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Latent Tuberculosis: Advances in Diagnosis and Treatment

Dimitrios Basoulis, Georgia Vrioni, Violetta Kapsimali, Aristeidis Vaiopoulos and Athanasios Tsakris
Medical School of the National and Kapodistrian University of Athens
Greece

1. Introduction

Tuberculosis (TB) is one of the oldest diseases known to affect humans. It is caused by bacteria belonging to the *Mycobacterium tuberculosis* complex and strains of these bacteria have been found in human bones dated from the Neolithic era. It was known to the ancient Greeks, Indians and the Inca, making it a disease with a global distribution even from ancient times. Latent tuberculosis infection refers to a time period where the host has been exposed and infected by the bacteria yet does not exhibit any signs or symptoms of infection. It is estimated that one third of the world, almost 2 billion people suffer from latent tuberculosis infection.

2. Epidemiology

Tuberculosis is a multisystemic infection with myriad presentations and manifestations. According to the World Health Organization (WHO) it is estimated that one third of the world’s population is currently infected by the bacillus and out of those people 5-10% will exhibit symptoms at some point during their life. WHO estimates that the largest number of new TB cases in 2008 occurred in the South-East Asia Region, which accounted for 35% of incident cases globally. However, the estimated incidence rate in sub-Saharan Africa is nearly twice that of the South-East Asia Region with over 350 cases per 100 000 population (WHO, 2011). Tuberculosis remains the most common cause of infectious disease related mortality worldwide. It is evident by this alone that latent tuberculosis is a serious public health problem, not only due to the possibility of the patients themselves eventually developing active tuberculosis, but also because of the public health risk that they impose.

*M. tuberculosis* is most commonly transmitted from a patient with infectious pulmonary tuberculosis via droplet nuclei, aerosolised by coughing, sneezing or even speaking. The tiny droplets dry rapidly, but the smallest of them (<10μm in diameter) can remain suspended in the atmosphere for several hours. When inhaled, these droplets can reach the terminal airspaces of the lung. Risk factors for transmission include the proximity of contact, the duration of contact, the degree of infectiousness of the case and the shared environment of the contact. It needs to be noted that patients that have sputum smear negative and culture positive tuberculosis are less infectious, whereas patients with culture negative
sputum pose essentially no risk for transmission. It is estimated that up to 20 people can be infected by a single patient before tuberculosis can be identified in high prevalence countries. Transmission is more common in tightly packed populations (i.e. overpopulated areas, military personnel etc.) in countries with a higher incidence.

It has been demonstrated that large clusters of TB are associated with an increased number of tuberculin skin test-positive contacts, even after adjusting for other risk factors for transmission. The number of positive contacts was significantly lower for cases with isoniazid-resistant TB compared with cases with fully-susceptible TB. This result has been interpreted to imply some connection between isoniazid resistance and mycobacterial virulence (Verhagen et al., 2011).

After exposure to the bacteria, the patient has a 5-10% chance of developing active tuberculosis. Risk factors that determine this progression include age, the individual's innate susceptibility to disease and level of function of cell-mediated immunity. Clinical illness directly following infection is classified as primary tuberculosis and is more common in children. The majority of patients infected will develop disease within a year while the rest will develop latent tuberculosis. Activation of tuberculosis bacilli at any point thereafter is termed secondary tuberculosis. Several diseases predispose the patient to develop active tuberculosis with chief amongst them HIV co-infection. It is estimated that nearly all of infected individuals that are HIV positive will at some point develop active tuberculosis; this risk depends on the level of immunosuppression and the CD4+ cell count of the infected patient. Patients with diabetes have 2-5 times increased risk for developing active disease, whereas the relative risk for patients with chronic renal failure climbs to 10-25.

3. Pathophysiology of tuberculosis infection

Two models for the pathophysiology of tuberculosis infection and the formation of granulomas have been suggested. The first one is the static model and it is considered to be the traditional one. The second was suggested a few years ago and it is the dynamic model of infection.

3.1 The static model

Mycobacteria belong to the family Mycobacteriaceae and the order Actinomycetales. The most important member of the Mycobacterium tuberculosis complex is the namesake organism, Mycobacterium tuberculosis. The complex also includes M. bovis (the bovine tubercle bacillus), M. africanum (isolated from cases in West, Central and East Africa). M. microti (a less virulent rarer bacillus), M. pinnipedii and M. canettii (very rare isolates). M. tuberculosis is a slow-growing, obligate aerobe and obligate pathogen. Most often, it is neutral on Gram's staining, however, once stained, the bacilli cannot be de-colorised by acid alcohol, hence the characterization as acid-fast and the reason they are best seen using the Ziehl-Neelsen stain. This ability of mycobacteria is derived from the high content of mycolic acids, long chain fatty acids and other lipids found in abundance in the cell wall of mycobacteria (Harada, 1976; Harada et al, 1977). In the mycobacterial cell wall, lipids are linked to underlying arabinolactan and peptidoglycan, which confers a high resistance to antibiotics due to low permeability of this structure. Another element of the cell wall structure is the lipoarabinomannan which is crucial to the mycobacterium's survival within
the host's macrophages. All of these proteins, characteristic of *M. tuberculosis* are included in the purified protein derivative (PPD, a precipitate of non-species-specific antigens obtained from filtrates of heat-sterilised, concentrated broth cultures.

