Transcriptional biomarkers for predicting development of tuberculosis: progress and clinical considerations

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Introduction
Achieving the ambitious targets for global tuberculosis (TB) control, will require an increased emphasis on preventing development of active disease in those with latent TB infection (LTBI) by preventative treatment or vaccination [1]. New shortened regimens of 1–3 months duration potentially allow much wider and more effective use of preventive therapy [2, 3]. A significant barrier is the limited ability to reliably identify those at high risk of disease progression, leading to high numbers needed to treat (NNT) to prevent a case of TB [4]. Hence, there is an interest in developing new diagnostic tests that better predict TB disease to allow more targeted preventive treatment and lower NNT [5]. These are often referred to as tests for incipient TB, stemming from the notion that TB prediction with low NNT most likely reflects detection of an early inflammatory response to multiplying Mycobacterium tuberculosis [6]. There are a number of promising biomarkers in development and undergoing evaluation as incipient TB tests [7]. Although it will be a few years before validated and approved diagnostics are ready for use in clinic, there is the potential for such tests to transform the TB management paradigm and provide a fresh approach to TB control. However, how such tests fit into future TB management and control algorithms is not fully defined and there remain limitations in their potential utility as well as a number of research gaps in this area.

How do predictive tests work?
Infection with M. tuberculosis results in an acquired immune response which enables effective granulomatous control of the organism and prevents clinical disease in approximately 90% of cases. Diagnosis of latent infection relies on detection of this antigen-specific response either in vivo with
A number of approaches have been utilised for diagnostic development which include 1) detecting low levels of *M. tuberculosis* antigens or nucleic acid, 2) identifying *M. tuberculosis*-specific immunophenotypes that correlate with disease risk and 3) detecting host RNA or protein signatures sensitive for early disease [14–16]. In contrast to the T-cell-based re-stimulation approach used in IGRA, a more global consideration of the host response in unstimulated blood using transcriptomics to evaluate mRNA abundance, have proved most successful for developing biomarkers of early disease. Over the past decade a large number of studies utilising RNA sequencing or microarray for biomarker discovery have identified mRNA transcript signatures in whole blood that distinguish those with clinical, active pulmonary TB from those with latent infection or other diseases [17]. These approaches characterise the differences in the global host response between disease states, reflecting both the changes in gene expression in circulating immune cells and their relative abundance in the blood sample. Of note, transcriptional changes have been shown to normalise with treatment of active TB [18]. In addition, and importantly for predicting disease, for a subset of the active TB-associated transcripts, changes in abundance were detectable in asymptomatic individuals in whom early TB pathology was demonstrated using PET/CT imaging, prior to clinical disease development [19]. Cohort studies following at-risk populations have also shown that transcriptional signatures can distinguish progressors from non-progressors from at least a year before clinical presentation [20]. While exact transcript signatures defined by these studies has differed with different study populations, aims and bioinformatic approaches, there is considerable overlap and co-correlation of transcripts, and the biological processes reflected by them is consistent [21, 22].

While in initial studies several signatures published met the WHO thresholds [7], they appear to perform less well in study populations outside those in which they were discovered. A recent meta-analysis evaluating 17 published candidate TB signatures against a pooled dataset from four studies containing 127 cases of incipient TB, identified eight signatures that performed equivalently. However, all failed to meet the minimal standard of 75% sensitivity and specificity for disease progression at 2 years proposed in the WHO TPP [22].

These limitations in part reflect two important features of disease progression, namely a significant proportion of individuals who develop incipient disease will self-heal and not progress registering a false-negative test. Secondly, progression rates of disease are heterogenous. While in some progression may take years, those that develop disease more rapidly will only test positive close to onset of clinical disease.
Indeed, the positive predictive value of the eight identified signatures for disease development was 11.1–14.3% over 3 months, 9.5–11.7% over 6 months and 6.8–9.4% over 24 months [22]. Assuming 60% treatment efficacy [23, 24] these values translate to NNTs of 12–15, 14–18 and 18–25, respectively, but it should be kept in mind that they are population-dependent (mainly TB incidence-dependent, if not considering host immune suppression) (figure 1).

In addition, there are inherent issues with the sensitivity and specificity of the transcriptomic approach. Gene expression is determined by the nature of the host response; however, not all forms of TB share a common disease pathogenesis. As a result, the transcriptional response may differ in miliary disease, in disease presentations in early childhood or in immunocompromised hosts. It is notable that sensitivity of these transcriptional signatures is ~90% even in confirmed active disease, which inherently limits the sensitivity for incipient disease [25]. Furthermore, other diseases share elements of the whole blood transcriptional responses found in TB, limiting specificity of the tests. In particular, the interferon signalling, which is the most enriched pathway in many of the TB transcriptional signatures, is also an important component of the transcriptional response in untreated HIV infection [19].

