial cell wall components, virulence genes, small guanosine triphosphatases etc.,\textsuperscript{32} and inhibitors of the aspartate kinases involved in bacterial cell wall synthesis.\textsuperscript{33} Mutations at hotspots are responsible for the development of extensively drug-resistant tuberculosis strains, as observed in a study involving 194 \textit{M. tuberculosis} strains isolated from patients.\textsuperscript{34} Early detection of these extensively drug-resistant cases could be done using a rapid test called MTBDRsI, which detects resistance to ethambutol, fluoroquinolones, second-line aminoglycosides (amikacin and kanamycin), and cyclic peptide (capreomycin), and this could be confirmed by drug susceptibility testing.\textsuperscript{35}

The principles of treatment (and poor prognostic indicators) for multidrug-resistant tuberculosis include:

- Use of as many first-line agents to which the organism is sensitive
- At least one parenteral drug should be part of the regimen
- At least three agents to which the isolate is sensitive should be used
- Administration of oral agents should be directly observed
- Bactericidal agents should be used whenever possible
- Single drugs should never be added to a failing regimen.\textsuperscript{36}

Poor prognostic indicators are:

- Category II failures given intermittent therapy
- Cavity at presentation
- HIV positivity

Other factors that predict progression to extensively drug-resistant tuberculosis in multidrug-resistant patients are bilateral cavitating lesions, prior exposure to a second-line tuberculosis. It has also been observed that this vaccine exerted therapeutic efficacy (100% survival in Cynomolgus monkeys infected with tuberculosis) and showed a synergistic effect after an initial priming Bacillus Calmette-Guérin dose. The implications of this study are that this vaccine might be useful against \textit{M. tuberculosis}, including multidrug-resistant and extensively drug-resistant tuberculosis in humans participating in therapeutic clinical trials.\textsuperscript{29} Other agents shown to be effective in animal models include a dry powder PA-824 cyclodextrin-lecithin aerosol suspension.\textsuperscript{30}

We need novel drugs and vaccines to be able to face the challenges in the treatment of multidrug-resistant and extensively drug-resistant tuberculosis. Some of the new drugs and vaccines having the potential to cure multidrug-resistant and extensively drug-resistant tuberculosis are linezolid,\textsuperscript{16,22,23} thoridizaine (a neuroleptic phenothiazine),\textsuperscript{24–26} myrtle leaf extracts (used as an antiseptic in Sardinian traditional medicine),\textsuperscript{27} medicinal plants having antituberculous activity (Acalypha indica, Adhatoda vasica, Allium cepa, Allium sativum, Aloe vera\textsuperscript{28}), and HSP65-IL-12/HVJ vaccine.\textsuperscript{29} HSP65-IL-12/HVJ is a combination of DNA vaccines expressing mycobacterial heat-shock protein 65 and interleukin-12, delivered by the hemagglutinating virus of Japan envelope and liposome.\textsuperscript{29} The HSP65-IL-12/HVJ vaccine has been shown to have protective and therapeutic efficacy in murine models and in Cynomolgus monkeys (which are considered to be the best animal model of human tuberculosis). It has also been observed that this vaccine exerted therapeutic efficacy (100% survival in Cynomolgus monkeys infected with tuberculosis) and showed a synergistic effect after an initial priming Bacillus Calmette-Guérin dose. The implications of this study are that this vaccine might be useful against \textit{M. tuberculosis}, including multidrug-resistant and extensively drug-resistant tuberculosis in humans participating in therapeutic clinical trials.\textsuperscript{29} Other agents shown to be effective in animal models include a dry powder PA-824 cyclodextrin-lecithin aerosol suspension.\textsuperscript{30}
injectable antibiotic, and each additional month in which a patient has failed to take at least 80% of their prescribed drugs. In a recent study, the success rate for treatment was 87% for multidrug-resistant but not extensively drug-resistant tuberculosis, and 41% for extensively drug-resistant cases. In this study, resistance to ciprofloxacin, ofloxacin, and amikacin is associated with an unsuccessful outcome. However, DOTS-Plus, which has been found to be an effective strategy for treatment of multidrug-resistant tuberculosis showed the best cure rates at 70%. Monitoring with serial cultures should be done at months 4, 6, 12, 18, and 24 of treatment. Treatment for 18–24 months is mandatory in multidrug-resistant tuberculosis, and the drugs used are costly and have higher toxicity rates. Important drugs used for treatment of multidrug-resistant tuberculosis and their main adverse effects are shown in Table 4.

### Prevention strategies

It has been noted worldwide that only 20% of patients with tuberculosis receive treatment according to current WHO recommendations. The WHO emphasizes and warns that if immediate action is not taken to persuade more public health authorities and doctors to use the recommended treatment, the window of opportunity to prevent the spread of drug-resistant strains will be missed.

On the basis of small-scale studies that have used the DOTS strategy to prevent the spread of resistance in tuberculosis patients, the WHO has been emphasizing and arguing for DOTS for years. Measures to prevent the spread of multidrug-resistant tuberculosis include:

- Cough hygiene
- Well-ventilated wards
- Segregation of suspected cases
- Negative pressure exhausts
- Air exchangers and high efficiency particulate arresting filters
- Personal protection with N95 masks

Apart from these, other measures important in the control of the spread of multidrug-resistant and extensively drug-resistant tuberculosis include major improvements in laboratory facilities and infection control, improving the performance of tuberculosis control programs, better treatment for both drug-susceptible and drug-resistant disease, massive scale-up in diagnosis and treatment of all types of tuberculosis, newer diagnostic tests, newer vaccines, early pulmonary resection combined with chemotherapy in multidrug-resistant and extensively drug-resistant tuberculosis cases, and commitment on a national scale by countries with the highest tuberculosis burden (ie, India, China, and Russia) by improving national policies and health systems that remove financial barriers to treatment, encourage rational drug usage, and create the relevant infrastructure.