The majority of inhaled bacilli are trapped at the level of the upper airways and expelled. A small fraction (<10%) will descend further down the bronchial tree. When the inhaled droplet nuclei reach the terminal airspaces of the lung, the bacilli, transported with the droplets, begin to grow for 2-12 weeks before any immune response from the host can be elicited. The host's immune system responds when the bacillary load reaches 1000-10,000 cells. Non-specifically activated alveolar macrophages will eventually begin to ingest the bacilli and sequester them from the host.

Phagocytes have 2 methods of dealing with the mycobacteria. Fusing the phagosomes containing the mycobacteria with lysosomes they create phagolysosomes. Phagolysosomes are the product of a fusion-fission process between the lysosomes, the phagosomes and other intracellular vesicles. The Ca$^{2+}$ signalling pathway and recruitment of vacuolar-proton transporting ATPase (vH+-ATPase) lead to a decrease in the pH of the phagolysosome, that in turn allows acid hydrolases to function efficiently for their microbicidal effect. Another way that phagocytes deal with the mycobacteria is through ubiquitination of mycobacterial cell wall and membrane components, which in turn leads to increased susceptibility to nitric oxide produced by the phagocytes. This process leads to phagocyte apoptosis (Beisiegel et al 2009; Bermudez & Goodman, 1996; Chan & Flynn, 2004; Cooper, 2009; Pieters, 2008; Ahmad, 2010).

This form of defence, however, proves inefficient as the bacilli have the ability to survive inside the macrophages by modulating the behaviour of its phagosome, preventing its fusion with acidic, hydrolytically-active lysosomes (Pieters, 2008; Russel et al 2009) The escape of *M. tuberculosis* from macrophage destruction is dependent on the 6-kDa early secreted antigenic target (ESAT-6) protein and ESX-1 protein secretion system encoded by the region of difference 1 (RD1). The ESAT-6 protein associates with liposomes containing dimyristoylphosphatidylcholine and cholesterol and causes destabilization and lysis of liposomes. It can also infiltrate the phagosome's membrane and cause lysis of the phagosome, enabling the mycobacteria to escape (Brodin et al, 2004; de Jonge et al, 2007; Derrick & Morris, 2007; Kinhikar et al, 2010).

In this initial stage of interaction, either the macrophages manage to contain the bacillary reproduction through sequestration and production of cytokines and proteolytic enzymes, or the bacilli manage to survive and multiply, leading to macrophage lysis. Through chemotaxis, monocytes arrive at the site of infection to ingest the bacilli after the macrophage lysis. Either through lysis or apoptosis the mycobacterial antigens are exposed and presented to T lymphocytes that will carry out the burden of the host's immune response orchestration.

Following these events, the host's immune system activates two more mechanisms to battle the invading bacteria: a tissue damaging response and a macrophage activating response. The tissue damaging response is a delayed-type hypersensitivity reaction to bacillary antigens leading to the destruction of “infected” macrophages. The macrophage activation focuses on activating specific macrophages to ingest and destroy the bacteria. Local macrophages are activated when the non-specific macrophages present bacillary antigens to
T lymphocytes, stimulating them to release lymphokines. Depending on which one of the two mechanisms is predominant, the subsequent form of tuberculosis is determined.

If the macrophage activation predominates, large numbers of activated macrophages arrive at the site of infection and granulomatous lesions begin to form. During this early stage and under the influence of a vascular endothelial growth factor (VEGF), the granuloma becomes highly vascularised which in turn will provide the pathway for the lymphocytes and macrophages to arrive at the site (Alatas F et al, 2004). Once there, the macrophages will further differentiate into different cells such as multi-nucleated giant cells, epithelioid cells and foamy macrophages. These cells will form the outer wall of the granuloma, now dubbed tubercle. The structure becomes much more stratified and a fibrous cuff forms outside the macrophage layer. Lymphocytes move away from the centre and aggregate outside this fibrous layer (Cáceres et al 2009).

The tissue damaging response on the other hand leads to destruction of macrophages that fail to contain the bacilli and in turn creates a necrotic area at the centre of the tubercle with dead macrophages. Due to low oxygen, presence of nitric oxide, nutrient deficiency and very acidic pH the mycobacteria cannot continue to multiply inside the tubercle centres, yet they can survive and remain dormant (Ahmad, 2010; Ohno et al, 2003; Voskuil et al, 2003). The central necrotic region resembles cheese in texture and has granted the name caseous necrosis to this process. At this point, some of the tubercles calcify and heal while others evolve further.

Two distinct types of granulomas have been identified. The classic caseous granulomas are composed of epithelial macrophages, neutrophils, and other immune cells surrounded by fibroblasts. *M. tuberculosis* resides inside macrophages in the central caseous necrotic region. The second type of granulomas (fibrotic lesions) is composed of mainly fibroblasts and contains very few macrophages. The exact location of viable *M. tuberculosis* in these lesions is not known (Barry et al, 2009). It needs to be noted that even the healed, fibrotic tubercles can still contain mycobacteria in a dormant state.

It has been suggested that the caseating centre of the granuloma is not the site where the host's immune response is organized and maintained, but rather that site is at the outer layers of the tubercle, where the macrophages can present their antigens to the lymphocytic population of the tubercle. This formation resembles a secondary lymphoid organ and is theorised to be better suited to orchestrate the host's immune response, as suggested by the high proliferative activity only observed in peripheral follicle-like structures (Ulrichs et al, 2004).

If the tissue damaging response predominates, due to a weak response from the macrophages, the initial lesion cannot be contained and continues to grow at the expense of the surrounding tissue. Bronchial walls and blood vessels are destroyed in this process (hence why haemoptysis is a chief symptom in rampant tuberculosis) and cavities are gradually formed (Zvi et al, 2008).

