REVIEW Diagnosis and management of miliary tuberculosis: current state and future perspectives

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disease, and it poses formidable challenges to global here h at the put ic health, scientific, and political level. Miliary TB is a potentially fatal form TB by results for massive lymphohematogenous dissemination of Mycobacterium tul arculosis bach. *The epidemiology of miliary* TB has been altered by the emergence of the hand nmunodeficiency virus (HIV) infection and widespread use of immunosuppressing drugs. Lingnost of miliary TB is a challenge that can perplex even the most experience clinical symptoms, and the chest radiographs do not always reveal consical miliary changes. Atypical presentations like cryptic miliary TB and ac the respiratory distress syndrome often lead to delayed diagnosis. High-resolution computed to pgraphy (HeCT) is relatively more sensitive and shows randomly distributed miliary nodules. In xtrapulp nary locations, ultrasonography, CT, and magnetic resonance imaging a ful in discerning the extent of organ involvement by lesions of miliary TB. Recently, pos on mission tomographic CT has been investigated as a promising tool d TB. Fundus examination for choroid tubercles, histopathological for evaluati uspel examine on of t sue bio sy specimens, and rapid culture methods for isolation of *M. tuber*s in m, bouy fluids, and other body tissues aid in confirming the diagnosis. Several el diagnos tests have recently become available for detecting active TB disease, screening ant *M. tuberculosis* infection, and identifying drug-resistant strains of *M. tuberculosis*. However, progress toward a robust point-of-care test has been limited, and novel biomarker discovery remains challenging. A high index of clinical suspicion and early diagnosis and timely instiction of antituberculosis treatment can be lifesaving. Response to first-line antituberculosis ags is good, but drug-induced hepatotoxicity and drug–drug interactions in HIV/TB coinfected patients create significant problems during treatment. Data available from randomized controlled trials are insufficient to define the optimum regimen and duration of treatment in patients with drug-sensitive as well as drug-resistant miliary TB, including those with HIV/AIDS, and the role of adjunctive corticosteroid treatment has not been properly studied. Research is going on worldwide in an attempt to provide a more effective vaccine than bacille Calmette-Guérin. This review highlights the epidemiology and clinical manifestation of miliary TB, challenges, recent advances, needs, and opportunities related to TB diagnostics and treatment.

Abstract: Tuberculosis (TB) remains one of the most import a cause of death from an infectious

Keywords: Mycobacterium tuberculosis, human immunodeficiency virus, diagnostic tests, biomarkers, antituberculosis drugs, vaccine

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Introduction

Tuberculosis (TB) is a leading cause of preventable morbidity and mortality worldwide. The latest World Health Organization (WHO) figures indicate that in 2010 there were 8.8 million incident cases of TB, with 13% of cases occurring among patients with

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human immunodeficiency virus (HIV) infection.^{1,2} The disease primarily involves the lungs, and at times distant bloodborne spread results in the development of extrapulmonary TB (EPTB). Miliary TB is a pathological name describing millet seed-sized (1–2 mm) granulomas in various organs affected by tubercle bacilli.³ It results from massive lymphohematogenous dissemination from a *Mycobacterium tuberculosis*-laden focus. In 1700, John Jacob Manget coined the term "miliary TB" (derived from the Latin word "miliarius," meaning related to millet seed) to denote this lethal form of disseminated TB.^{4–6} In order to clarify the difference between clinical and pathological diagnoses, it has been proposed that the term miliary TB should be restricted to disseminated TB with miliary shadows on chest radiograph.⁷

Miliary TB has a spectrum of manifestations that still perplex the most erudite and experienced clinicians and are a diagnostic and therapeutic challenge. Despite effective therapy being available, mortality from this disease has remained high. The myriad clinical manifestations, atypical radiographic findings, and difficulties in establishing TB as the etiological diagnosis are challenges in the diagnosis and treatment of miliary TB.

Miliary TB is diagnosed by the presence of a diffuse miliary infiltrate on chest radiograph or high-resolution computed tomography (HRCT) scan, or evidence of miling, tubercles in multiple organs at laparoscopy, open surgery, a autopsy. The clinical and morbid anatomic picture even to be confirmed by bacteriology, histopathology, and or a drumatic chemotherapeutic response. The disease is charterized by high mortality, reported to be between 8% and 5.%. The diagnosis is frequently missed, and more intersive investigations are often required.

In this review, we first provide an overview regarding the epidemiology, current orders inding a key pathogenetic mechanisms, and the varied minical confestations in miliary TB, and then the available arage, sic modalities with recent advances and current the tment guidelines of miliary TB are addressed in detail.

Burden of the problem

TB remains a major worldwide health problem, causing almost 2 million deaths every year. The epidemic of TB, fuelled by HIV coinfection and bacillary resistance to current antimycobacterial drugs, continues to plague low-income countries particularly. India bears the highest burden of TB (1.96 million cases annually),⁸ and also a significantly high number of HIV patients (2.3 million prevalent cases).⁹ It is estimated that miliary TB accounts for about less than 2% of all cases of TB in immunocompetent persons and up to 20% of all EPTB cases. Of 11,182 incident cases reported in the United States in 2010, EPTB accounted for approximately 22% of cases; miliary disease was reported in 299 (2.7%).¹⁰ Immunocompromised patients have a significantly higher prevalence of TB than the general population. The disease is more frequently encountered in immunosuppressed individuals. EPTB accounts for more than 50% of all cases of TB in late HIV infection.^{11–18} This disease has shown a high mortality, despite effective therapy being available. Worldwide, estimates of its incidence are hampered largely by incomplete reporting and imprecise diagnostic criteria.

Since its first description Jon Jacob Manget, the clinical presentation has changed dramate ally. Miliary TB has been considered to be a child odd bease for a long time. However, during the last the decrees, it is increasingly being recognized in club, 10.19-21 Leas been noticed recently that there is a hor hor hor hor hor hor miliary TB owing to the MV ep. emic, and the increasing list of causes of and suppression, such as introduction of biological and im munosuppressive drugs for treatment of various medical reasing occurrence of organ transplantation, and orders, in d modialysis programs. Bacille Calmette–Guérin chron vaccination has resulted in substantial reduction in miliary TB and TB meningitis (TBM) among young vaccines. Increasing use of CT scans and wider application of invasive diagnostic methods are likely to have contributed to the demographic shift. At present, two additional peaks are evident: one involving adolescents and young adults, and the other later in life among elderly persons.^{5,22,23–36} Males appear to be more frequently affected by miliary TB in pediatric as well as adult series.21-38

Pathogenesis

The central event in the development of miliary TB is a massive lymphohematogenous dissemination of *M. tuberculosis* from a pulmonary or extrapulmonary focus and embolization to the vascular beds of various organs. It most commonly involves the liver, spleen, bone marrow, lungs, and meninges. The most likely reason for this distribution is that these organs have numerous phagocytic cells in their sinusoidal wall. Sometimes, simultaneous reactivation of multiple foci in various organs can result in miliary TB. This reactivation can occur either at the time of primary infection or later during reactivation of a dormant focus. When miliary TB develops during primary disease (early generalization), the disease has

an acute onset and is rapidly progressive. Late generalization during postprimary TB can be rapidly progressive (resulting in acute miliary TB), episodic, or protracted, leading to chronic miliary TB. Reinfection also has an important role, particularly in highly endemic areas with increased transmission of *M. tuberculosis*.

The inadequacy of effector T-cell response in suppression of *M. tuberculosis* is thought to be responsible for the development of miliary TB.39-42 The abundance of T-helper 1 and 2 polarized effector T (Teff) cells in the peripheral blood as well as at local disease site(s) of patients with miliary TB suggests that miliary TB possibly represents the T-helper 2 end of the spectrum.^{41,42} Interleukin-4 (IL-4), with its ability to downregulate inducible nitric oxide synthase, toll-like receptor 2, and macrophage activation, may play a crucial role in determining whether the infection becomes latent or progressive.^{39,40} M. tuberculosis can either fail to induce the protective response or can drive the protective mechanisms and then deliberately "sabotage" them, resulting in progressive disease.^{40–42} In miliary TB, the selective recruitment of the Teff cells at the pathologic site, however, fails to provide an adequate level of effector immunity at the disease site due to efficient and comparable homing of regulatory T (Treg) cells, which inhibit the function of the Teff cells that have infiltrated the disease site. It has been postulated that when the bala of homing shifts toward the Treg cells, there occurs a state local immunosuppression leading to disease diss tion.

Clinical presentation

The clinical manifestations of miliary of are proton, non-specific, and can be obscure till late in the clease.

Constitutional symptoms

Presentation with fever a freever a weeks churation, anorexia, weight loss, lassitude and olugh is requent. Occurrence of daily morning templeature spins is reported to be characteristic of miliary TB.⁴ However, fever may be absent and the patients may present with progressive wasting strongly mimicking a metastatic carcinoma (cryptic miliary TB).^{21,44,45} Previously, cryptic miliary TB, which was often diagnosed only at autopsy, is now being increasingly diagnosed with the advent of HRCT. Chills and rigors, described in patients with malaria, or sepsis and bacteremia, have often been described in adult patients with miliary TB.⁴⁶ Night sweats are common.

Systemic involvement

Since miliary TB can involve many organs, patients present with symptoms and signs referring to various organ systems. TBM has been described in 10%–30% of adult patients with miliary TB.^{23–38} On the contrary, about one-third of patients presenting with TBM have underlying miliary TB.⁴⁷ A recently published study⁴⁸ found TBM with and without tuberculomas and thoracic transverse myelopathy as the most frequent neurological complication in patients with miliary TB.

Choroidal tubercles occur less commonly in adult patients with miliary TB than children. If present, choroidal tubercles are pathognomonic of miliary TB and offer a valuable clue to the diagnosis (Figure 1A and B). Choroidal tubercles are bilateral, pale, gray-white, or yellowish lesions usually less than one-quarter of the size of the optic disk and are located within 2 cm of the optic nerve. Therefore, a systematic ophthalmoscorie examination after mydriatic administration is recommended in all patients with suspected miliary TB.

