HIV-TB CO-INFECTION: PATHOGENESIS, DIAGNOSIS AND MANAGEMENT IN ADULTS

Salami, A.K

Department Of Medicine, College Of Medicine, University Of Ilorin, PMB 1515, Ilorin

Correspondence to: Dr. Salami. AK, E-mail: salkaz2000@yahoo.com

ABSTRACT
There is a looming epidemic of HIV/TB co-infection in tandem to the prevailing wave of HIV infection in sub-Saharan Africa and South East Asia where TB is endemic. HIV positive patients become susceptible to Mycobacterial infection following depletion of the immune cells that usually resist mycobacterial infection i.e. CD4+ and macrophages. Host's TH1 immune response to TB produces several cytokines some of which further enhance local and systemic HIV replication. DOTS remain the best option for treating TB in HIV patients. This is sometimes co-administered with antiretroviral (ARV) drugs; however, this is fraught with complex bidirectional drug-drug interactions between rifamycin component of anti-TB regimen and the protease inhibitors (PI) and non-nucleoside reverse transcriptase inhibitors (NNRTI) components of ARV. Some patients could also develop paradoxical reactions. Cytokine inhibitors could have an adjunct role to Anti-TB therapy when it roles in the pathogenesis of TB in HIV/TB is fully understood.

KEYWORDS: HIV/TB co-infection, pathogenesis, DOTS, HAART, Drug interactions

INTRODUCTION
By estimate one third of the world population is infected with Mycobacterium tuberculosis (MTB) and every day 23,000 people develop active TB, i.e. about 8.7million cases per year (1). Averagely, TB kills 2million people every year (1). Also about 40million people are HIV infected worldwide with 16,000 people being infected every day (1). TB and HIV have formed a synergy and this is facilitating the spread of a co-epidemic of HIV-TB especially in sub-Saharan Africa and South-East Asia (23). HIV positive individuals are 50 times more susceptible to MTB infection, and once infected, are 800 times more likely to develop an active TB (45). About 70% of all TB cases in sub Saharan Africa are co-infected with HIV (1), the highest in the world. This has impacted heavily on the regional TB control programme with a threat of MDR-TB especially in countries with ineffective DOTS implementation (1). The prevalence of HIV infection in Nigeria is about 5%; this is 4times lower than the estimated 20% mark of HIV prevalence at which the annual percentage increase in TB will be high at over 10% (6). However, Nigeria still has the highest annual estimate of new TB cases in Africa with about 27% of its adults infected with TB/HIV (7) while South Africa has TB as the leading cause of mortality amongst her HIV infected persons (8). According to WHO, TB accounts for 11% of all deaths in AIDS patients (9); this makes it one of the leading causes of mortality in HIV infected patients. To effectively control this growing epidemic of HIV/TB, both
HAART and DOTS should have a wider coverage. These medications will provide lifelines to millions of co-infected people. Clinicians have to be educated and updated on the complex interactions between TB and HIV as well as the dual drug-drug interactions that exist between anti-retroviral and anti-TB drugs.

**PATHOGENESIS**

*MTB* is the infectious agent of TB and it is acquired through inhalation of aerosolized droplets nuclei produced by patients with pulmonary, bronchial or laryngeal TB when they cough, sneeze, speak, or sing (10). These are the main sources of mycobacterial transmission, but smear negative PTB and extra pulmonary TB (EPTB) could also transmit the bacilli (11), especially during cough induction (12&13), irrigation of TB abscesses (14) or changing of wound dressings (15). In most infected individuals the tubercle bacilli remain dormant for years after infection before entering a phase of exponential multiplication to give active disease (16). Development of active TB is often prevented by the host's intact immune system (17), specifically the cell-mediated type (18), but this is the target of HIV infection. In TB/HIV co-infected patients therefore, there will be a steady deterioration in this protective capacity of the cell-mediated immunity till a critical point at which tubercle bacilli begin to proliferate and cause clinical disease (17). That is the balance between quiescent TB focus and the host immunity has broken down and resulted in endogenous reactivation. This occurs with the CD4+ count around 500 cells/ul or slightly higher. The ensuing reactivation often manifest as a localized pulmonary form, however, in advanced state of immunosuppression poor containment of the resulting infection could occur with resultant widespread of the bacilli causing extra pulmonary, disseminated or miliary TB (18). When the CD4+ count is less than 500 cells/ul (19) active TB could progress rapidly from primary mycobacterial infection (20&21) and some patients could have exogenous re-infection (22). The estimated annual risk of reactivation among HIV/TB patients is about 5 to 8% with a cumulative lifetime risk of 30% or more compared to a cumulative lifetime risk of 5 to 10% in HIV-negative patients (9). Early in the course of HIV infection reactivation is on the upper lobes or the upper part of the lower lobes of the lungs where ventilation is greatest; $P_{a}O_2$ of 140 mmHg (23) and lymphatic drainage is relatively impaired (24&25) but as the disease progresses it could occur at any earlier seeded site in the body.