The mycobacterial cell wall components are recognized by host receptors that include toll-like receptors (TLRs), nucleotide-binding oligomerisation domain (NOD)-like receptors (NLRs), and C-type lectins, including mannose receptor (MR), the dendritic cell-specific intercellular adhesion molecule grabbing nonintegrin (DC-SIGN), macrophage inducible C-type lectin (Mincle) and dendritic cell-associated C-type lectin-1 (Dectin-1). The TLR
signalling is the main arm of the innate immune response and *M. tuberculosis* phagocyted through different receptors may have a different fate (Harding & Boom, 2010; Ishikawa E et al, 2009; Jo, 2008; Jo et al, 2007; Noss et al, 2001).

Cell mediated immunity, more specifically macrophages and CD4+ T lymphocytes, plays a very important role in the above process. The infected macrophages produce a host of cytokines: Interleukin 1 (IL1) which leads to the development of fever, interleukin 6 (IL6) which leads to hyperglobulinemia and tumour necrosis factor α (TNF-α) that contributes to the killing of mycobacteria, the formation of caseating granulomas, fever and weight loss. As mentioned earlier, non-specific macrophages are also responsible for presenting the bacillary antigens to the T cells and eliciting their response (Khader & Cooper, 2008; Kursar et al 2007). Activated T helper Type 1 lymphocytes participate in the destruction of infected cells through an MHC class II restricted process. They also produce interferon γ (IFN-γ) and interleukin 2 (IL2) and promote cell-mediated immunity. Once the bacillary growth is stabilized, the presence of CD8+ T cells appears to gain importance, both for the production of IFN-γ and an increase in the cytototoxic activity. This is a period of stalemate where the bacillary load remains relatively constant and the infection is in a state of latency (Bodnar et al, 2001; Russel et al, 2009).

More recently, it was demonstrated that IL1-beta, a subset of interleukin 1, which plays an important part as mediator in the host’s immune response, is induced when ESAT-6 is secreted from the bacilli. IL1-beta is activated through the inflamasome, a caspase activating protein complex. Caspases are cysteine-aspartic proteases that play a part in inflammation response and apoptosis. Mycobacteria have developed the ability to halt the inflamasome’s formation by secreting a Zn²⁺ metalloprotease, encoded by the zmp1 gene. Mycobacteria genetically modified for zmp1 deletion and through the secretion of ESAT-6 lead to IL1-beta activation and elicit a stronger immune response from the host leading to improved mycobacterial clearance by macrophages, and lower bacterial burden in the lungs of aerosol-infected mice (Danelishvili, 2010; Lalor, 2011; Master 2008; Mishra, 2010). Mycobacteria secrete their own enzymes (Rv3654c and Rv3655c) within the macrophage cytoplasm with the ability to cleave caspase-8. In this manner, the bacilli prevent macrophage apoptosis by preventing the inflamasome’s formation and promote cellular lysis (Danelishvili, 2010). It has been demonstrated that it is more beneficial to bacterial growth if the macrophages are steered towards lysis as opposed to apoptosis. Necrosis was correlated with Caspase 3 activity and bacterial growth, whereas activation of calcium, TNF-alpha and Caspase 8 was associated with apoptosis and decreased bacterial load (Arcila et al, 2007).

Humoral immunity seems to play a much lesser role if any. The evidence that B-cells and *M. tuberculosis*-specific antibodies can mediate protection against extracellular *M. tuberculosis* is highly controversial as their contribution is probably of minor importance (TBNET, 2009).

The host’s immune response can eventually cause more problems through tissue destruction and uncontrolled activation of macrophages and lymphocytes. For this reason there is a negative feedback mechanism in place, to control the extent of the response. A family of receptor tyrosine kinases provide this negative feedback mechanism to both, TLR-mediated and cytokine-driven proinflammatory immune responses (Liew, 2005). Again, the mycobacteria have developed mechanisms to take advantage of this process in order to halt the immune response to their benefit. Several *M. tuberculosis* cell wall components or protein
products such as 19-kDa lipoprotein, glycolipids (particularly Man-LAM), trehalose dimycolate (cord factor) can modulate antigen-processing pathways by MHC class I, MHC class II and CD1 molecules, phagolysosome formation and other macrophage intracellular signalling pathways (Ahmad, 2010; Bowdish et al, 2009; Gehring et al, 2004; Harding & Boom, 2010; Jo et al, 2007; Nigou et al, 2001; Noss et al, 2001; Pecora et al; 2006). This results in a subset of macrophages that are unable to present mycobacterial antigens to T lymphocytes.

It is hypothesized that the infection sustains itself not through replicating bacilli forming equilibrium with those being destroyed by the host's immune system, but through a population of non-replicating bacilli that can withstand the immune response. The evidence to this is indirect, suggested by the lack of cellular debris in the granuloma centres of infected mice (Rees & Hart, 1961). It is believed that the host's immune response is driven by antigens produced during active multiplication of the bacilli and thus, those that remain dormant would not sustain that response to its maximum potential (Andersen, 1997).

### 3.2 The dynamic model

More recently a dynamic model of infection was proposed able to give some logical explanations to some short-comings of the static model. The first question posed was how it is possible for the mycobacteria to remain dormant in the tubercle environment when the host is trying to re-structure the damaged tissues. The alveolar macrophages have a lifespan of 3 months, yet according to the static model, they exist in stalemate with the mycobacteria for a much longer period of time, whether as part of the middle layer of the granuloma or as part of the caseous centre having phagocyted bacilli and sustaining them in their dormant state (Cardona, 2009).