Cutaneous le ions r ay off, a valuable clue to the diagnosis of hilk B. She involvement in the form of erythy tosus may and papules has also been described.³⁻⁶ Sons of hepatic involvement may be evident Torm of ictorys and hepatosplenomegaly. Before the in vent of nodern imaging modalities, such as CT, MRI, echocatiography, clinically evident cardiac or renal al invol nt was seldom documented in patients with mil-**TB**.³⁻⁶ Overt adrenal insufficiency at presentation or during treatment has also been described in miliary TB.49 Atypical presentations^{21,25–38,44,48–66} can delay the diagnosis, and miliary TB is often a missed diagnosis. Patients with occult miliary TB can present with "pyrexia of unknown origin" without any localizing clue. Clinical presentation such as absence of fever and progressive wasting strongly mimicking a metastatic carcinoma can occur, especially in the elderly. Proudfoot et al²¹ suggested the term "cryptic miliary TB." Few studies have highlighted the important differences between classical and cryptic forms of miliary TB.^{21,44,45}

Children

By contrast with adults, fewer published series are available on childhood miliary TB.^{67–71} Clinical presentation of miliary TB in children is similar to that observed in adults. In children with miliary TB, chills, night sweats, hemoptysis, and productive cough have been reported less frequently, while peripheral lymphadenopathy and hepatosplenomegaly are more common, compared with adults. A higher proportion of children with miliary TB (20%–40%) suffer from TBM^{67–71} compared with adults.

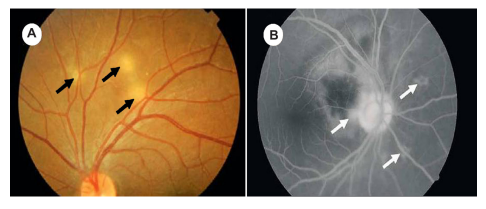


Figure I (A) Ophthalmoscopic pictures showing multiple choroidal tubercles (black arrows); (B) choroidal tubercles (white arrows): fluorescein angiogram.

Immunosuppressed individuals

The clinical presentation of miliary TB in early HIV infection (CD4⁺ cell counts > 200 cells/ μ L) is similar to that observed in immunocompetent individuals.72-74 With progression of immunosuppression in late, advanced HIV infection (CD4+ cell counts < 200 cells/ μ L), disseminated and miliary TB are seen more often.^{15,75} A number of studies have addressed the comparison of various aspects of miliary TB in the late advanced stage of HIV infection and in immunocompetent individuals.^{15,72,75–77} Cutaneous involvement is unusual in miliary TB, but is more commonly seen in HIV-infected patients with severe immunosuppression.78 Typically, cutaneous lesions are few in number and appear as til papules or vesiculopapules,79 described as tuberg cuti miliaris disseminata, tuberculosis cutis acut gener lisita, and disseminated TB of the skin. Sometizes, h dar, pustular, or purpuric lesions, indurated ut ating play es, and subcutaneous abscesses have been reported

In miliary TB patients coincected with FeV, intrathoracic lymphadenopathy an etuber alin anergy are more common; sputum smear are sodom rusitive, and blood culture may grow *Metubercolosis*, exercially with profound immunosuppression.^{2–74}

Immune reconstitution inflammatory syndrome (IRIS) has been implicated as the cause of paradoxical worsening of lesions in patients with TB. IRIS has been reported to occur in about one-third of patients with HIV/TB coinfection within days to weeks of the initiation of highly active antiretroviral therapy. IRIS can be brief or prolonged with multiple recurrences. Manifestations of IRIS range from isolated instances of fever to increased or initial appearance of lymphadenopathy, new or worsening pulmonary infiltrates, serositis, cutaneous lesions, and new or expanding central nervous system (CNS) mass lesions.⁸⁰ Consequently, HIV/miliary TB coinfected patients may

develop acute renal failure⁸¹ or acute respiratory distress syndrome (ARDS).⁸²

Uncommon clinical manifestations and complications

Several uncommon mical manifestations and complications have been observed in patients with miliary TB (Table 1).^{21,25} 14:50–66 Atypical clinical presentation often debys diagnosis and treatment, and miliary TB is often a "hissed diagnosis."

 Table 1 Oncommon clinical manifestations and complications in

 mina.
 tuberculosis

Systemic manifestations

- Cryptic miliary tuberculosis
- Pyrexia of unknown origin
- Shock, multiorgan dysfunction

Incidental diagnosis on investigation for some other reason
Pulmonary

- Acute respiratory distress syndrome
- "Air leak" syndrome (pneumothorax, pneumomediastinum)
- Acute empyema
- Cardiovascular
- Pericarditis with or without pericardial effusion
- Sudden cardiac death
- Mycotic aneurysm of aorta
- Native valve, prosthetic valve endocarditis
- Renal
- Overt renal failure due to granulomatous destruction of the interstitium
- Immune complex glomerulonephritis
- Hematological
 - Myelopthisic anemia
 - Immune hemolytic anemia
 - Endocrinological
 - Thyrotoxicosis
 - Hepatic
 - Cholestatic jaundice
 - Others
 - Presentation as focal extrapulmonary tuberculosis

Complications are often self-limited and improve with antituberculosis therapy (ATT) alone. However, at times they can be life-threatening, necessitating prompt recognition and treatment. Important complications in patients with miliary TB include air-leak syndromes (eg, pneumothorax, pneumopericardium), ARDS, antituberculosis drug-induced hepatotoxicity, and renal failure. Rarely, cardiovascular complications and sudden cardiac death have been described in miliary TB.^{61–65}

Diagnosis

Even in the endemic area, the diagnosis of miliary TB can be difficult, as the clinical symptoms are nonspecific, the chest radiographs do not always reveal the classical miliary changes, and atypical presentations are commonly encountered. Therefore, a high index of clinical suspicion and a systematic approach to diagnostic testing is required to establish the diagnosis of miliary TB.

The following criteria have been proposed for the diagnosis of miliary TB:³⁵ (1) clinical presentation consistent with the diagnosis of TB – like pyrexia with evening rise of temperature, night sweats, anorexia, and weight loss of greater than 6 weeks in duration – responding to antituberculosis treatment; (2) typical miliary pattern on chest radiograph; (3) bilateral, diffuse reticulonodular lung lesion on a background of miliary shadows demonstrable either of chest radiograph or HRCT scan; and (4) microbiological of histopathological evidence of TB.

A high index of clinical suspicion with en towards confirming the diagnosis by demonstrating M. tub. culosis early in the course of disease is imperative. Smear and culture examination of spontaneously expectorate or induced sputum, gastric lavage, plenal, portoneal, or pericardial fluid, cerebrospinal fluid urine pus form cold abscess, bronchoscopic secretions, and periperal blood is helpful in the diagnosis of milk v T. Methological and histopathological examination of one marrow, liver and peripheral lymph node, and transbronchial lung biopsy specimens have all been used to confirm the diagnosis of miliary TB, with varying results.^{25,32–36,83} Whenever possible, efforts should be made at procuring tissue/fluid for mycobacterial culture and sensitivity testing. Rapid-culture methods such as the Bactec 460 radiometric method or Bactec Mycobacterial Growth Indicator Tube (MGIT) 960 system may be useful for rapid drug-susceptibility testing.^{17,84} In the published reports, no systematic pattern of diagnostic approach is available. A standard diagnostic approach to a patient with suspected miliary TB is shown in Figure 2.

Laboratory findings

Hematological and biochemical

A number of hematological and biochemical abnormalities are known to occur in patients with miliary TB.^{23–30,33–38,67–69} Anemia of chronic disease, leukocytosis, leucopenia, leukamoid reactions, and thrombocytopenia are some of the common abnormalities found. Erythrocyte sedimentation rate is usually elevated in patients with miliary TB. Disseminated intravascular coagulation has been described in patients with miliary TB in the setting of ARDS and multiple organ dysfunction syndrome and is associated with a high mortality.⁸⁵ Immune mechanisms have been implicated to cause bone marrow suppression and resulting pancytopenia or hypoplastic anaemia.⁵⁶

Hyponatremia in miliarur B cale ccur due to an acquired disturbance of neurohypophyseal function resulting in unregulated antidiurete horme e release. Hyponatremia may indicate the presence of 7 BM³⁶ and may also be a predictor of mortality²⁰⁻⁵ Hypotcalcenta has also been described in miliary TP, but is uncentron.

Tylerculin skin test

A higher proportion of patients with miliary TB manifest therculin a ergy than those with pulmonary TB or EPTB. Tubercum skin test (TST) conversion may occur following a ccessful treatment. In various published pediatric^{67–71} and adult series,^{4,24–29,32–34,37} tuberculin anergy has ranged from 35% to 74% and 20% to 70%, respectively. Because of tuberculin anergy, cross-reactivity with environmental mycobacteria and tuberculin positivity due to BCG vaccination, the TST is not useful as a diagnostic test in patients with miliary TB. Tuberculin test positivity suggests infection, but it does not distinguish between latent TB infection and active disease. Although a positive TST signifies a possible diagnosis of miliary TB, a negative test does not exclude it.

Interferon-gamma release assays

Currently, two commercial interferon- γ release assays (IGRAs), the Quantiferon-TB Gold (QFT-G) and the T-Spot-TB, are approved. They measure interferon- γ released following incubation of patient blood with antigens specific to *M. tuberculosis*, namely early secretory antigenic target-6 (ESAT-6) and culture filtrate protein 10 (CFP-10). The QFT-G test is now available as an "in-tube" version, which also includes, in addition to ESAT-6 and CFP-10, the antigen TB7.7.⁸⁶ IGRAs do not differentiate latent TB infection from active TB disease and are not significantly superior to TST,

Suspected miliary TB ↓

No single symptom or sign is diagnostic of miliary TB. Clinicians should look for a constellation of symptoms and signs suggestive of miliary TB, such as

- Peripheral lymphadenopathy
- Skin lesions
- Hepatosplenomegaly
- Signs of meningeal irritation
- Pleural, pericardial effusions, ascites
- Choroid tubercles (fundoscopy)

Diagnostic evaluation should focus on confirming miliary TB

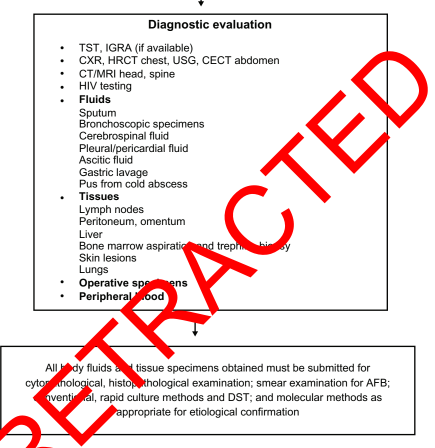


Figure 2 Algorithm for the gno even of a patient with suspected miliary tuberculosis (TB).