HIV and *MTB* influence each other in a synergistic and bidirectional ways (26). Host immune response to tubercle bacilli enhances both systemic and local HIV replication and tends to accelerate the course of progression of HIV infection (26&27). Mechanisms of these interactions are now being understood. The initial contact between the host immune system and *MTB* occurs in the alveolar macrophages that present mycobacterial antigens to antigen-specific CD4+ cells (18). This is via several cytokines, which are inflammatory mediators produced by macrophages, monocytes, and lymphocytes. When these cells are sensitized by prior exposure to MTB, and then re-exposed to the same antigens, they produce several cytokines such as interferon-gamma (IFN-γ), interleukin 6 (IL-6), IL-12, and IL-18 (28,29). This is a Th1 cell-mediated immune response; it is typical for TB and other intracellular pathogens. This is in contrast to asthma and other atopic diseases whose response is Th2 cell-mediated with different cytokines, like IL-4 and IL-5 (30). Mycobacterial infected macrophage releases IL-12 and 18, these cytokines stimulate CD4+...
lymphocytes to release IFN-γ (31&32), which in turn activate more macrophages to enhance their ability to contain mycobacterial infection. The activated macrophages also release tumor necrosis factor (TNF-α), IL-1 and IL-6, it is these set of cytokines that enhance viral replication (26,33-36). Tubercle bacilli and their products also enhance viral replication by inducing nuclear factor kappa-B (NF-κB), this cellular factor binds to promoter regions of HIV (37&38) and TNF-α-induced-HIV replication is mediated predominantly through the increased activation of this factor (39&40). The long terminal repeat (LTR) of HIV contains 2 NF-κB sites, and NF-κB, either alone (41) or in concert with other transcription factors (42), is critical to the transcriptional activation of HIV. Activation of mitogen-activated protein (MAP) kinase pathway has also been implicated in the increased HIV replication (43). In particular, the p38 MAP kinase pathway has been found to be critical in HIV replication in both CD4+ cells (43) and macrophages (44). IL-1 β and TNF-α activate p38 MAP kinase and the HIV-1 LTR (45), and these cytokines are up-regulated by the MTB infection of mononuclear phagocytes (46&47). β-chemokine, monocyte chemotactic protein (MCP-1) are also known to play active roles in enhancing HIV replication (48). The recovered broncho-alveolar fluid from TB affected lung has demonstrated local increase in HIV replication by containing higher level of viral load compared to the unaffected segment of the lung and this correlated with TNF-α suggesting local production of the virus (49). The local immune activation against TB also favours the development of latent HIV infection in the macrophages and dendritic cells, thereby potentially enhancing dissemination of HIV (50&51). Thus in HIV-infected persons with active TB, the active sites of TB infection act as epifoci of increased HIV replication and evolution independent of systemic HIV disease activity (50). The resulting HIV viraemia will deplete immune cells that play a central role in anti-mycobacterial defenses (52); such CD4+ lymphocytes, macrophages and monocytes. MTB is characterized by delayed-type hypersensitivity reaction and granuloma formation in the infected tissues. Resolution of this granuloma is controlled by both cell-mediated immunity and delayed-type hypersensitivity reaction, both of which are often accompanied by some level of tissue destruction (24). Cell-mediated immunity controls TB by activating macrophages to kill ingested bacilli while delayed-type hypersensitivity causes caseous necrosis that result in killing of bacilli-laden macrophages (53). Some of the granulomas may undergo necrosis and sloughs off forming cavities others may heal with fibrosis and some may calcify (54). The extent of necrosis and cavitation in HIV/TB patients is dependent on the relative efficacy of each of these two immunologic processes in inhibiting multiplication of MTB. However, both processes are reduced in this circumstance.