The second question was how did the bacilli reactivate themselves from their dormant state, as it has been demonstrated that the resuscitation factors necessary for this are only produced by active bacilli (Cardona, 2009; Shleeva et al, 2002).

The third question posed seeks an explanation based on a physiological model regarding the ability of isoniazid to treat latent tuberculosis when it is known that isoniazid can only take effect on actively multiplying bacilli (Cardona, 2009; TBNET, 2009).

According to the dynamic model that has been suggested, the granulomas are not static formations but rather, inside the granuloma, there exists a balance between inactive dormant bacilli, rapidly multiplying ones, dying bacilli and cellular debris constantly being removed from the site (TBNET, 2009). The exact nature of the metabolic state of mycobacteria within the macrophages in the granuloma is a matter of great debate and investigation.

The size of the actively multiplying mycobacterial load in the granuloma determines the antigen-specific re-stimulation of memory T lymphocytes. On the other hand, if the mycobacteria are mostly contained within macrophages in their dormant state, it is more likely that T cell immunity will begin to decline. This in turn would explain why a tuberculin skin test can revert to negative after exposure at a rate of about 5% per year (TBNET, 2009).
Perhaps the most important element in this proposed model is the role of the foamy cell, i.e. alveolar macrophages at the end of their life cycle and filled with lipids, due to phagocytosis of extracellular debris, mostly consisting of lipid-rich cellular membrane remains. The mycobacteria phagocytosed by these cells can survive through the mechanisms explained earlier. The dynamic model suggests that the mycobacteria can continue to grow albeit at very slow rates instead of becoming dormant. The slower metabolic rate provides resistance to stress and reduces the nutritional needs of the bacilli, thus allowing their survival (Cardona, 2009; Muñoz-Elias et al, 2005). It has not been fully researched but evidence suggests that mycobacteria can escape the phagosomes of the foamy cells and reach the bronchial tree and become aerosolised.

Foamy cells provide a stressful environment that conditions the bacilli to become more resistant. This in turn, confers them the ability to better survive in the open air and according to some studies explains why they are more virulent. Moreover, the high lipid content of the foamy cells also provides triglycerides to the bacilli that will in turn provide them with nutrients in new infection sites in the event of starvation. In fact the highly aggressive Beijing strains have also been found to contain large amounts of lipids, which would at least partly account for the greater virulence (Garton et al, 2002; Neyrolles et al, 2006; Peyron et al, 2008). Finally the high lipid content of foamy cells when exposed to the alveolar spaces will contribute to increased surfactant concentration and thus will make aerosolisation of the bacteria easier (Cardona, 2009).

Growing bacteria are easy to combat since they cannot survive in stressful environments. The dynamic model offers a different explanation of the mechanism, with which the host's immune system focuses on the non-replicating bacteria. The phagocytosed bacilli, as explained in the static model, will eventually lead to lysis or apoptosis of the macrophages. This cellular debris and the extracellular bacteria will form the population of the non-replicating bacteria at the caseous centre. The attraction of specific macrophages and neutrophils will provide a new breeding ground for the active bacteria and also material for the formation of the foamy cells, as they will phagocyte cellular membrane remnants to clear the debris from the caseous centre of the granuloma. The bacilli, inside the foamy cells, under these circumstances, will eventually find themselves within the bronchial spaces and after they are aerosolised they will reinfect the host at new sites. Due to their higher virulence they will manage to overcome the initial immune response and form a new granuloma to repeat the same sequence of events (Cardona, 2009). At the new site of infection the bacilli are actively multiplying again and thus are susceptible to isoniazid. This would explain why a single-drug nine month treatment is effective in most cases of latent tuberculosis.

4. Latent tuberculosis and reactivation

Mycobacteria are completely eradicated only in about 10% of the cases, while in the remaining, the bacilli survive for years to come, through the processes explained. This state has been termed latent tuberculosis infection. In any event where the host's immune response dwindles, there is a risk for the bacilli to reactivate themselves and lead to active tuberculosis infection. Most of the new cases of tuberculosis in low incidence countries are the result of such reactivation of latent tuberculosis infections. It is of interest to note that expression of DosR-regulated dormancy antigens continues even in this latent stage of infection, providing a promising new target for vaccines that would help battle latent TB
infections in the future (Leyten et al, 2006; Lin & Ottenhoff, 2008). It is also probable that \textit{M. tuberculosis}, during the latent stage of infection can form spore-like structures, typically seen with other mycobacteria, in response to prolonged stationary phase or nutrient starvation, for its survival (Ghosh et al, 2009).

The reactivation of latent infection requires \textit{M. tuberculosis} to exit dormancy. This is mainly achieved through the effects of a family of five proteins, dubbed resuscitation promoting factors (Rpfs), that have the effect of a lytic transglycosylase. These molecules were found to be able to cause degradation to cell wall components of the mycobacteria. It is not exactly known how this activity relates to the resuscitation process, it is however theorised that the end result of this enzymatic activity is changes to the mycobacterial cell-wall, overcoming the environmental restraints to the bacterial multiplication. Another theory states that the changes brought to the cell wall, lead to production and secretion of peptidoglycans with the ability to modulate the environment and the host's immune response (Hett et al, 2007; Tufariello et al, 2006). It needs be noted that \textit{M. tuberculosis} bacilli found in the sputum of patients with latent infection and after deletion of the Rpfs encoding genes, can only be cultured when Rpfs are introduced to the growth material and thus resuscitation is possible (Mukamolova et al, 2010), however for non-dormant mycobacteria it seems that the Rpfs are not important for their multiplication (Kana et al, 2008).