Abbreviations: AFB, acid-fast cilli; CECT, contrast enhanced computed tomography; CXR, chest radiograph; DST, drug-susceptibility testing; HRCT, high resolution computed tomography; IGRA, interview on-y release assays; MRI, magnetic resonance imaging; TST, tuberculin skin test; USG, ultrasonography.

albeit they have the ability to identify latent TB infection in HIV-infected individuals.^{86,87} The WHO advises against the use of IGRAs over TST as a diagnostic test in low- and middle-income countries with typically high TB and/or HIV burdens.⁸⁸

Imaging studies

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Miliary pattern on the chest radiograph is often the first clue suggestive of miliary TB. Several other imaging modalities, such as ultrasonography, CT, MRI, and positron-emission tomography (PET), help to assess the extent of organ involvement and are also useful in evaluating response to treatment.

Chest radiograph

The radiographic hallmark of miliary TB is the miliary pattern on chest radiograph (Figure 3A). The term miliary refers to the "millet seed" size of the nodules (<2 mm) seen on classical chest radiograph. Subtle miliary lesions are best delineated in slightly underpenetrated films, especially

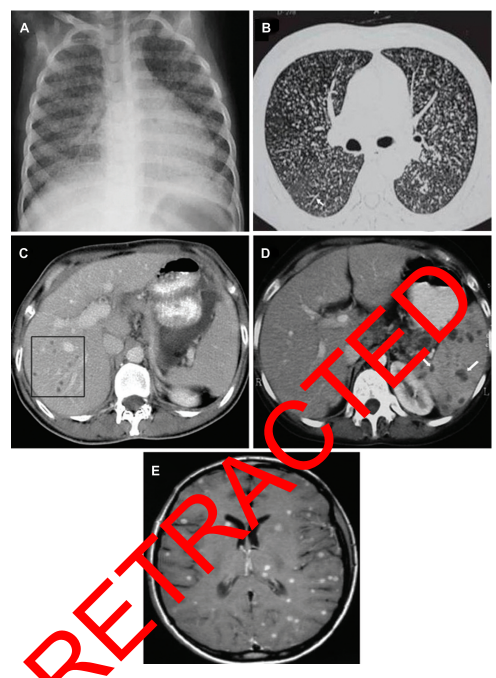


Figure 3 (A) Chest radiograph, esteroanterior view) showing classical miliary pattern. (B) High-resolution computed tomography image (1.0 mm section thickness) shows uniform-sized small nodules rando by distributed throughout both lungs. Note the classical "tree-in-bud" appearance (white arrow). (C) Contrast-enhanced computed tomography of the abdomen, showing ocal miliary lesions in the liver (square) and (D) spleen (white arrows). (E) Miliary central nervous system tuberculosis. Note: Axial contrast-enhanced T_1 -weighted magnetic resonance image shows multiple small foci within both cerebral hemispheres.

when the areas of the lung in between the ribs are carefully scrutinized.^{89,90} The chest radiographic abnormalities in miliary TB are described in Table 2.⁴ In about 10% of cases, the nodules may be greater than 3 mm in diameter.⁷⁸ Chest plain films are usually normal at the onset of symptoms, and the earliest finding, seen within 1–2 weeks, may be hyperinflation. As the typical changes evolve over the course of disease, obtaining periodic chest radiographs in patients

presenting with pyrexia of unknown origin may be rewarding. In the pre-CT scan era, diagnosis of miliary TB was frequently missed on the chest radiographs and was evident only at autopsy. Evidence from published studies indicates that the classic miliary pattern may not be evident in up to 50% of patients with miliary TB.^{23–26,90}

A classical miliary pattern on the chest radiograph represents the summation of densities of tubercles that

Classical presentation (50%)	Nonmiliary pulmonary manifestations (10%–30%)	Other associated findings (<5%)
Miliary pattern	Asymmetrical nodular pattern	 Pulmonary Parenchymal lesions and cavitation Segmental consolidation Thickening of interlobular septae
	Coalescence of nodules	Pleural Pleural effusion Empyema Pneumothorax Pneumomediastinum
	Mottled appearance	Others Intrathoracic Lymphadenopathy Pericardial effusion
	"Snowstorm" appearance Air-space consolidation	

Table 2 Chest radiographic abnormalities in miliary tuberculosis

are perfectly aligned, whereas curvilinear densities and reticulonodular pattern result from imperfectly aligned tubercles.⁹¹ The histopathological composition of the tubercles, their number, and their size have been proposed to be the determinants of radiographic visibility of the nodules.⁹¹ Rarely, lymphatic obstruction or infiltration can result it ground-glass appearance.⁹² The diagnosis of milia / TB becomes easier when a patient presents with typical and shadows on chest radiograph in an appropriate utting, as compared to those who do not show the plassical pattern. Thus, if there is a high index of subjicion of n iliary TB and the chest radiograph is atypical, it ispuggested that HRCT be done to support the diagnostic

Ultrasonography

In patients with miliary, B, ultrasonography is a useful tool in detecting associated lesions, such as loculated ascites, focal hepatic and splenic lesions, adnexal mass, intra-abdominal lymphadenopathy, and cold abscess. Ultrasonography guidance also facilitates diagnostic thoracic or abdominal paracentesis to procure pleural or peritoneal fluid for diagnostic testing, especially if the fluid is loculated.

Computed tomography and magnetic resonance imaging

In comparison with the pre-CT era, HRCT scans have considerably improved the antemortem diagnosis of miliary TB and may demonstrate miliary disease before it becomes radiographically apparent. On a thin-section CT, a mixture of both sharply and poorly defined 1-4 mm nodules are seen in a diffuse, random distribution often associated with intra- and interlobular septal thickening (Figure 3B).93 The interlobular septal thickening or intralobular fine network that is evident on HRCT scans in miliary TB seems to be caused by the presence of tubercles in the interlobular septa and alveolar walls. Sometimes, in subjects with active postprimary disease, centrilobular nodules and branching linear structures with a "tree-in-bud" appearance may be evident.94 Contrastenhanced CT scans are better for detecting additional findings, such as intrathoracic lymphadenopathy, calcification, and pleural lesions. A higher prevalence of interlobular septal thickening, necrotic implemented and extrathoracic involvement has been observed in HIL seropositive patients with miliary TB.76

Miliary TB is an intertitial Lag disease (ILD), having clinical, radio ogn 1 and physiological similarities with other ILD is a result of miliary TB with other ILDs, it pses diagnostic and therapeutic challenges nysicians. It is to be emphasized that an early and to finite diagonosis of miliary TB is of paramount importance table condition, whereas most other ILDs do t is a tre not ha a specific treatment. On this issue, Pipavath and on gues⁹⁵ describe the HRCT findings and correlation of these findings with pulmonary function and gas-exchange parameters in miliary TB. In addition to the demonstration of miliary nodularity in HRCT, this study has demonstrated other radiological features (consolidation, ground-glass, and focal cystic abnormalities), which cannot be seen in chest radiographs. Another important HRCT finding from this study is the demonstration of emphysematous changes following treatment. They have also demonstrated that HRCT findings correlate with restrictive physiology and impaired gas exchange, as in other interstitial lung diseases.95

Contrast-enhanced CT and MRI have been useful in identifying miliary lesions at occult extrapulmonary sites, an exercise that was earlier possible only at postmortem examination. Abdominal CT is useful in identifying lesions in the liver, spleen, mesentery, peritoneum, and intra-abdominal lymphadenopathy, and also detects cold abscesses.^{17,96} Unlike HRCT scans of the chest, where the classic nodular lesions are less than 2 mm, miliary lesions in the liver and spleen may appear as confluent or discrete hypodense lesions (Figure 3C and D), sometimes with peripheral rim enhancement.^{17,96}

Miliary CNS TB is usually associated with TBM and appears at MRI as multiple tiny, hyperintense T_2 foci that

homogeneously enhance on contrast enhanced T_1 -weighted images (Figure 3E). The MRI is particularly helpful in identifying and delineating the extent of tuberculomas and cold abscesses and monitoring the response to treatment.

Pelvic evaluation with all imaging modalities should be routinely done in all female patients for defining the extent of involvement. Image-guided radiological procedures such as fine-needle aspiration for cytological examination and biopsy under CT or MRI guidance are useful for procuring tissue/body fluids for diagnostic testing.

Positron-emission tomography

PET-CT using the radiopharmaceutical ¹⁸F fluorodeoxyglucose has the potential to play a role in assessing the activity of various infectious lesions, including TB.^{97,98} The PET-CT is suitable for defining the extent of disease at the time of initial presentation (Figure 4A and B). Though ¹⁸F fluorodeoxyglucose PET/CT is not specific for TB, it plays an important role in the evaluation of known or suspected TB cases. It can determine the activity of lesions, guide biopsy from active sites, detect occult distant foci, and evaluate response to therapy. In the future, labeling antituberculous drugs like isoniazid and rifampicin with positron-emitting isotopes may culminate in the development of TB-specific PET radiopharmaceuticals.