PATTERN OF TB IN HIV INFECTION: The location and pattern of distribution of TB in HIV/TB patients is a measure of their level of immunity (18). In the earlier stages of HIV disease the clinical features is typical and similar to that seen in HIV negative patients. The manifestation is often pulmonary with infiltrates and cavitations in the apical posterior segments of the upper lobe and the superior segment of the lower lobe of the lungs. As the level of immunosuppression increases the presentation becomes atypical resembling primary TB with interstitial non-cavitary lesions because of poor granuloma formation and these involve more of the lower lung fields (55). At the terminal stage of HIV/AIDS extra pulmonary presentation
involving single or multiple sites are commoner along with miliary and disseminated disease i.e., involvement of two or more non-contagious sites.

**DIAGNOSIS**

Definitive diagnosis of TB in HIV infected patients requires the isolation and identification of *MTB* from the culture of the infected tissue or fluid; a presumptive diagnosis is often made from microscopic observation of acid-fast bacilli (AFB) in the stained smear of sputum (56). AFB are rod shaped organisms with large amount of lipid in their cell walls making them difficult to stain but once stained resist decolourization even when washed with 95% alcohol containing 3% hydrochloric acid, thus the characteristic acid-fast property (57&58). Laboratories diagnosis is by Ziehl-Neelsen (ZN) or Kinyoun or Tan Thiam Hok staining procedures all of which utilize carbol-fuschin (57). The former is heat fixed while the latter two are cold staining methods that require increased concentration of phenol in the staining solution. ZN staining with light microscope is the most commonly used methods of the three, it is however, time consuming and have low sensitivity requiring at least 10 (5) of tubercle bacilli per ml of specimen for reliable routine diagnosis (58). However, auramine-rhodamine staining technique with fluorescence microscope is a much faster and sensitive alternative. Properly collected sputum smears that fail to demonstrate AFB do not exclude the diagnosis of TB; because post-primary TB, the main source of infection, or re-infection in a given population is smear positive in about 50% of cases (3&59) while primary and miliary TB are smear positive in less than 25% of the cases (60&61). These percentages decrease further in the HIV-seropositive population because of their lower propensity to develop cavitary disease. Invasive procedures such as bronchoscopy with transbronchial biopsy may be necessary to establish the diagnosis of TB in them because of this high rate of smear negative PTB and increasing cases of extra pulmonary TB (EPTB) as well as other opportunistic diseases that may resemble TB in presentation (62-64). *MTB* can be differentiated from other mycobacteria that could equally infect HIV/AIDS patients by culture (65), but the commonly used Lowenstein-Jensen culture agar requires 4–8 weeks for adequate growth to allow identification (66). Recent improvement in methods of mycobacterial specification is with the use of radiometric technique (BACTEC method). The technique uses radio labeled palmitic acid, a substrate that is metabolized to released (14) CO₂, which is quantified to identify presence and growth of the mycobacteria (66&67). The BACTEC system allows detection of *MTB* growth with a mean detection time of 7–13 days for smear-positive and 14–22 days for smear-negative sputum specimens (66). Rapid diagnosis of TB is also possible with molecular amplification and identification of *MTB* specific DNA or ribosomal RNA sequences by polymerase chain reaction (68). For epidemiologic purposes, patterns of infection within a population could be studied, with identification of the points of transmission by restriction fragment length polymorphism also referred to as "DNA fingerprinting," this is a molecular biology technique that allows differentiation of unrelated strains of *MTB* by demonstration of nucleotide sequence differences at selected sites in their DNA genome (69).