Finally, for these biomarker signatures to have a role in clinical care, several steps are needed to translate these into assays for routine use [26]. Whole genome discovery approaches need to be translated to more tractable, such as multiplex PCR or NanoString’s Counter technology, for the most parsimonious sets of predictive transcripts. Scoring methods must be determined to provide diagnostic cut-off values and labour-intensive RNA extraction methods need to be streamlined and ideally integrated into the diagnostic platform. Progress is being made, with companies such as Cepheid developing cartridges to monitor host mRNA abundance by real time PCR on the GeneXpert platform. These assays, once developed, should then be evaluated in prospective studies in populations of their intended use for their clinical and public health utility [10] However, it is also important to reflect that the challenges in overcoming all these steps means that numerous promising biomarkers fail to progress to the clinic. Transcriptional biomarkers are slowly making their way into clinical practice primarily in the cancer field where a number of commercial products are available to measure transcript abundance, mainly in tissue samples. Additionally, although at this stage transcriptional biomarkers for the purpose of short term prediction of disease progression look most promising, it may be that other approaches, particularly T-cell activation markers or biomarkers

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**FIGURE 1** In this illustrative scenario, 480 individuals are tested for latent tuberculosis (TB) and incipient TB at baseline. Among them, 13 (2.7%) develop incident TB over 24 months. At baseline 50% have a positive test for latent TB whereas 15% have a positive test for incipient TB. At 24 months, the latent TB test accuracy for predicting incident cases has a sensitivity of 62%, specificity 50%, positive predictive value (PPV) 3.3% and negative predictive value (NPV) 98%. In contrast, the incipient TB test for predicting incident cases has a sensitivity 39%, specificity 86%, PPV 6.9% and NPV 98%. The number needed to treat to prevent disease within 24 months respectively would be 50 for the latent TB test (with five missed cases) and 24 for the incipient TB test (with eight missed cases). Note this does not take into account reinfection over the 24-month period.
derived from proteomic studies or other approaches that are in early development, may be more fruitful in the longer term.

**Potential clinical use of tests**

Whole blood transcriptional biomarkers mark clear progress in predicting clinical progression of disease. If the minimal test performance recommended by WHO can be attained and technical challenges overcome, transcriptional biomarker-based tests hold promise for selecting high TB risk individuals for preventive therapy. In figure 1 we show a potential use of these assays. In low-incidence settings these could include household, school and occupational contacts of infectious TB patients, healthcare workers and recent immigrants from high-incidence countries. In high-incidence settings, incipient TB tests could additionally be used for community screening [27], for example in (bi-)annual campaigns. Because of a time-lag effect in which an inflammatory response may commence only months after *M. tuberculosis* infection, repeat testing may be warranted. As a result, we may need to reconsider our approach to evaluating and testing risk groups such as household contacts, who are typically only assessed for evidence of TB infection (with IGRA or TST) and consideration for preventive therapy at baseline. A new algorithm, incorporating novel predictive diagnostics testing, may require repeated testing at 6 monthly or yearly intervals for a period of 2–3 years. Comparison of diagnostic and treatment algorithms should be evaluated in clinical trials, but patient and provider views on the different approaches will also need to be explored. Additionally, technical feasibility and affordability will be important considerations, and cost-effectiveness analyses will be needed to inform policy decisions with regard to target groups and frequency of testing. Transcriptional biomarker-based tests can likely also be used for triaging symptomatic patients or those with abnormal CXR being evaluated for TB, by more precisely identifying those who need further diagnostics for bacteriological confirmation [28], this also being important for cost considerations. As there is no time-lag effect, sensitivity for this use case is expected to be higher than for prediction, whereas specificity may be lower because alternative causes of disease may elicit a similar transcriptional response.

It is as yet unclear whether transcriptional biomarker-based tests should be used to guide preventive therapy in individuals with impaired cell-mediated immunity, such as people living with HIV, pre-transplant patients and those using tumour necrosis factor-α blocking agents [28]. In these target groups contained *M. tuberculosis* infection may quickly progress to severe disease, with a narrow time window to test for incipient TB. Therefore, current approaches to risk stratification, combining epidemiological factors with or without IGRA, may remain appropriate.

Another question that these new assays raise is how incipient TB, defined by a positive prediction test but negative molecular assay or *M. tuberculosis* culture, should be treated. Clinical trials will be required to establish non-inferiority of preventive treatment to full-course chemotherapy, with a potential role for shortened (e.g. 2 months) four-drug regimens [29].

Acknowledgements: This project is part of the Global TB Network activities carried out for the latent tuberculosis group. Moreover, this work is a part of the research performed thanks to the grant of Ricerca Corrente from the Italian Ministry of Health Linea 4, program 2.

Support statement: This work was supported by Ricerca Corrente, Italian Ministry of Health. Funding information for this article has been deposited with the Crossref Funder Registry.

Conflict of interest: H. Esmail has nothing to disclose. F. Cobelens has nothing to disclose. D. Goletti reports grants and personal fees from Quidel, personal fees from Qiagen, Janssen and Diasorin, outside the submitted work.

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https://doi.org/10.1183/13993003.01957-2019
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