Pulmonary functions, gas-exchange abnormalities

Miliary TB is associated with abnormalities of admonary function typical of diffuse interstitial diffuse of the heres.^{99,100} Impairment of diffusion has been the monefrequent and severe abnormality encountered.¹⁰⁰ Additionally, a mild reduction in flow rates suggestive of peripheral airways involvement may be observed. During the acute stage, arterial hypoxemia are to videning of the alveolar–arterial oxygen gradient and hepocapina due to tachypnea are



Figure 4 Coronal plain computed tomography (A) and positron-emission tomography (B) images showing diffuse increased ¹⁸F fluorodeoxyglucose uptake in spleen and multifocal uptake in liver, mediastinal node (black arrow).

also observed. These patients have abnormal cardiopulmonary exercise performance, with lower maximum oxygen consumption, maximal work rate, anaerobic threshold, peak minute ventilation, breathing reserve, and low maximal heart rate.^{102,103} Some of these patients manifest a demonstrable fall in oxygen saturation (to 4% or more) with exercise. Following successful treatment, most patients reveal reversal of abnormalities. However, some of these abnormalities may persist following treatment.^{102,103}

Sputum examination – staining and culturing

Though not all patients with miliary TB manifest productive cough, when available, sputum must be subjected to smear and mycobacterial autum examination. Sputum smear microscopy using Kiehl–Neel en staining is useful in detecting acid-fact bacilla Fluore cence microscopy is credited with increased sensitivity and lower work effort, but has a rider of acreated cost and technical complexity. Various developments are using made in the field of fluorescent microscopy, including light-emitting diode-based fluorescent microscopy, mobile phone-based microscopy, and automated of tection systems using image processing.¹⁰⁴

Culture emains the gold standard for the laboratory contr. on of TB. Although culture-based diagnosis of recommended in the International Standards of Tuberculosis Care,105 lack of resources and technical expertise poses a major limitation in most of the high-prevalence countries. Traditionally, primary isolation and culture of mycobacteria is performed on Löwenstein-Jensen medium, which takes at least 21 days for a result. Liquid culturing with radioisotopic detection or with the incorporation of fluorescent dyes was introduced in the past as a confirmatory method (Bactec 460, Bactec MGIT 960 system, MB/ BacT, and Versa Trek system). The mean turnaround time for mycobacterial growth in smear-positive specimens is 9 days for MGIT 960 and 38 days for Löwenstein-Jensen medium, whereas in smear-negative specimens it is 16 and 48 days, respectively.¹⁰⁶ Microscopic observation drug susceptibility testing developed recently allows both rapid and low-cost TB diagnosis in liquid culture with the simultaneous determination of drug susceptibilities.107 Some other unconventional methods, like thin-layer agar and the direct nitrate reductase assay, have attempted to address the problem of multiple-point processing and hence the generation of aerosols by incorporating visual inspection of results in the form of typical colony morphology or color change to identify TB growth.108

Bronchoscopy

Fiberoptic bronchoscopy, bronchoalveolar lavage (BAL), bronchoscopic aspirate, brushings, washings, and transbronchial lung biopsy are useful in confirming the diagnosis of miliary TB. The cumulative diagnostic yield for various bronchoscopic specimens by smear and culture methods in published studies has been found to be 46.8%.^{32–36} In patients with dry cough, BAL fluid obtained through fiberoptic bronchoscopy should be submitted for mycobacterial smear, culture, and molecular methods.

Body-fluid and tissue examination

In patients with suspected miliary TB, depending on the extent of organ-system involvement, appropriate tissue and body-fluid samples must be obtained to confirm histopathological microbiological diagnosis. Elevated serum alkaline phosphatase levels indicate diffuse liver involvement; needle biopsy of the liver can be useful in confirming the diagnosis. Bone marrow aspiration and needle biopsy have also been found to be useful for the diagnosis of miliary TB. Pleural fluid, pericardial fluid, ascitic fluid, cerebrospinal fluid (CSF), urine, bronchoscopic secretions, blood and tissue biopsy specimens have all been employed to confirm the diagnosis of disseminated and miliary TB. The diagnostic yield of various tissue and body-fluid specimens has b yravariable.^{23-30,32-37,67-69}

Immunological abnormalities

A limited number of reports on the cellular connecteristics of BAL in patients with miliary TB line been published, with conflicting results.^{100,101,109} Patients with TB had a significantly higher total cell count and increased proportion of lymphocytes and CLC⁴ and 2D4⁺ T lymphocytes in the BAL fluid.¹⁰¹ In patients who miliare TB, BAL showed lymphocytic alveolits.^{101,104} The finding of increased CD4⁺ lymphocytes in the BAL fluid.¹⁰¹ The finding of increased CD4⁺ lymphocytes in the BAL fluid.¹⁰¹ The finding of lymphocytes at the site of inflammation.

Polyclonal hypergammaglobulinemia with increase in immunoglobulin (Ig) G, IgA, and IgM was observed in peripheral blood and BAL fluid in one study.¹⁰¹ These findings probably result from increased local synthesis by activated B lymphocytes. Increased BAL fluid fibronectin^{101,110} and serum complement (C3)¹⁰¹ have also been described in patients with miliary TB. The increase in serum C3 has been thought to be the result of "acute-phase response" to ongoing inflammation and elevated BAL fluid fibronectin compared with peripheral blood suggest local synthesis in the lung.

Serodiagnostic and molecular methods

When ascitic or pleural fluid is present, adenosine deaminase (ADA) and interferon- γ estimations can be useful adjuncts in the diagnosis, especially in areas where TB is highly prevalent.^{17,111–113} A recent study¹¹⁴ has shown that that CSF-ADA is a more sensitive indicator than polymerase chain reaction (PCR) for the diagnosis in patients with TB meningitis. As ADA estimation is a cheap, cost-effective test, the utility of CSF-ADA estimation in the diagnosis of TB meningitis merits further study. PCR of blood (especially in HIVinfected patients), CSF fluid, and tissue biopsy specimens may be useful for confirmation of diagnosis.¹⁷ PCR has been found to be most useful when applied to clean specimens such as CSF fluid, where its sensitivity and specificity have been reported to be 50%-50% and 100%, respectively.¹⁷ In patients with suspect miliary 1, wherever possible, automated molecular tests for *M. tube culosis* detection and drug-resistance tering my be use for early confirmation of diagnosis.¹¹⁵ The PUP based supplification of various target nucleic acts has been trind extensively that allows rapid and sensitive detection of target DNA sequences. The PCR mution of the entire 16S–23S rRNA spacer region am d use of secondary technique of randomly amplified DNA fingerprinting to differentiate strains vmorphi p belong to the Mycobacterium genus has been reported.¹¹⁶ targets include the 16S rRNA gene, the 16S–23S internal transcribed spacer, the 65 kDa heat-shock protein, recA, rpoB, and gyrB.

The most significant advance toward a point-of-care (POC) test for TB has come in the field of nucleic acid amplification with the launch of the GeneXpert MTB/RIF assay.117 The assay is capable of detecting the *M. tuberculosis* complex while simultaneously detecting rifampicin resistance within 2 hours. When testing a single sputum sample, the assay detects 98%-100% of sputum smear-positive disease and 57%-83% of smear-negative disease among prospectively studied TB suspects.¹¹⁸ Based on currently available evidence and expert opinion, molecular assays to detect gene mutations that signal drug resistance have been endorsed by the WHO as being most suited for rapid diagnosis.115 Urine represents a clinical sample that is easy to collect from both adults and children, and has been used extensively to evaluate several antigen and DNA detection assays.¹¹⁹ Commercially available assays are able to detect lipoarabinomannan (LAM) in the urine of patients with TB. A cheap POC lateral flow (Determine TB-LAM Ag urine dipstick test) has now been developed, which provides a qualitative (yes/no) readout of a TB diagnosis.120

The ideal TB test would be a POC device capable of providing an on-the-spot accurate diagnosis of active TB in HIV-infected and -uninfected adults and children with pulmonary and EPTB; it should also be able to detect resistance to the first-line TB drugs to avoid initial treatment failure.¹²¹ Table 3 summarizes the strengths and limitations of the currently available tests for TB.

In geographical areas where the prevalence of TB is high, when a patient presents with a compatible clinical picture and a chest radiograph suggestive of classical miliary pattern, it is common practice to start the ATT straight away, keeping in mind the potential lethality of the condition. Measures to confirm the diagnosis are initiated simultaneously.

The Indian perspective

Scientific efforts have been put in by academia and research institutes in India for the development of better diagnostic tools. India has been a big market for in vitro diagnostics, but has been dominated by imported and generic products,

mostly serological, with virtually no innovations. The Revised National Tuberculosis Control Programme (RNTCP), being an official caretaker in India for TB control, has been very active in the recent past. In line with the WHO twelve-point policy package, RNTCP has also adopted strategies to diagnose and manage TB in HIV-infected patients. The program has immediate priorities of restricting TB infection by providing treatment to all infected individuals. For diagnosis, there exist the guidelines for intensive case-finding at the community level, but for early diagnosis of TB in the Indian population, not many efforts could be made. This is very justifiable in the light of huge numbers of already existing cases of TB. The Indian Council of Medical Research (ICMR) has also been working extensively on disease-control programs with the support of the continued exploitation of sometific and technological advances from basic to applied research, from biomedical to health sciences, and for laberatory field research. ICMR is providing significant information through its laboratories and al provides funding to various engaged in TP rese academic research ast ations for research in this area.

Table 3	Key	features	of tests	for T	В
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Test	Pros	Cons
Tuberculin skin test	High specificity in non-BCG-varinated provulations	Training required for administration
	Cost-effectiveness	and interpretation
		Return visit required in 48–72 hours
		for test result
		Possible false-positive and false-
		negative results
Interferon-γ release assay	High spectrucy	Blood withdrawal required
	Only one putent visit required	Indeterminate results in those who are
•	Nults available in 16–24 hours	immunosuppressed
	No confounding by BCG vaccination	No capacity to differentiate between
		latent and active TB
		High cost
Chest radiography	ady availability	Low sensitivity and specificity
	Capacity to differentiate latent infection	Not confirmatory
	from active TB	
Smear microscopy	Ease, speed, and cost-effectiveness of the technique	Low sensitivity
	Quantitative estimate of the number of bacilli	No capacity to differentiate from
	Usefulness in determining infectiousness and in monitoring treatment progress	nontuberculous mycobacteria
Conventional culture using solid media	Examination of colony morphology possible	Wait of 3–8 weeks for result
	Quantitative results	
Automated liquid-culture systems	Sensitivity greater than culture in solid media	Contamination-prone
	Faster results (1–3 weeks)	Stringent quality-assurance systems required
		Expensive equipment required
Nucleic acid amplification test (NAATs)	High specificity	Low sensitivity with smear-negative TB
	Higher sensitivity than smear microscopy	Contamination-prone
	Rapid (1–2 days) diagnosis	Technical skill and expertise required
	Capacity to differentiate TB from other	High cost
	mycobacteria	

Abbreviations: TB, tuberculosis; BCG, bacille Calmette–Guérin.

An international symposium on TB diagnostics held at the International Centre for Genetic Engineering and Biotechnology, New Delhi, India, in December 2010 titled "Innovating to Make an Impact" discussed multiple aspects regarding the challenges in TB diagnostics. A very positive feel for support in the field of diagnostic development came out of this.122 A consultative meeting held in January 2011 at the National AIDS Research Institute, India, - "Galvanizing Evidence for HIV Management" - also incorporated a full session on TB supported by WHO. Exclusive discussions on diagnosing EPTB, childhood TB, and HIV-TB were conducted, as these pose serious challenges to developing universally applicable diagnostic tools for TB. The willingness and determination for better diagnosis and management of TB from laboratory workers to the policy-makers have further shown a promising future.