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**Fig. 1.** Natural progression of tuberculosis, adapted from Ahmad, 2010
It has also been demonstrated that amongst the Rpfs, those that seem to be the most important are RpfA and RpfB. Infected mice with strains of mycobacteria with deletion of the genes encoding these specific Rpfs, were found to be more resistant to TB reactivation and also their macrophages were found to produce larger quantities of TNF-α and IL6 (Russel-Goldman et al, 2008). These resuscitation factors are another possible target for future vaccines against latent TB (Zvi et al, 2008).

5. Latent tuberculosis diagnosis

Diagnosis of latent tuberculosis is a matter of active current research due to the difficulties presented in identifying patients with latent infection. There is no question that controlling contacts and identifying people who are carrying the bacilli would be the best prevention plan. However, due to the lack of any physical signs or symptoms and the fact that all or most of the bacilli during this state remain dormant, it is very difficult to elicit an immune system response that would be evident to the observer. This in turn means that it is difficult to identify individuals with latent infection. An ideal test for latent tuberculosis infection diagnosis should meet the following criteria:

- High sensitivity in all populations at risk.
- High specificity regardless of BCG vaccination and infection with environmental mycobacteria.
- Reliability and stability over time.
- Objective criteria for positive result, affordability and easy administration.
- Ability to distinguish recently infected individuals with increased risk of progression to active tuberculosis.

There are currently two groups of tests for latent tuberculosis infection diagnosis: tuberculin skin tests (TST) and interferon-γ release assays (IGRA).

5.1 The tuberculin skin test

Historically, the most accurate method for detecting if an individual had come in contact with *M. tuberculosis* was the tuberculin skin test (TST). This test measures the hosts' *in vivo* immune response in the form of a cell-mediated delayed hypersensitivity reaction to a mixture of more than 200 *M. tuberculosis* antigens, termed as purified protein derivative (PPD). The PPD is a crude mixture of antigens, not specific to *M. tuberculosis*, but also found in other mycobacteria such as the BCG bacillus, *M. bovis* and even non-tuberculous mycobacteria. This mixture is intradermally injected, usually at the inner side of the forearm and the test result is read as an induration on the site of injection after 48-72 hours (Huebner et al 1993). This reaction may last for up to 1 month, depending on the quality and quantity of the initial reaction. Strong reactions may result in tissue necrosis, which is the only absolute contraindication to the TST (TBNET, 2009). The induration is caused due to the introduction of the antigens that causes non-specific neutrophils and antigen-specific T lymphocytes to arrive at the site and spark an inflammatory cascade of cytokine production. The migration of immune cells to the site seems to have a biphasic distribution: an initial nonspecific infiltration where the neutrophils arrive at the site, taking place in the first 4-6 hours and which is an event that also occurs in nonsensitised subjects and a second specific peak, where the specific T cells arrive at the site (Kenney et al, 1987; Platt et al, 1983;
Poulter et al, 1982; TBNET, 2009). The lymphocyte population is a mix of CD4+ and CD8+ cells with the former being always greater in number (Gibbs et al, 1984). The lymphocytic infiltration is at first perivascular and under the influence of early cytokines, such as IFN-γ, TNF-α and TNF-β, the endothelium is stimulating into expressing adhesion molecules (E-selectin), increasing the permeability of the vascular walls and enabling the cells to migrate to the dermis. Regulatory T-cells influence the size of the induration of the tuberculin skin test. Cutaneous CD4 T-cells accumulating after tuberculin PPD stimulation in the skin are predominantly of a CD45 RO memory phenotype (Sarrazin et al, 2009). The criteria for the test’s interpretation vary considerably and depend on the nature of the population being tested. They are arbitrary and the result of international consensus.

In the United States, according to the Center for Disease Control (CDC), 5 tuberculin units (TUs) are used and a test is considered positive for the general population with no known TB contacts when the induration measures 15mm or more. An induration of 10 or more millimetres is considered positive in recent immigrants (<5 years) from high-prevalence countries, injection drug users, residents and employees of high-risk congregate settings, mycobacteriology laboratory personnel, persons with clinical conditions that place them at high risk, children < 4 years of age, infants, children, and adolescents exposed to adults in high-risk categories. Finally, an induration of 5 or more millimetres is considered positive in HIV-infected persons, a recent contact of a person with TB disease, persons with fibrotic changes on chest radiograph consistent with prior TB, patients with organ transplants, persons who are immunosuppressed for other reasons (e.g., taking the equivalent of >15 mg/day of prednisone for 1 month or longer, taking TNF-α antagonists, etc.) (CDC, 2011).

In Europe, the situation differs from country to country depending on the incidence and prevalence of TB. In countries with high incidence, such as former Soviet Union countries, a 10mm induration is considered positive. In most European countries 2 TUs are used and interpretation of the results follows the same guidelines as in the US (ECDC, 2011).

As with every screening test, TST has a chance of false positive and false negative results. Possible false positive reactions are caused due to infections with non-tuberculous mycobacteria, previous vaccination with BCG, incorrect method of TST administration (including wrong amount of PPD injected as well as injecting it subcutaneously rather than intradermally), incorrect interpretation of reaction (more often than many would assume, doctors and/or nurses measure the erythema caused by the immune response rather than the induration leading to overestimation of the reaction caused), incorrect bottle of antigen used. False negative results are caused by cutaneous anergy (anergy is the inability to react to skin tests because of a weakened immune system, such as in HIV patients or patients under immunosuppression, particularly those taking anti-TNF-α medications for autoimmune conditions), recent TB infection (within 8-10 weeks of exposure), very old TB infection (many years), very young age (less than 6 months old), recent live-virus vaccination (e.g., measles and smallpox), overwhelming TB disease (tuberculosis by itself is thought to cause a degree of immunosuppression to the host in these advanced cases), some viral illnesses (e.g., measles and chicken pox), incorrect method of TST administration, incorrect interpretation of reaction (ECDC, 2011; CDC, 2011).