WHO recently advocated the screening of all TB patients for HIV infection, but this has not been universally accepted by the clinicians especially in the developing countries for reasons that include; increase in the cost of patients care (70), regional variation in the prevalence of the TB/HIV co-infection (7) and the fact that DOTS if properly
implemented is effective in curing TB in most patients regardless of their HIV status (17). However, offering voluntary counseling and HIV testing (VCT) to TB patients is beneficial because early diagnosis of HIV infection in TB patients has been associated with a good prognosis in terms of TB cure and it minimizes the negative effect of TB on the course of HIV (71). Patients could as well plan for the future. HIV co-infection should therefore be suspected in TB patients with history of risky life style, or in a TB patient that does not show prompt sputum conversion while on standard anti-TB regimen. It should also be considered in TB patients with chronic diarrhea or mucocutaneous lesions such as or oral thrush, multidermatomal herpes zoster, non specific generalized dermatitis or co-existing sexually transmitted disease. Patients with extra pulmonary or disseminated TB should also be evaluated for dual infection. These categories of TB patients should be offered VCT.

**TREATMENT**

Short course of anti-TB chemotherapy is as effective in HIV positive TB patients as it is in the HIV negative ones, sputum conversion is rapid and cure rate is good (72). However, drug compliance is often very poor (73) and this could encourage emergence of potentially incurable multidrug-resistant TB (MDR-TB). This is however, common in poorly managed TB control programmes, especially those that lack the basic elements of good control (74). (prevalence of MDR-TB in Nigeria is 1.7% (7)). Successful treatment of TB in HIV patients therefore, depends on early diagnosis and effective application of DOTS where anti-TB will be free and patients observed to swallow each pill. In order to increase patients’ easy access to free ARV drugs the existing local DOTS infrastructure could be utilized to deliver both medications in a DOT-HAART programme (75).

However, where DOTS is not fully feasible, self-supervised patient-centered care should be encouraged to forestall non-compliance. Current guidelines (74&76) recommend that HIV infected patients with drug susceptible TB be started on the standard six-month regimen of four drugs and that treatment should be initiated with isoniazid (INH), rifampicin/ rifabutin, pyrazinamide and ethambutol, all in mg/kg body weight for the first 2 months followed by rifampicin and INH for the subsequent 4 months. Intermittent anti-TB drug administration is not advisable in the management of HIV/TB and if it is to be adopted at all, the thrice weekly regimen should only be tried during the continuation phase of therapy (77). However, daily drug intake is preferred to the intermittent regimen so as to prevent development of MDR-TB (74&76). Response of HIV/TB patients to this 6-month therapy has been found to be good with similar recurrent rate to that of HIV-negative patients (78&79): A higher recurrence rate was however, found in one report (80), but this was ascribed to re-infection rather than treatment failure. A multi-centered study may be needed to compare the outcome of short course anti-TB management in HIV patients with early and well advanced disease (81). At the interim, prolongation of the continuation phase to 7 months to make a total of 9 months of treatment has been suggested if there is evidence of a delayed clinical or bacteriological response to therapy (74&76). The practice of extended post-treatment INH therapy to prevent recurrence of TB in HIV patients has not been widely accepted though some have found it effective (82). The same principles of treatment for PTB in HIV infected adults also apply to EPTB. The drug regimens and treatment durations are the same (76&83). However, for certain forms of EPTB, such as TB meningitis, bone, or joint, using rifamycin-based regimens for at least 9 months is
generally recommended (76&83). A non-rifampicin based anti-TB regimen comprising INH, pyrazinamide and ethambutol is generally not recommended for treatment of HIV-related TB, but in cases of rifampicin intolerance, severe allergy or toxicity, this regimen should be administered daily for 18 months (83).