Treatment

Miliary TB is uniformly fatal within 1 year if untreated.³⁻⁶ ATT is the cornerstone of management. Delay in diagnosis often leads to late institution of specific treatment and significantly contributes to mortality. A greater vigilance with efforts towards confirming the diagnosis by demonstrating *M. tuberculosis* early in the course of disease is imperative. There is no consensus regarding the optimum duration treatment in patients with miliary TB. Moreover, publish randomized controlled trials assessing the efficient of the standard WHO treatment regimens that hav been idely used in national TB-control programs are tso .ng.^{125,} We will discuss the treatment of milizer TB as per the current recommendations by different author tive bodies in the following sections.

Guidelines from professional organizations

According to the WED grademed ²³ patients are categorized as "new patients" or "proviously treated patients." Miliary TB is classified as pulmonary TB because there are lesions in the lungs. New patients with miliary TB receive 6 months of daily or intermittent treatment. The guidelines mention that some experts recommend 9–12 months of treatment when TBM is present given the serious risk of disability and mortality, and 9 months of treatment when bone and joint TB is also present.

In the absence of associated meningeal involvement, the American Thoracic Society, the Centers for Disease Control and Prevention (CDC), the Infectious Disease Society of America,¹²⁵ and the National Institute for Health and Clinical Excellence (NICE) TB guidelines¹²⁶ suggest 6 months of treatment (2-month intensive phase with isoniazid, rifampicin, pyrazinamide, and ethambutol or streptomycin, followed by a 4-month continuation phase with isoniazid and rifampicin) to be adequate in miliary TB, whereas the American Academy of Pediatrics¹²⁷ advocates 9 months of treatment. In the presence of associated TBM, treatment needs to be given for at least 12 months. The NICE TB guidelines¹²⁶ suggest that all patients with disseminated (including miliary) TB should be tested for CNS involvement by CT or MRI of the brain and/or lumbar puncture for those without CNS symptoms and signs. They recommend starting ATT even if initial liver functions are abnormal and careful monitoring during follow-up. Appropriate modification of drug treatment should be de le 11 h patient's liver function deteriorates significantients with miliary TB get treated undernation TB gntrol programs, with the Directly Observed Teatment, Short-course (DOTS) crmittee, thrice-weekly treatment in using short-course, low-econe c-resource countries.¹²³

These observations highlight the importance of accuracy assessing the extent of involvement clinically and radiologically. Thus, if underlying TBM remains undiagnosed in a patient with miliary TB, ATT for only 6 monocuracy be suboptimal. Therefore, though the standard and ion of treatment may be sufficient for many, each patient needs to be assessed individually, and wherever indicated, treatment duration may have to be extended.

Patients with HIV/tuberculosis coinfection

Sparse data are available regarding the efficacy of standard treatment regimens in the treatment of HIV/miliary TB coinfection. The WHO recommends all patients of suspected or confirmed military TB should be tested for HIV status, and in HIV-infected patients with TB, for antiretroviral treatment to be started after the completion of ATT.^{128,129} The strategy for initiation of treatment for both TB and HIV infection is shown in Table 4. In cases of HIV/MTB coinfected children, the CDC recommends 12 months' ATT, including HREZ for 2 months followed by HR for 10 months.¹²⁵ For children already receiving antiretroviral treatment (ART) in whom TB is diagnosed, the ART regimen should be reviewed and optimized for treating HIV/TB coinfection and to minimize potential toxicities and drug-drug interactions. Treatment of miliary TB in patients coinfected with HIV requires careful consideration of drug-drug interactions between antituberculosis and antiretroviral drugs.^{128,130}

 Table 4
 Strategy for initiation of treatment for both TB and HIV infection

Criteria	TB treatment	ART
Extrapulmonary TB	Start	Start ART as soon as TB
(regardless of CD4	immediately	treatment is tolerated
count)		(between 2 weeks and
Pulmonary TB	Start	2 months)ª.
$CD4 < 200 \text{ cells/mm}^3$	immediately	
Pulmonary TB	Start	Start ART after completion
CD4 = 200-350 cells/mm ³	immediately	of initial TB treatment
		phase (start earlier if
		severely compromised).
Pulmonary TB	Start	Monitor CD4 count.
$CD4 > 350 \text{ cells/mm}^3$	immediately	Consider ART if CD4 cell
		count drops below
		350 cells/mm ³ .

Note: ^aThe decision to start ART should also be based on clinical evaluation of other signs of immunodeficiency.

Abbreviations: TB, tuberculosis; ART, antoretroviral therapy.

Coadministration of rifampicin may result in dangerously low levels of antiretroviral agents by inducing the hepatic cytochrome P450 pathway. Rifabutin is preferred over rifampicin, especially when protease inhibitors are used, but it is costly. Efavirenz is preferred over nevirapine, but should be avoided during pregnancy. Recently, there has been a change in the WHO revised recommendations¹²⁸ based on the Grading of Recommendations Assessment, Development and Evaluation system¹³¹ regarding the time of starting antiretroviral drug the choice of drugs, and the time of initiation in conjon to institution of ATT.

In peripheral hospitals in endemic areas who and IV and TB are common, quality-assured laboratory facilities for HIV enzyme-linked immunosorbent assay, CDD T-lymphocyte counts and plasma HIV viral load estimation may not be available. Timing of initiation and ART, choice of ART and ATT regimens, and thug-ong interactions all require careful consideration

Role of corticos, roids

Published data regarding the role of adjunct corticosteroid treatment in patients with miliary TB are few and with conflicting results.¹³² A beneficial response was observed in one study,¹³³ although such benefit could not be documented by another study.¹³⁴ Presence of associated adrenal insufficiency is an absolute indication for corticosteroid administration. Adjunctive corticosteroid treatment may be beneficial in miliary TB with meningeal involvement, large pericardial or pleural effusion, endobronchial TB, IRIS, ARDS, immune complex nephritis, and histiocytic phagocytosis syndrome.^{3–6,81,82} The benefit of corticosteroid administration in patients with miliary TB merits further evaluation in future studies, especially in the setting of pulmonary function abnormalities.

Prevention

Evidence from published studies indicates that BCG vaccination is effective in reducing the incidence of miliary TB, especially in children.¹³⁵ However, it is not effective in individuals who are already infected and should not be administered to immunosuppressed hosts. BCG fails to induce immune responses to RD1 antigens, including ESAT6 and CFP10, which are genetically absent from BCG, but also against a new series of *M. tuberculosis* dormancy (DosR) regulon antigens that are expressed by M. tuberculosis under conditions of intracellular stress (eg, hypoxia), and which may be important in host control of aten. fection.¹³⁶ BCG is also a powerful inducer of Tike which may dampen immunity to *M. tuberculosis* as well as a oster decines. These factors might also explain - at less in the why BCG revaccination does not force, added value against TB.137 Targeted tuberculing uting and that then of latent TB infection is often practiced in co., tries with low prevalence of TB,125 but drughepatitis a potential risk with this intervention. ind ngoing remarch^{138,139} is likely to provide a more effective cine than BCG.

tality and prognostic factors

The mortality related to miliary TB is about 15%–20% in children^{67–71} and 25%–30% in adults.^{23–38} Mortality is strongly associated with age, mycobacterial burden, the delay in initiation of chemotherapy, and laboratory markers indicative of severity of disease, such as lymphopenia, thrombocytopenia, hypoalbuminemia, and elevated hepatic transaminases.^{7,34,140}

Several factors have been identified as predictors of poor outcome in patients with miliary TB.^{23–30,32–37,71} Recognition of these factors can alert clinicians managing patients with miliary TB. A 4-point nutritional risk score was defined according to the presence of four nutritional factors: low body mass index (<18.5 kg/m²), hypoalbuminemia (serum albumin < 30 g/L), hypocholesterolemia (serum cholesterol < 2.33 mmol/L), and severe lymphocytopenia (<7 × 10⁵ cells/L). Each risk factor was assigned a value of 1 if present or 0 if absent. Patients with 3 or 4 points were classified to have a high nutritional risk score.¹⁴¹

Challenges in the treatment of miliary TB

TB is unique among the major infectious diseases in that it lacks accurate rapid POC diagnostic tests. Failure to

control the spread of TB is largely due to our inability to detect and treat all infectious cases of pulmonary TB in a timely fashion, allowing continued *M. tuberculosis* transmission within communities. Challenges to effective solutions include lack of access to diagnosis and treatment, the frequent coexistence of epidemics of TB and HIV, and the increasing prevalence of drug-resistant TB.

Miliary TB is a challenge for clinicians. The key practical issues that may pose difficulties while treating a case of miliary TB are listed below.

- Choice of the right antituberculosis drug regimen, adding steroids, duration of ATT, inadequacy of laboratory monitoring facilities, and difficulties in managing complications (especially in peripheral centers due to lack of expertise) are all therapeutic challenges. Failure to assess the extent of organ-system involvement initially (eg, TBM) may result in suboptimal duration of therapy.
- While treating TB, drugs should be genuine with good bioavailability, which may not be the case in resource-limited nations despite having high disease prevalence.
- In HIV-coinfected patients, even with regular antituberculosis drug intake, adequate plasma levels may not be achieved because of malabsorption problems.
- Regarding ART and ATT, several issues are still unclear, like sufficient staff training for recognition of adverse effects and close monitoring of codrug toxicities, lace of quality-assured laboratory facilities where the lisease is common, and IRIS diagnosis (proper education of patients for recognition of drug toxicities, dreamanerence issues).