### Multidrug-resistant tuberculosis in general practice

General practitioners and primary care physicians have key roles in the successful management of tuberculosis and thus preventing emergence of multidrug resistance in patients. Implementation of DOTS in India should be mainly done in the primary care setting because most of the population reside in rural areas. Timely identification and prompt referral of suspected cases to specialty care facilities for DOTS-Plus are the main roles of primary care doctors in the management of multidrug-resistant tuberculosis. Default from drug treatment is one of the major causes of multidrug resistance, and hence vigilance should be exercised in handling such cases. Figure 1 shows the approach to cases with interrupted category II treatment to prevent emergence of multidrug-resistant tuberculosis.

Factors that are important in making treatment decisions for defaulters include duration of treatment interruption (less than two weeks, 2–7 weeks, or more than eight weeks), duration of treatment before interruption (less than one month, 1–2 months, or more than two months), and the sputum report (positive or negative). In all sputum-negative cases, irrespective

### Table 4 Drugs used in the treatment of multidrug-resistant tuberculosis

<table>
<thead>
<tr>
<th>Group</th>
<th>Drugs</th>
<th>Important adverse effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aminoglycosides</td>
<td>Streptomycin, Kanamycin, Amikacin</td>
<td>Nephrotoxicity, Ototoxicity, Capreomycin is hepatotoxic too</td>
</tr>
<tr>
<td>Thioamides</td>
<td>Ethionamide, Prothionamide</td>
<td>Psychosis, Hepatotoxicity</td>
</tr>
<tr>
<td>Fluoroquinolones</td>
<td>Ofloxacin, Levofloxacin</td>
<td>Gastrointestinal disturbances, Central nervous system symptoms</td>
</tr>
<tr>
<td>Others</td>
<td>Moxafloxacin, Pyrazinamide, Cycloserine, Para-amino salicylic acid, Ethambutol</td>
<td>Hepatotoxicity, hyperuricemia, Convulsions, suicidal tendency, Hepatotoxicity, hypersensitivity, Optic neuritis, peripheral neuritis</td>
</tr>
</tbody>
</table>
of the length of interruption and duration of treatment before interruption, it is advisable to continue the remaining months of WHO category II treatment. Patients who are sputum-positive during treatment of 1–2 months and who had 2–7 weeks of interrupted treatment, should take an extra month of intensive WHO category II treatment. All remaining sputum-positive cases, while on treatment, irrespective of length of interruption and/or duration of treatment before interruption, should receive a fresh course of WHO category II treatment.

Conclusion

Emergence of multidrug-resistant and extensively drug-resistant tuberculosis is a major threat to successful control of tuberculosis worldwide. Prompt implementation of DOTS reduces the chances for development of multidrug-resistant tuberculosis. Early identification and timely referral of suspected multidrug-resistant tuberculosis cases by primary care doctors for DOTS-Plus treatment will help in the appropriate management of multidrug-resistant tuberculosis and thereby prevent development of life-threatening extensively drug-resistant disease.

Also, because tuberculosis is compounded by HIV coinfec-
tion, which hastens the emergence of multidrug-resistant and extensively drug-resistant tuberculosis, novel antituberculous drugs are urgently required. In this context, the creation of new drugs that target metabolic pathways essential for survival of the bacteria, without damaging the host, will prove to be vital. It has recently been found that the 2C-methyl-d-erythritol 4-phosphate (MEP) pathway of M. tuberculosis is one of several pathways that are essential for viability of the organism, and that the human host lacks this particular enzyme, and hence treatment targeted to this enzyme would be less toxic to humans. Thus, the MEP pathway represents a bacterium-specific drug target and, indeed, fosmidomycin is now known to inhibit the second step in the MEP pathway and may lead us to find other novel antituberculous drugs.

Disclosure

The authors report no conflicts of interest in this work.

References


Continuing medical education questions

1. Which of the following statements is/are true regarding WHO treatment categories for tuberculosis?
   a. New, sputum smear-positive sample is WHO category I
   b. Seriously ill, sputum smear-negative is WHO category I
   c. Seriously ill, extrapulmonary is WHO category I
   d. Sputum smear-positive relapse is WHO category I.

2. Which of the following statements is/are true regarding methods used to diagnose multidrug-resistant tuberculosis?
   a. Mycobacteriophage method is conventional culture method
   b. The mycobacterial growth indicator tube is a conventional culture method
   c. Polymerase chain reaction is a molecular method
   d. BACTEC-460 is a molecular method.

3. Which of the following statements is/are true regarding the definition of multidrug-resistant tuberculosis?
   a. It is resistant to isoniazid, rifampicin ± other first-line drugs
   b. It is resistant to isoniazid, rifampicin ± other first-line drugs + an injectable form
   c. It is resistant to isoniazid, rifampicin ± other first-line drugs + a fluoroquinolone
   d. It is resistant to a fluoroquinolone and rifampicin.

4. Which of the following statements best describes the purpose of DOTS-Plus?
   a. To cure tuberculosis and prevent multidrug resistance
   b. To monitor, report, and evaluate treatment outcome
   c. To make the diagnosis by sputum microscopy
   d. Duration of intensive phase of treatment is 2–3 months.

5. Which of these statements predicts poor prognosis in multidrug-resistant tuberculosis WHO category II failures who received intermittent therapy?
   a. Cavity at presentation
   b. HIV positivity
   c. Sputum culture-positive at three months on optimal therapy.

Answers
True: 1.a, 1.b, 1.c, 2.c, 3.a, 4.b, 5.a, 5.b, 5.c.
False: 1.d, 2.a, 2.b, 2.d,