Epigenetics and Proteomics Join Transcriptomics in the Quest for Tuberculosis Biomarkers

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ABSTRACT  An estimated one-third of the world’s population is currently latently infected with Mycobacterium tuberculosis. Latent M. tuberculosis infection (LTBI) progresses into active tuberculosis (TB) disease in ~5 to 10% of infected individuals. Diagnostic and prognostic biomarkers to monitor disease progression are urgently needed to ensure better care for TB patients and to decrease the spread of TB. Biomarker development is primarily based on transcriptomics. Our understanding of biology combined with evolving technical advances in high-throughput techniques led us to investigate the possibility of additional platforms (epigenetics and proteomics) in the quest to (i) understand the biology of the TB host response and (ii) search for multiparameter biosignatures in TB. We engaged in a pilot study to interrogate the DNA methylation, transcriptome, and proteome in selected monocytes and granulocytes from TB patients and healthy LTBI participants. Our study provides first insights into the levels and sources of diversity in the epigenome and proteome among TB patients and LTBI controls, despite limitations due to small sample size. Functionally the differences between the infection phenotypes (LTBI versus active TB) observed in the different platforms were congruent, thereby suggesting regulation of function not only at the transcriptional level but also by DNA methylation and microRNA. Thus, our data argue for the development of a large-scale study of the DNA methylome, with particular attention to study design in accounting for variation based on gender, age, and cell type.

IMPORTANCE  DNA methylation modifies the transcriptional program of cells. We have focused on two major populations of leukocytes involved in immune response to infectious diseases, granulocytes and monocytes, both of which are professional phagocytes that engulf and kill bacteria. We have interrogated how DNA methylation, gene expression, and protein translation differ in these two cell populations between healthy individuals and patients suffering from TB. To better understand the underlying biologic mechanisms, we harnessed a statistical enrichment analysis, taking advantage of predefined and well-characterized gene sets. Not only were there clear differences on various levels between the two populations, but there were also differences between TB patients and healthy controls in the transcriptome, proteome, and, for the first time, DNA methylome in these cells. Our pilot study emphasizes the value of a large-scale study of the DNA methylome taking into account our findings.

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Tuberculosis (TB) is a threatening disease, with currently 9 million new cases and 1.5 million deaths per year (1). One-third of the global population is latently infected (latent TB infection [LTBI]) with Mycobacterium tuberculosis, thus facing the risk of developing active TB during their lifetime. Effective drug treatment regimens exist, albeit they have been challenged by increasing multidrug-resistant, extensively drug-resistant, and totally drug-resistant forms of TB (2). In addition, TB diagnosis and control are still hampered by the unavailability of vaccines capable of preventing TB. The fact that only an estimated 5 to 10% of people with LTBI develop active TB disease combined with the high occurrence of infection in household contacts emphasizes that the underlying biological mechanisms remain poorly understood. A better understanding of the biological processes involved in progression from LTBI to active TB will contribute toward better intervention measures. To this end, sufficiently validated bio-
markers to support development of TB vaccines, diagnostics, and drugs are needed (3, 4). To add further complexity to this scenario, the biology of and interactions between the host, microbe, and environment are highly complex and variable (5). Some factors are known to affect the host response to M. tuberculosis infection, including inherent host genetics, variability among hosts, status of the immune system, and external factors, such as nutrition, pollution, coinfection, frequency of previous infections and coinfections, stress levels, and adaptations by bacterial strains in specific host populations (6–8). External factors are also known to affect host epigenetics, prompting us to include the epigenome in future TB biomarker research. In particular, stable marks such as those derived from DNA methylation patterns are currently absent from studies of this kind, while more dynamic epigenetic marks (histone modifications) have uncovered matters relating to “trained immunity,” even in the case of recipients of the only current TB vaccine, M. tuberculosis BCG (bacillus Calmette-Guérin) (9, 10).

To date, transcriptomic profiles from peripheral blood cells have been the main focus in the exploration for biomarkers (11). In addition to easy access, blood also represents a site of dynamic exchange of chemokines, cytokines, and cells trafficking between foci of active disease and the lymphoid system (12). Within the immune system, cells have their own discrete functions, but the system as a whole exerts a concerted function with remarkable plasticity. Hence careful consideration of different cell types is required when describing either predispositions for disease development or the resultant functions following infection. Understanding how each specific cell contributes to maintenance of LTBI instead of progressing to active TB is essential. Professional phagocytes play a central role in these processes (13). Phagocytes comprise dendritic cells, monocytes (differentiating into macrophages), and granulocytes, predominantly neutrophils. Monocytes and granulocytes have epigenomes distinct from other cell types (14, 15). Due to its inherent stability, disruption of the normal DNA methylome can produce stable cell populations with prolonged aberrant phenotypes and thereby contribute to disease, which suggests a useful source of biomarkers for risk stratification and disease diagnosis (16). In primary human leukocyte subsets, single nucleotide polymorphisms (SNPs), which are associated with immune-mediated disease, preferentially map to cell-specific regulatory hypomethylated regions (HMRs) (14). Such loci, in combination with DNA methylation sites, offer hypotheses toward depicting cellular subsets in which specific epigenetic changes may drive disease.