In order to limit mortality from HIV/TB co-infection anti-TB drugs may have to be co-administered with antiretroviral (ARV) drugs in those with advanced disease (84). However, treatment of TB should be given priority if possible to avoid drug induced hepatic reactions from the dual potential hepatotoxic combinations. Initiation of ARV should be based on CD4+ count and risk of disease progression (74&76). There is yet to be a consensus on the time to introduce ARV. Tables 1 and 2, show that commencement of ARV in HIV/TB should be individualized and balanced between potential overlapping drug toxicities, drug-drug interactions and increasing TB morbidity from immune reconstitution reactions, against the reciprocal beneficial effect each therapy have on each other, which is to the benefit of the patients (74&76). HAART improve immune responses to TB and reduces the risk of relapse and re-infection, while anti-TB drugs results in quick lowering of the viral load thereby reducing the rate of CD4+ cells loss (85&86). Good as this combination therapy may sound, it is associated with bi-directional pharmacokinetic interactions between the rifamycin components of anti-TB regimen and the protease inhibitors (PIs) and non-nucleoside reverse transcriptase inhibitors (NNRTIs) components of antiretroviral therapy (87). Therefore diligent consideration of choice of drugs becomes imperative to prevent or minimize these drug-drug interactions. PIs and NNRTIs may inhibit or induce hepatic cytochrome (CYP-450) isoenzymes and thus alter the serum concentration of rifamycins (87). Rifamycins in turn can induce CYP-450 and therefore substantially decrease blood levels of the ARV drugs too Rifampicin is the most potent CYP-450 inducer of all the rifamycins (88), followed by rifabutin, rifapentine has an intermediate activity (89&90). Only rifampicin is currently available in Nigeria. All the PIs inhibit CYP-450 (91&92) with ritonavir as the most potent and saquinavir as the least potent while indinavir, nelfinavir have intermediate inhibitory properties. The available NNRTIs have diverse effects on CYP-450: nevirapine is an inducer, delavirdine is an inhibitor, and efavirenz is both an inducer and an inhibitor (92&93). In contrast to the PIs and the NNRTIs, the other class of ARV drugs, NRTIs; zidovudine, didanosine, zalcitabine, stavudine, and lamivudine are not metabolized by CYP-450, therefore, concurrent use of NRTIs and rifamycins is not contraindicated. Also, no contraindication exists for the use of NRTIs, NNRTIs, and PIs with INH, pyrazinamide, ethambutol, or streptomycin. These first-line anti-TB medications, in contrast to the rifamycins, are not CYP-450 inducers or inhibitors. Commonly prescribed ARV for co-infected patients in Nigeria includes zidovudine or stavudine with lamivudine and efavirenz, other approved combinations are also possible as most of the HAART drugs are now available in the country. Efavirenz is contraindicated in pregnant women because of it teratogenicity. Squanvir / ritonovir or abacavir with stavudine or zidovudine and lamivudine are the alternative to it.

Adjunct therapy with cytokine inhibitors may have a role in the management of HIV/TB co-infection (94) to limit HIV replication before initiation of ARV. Thalidomide, a specific TNF-α inhibitor and pentoxifylline a nonspecific inhibitor have been tried (95). However, inhibition of TNF-α was associated with profound immune defects
akin to that of advanced AIDS and these predisposed patients to reactivation of old TB, which progressed to disseminated form (95). Full understanding of the role of cytokines inhibition in the pathogenesis of reactivation of TB in HIV/TB is therefore required. Good nutrition, including food supplements will serve as essential adjuncts to anti-TB chemotherapy.

PATIENTS MONITORING. All HIV/PTB patients should be monitored by sputum smear microscopy during treatment and if available, sputum culture and susceptibility testing. HIV-EPTB should also be monitored, but the frequency and types of evaluations will depend on the involved sites and the ease with which specimens can be obtained from these sites (81). In resource limited countries, a monthly clinical assessment has been recommended for three months followed by 3-6 monthly clinical evaluation (96) to monitor improvement, ARV adherence and to identify possible drug reaction (96). Clinical and laboratory assessment should be more frequent for patients with underline liver disease. In most developing countries viral load measurement is unavailable and monitoring of therapy is by regular clinical assessment for signs of disease regression, increasing body weight and rising CD4+ count (97). Laboratory monitoring is prioritized by observing the trimmed down WHO guidelines for HIV treatment in poor countries (98). These are inexpensive tests that have been divided into basic and desirable tests and include haemoglobin, white cells and total lymphocyte count, liver enzymes and CD4+ cells. These should be done every 3-6 months. Serum amylase, bilirubin and hoids are also desirable but optional (98). Some HIV/TB patients could experience temporary exacerbation of symptoms and signs of TB, and some may show worsening of the radiographic features of TB at the beginning of anti-TB treatment. This phenomenon is termed a paradoxical (or immune reconstitution) reaction (96). It could also occur among HIV negative TB patients, but it is commoner amongst HIV/TB co-infected while on HAART (99). Features of a paradoxical reaction include high fevers, increase in size and inflammation of involved lymph nodes, new lymphadenopathy, expanding central nervous system lesions, worsening of pulmonary parenchymal infiltrations, and increasing pleural effusions (99). A reaction that is not severe should be treated symptomatically with non-steroidal anti-inflammatory agents without a change in anti-TB or ARV therapy (100&101). Those with severe reactions (e.g. airway compromise from enlarging lymph nodes, enlarging serosal fluid collections, and sepsis syndrome) may benefit from the prednisone or methylprednisolone 1mg/kg body weight and gradually reduced after 1-2 weeks (100&101).