Besides these challenges, heating in TB following "successful" treatment results in fibrosis and consequent anatomical and physiological alterations of the involved organs.^{142–144} The persistence corphysiological, immunological, and radiological delects in antilary TB in spite of treatment and the observation corpulate in treated cases of pulmonary TB patients point out that these patients will not regain optimal health despite achieving a microbiological cure.¹⁴⁵

Conclusion and future direction

Miliary TB is a potentially lethal disease that still perplexes even the most experienced clinicians. Newer technological tools should be used to unravel the immunopathologic phenomenon that results in this form of TB. The role of new interferon- γ assays in the diagnosis of miliary TB needs to be explored in the field. An attempt should be made for systematic data collection and reporting to study the global epidemiology of miliary TB through national TB-control programs to ensure that the proposed diagnostic criteria are strictly adhered to. Miliary TB has shown a high mortality despite the availability of effective treatment. The cause of death in patients with miliary TB merits further study. Appropriately designed randomized controlled trials are needed to define the optimum regimen and duration of treatment in miliary TB patients, including those with HIV/AIDS. The role of adjunctive corticosteroid therapy in the treatment of miliary TB to prevent physiological and radiological abnormalities has not been properly studied in controlled clinical trials and needs to be elucidated in future studies. The scope and utility of PET-CT in assessing the activity of posttreatment residual lesions in miliary TB needs to be ascertained.

Our tools to combat TB re day erously out of date and ineffective. Besides new to ls, we also need new strategies to identify key *M. tuber dlosis*, man best interactions, where we can most like of find M. tus culosis's Achilles' heel. Equally important that we boild high-quality clinical trial capacity a biobanks br 3 biomarker identification. The attempt to preserve an ultimate TB vaccine is still a reverie a mere rell-targeting vaccine may not be sufbe ient; rather; other (innate immune-related) cells, such as vral-kille cells, $\gamma\delta$ T cells, DC, or macrophages need to n. be act. and triggered in a timely fashion.¹⁴⁶ Translational sh into better TB diagnostics, drugs, and vaccines has increased globally, but an improved understanding of the basic infection biology of this complex disease is required before radically new interventions can be designed. The search for a better vaccine than BCG is still on, and more data on the candidate vaccines that are currently being evaluated are expected to emerge.

The precise immunopathogenesis of pulmonary fibrosis is not adequately understood, and drugs are not presently available to reverse the process. Nevertheless, there are promising results from basic science research that stem cell therapy in the lung may facilitate lung regeneration and repair.¹⁴⁷ Research, therefore, should be aimed at unraveling the mystery of the immunopathogenesis of fibrosis and discovering drugs that can avert the incidence of fibrosis and reverse fibrosis once it has developed.

In response to the global emergency of the TB pandemic, the Stop TB Partnership was established by the World Health Assembly in May 2000, which consists of a partners' forum, a coordinating board, and a partnership secretariat currently hosted by the WHO in Geneva, Switzerland. A promising and important portfolio of new TB diagnostics, new TB drugs and vaccines has been endorsed by the Stop TB Partnership.

The challenge now is to complete development and validation of these in high-TB and high-TB/HIV-burden countries and then translate them into clinical practice at peripheral points of health care.¹⁴⁸ With limited finances, priority must be given to the development of technologies that will reach those not being served by current diagnostic provision. In the future, therapeutic interventions based on the results of novel diagnostic procedures can be made earlier, leading to improvements in patient care.

Disclosure

The authors report no conflicts of interest in this work.

References

- 1. World Health Organization. *WHO Report: Global Tuberculosis Control*. Geneva: World Health Organization; 2011.
- Murray CJ, Lopez AD. Global mortality, disability, and the contribution of risk factors: Global Burden of Disease Study. *Lancet*. 1997;349: 1436–1442.
- Sahn SA, Neff TA. Miliary tuberculosis. Am J Med. 1974;56: 495–505.
- Sharma SK, Mohan A. Disseminated and miliary tuberculosis. In: Sharma SK, Mohan A, editors. *Tuberculosis*, 2nd ed. New Delhi: Jaypee Brothers; 2009:493–518.
- Baker SK, Glassroth J. Miliary tuberculosis. In: Rom WN, Garay SM, editors. *Tuberculosis*. Philadelphia: Lippincott Williams & Wilkins; 2004:427–444.
- Divinagracia R, Harris HW. Miliary tuberculosis. In: Schlossberg D, editor. *Tuberculosis and Nontuberculous Mycobacterial Infection*. Philadelphia: WB Saunders; 1999:271–284.
- Matsushima T. Miliary tuberculosis or disseminated tuberculosis. *Med.* 2005;44:687.
- World Health Organization. Global Tuberculosis Control Surve lance Planning, Financing. Geneva: WHO; 2008.
- 9. National AIDS Control Organisation. *HIV Sentrel & even ance and HIV Estimation in India 2007: A Technical Brit /* 2008. As dable from: http://www.nacoonline.org/upload/Publicat.on/M&ESurvillance, Research/HIVSentinelSurveillanceandHIV etimation2007_A Technical Brief.pdf. Accessed November 15, 2012.
- 10. Centers for Disease Control and revention. *Reported Suberculosis in the United States, 2010.* Atlant, GA: C. C; 2011.
- Directorate General of Health Service. Ministroof Health and Family Welfare. RNTCP Performance report, edian with quarter. New Delhi: Ministry of Health and Family Welfare; 104.
- 12. Wares F, Balasubrah, and F, Wares F, Sharma SK. Extrapulmonary tuberculosis: management and control. In: Agarwal SP, Chauhan LS, editors. *Tuberculosis Control in India*. New Delhi: Elsevier; 2005: 95–114.
- [No authors listed.] National survey of notifications of tuberculosis in England and Wales in 1988. Medical Research Council Cardiothoracic Epidemiology Group. *Thorax*. 1992;47:770–775.
- Shafer RW, Kim DS, Weiss JP, Quale JM. Extrapulmonary tuberculosis in patients with human immunodeficiency virus infection. *Medicine* (*Baltimore*). 1991;70:384–397.
- Jones BE, Young SM, Antoniskis D, Davidson PT, Kramer F, Barnes PF. Relationship of the manifestations of tuberculosis to CD4 cell counts in patients with human immunodeficiency virus infection. *Am Rev Respir Dis.* 1993;148:1292–1297.
- Lee MP, Chan JW, Ng KK, Li PC. Clinical manifestations of tuberculosis in HIV-infected patients. *Respirology*. 2000;5:423–426.
- Sharma SK, Mohan A. Extrapulmonary tuberculosis. *Indian J Med Res.* 2004;120:316–353.

- Shao C, Qu J, He L. A comparative study of clinical manifestations caused by tuberculosis in immunocompromised and nonimmunocompromised patients. *Chin Med J.* 2003;116:1717–1722.
- Jacques J, Sloan JM. The changing pattern of miliary tuberculosis. *Thorax.* 1970;25:237–240.
- [No authors listed.] Miliary tuberculosis: a changing pattern [editorial]. Lancet. 1975;1:985–986.
- Proudfoot AT, Akhtar AJ, Doughs AC, Home NW. Miliary tuberculosis in adults. *Br Med J.* 1969;2:273–276.
- Braun MM, Cote TR, Rabkin CS. Trends in death with tuberculosis during the AIDS era. JAMA. 1993;269:2865–2868.
- Slavin RE, Walsh TJ, Pollack AD. Late generalized tuberculosis: a clinical pathologic analysis and comparison of 100 cases in the preantibiotic and antibiotic eras. *Medicine (Baltimore)*. 1980;59:352–366.
- Long R, O'Connor R, Palayew M, Hershfield E, Manfreda J. Disseminated tuberculosis with and without a miliary pattern on chest radiograph: a clinical-pathologic-radiologic correlation. *Int J Tuberc Lung Dis.* 1997;1:52–58.
- 25. Biehl JP. Miliary tuberculosis; a review of sixty-eight adult patients admitted to a municipal general posital. *Am Rev Tuberc*. 1958;77: 605–622.
- Munt PW. Miliary tuber closis in the memotherapy era: with a clinical review in 69 American clults. *Medvine (Baltimore)*. 1972;51: 139–155.
- 27. Campbell IG. Malary tubecculosis a British Columbia. *Can Med* Assoc J. 1973-106, 517-1519.
- Gelb AF, effler C, vrewing , Mascatello V, Lyons HA. Miliary tubercurs. Am Rev Rev 5 Dis. 1973;108:1327–1333.
- Griece MH, et al. Acule disseminated tuberculosis as a diagnostic patham. A clinic study based on twenty-eight cases. *Am Rev Respir Dis.* 1974;109:554–660.
- Onadeko (D), Dickinson R, Sofowora EO. Miliary tuberculosis of the lung in Nigerian adults. *East Afr Med J.* 1975;52:390–395.
- 31. Juliu B. Jutler J, Ostrow JH. Miliary tuberculosis. A review of 83 cases treated between 1950 and 1968. *Ethiop Med J*. 1977;15:39–48.
- 2 Just S, Benatar SR. Disseminated tuberculosis. A study of 62 cases. *SAfr Med J*. 1980;58:835–842.
- Kim JH, Langston AA, Gallis HA. Miliary tuberculosis: epidemiology, clinical manifestations, diagnosis, and outcome. *Rev Infect Dis.* 1990;12: 583–590.
- Maartens G, Willcox PA, Benatar SR. Miliary tuberculosis: rapid diagnosis, hematologic abnormalities, and outcome in 109 treated adults. *Am J Med.* 1990;89:291–296.
- Sharma SK, Mohan A, Pande JN, Prasad KL, Gupta AK, Khilnani GC. Clinical profile, laboratory characteristics and outcome in miliary tuberculosis. *QJM*. 1995;88:29–37.
- Al-Jahdali H, Al-Zahrani K, Amene P, et al. Clinical aspects of miliary tuberculosis in Saudi adults. *Int J Tuberc Lung Dis*. 2000;4:252–255.
- Mert A, Bilir M, Tabak F, et al. Miliary tuberculosis: clinical manifestations, diagnosis and outcome in 38 adults. *Respirology*. 2001;6:217–224.
- Hussain SF, Irfan M, Abbasi M, et al. Clinical characteristics of 110 miliary tuberculosis patients from a low HIV prevalence country. *Int J Tuberc Lung Dis.* 2004;8:493–499.
- Collins HL, Kaufmann SH. The many faces of host responses to tuberculosis. *Immunology*. 2001;103:1–9.
- Rook GA, Hernandez-Pando R, Dheda K, Teng Seah G. IL-4 in tuberculosis: implications for vaccine design. *Trends Immunol*. 2004;25: 483–488.
- Sharma PK, Saha PK, Singh A, Sharma SK, Ghosh B, Mitra DK. FoxP3+ regulatory T cells suppress effector T-cell function at pathologic site in miliary tuberculosis. *Am J Respir Crit Care Med.* 2009;179: 1061–1070.
- Sharma SK, Mitra DK, Balamurugan A, Pandey RM, Mehra NK. Cytokine polarization in miliary and pleural tuberculosis. *J Clin Immunol*. 2002;22:345–352.
- Cunha BA, Krakakis J, McDermott BP. Fever of unknown origin (FUO) caused by miliary tuberculosis: diagnostic significance of morning temperature spikes. *Heart Lung*. 2009;38:77–82.