Of special consideration is the so-called booster effect after TST testing. In certain people, who have been exposed to M. tuberculosis, the ability of their immune system to react to the PPD antigens might have diminished over the course of time. These patients when tested
with the TST would have a negative reading. However, reintroducing the tuberculosis antigens to their immune system by the test itself stimulates their immune system to react more fiercely to these antigens. Subsequent tests in these individuals would result as positive even though they haven't been exposed to the bacilli in the time between the two tests. In a sense, the first TST “boosted” the results of the second one. In certain populations, the CDC suggests performing a two-step test in order to identify possible false negative first tests and prevent unnecessary treatment. Such populations include health-care workers, doctors, nurses or nursing home residents, whose status with regards to tuberculosis exposure and/or infection is important to know.

It is evident that the TST has several limitations to its use, which in turn sparked the interest in developing new diagnostic tools such as the IGRAs. Such limitations include a high proportion of false positive and false negative results, difficulty in separating true infection from the effects of BCG vaccination and NTM infection, technical problems in administration, immune response boosting after repeated TST, complicated and subjective interpretation and a need for a second visit for the interpretation of the test's result.

5.2 The interferon-γ release assays

Interferon-γ release assay kit tests were developed the past decade as an alternative to the TST. They are whole-blood tests that can aid in diagnosing *M. tuberculosis* infection, including both latent tuberculosis infection and active disease. They are indirect *in vitro, ex vivo* tests that measure the production of interferon-γ by a patient's T lymphocytes after the latter are incubated with specific *M. tuberculosis* antigens *in vitro* (Andersen et al, 2000; Harboe et al, 1996; Mahairas et al, 1996). To conduct the test, fresh blood sample from the patient is mixed with the antigens and the response is measured either by measuring the produced interferon through enzyme-linked immunosorbent assay (ELISA), rapid enzyme-linked immunospot assay or by measuring the number of activated T cells through flow cytometry. The difference in method used is what distinguishes the two commercially available kits. QuantiFERON-TB Gold In-Tube (QFT-GIT, by Cellestis Limited, Carnegie, Victoria, Australia) uses the ELISA method and the T-SPOT (by Oxford Immunotec Limited, Abingdon, UK) uses the ELISPOT. It is interesting to mention that initially IGRAs would use the PPD as antigen but still follow the same principle and in an interesting twist of fate, it has been suggested to use the specific IGRA antigens for TST, as these antigens have been found to elicit a distinctive immune response with induration on animals. IGRAs are performed on fresh blood specimens.

The antigens used in these methods are peptides derived from ESAT-6, CFP-10 and for the Quantiferon method TB7.7 proteins of the mycobacteria. The first two are encoded at the region of difference (RD) 1 genetic locum whereas the third at the RD11, regions that are deleted from the *M. bovis* BCG genome and are absent in most environmental mycobacteria, with the exception of *M. kansasii, M. szulgai* and *M. marinum* (TBNET, 2009). During earlier stages of the method's development, the entire protein product was used. The early secretory antigenic target (ESAT) is a 6kDa protein and the culture filtrate protein (CFP-10) is a 10kDa protein. Together they form an heterodimeric complex and depend on each other for stability. They are secreted through the ESX1 secretion system and are considered to be an indication of virulence. Their role is not fully understood but they seem to induce lysis through integration on the macrophage cellular membrane (Brodin et al, 2004; de Jonge et al,
2007; Derrick & Morris, 2007; Kinhikar et al, 2010; Renshaw et al, 2005). Even less is known regarding TB7.7. IGRA techniques support the dynamic model for latent TB since they detect IFN-γ produced by T cells, with a short lifespan that have been activated by macrophages that presented to them the tuberculosis antigens (Cardona, 2009).

For the QFT-GIT (Table 1), 1 ml of blood is drawn into one of each of three special testing tubes. These are precoated and heparinised by the manufacturer. Within 16 hours the tubes must be incubated for another 16 to 24 hours at 37 °C. After centrifugation, the plasma is harvested to be further processed. QFT-GIT collection tubes contain a gel plug that separates the plasma from the cells when centrifuged. The plasma can be used immediately or at a later point in time. Results are interpreted according to the manufacturer’s recommendations (ECDC, 2011).

<table>
<thead>
<tr>
<th>Result</th>
<th>IFN-γ concentration (International Units per ml, IU/ml)</th>
</tr>
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<tbody>
<tr>
<td>Positive</td>
<td>≥ 0.35 IU/ml and ≥ 25% over nil</td>
</tr>
<tr>
<td>Negative</td>
<td>&lt; 0.35 IU/ml or &lt; 25% over nil</td>
</tr>
<tr>
<td>Indeterminate</td>
<td>&lt; 0.35 IU/ml or &lt; 25% over nil</td>
</tr>
<tr>
<td></td>
<td>Any</td>
</tr>
</tbody>
</table>

Table 1. Quantiferon results interpretation, adapted from ECDC, 2011

For the T-SPOT assay (Table 2), 8 ml of blood are required and the assay must be performed within eight hours of blood collection. Alternatively, the manufacturer also provides a reagent (T-Cell Xtend) which extends processing time to 32 hours after blood collection. The T-cell-containing peripheral blood mononuclear cell fraction is separated from whole blood and distributed to the microtitre plate wells (250,000 cells/well) provided in the assay kit. Following 16 to 20 hours incubation, the number of IFN-γ-secreting T-cells (represented as spot-forming units) can be detected by ELISPOT assay. As with QFT-GIT the test's results are interpreted according to the manufacturer’s recommendations (ECDC, 2011).