CHEMOPROPHYLAXIS.
Chemoprophylaxis against TB in HIV positive patients may not be strongly advocated in a developing country like Nigeria where TB is an endemic problem. The value of a positive tuberculin test may also be difficult to determine because BCG is routinely administered at birth. In addition INH prophylaxis may be abuse and this could encourage INH-resistance when decision is finally made to treat active TB.
### Table 1. WHO. Suggested timing of HAART in HIV/TB co-infection (WHO- Dec 2003)

<table>
<thead>
<tr>
<th>CD4+ count</th>
<th>Recommended regimen</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD4+&lt;200/mm³</td>
<td>Start TB treatment as soon as possible</td>
<td>Start ARV as soon as anti-TB is tolerated. EFV is contraindicated in pregnant women.</td>
</tr>
<tr>
<td>CD4+ 200-350/mm³</td>
<td>Start TB treatment</td>
<td>Start ARV after intensive phase of anti-TB.</td>
</tr>
<tr>
<td>CD4+&gt;350/mm³</td>
<td>Start TB treatment</td>
<td>Defer ARV*</td>
</tr>
<tr>
<td>CD4+ not available</td>
<td>Start TB treatment</td>
<td>Consider ARV**</td>
</tr>
</tbody>
</table>

*Unless non-TB stage IV conditions are present. Otherwise start ART upon completion of TB treatment.**If no other signs of immunodeficiency are present is improving on TB treatment, ART should be started upon completion of TB treatment.

### Table 2. BHIVA. Suggested timing of HAART in HIV/TB co-infection

<table>
<thead>
<tr>
<th>CD4+ count cells/uL</th>
<th>When to treat with HAART</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100</td>
<td>As soon as possible-dependent on physician assessment, [Some physicians delay up to 2 months]</td>
</tr>
<tr>
<td>100-200</td>
<td>After 2 months of TB treatment</td>
</tr>
<tr>
<td>&gt;200</td>
<td>After completing 6 months TB treatment*</td>
</tr>
</tbody>
</table>

http://www.bhiva.org * BHIVA treatment guidelines for TB/HIV infection – February 2005
REFERENCES


4. WHO. TB/HIV. Available at: http://www.who.int/tb/hiv/en/

5. WHO. TB/HIV co-infection. Available at: http://www.cdc.gov/nchstp/tb/pubs/MTB_HIVco_infection/default.htm


7. Global TB control-surveillance, planning, financing. WHO. 2005


37. Lederman MM, Georges DL, Kusner DJ et al. MTB and its purified protein derivative activate expression of the human


55. Perlman DC, el-Sadr WM, Nelson ET et al. Variation of chest radiographic patterns in pulmonary TB by degree of human immunodeficiency virus-related


77. CDC. Acquired rifamycin resistance in persons with advanced HIV disease being treated for active TB with intermittent rifamycin-based regimens. MMWR 2002;51:214-215.


81. CDC. 2004: Treatment of Pulmonary Tuberculosis Among HIV-Infected Adults and Adolescents; available at: http://www.cdc.gov/mmwr/preview/mmwrhtml/mm5315a1.htm


90. Durand DV, Hampden C, Boobis AR et al. Induction of mixed function oxidase activity in man by rifapentine (MDL 473), a long-acting


98. WHO. Antiretroviral newsletter Issue no.4. Clinical and laboratory monitoring of antiretroviral therapy in resource-limited and unlimited settings. Available at: http://www.wpro.who.int/NR/rdonlyres/DFC8 8S4E-1EA8-4B06-B04A- 4E05797F17EE0/ART. Newsletter Issue 4.pdf