- Yu YL, Chow WH, Humphries MJ, Wong RW, Gabriel M. Cryptic miliary tuberculosis. *QJM*. 1986;59:421–428.
- Ormerod LP. Respiratory tuberculosis. In: Davies PDO, editor. Clinical Tuberculosis. London: Chapman and Hall Medical; 1997:76.
- Sharma SK, Mohan A. Miliary tuberculosis. In: Schlossberg D, editor. *Tuberculosis and Nontuberculous Mycobacterial Infections*, 6th ed. Washington: American Society for Microbiology; 2011:415–435.
- Thwaites GE, Nguyen DB, Nguyen HD, et al. Dexamethasone for the treatment of tuberculous meningitis in adolescents and adults. *N Engl J Med.* 2004;351:1741–1751.
- Garg RK, Sharma R, Kar AM, et al. Neurological complications of miliary tuberculosis. *Clin Neurol Neurosurg*. 2010;112:188–192.
- Braidy J, Pothel C, Amra S. Miliary tuberculosis presenting as adrenal failure. *Can Med Assoc J.* 1981;124:748–751.
- Penner C, Roberts D, Kunimoto D, Manfreda J, Long R. Tuberculosis as a primary cause of respiratory failure requiring mechanical ventilation. *Am J Respir Crit Care Med.* 1995;151:867–872.
- Mohan A, Sharma SK, Pande JN. Acute respiratory distress syndrome in miliary tuberculosis: a 12-year experience. *Indian J Chest Dis Allied Sci.* 1996;38:157–162.
- 52. Sharma N, Kumar P. Miliary tuberculosis with bilateral pneumothorax: a rare complication. *Indian J Chest Dis Allied Sci.* 2002;44:125–127.
- Das M, Chandra U, Natchu M, Lodha R, Kabra SK. Pneumomediastinum and subcutaneous emphysema in acute miliary tuberculosis. *Indian J Pediatr.* 2004;71:553–554.
- Singh KJ, Ahluwalia G, Sharma SK, Saxena R, Chaudhary VP, Anant M. Significance of haematological manifestations in patients with tuberculosis. *J Assoc Physicians India*. 2001;49:788, 790–794.
- Kuo PH, Yang PC, Kuo SS, Luh KT. Severe immune hemolytic anemia in disseminated tuberculosis with response to antituberculosis therapy. *Chest.* 2001;119:1961–1963.
- Runo JR, Welch DC, Ness EM, Robbins IM, Milstone AP. Miliary tuberculosis as a cause of acute empyema. *Respiration*. 2003;70:529–532.
- Sydow M, Schauer A, Crozier TA, Burchardi H. Multiple organ failure in generalized disseminated tuberculosis. *Respir Med.* 1992 10: 517–519.
- Nieuwland Y, Tan KY, Elte JW. Miliary tuberculosis presenting win thyrotoxicosis. *Postgrad Med J.* 1992;68:677–679.
- Mallinson WJ, Fuller RW, Levison DA, Baker LR, Catell WR, Diffuse interstitial renal tuberculosis – an unusual cause of ren Ufai are. 2011. 1981:50:137–148.
- Shribman JH, Eastwood JB, Uff J. Immute: unplex nepholis complicating miliary tuberculosis. *Br Med J (Clin. ves Ed)*. 1983;287: 1593–1594.
- Wallis PJ, Branfoot AC, Emerson A. Sudden death due to myocardial tuberculosis. *Thorax*. 1984;39: 5–156
- Felson B, Akers PV, Hall GS, Schupper JT, Grene RE, Pedrosa CS. Mycotic tuberculous an arysmof the corary aorta. *JAMA*. 1977;237: 1104–1108.
- Cope AP, Heber M, Vilkie Leo, Jular tuberculous endocarditis: a case report and review of the literature. *J Infect*. 1990;21:293–296.
- Wainwright J. Tuberculou endocarditis: a report of 2 cases. S Afr Med J. 1979;56:731–733.
- Rose AG. Cardiac tuberculosis. A study of 19 patients. Arch Pathol Lab Med. 1987;111:422–426.
- Asada Y, Hayashi T, Sumiyoshi A, Aburaya M, Shishime E. Miliary tuberculosis presenting as fever and jaundice with hepatic failure. *Hum Pathol.* 1991;22:92–94.
- 67. Hussey G, Chisholm T, Kibel M. Miliary tuberculosis in children: a review of 94 cases. *Pediatr Infect Dis J*. 1991;10:832–836.
- Kim PK, Lee JS, Yun DJ. Clinical review of miliary tuberculosis in Korean children. 84 cases and review of the literature. *Yonsei Med J*. 1969;10:146–152.
- Gurkan F, Bosnak M, Dikici B, et al. Miliary tuberculosis in children: a clinical review. *Scand J Infect Dis.* 1998;30:359–362.
- Aderele WI. Miliary tuberculosis in Nigerian children. *East Afr Med J*. 1978;55:166–171.

- Rahajoe NN. Miliary tuberculosis in children. A clinical review. Paediatr Indones. 1990;30:233–240.
- Sharma SK, Mohan A. Co-infection of human immunodeficiency virus (HIV) and tuberculosis: Indian perspective. *Indian J Tuberc*. 2004;51: 5–16.
- Haas DW, Des Prez RM. Tuberculosis and acquired immunodeficiency syndrome: a historical perspective on recent developments. *Am J Med*. 1994;96:439–450.
- Theuer CP, Hopewell PC, Elias D, Schecter GF, Rutherford GW, Chaisson RE. Human immunodeficiency virus infection in tuberculosis patients. *J Infect Dis.* 1990;162:8–12.
- Lado Lado FL, Barrio Gomez E, Carballo Arceo E, Cabarcos Ortiz de Barron A. Clinical presentation of tuberculosis and the degree of immunodeficiency in patients with HIV infection. *Scand J Infect Dis.* 1999;31:387–391.
- Kim JY, Jeong YJ, Kim KI, et al. Miliary tuberculosis: a comparison of CT findings in HIV-seropositive and HIV-seronegative patients. *Br J Radiol*. 2010;83:206–11.
- Harries A, Maher D, Graham S. *TB/HIV: A Clinical Manual*, 2nd ed. Geneva: World Health Organization 2004.
- Daikos GL, Uttamchandani Ko, Tuda vet al. Disseminated miliary tuberculosis of the skin in prents with Al. 3: report of four cases. *Clin Infect Dis.* 1998;27:205-208.
- del Giudice P, Berrand E, Perrin C, et al Janusual cutaneous manifestations of miliary to erculor. *Clin In. A Dis.* 2000;30:201–204.
- Crump JA, Tyre, Mic Lloyd-Owen SJ, Han LY, Lipman MC, Johnson MA. Milian tuberer osis with paradoxical expansion of intracretical uberculomatic plicating human immunodeficiency virus infection in a plicent receiving highly active antiretroviral therapy. *Clin L* for Dis. 1998, 11008–1009.
- Jehle AW, Khanna A, Sigle JP, et al. Acute renal failure on immune reconstitution in an HIV-positive patient with miliary tuberculosis. *Clin Infect Dis.*, 004;38:e32–e35.
- boldsack, R, Allen S, Lipman MC. Adult respiratory distress syndrome as a severe immune reconstitution disease following the commencement Shighly active antiretroviral therapy. *Sex Transm Infect.* 2003;79: 337–338.
- Willcox PA, Potgieter PD, Bateman ED, Benatar SR. Rapid diagnosis of sputum negative miliary tuberculosis using the flexible fibreoptic bronchoscope. *Thorax.* 1986;41:681–684.
- Rodrigues C, Shenai S, Sadani M, et al. Evaluation of the bactec MGIT 960 TB system for recovery and identification of *Mycobacterium tuberculosis* complex in a high through put tertiary care centre. *Indian J Med Microbiol*. 2009;27:217–221.
- Rosenberg MJ, Rumans LW. Survival of a patient with pancytopenia and disseminated coagulation associated with miliary tuberculosis. *Chest.* 1978;73:536–539.
- Cattamanchi A, Smith R, Steingart KR, et al. Interferon-gamma release assays for the diagnosis of latent tuberculosis infection in HIV-infected individuals: a systematic review and meta-analysis. *J Acquir Immune Defic Syndr*. 2011;56:230–238.
- Mori T. Usefulness of interferon-gamma release assays for diagnosing TB infection and problems with these assays. *J Infect Chemother*. 2009;15:143–145.
- World Health Organization. Use of Interferon-γ Release Assays (IGRAs) in TB Control in Low and Middle-Income Settings. Geneva: WHO; 2010.
- Steiner PE. The histopathological basis for the X-ray diagnosability of pulmonary miliary tuberculosis. *Am Rev Tuberc*. 1937;36:692–705.
- Kwong JS, Carignan S, Kang EY, Müller NL, FitzGerald JM. Miliary tuberculosis. Diagnostic accuracy of chest radiography. *Chest.* 1996;110: 339–342.
- Jamieson DH, Cremin BJ. High resolution CT of the lungs in acute disseminated tuberculosis and a pediatric radiology perspective of the term "miliary." *Pediatr Radiol.* 1993;23:380–383.
- Price M. Lymphangitis reticularis tuberculosa. *Tubercle*. 1968;49: 377–384.