<table>
<thead>
<tr>
<th>Result</th>
<th>Spot count</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M. tuberculosis antigens</td>
</tr>
<tr>
<td></td>
<td>ESAT-6</td>
</tr>
<tr>
<td>Positive</td>
<td>≥ 6 over nil and/or ≥ 6 over nil</td>
</tr>
<tr>
<td>Negative</td>
<td>≤ 5 over nil and/or ≤ 5 over nil</td>
</tr>
<tr>
<td>Borderline</td>
<td>If for any antigen highest is 5 - 7 over nil</td>
</tr>
<tr>
<td>Indeterminate</td>
<td>≤ 6 over nil and/or ≤ 6 over nil</td>
</tr>
<tr>
<td></td>
<td>Any</td>
</tr>
</tbody>
</table>

Table 2. T-Spot results interpretation, adapted from ECDC, 2011
The presence of negative and positive controls ensures that IGRAs are correctly performed. The three testing tubes contain the mycobacteria antigens (Mtb), no antigens (Nil) and phytohaemagglutinin A (PHA), a T-cell activating mitogen. The Nil vial serves as the negative control for the process whereas the PHA as the positive one. If there is IFN-γ production in the Mtb tube, none in the Nil and any amount in the PHA, it means that the result is a positive one because it would imply that the sample's lymphocytes reacted to the antigens as expected and did not react to any other antigens that might have contaminated the sample. If on the other hand there is no IFN-γ production in the Mtb tube and the Nil tube but there is in the PHA one, it implies that the lymphocytes react normally to the PHA antigen yet they do not react when exposed to the bacilli antigens and therefore these lymphocytes haven't met these antigens before. Finally, the results are indeterminate if at any point there is IFN-γ production in the Nil tube, which might imply contamination or there is increased baseline interferon production or if there is no sufficient production in the PHA tube, which might imply anergy. Technical factors (sample collection, storage and transportation) might also contribute to returning indeterminate results (ECDC, 2011).

There is a lot of debate on whether IGRAs are indeed more reliable than the traditional TST. In Germany, Denmark and Switzerland, IGRAs have substituted TST when screening populations receiving anti-TNF-α therapies. The US, Australia, France and Denmark use either TST or IGRAs, whereas Canada, the United Kingdom, Italy, Spain, Australia and Slovakia to name a few, support a 2-step approach using both TST and IGRAs in an attempt to increase sensitivity and specificity of both methods. The two-step approach seems to be the most favoured strategy for IGRA use, especially in BCG vaccinated contacts.

IGRAs have some distinct advantages over TST with regards to diagnosing latent tuberculosis infection. IGRA testing requires a single patient visit to conduct the test and results can be available within the day. Moreover there is no “booster” effect associated with IGRAs since they are \textit{ex vivo, in vitro} tests. Finally, due to the specificity of the \textit{M. tuberculosis} antigens used, BCG vaccination does not cause false positive results. Due to the positive control, IGRAs are able to differentiate between immunocompromised hosts and negative results with more accuracy. In the TBNET/ECDC systematic review and meta-analysis (Sester et al. 2010) IGRAs were also found to have greater sensitivity in diagnosing active TB infection compared to the TST, 80% for QFT-GIT, 81% for T-Spot compared to only 65% for the TST. In the same review, specificity was found to be 79% (75-82%) for QFT-GIT, 59% (56-62%) for T-spot and 75% (72-78%) for TST. Sensitivity to diagnose latent TB infection was found 67%, 87% and 71% for QFT-GTI, T-Spot and TST respectively, whereas specificity for latent TB infection was 99%, 98% and 88% respectively (Diel et al, 2011; Menzies et al, 2007; Pai et al, 2008; Sester et al, 2010).

Current consensus amongst the European countries is that IGRAs can be included in screening for latent TB infection, albeit there is not enough evidence yet to provide a clear picture. Nonetheless it can provide an extra step in establishing a diagnosis. On the other hand, due to their high negative predictive value for immunocompetent patients, negative IGRA results can safely exclude progression to active disease, albeit it does not rule out the possibility of latent infection (Diel et al, 2011). Applying the IGRAs to specimens from possible infection sites (i.e. Bronchoalveolar Lavage) as opposed to blood samples, especially in immunodeficient individuals can help distinguish between active and latent TB (Jafari et al, 2009). In diagnosing active tuberculosis we mention for completeness, that
current consensus is that IGRAs do not have a place in routine screening, yet in certain cases when there is a strong clinical suspicion yet no laboratory proof, they can contribute. Neither IGRAs nor TST can replace the standard laboratory tools for diagnosing active tuberculosis (ECDC, 2011).

As with the TST, IGRAs also have some shortcomings. Perhaps most importantly IGRAs, just like TST are unable to distinguish between latent and active infection when limited to blood testing. Moreover, blood samples need to be processed within 8-30 hours after collection; otherwise the white blood cells will gradually become non-responsive to the antigenic stimulation. Errors in collecting or transporting blood specimens or in running and interpreting the assay can decrease the accuracy of IGRAs. Since these techniques are relatively new, there is still limited data on the use of IGRAs in certain population groups such as children younger than 5 years of age, HIV patients, anti-TNF-α treated patients or in general immunocompromised patients. Finally there is a significant cost to this process as opposed to the fairly cheap TST method.