- Van Dyck P, Vanhoenacker FM, Van den Brande P, et al. Imaging of pulmonary tuberculosis. *Eur Radiol*. 2003;13:1771–1785.
- McGuinness G, Naidich DP, Jagirdar J, Leitman B, McCauley DI. High resolution CT findings in miliary lung disease. *J Comput Assist Tomogr.* 1992;16:384–390.
- Pipavath SNJ, Sharma SK, Sinha S, Mukhopadhyay S, Gulati MS. High resolution CT (HRCT) in miliary tuberculosis (MTB) of the lung: correlation with pulmonary function tests and gas exchange parameters in north Indian patients. *Indian J Med Res.* 2007;126: 193–198.
- Yu RS, Zhang SZ, Wu JJ, Li RF. Imaging diagnosis of 12 patients with hepatic tuberculosis. World J Gastroenterol. 2004;10: 1639–1642.
- Ichiya Y, Kuwabara Y, Sasaki M, et al. FDG-PET in infectious lesions: the detection and assessment of lesion activity. *Ann Nucl Med.* 1996;10: 185–191.
- Goo JM, Im JG, Do KH, et al. Pulmonary tuberculoma evaluated by means of FDG PET: findings in 10 cases. *Radiology*. 2000;216: 117–121.
- Williams NH Jr, Kane C, Yoo OH. Pulmonary function in miliary tuberculosis. *Am Rev Respir Dis.* 1973;107:858–860.
- 100. Ainslie GM, Solomon JA, Bateman ED. Lymphocyte and lymphocyte subset numbers in blood and in bronchoalveolar lavage and pleural fluid in various forms of pulmonary tuberculosis at presentation and during recovery. *Thorax*. 1992;47:513–518.
- 101. Sharma SK, Pande JN, Singh YN, et al. Pulmonary function and immunologic abnormalities in miliary tuberculosis. *Am Rev Respir Dis.* 1992;145(5):1167–1171.
- Sharma SK, Ahluwalia G. Exercise testing in miliary tuberculosis some facts. *Indian J Med Res.* 2007;125:182–183.
- 103. Sharma SK, Ahluwalia G. Effect of antituberculosis treatment on cardiopulmonary responses to exercise in miliary tuberculosis. *Indian J Med Res.* 2006;124:411–418.
- Lange C, Mori T. Advances in the diagnosis of tuberculosis Respirology. 2010;15:220–240.
- Hopewell PC, Pai M, Maher D, Uplekar M, Raviglione M International standards for tuberculosis care. *Lancet Infect Pie* 2006; 710–725.
- 106. Moore DA, Mendoza D, Gilman RH, et al. Microsoppic observation drug susceptibility assay, a rapid, reliable diagnostic to of or authorugresistant tuberculosis suitable for use in resource-poor storings. *J Clin Microbiol.* 2004;42:4432–4437.
- 107. Moore DA, Evans CA, Gilman RH, etval. Microscopic-observation drug-susceptibility assay for the ragnosis of TLON *Engl J Med*. 2006;355:1539–1550.
- Shikama ML, Ferro e Silva & Villeir G, et al. Multicentre study of nitrate reductase assay for upid detration of rifampicinresistant *M. tubergrosts*, *Int J. Subres Lung Dis.* 2009;13: 377–380.
- 109. Sharma SK, Pando N, Vana A, Onchoalveolar lavage (BAL) in miliary tuberculosis. A *ercle*. 1988;69:175–178.
- 110. Prabhakaran D, Sharman K, Verma K, Pande JN. Estimation of fibronectin in bronchoalveolar lavage fluid in various diffuse interstitial lung diseases. *Am Rev Respir Dis.* 1990;141:A51.
- 111. Sharma SK, Suresh V, Mohan A, et al. A prospective study of sensitivity and specificity of adenosine deaminase estimation in the diagnosis of tuberculosis pleural effusion. *Indian J Chest Dis Allied Sci.* 2001;43: 149–155.
- Sharma SK, Banga A. Diagnostic utility of pleural fluid IFN-gamma in tuberculosis pleural effusion. *J Interferon Cytokine Res.* 2004;24: 213–217.
- Sharma SK, Banga A. Pleural fluid interferon-gamma and adenosine deaminase levels in tuberculosis pleural effusion: a cost-effectiveness analysis. J Clin Lab Anal. 2005;19:40–46.
- Rana SV, Chacko F, Lal V, et al. To compare CSF adenosine deaminase levels and CSF-PCR for tuberculous meningitis. *Clin Neurol Neurosurg*. 2010;112:424–430.

- 115. World Health Organization. Policy statement. Molecular line probe assays for rapid screening of patients at risk of multidrug-resistant tuberculosis (MDR-TB). 2008. Available from: http://www.who. int/tb/features_archive/policy_statement.pdf. Accessed July 19, 2012.
- 116. Kanduma E, McHugh TD, Gillespie SH. Molecular methods for *Mycobacterium tuberculosis* strain typing: a users guide. J Appl Microbiol. 2003;94:781–791.
- Boehme CC, Nabeta P, Hillemann D, et al. Rapid molecular detection of tuberculosis and rifampin resistance. *N Engl J Med.* 2010;363: 1005–1015.
- Helb D, Jones M, Story E, et al. Rapid detection of *Mycobacterium* tuberculosis and rifampin resistance by use of on-demand, near-patient technology. J Clin Microbiol. 2010;48:229–237.
- Minion J, Leung E, Talbot E, Dheda K, Pai M, Menzies D. Diagnosing tuberculosis with urine lipoarabinomannan: systematic review and meta-analysis. *Eur Respir J.* 2011;38:1398–1405.
- 120. Lawn SD, Kerkhoff AD, Vogt M, Wood R. Diagnostic accuracy of a low-cost, urine antigen, point-of-care screening assay for HIVassociated pulmonary tubercularity before antiretroviral therapy: a descriptive study. *Lancet Vect Dis.* 11;12:201–209.
- Weyer K, Carai S, Nunn, TB diagnostes what does the world really need? J Infect Pis. 20, 204 Supp. :S1196–S1202.
- Ghanashyam B. To erculosis the generative section in the section of the section of
- 123. World Health Organization. *Treasurent of Tuberculosis: Guidelines for* Nationale rogrammer 3rd efficienceva: WHO; 2003.
- 124. World and the Organization *Treatment of Tuberculosis: Guidelines*, 4th 6d. Genery: WHO; 2009.
- 125. Pit uberg HM, Aurman WJ, Chaisson RE, et al. American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America: treatment of tuberculosis. *Am J Respir Crit Care Med* 2003;167:603–662.
- 120 National Institute for Health and Clinical Excellence, National Conductating Centre for Chronic Conditions. Management of nonrepiratory tuberculosis. In: *Tuberculosis: Clinical Diagnosis and Management of Tuberculosis, and Measures for Its Prevention and Control.* London: Royal College of Physicians; 2006:63–76.
- 127. [No authors listed.] American Academy of Pediatrics Committee on Infectious Diseases: Chemotherapy for tuberculosis in infants and children. *Pediatrics*. 1992;89:161–165.
- World Health Organization. Rapid Advice: Antiretroviral Therapy for HIV Infection in Adults and Adolescents. Geneva: WHO; 2009.
- Sharma SK, Mohan A, Kadhiravan T. HIV-TB co-infection: epidemiology, diagnosis and management. *Indian J Med Res.* 2005;121:550–567.
- Pozniak AL, Coyne KM, Miller RF, et al. BHIVA treatment guidelines for (TB)/HIV infection 2005. *HIV Med*. 2011;12:517–524.
- Guyatt GH, Oxman AD, Vist GE, et al. GRADE: an emerging consensus on rating quality of evidence and strength of recommendations. *BMJ*. 2008;336:924–926.
- Dooley DP, Carpenter JL, Rademacher S. Adjunctive corticosteroid therapy for tuberculosis: a critical reappraisal of the literature. *Clin Infect Dis.* 1997;25:872–887.
- 133. Sun TN, Yang JY, Zheng LY, Deng WW, Sui ZY. Chemotherapy and its combination with corticosteroids in acute miliary tuberculosis in adolescents and adults: analysis of 55 cases. *Chin Med J (Engl)*. 1981;94:309–314.
- Massaro D, Katz S, Sachs M. Choroidal tubercles. A clue to hematogenous tuberculosis. *Ann Intern Med.* 1964;60:231–241.
- Rodrigues LC, Diwan VK, Wheeler JG. Protective effect of BCG against tuberculous meningitis and miliary tuberculosis: a meta analysis. *Int J Epidemiol.* 1993;22:1154–1158.
- 136. Leyten EM, Lin MY, Franken KL, et al. Human T-cell responses to 25 novel antigens encoded by genes of the dormancy regulon of *Mycobacterium tuberculosis*. *Microbes Infect*. 2006;8: 2052–2060.

- 137. Murray RA, Mansoor N, Harbacheuski R, et al. Bacillus Calmette Guerin vaccination of human newborns induces a specific, functional CD8+ T cell response. *J Immunol.* 2006;177:5647–5651.
- Kaufmann SH, Hussey G, Lambert PH. New vaccines for tuberculosis. Lancet. 2010;375:2110–2119.
- Parida SK, Kaufmann SH. Novel tuberculosis vaccines on the horizon. Curr Opin Immunol. 2010;22:374–384.
- Miyoshi I, Daibata M, Kuroda N, Taguchi H, Enzan H. Miliary tuberculosius not affecting the lungs but complicated by acute respiratory distress syndrome. *Intern Med.* 2005;44:622–624.
- Kim DK, Kim HJ, Kwon SY, et al. Nutritional deficit as a negative prognostic factor in patients with miliary tuberculosis. *Eur Respir J*. 2008;32:1031–1036.
- Wilcox PA, Ferguson AD. Chronic obstructive airways disease following treated pulmonary tuberculosis. *Respir Med.* 1989;83:195–198.
- 143. Hnzido E, Singh T, Churchyard G. Chronic pulmonary function impairment caused by initial and recurrent pulmonary tuberculosis following treatment. *Thorax*. 2000;55:32–38.

- 144. Ohi M, Chin K, Tsutsui T, Fukunaga T, Kuno K. Tuberculosis sequelae: pathophysiological aspect (ventilation). *Kekkaku*. 1990;65:847–854. Japanese.
- 145. Pasipanodya JG, Miller TL, Vecino M, et al. Pulmonary impairment after tuberculosis. *Chest.* 2007;131:1817–1824.
- Kuijl C, Savage ND, Marsman M, et al. Intracellular bacterial growth is controlled by a kinase network around PKB/AKT1. *Nature*. 2007;450: 725–730.
- Loebinger MR, Janes SMJ. Stem cells for lung diseases. Chest. 2007; 132:279–285.
- 148. Mwaba P, McNerney R, Grobusch MP, et al. Achieving STOP TB Partnership goals: perspectives on development of new diagnostics, drugs and vaccines for tuberculosis. *Trop Med Int Health*. 2011;16: 819–827.

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