Finally, another method is being developed for use that employs flow cytometry for the detection of interferon producing lymphocytes. This method is not yet commercially available and due to the high cost of the process it is not known yet if it will contribute to latent tuberculosis diagnosis (Fuhrmann et al 2008). There are experimental methods detecting antibodies against tuberculosis antigens, but as mentioned already humoral immunity plays a small part in tuberculosis if any at all and thus these methods so far have no clinical application (El-Shazly, 2007). Most recently the WHO issued a statement asking countries to ban antibodies based tests for the diagnosis of tuberculosis (WHO, 2011).

6. Latent tuberculosis treatment

Individuals with known contacts with patients suffering from active tuberculosis and who test positive with the aforementioned methods are considered, given reasonable clinical suspicion, to have latent infection. They are eligible to receive treatment in order to prevent them from developing an active infection. In some cases (i.e. children, HIV patients) even without TST or IGRAs supporting, clinical suspicion alone is enough to start treatment and re-test the patient at a later time to verify the result of the diagnostic tests. Treatment for latent tuberculosis is less expensive than for active and preventing the disease provides overall a great economic benefit for the health-care system.

Current guidelines (American Thoracic Society & CDC, 2000, revised 2005) in the US, suggest a 9-month daily treatment with isoniazid (INH) 5mg/kg up to 300mg. This can be reduced to only 6 months, for adults seronegative for HIV co-infection. In most cases the 9 month treatment plan is followed since it has been show to achieve better results (70% complete remission vs. 60% for the 6 month regimen). In very few cases a 12-month regimen is recommended, particularly for populations with a higher incidence of active tuberculosis (TBNET, 2009).

As is the problem with most tuberculosis therapies there is a high amount of non-compliant patients contributing to failure of treatment. One solution would be to enforce Directly Observed Treatment (DOT) for patients taking isoniazid for latent tuberculosis, but such a decision comes with a high financial cost. Under these circumstances, treatment can be modified to a 2/week regimen at a dose of 15mg/kg up to 900mg. Isoniazid side-effects
include polyneuropathy, preventable with administration of B6 vitamin and hepatic toxicity that remains a prime reason for discontinuation of treatment. Studies have shown that 10-20% of patients will have an increase in liver transaminases and about 2% will have clinically significant hepatitis, with that percentage increasing in the present of co-morbidity factors (Nolan et al, 1999).

Due to these problems the ATS and CDC have suggested alternative treatment options. One such option is a daily dose of rifampicin (RMP) 4-month single-drug regimen or a daily dose of pyrazinamide (PZA)-rifampicin 4-month regimen. The RMP treatment is not recommended for HIV positive patients due to interactions with HAART treatment, but otherwise it has shown promising results for patients intolerant of INH or for those cases where INH resistance is verified or suspected. Benefits of this shorter regimen include a lower cost and also higher degrees of compliance (Jasmer et al, 2002; Menzies et al, 2004, 2008; Polesky et al, 1996; Reichman et al, 2004; Villarino et al, 1997).

Initially the PZA-RMP regimen was designed to be administered for 2 months, but due to adverse effects (serious hepatotoxicity and death) it is no longer recommended, but for some rare cases (CDC, 2001; Leceouer, 1989; Gao, 2006) Other possible regimens that are under evaluation include a 3 month daily treatment with INH-RMP and a 3 month weekly INH-rifapentin regimen. The former has been tested in the UK and exhibiting satisfactory results in terms of adverse effects and success of treatment (Ena & Valls, 2005). The latter is under study in the US, the CDC recently made public that patients on this regimen have higher compliance, satisfactory remission results compared to INH but it seems that they have increased adverse effects and also the cost of treatment is higher than the RMP regimen.

7. Conclusion

Latent tuberculosis is a field of great scientific interest and research possibilities. We have investigated the granuloma and its formation and 2 theories exist, a lot of the secrets still remain hidden and more evidence is needed to support either theory. In the field of diagnosis new tools are available and it remains to be seen how they will fare when tested against special populations (i.e. HIV patients which is the field of our own research as well). New guidelines for treatment are issued and those are under evaluation. Latent tuberculosis is an important public health issue, an insidious infection that can persist for years; above all, clinical suspicion is paramount for its diagnosis.

8. References


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and CD4 T cells that mediate autoimmunity. *J Immunol.* Vol 186 No 10 May 2011 pp5738-48


Pulmonary infections are notorious in causing considerable morbidity and mortality. Caused by bacteria, viruses or fungi, respiratory infections require distinct knowledge of recent advances in pathogenesis. Progress in the understanding of immunopathogenesis of Acinetobacter baumannii infection will explain how an atypical organism establishes infection. The chapter regarding pulmonary nontuberculous mycobacterial infections in the State of Para depicts a unique study in an endemic region for tuberculosis in North of Brazil. The diagnosis and treatment of latent tuberculosis is a formidable challenge. Thus, new developments in diagnosis and treatment of latent tuberculosis are included in this book. Challenging in their diagnosis, nontuberculous mycobacterial pulmonary diseases require special education for management. The problems of respiratory infections in the immunocompromised host are increasing in numbers and in resilience to treatment. Therefore, the chapter describing the host immune responses against pulmonary fungal pathogens comes as a necessary section in this book. The insight brought forth from this book can be valuable for both clinicians and scientists.

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