This pilot study aims to investigate the level of differentiation between TB patients and genetically unrelated LTBI household contacts in a cell-specific manner. Isolated monocytes and granulocytes from peripheral blood were investigated to determine the extent and biological functions of differences between the DNA methylome, transcriptome (mRNA and microRNA [miR]), and proteome at a global level using high-throughput techniques.

Toward this end, we studied each platform in a descriptive manner, followed by differential analyses between LTBI and TB at each platform in monocytes and granulocytes alone and in combination. Next, we determined if these differences point toward specific functions. Finally, we analyzed these data to determine which, if any, correlations of differences between LTBI and TB exist in functions from one platform to another. Specifically, we wanted to ascertain in this very-limited-sample-size study whether a portion of the differences between LTBI and TB in global DNA methylation and miR platforms reveal gene regulation that will be reflected by the mRNA data set and in turn is translated into the proteome.

At the levels of DNA methylation, miR, mRNA, and proteins, the data allowed differentiation between LTBI and TB. Moreover, unique functions were congruent from one platform to another, suggesting a global regulation of function (or loss thereof) during TB.

RESULTS AND DISCUSSION

Global DNA methylation in LTBI and TB. The DNA methylation status of over 485,000 CpGs was interrogated in isolated monocytes and granulocytes from LTBI and TB participants. For both cell types, a clear bimodal data distribution was evident, with no major variation of these distributions between LTBI and TB (see Fig. S1A and B in the supplemental material) either together or per cell type. Frequency distributions illustrate similar levels of hypomethylated (0 to 30%) and hypermethylated (70 to 100%) CpGs in both study groups. This observation differs from the overall spread of methylated CpGs in precursor cells, such as hematopoietic stem cells (15).

We tested whether methylation of CpGs in professional phagocytes was related to known functions in these cell types. The list of genes with fully hyper- or hypomethylated CpGs associated with promoter regions was tested for enrichment in blood transcription modules (BTMs) using the hypergeometric test, in which genes with fully hyper- or hypomethylated CpGs associated with promoter regions constituted the foreground set, while genes lacking such sites constituted the background (17, 18). Genes with promoter-associated hypermethylated sites (i.e., putatively silenced in the analyzed cell subsets) were not significantly enriched in BTMs. However, significant enrichment in BTMs for genes with hypomethylated CpGs associated with promoter regions (i.e., potentially activated genes) was observed with statistically significant enrichment in modules involved in cell cycling and transcription, as well as those involved in immune activation (see Table S1 in the supplemental material).

To determine which known sources of variation (that is, disease phenotype, gender, and cell type) explain the total variance of the data set, we applied principal component analysis (PCA) regression of principal components (PCs) over the independent variables. Both gender and cell type explained a substantial portion of the variance of the first two PCs, while disease phenotype explained a smaller fraction of the variance (Fig. 1A). Disease phenotype explained the majority of the variance of PC6 (Fig. 1A and B). When the global DNA methylation status of these and other cell types in peripheral blood was compared in another study of similar design, PCA indicated that DNA methylation patterns differ more profoundly between cell types than between individuals (19). These results, in combination with our findings, indicate that M. tuberculosis infection affects DNA methylation in monocytes and granulocytes to a lesser extent than differences observed in cell type at the level of the global DNA methylome.

Given the clear binomial distribution both in cell types and in disease phenotypes, a nonspecific filtered data set was defined to exclude uninformative data. This set excluded (i) CpGs for which no link to functional genes had been established thus far and (ii) CpGs that were either hypermethylated or hypomethylated in every sample. The resulting filtered data set contained 80,198 CpG
sites and showed an even more concise discrimination between genders (Fig. 1C and D). Unsupervised hierarchical clustering of the methylation profiles from 36 samples confirms gender differences in these two cell types (Fig. 1E). Thus, gender accounted for higher variation between individuals than TB disease phenotype in this data set. However, when the filtered set of CpGs was evaluated for differences between LTBI and TB, a clear distinction between groups was observed in PC5 and PC6 of the PCA (Fig. 1D), suggesting that some of the epigenetic changes in monocytes and granulocytes were either the result or the cause of TB.

Second, differential analyses revealed <1% of the CpGs in the filtered set to be statistically differentially methylated between TB and LTBI (Wilcoxon \( q < 0.05 \), adjusted for multiple testing using the Benjamini-Hochberg procedure) (see Table S2 in the supplemental material). Of these, the CpGs showing most differentially methylated targets occurred in CpG islands and shores (see Fig. S1C in the supplemental material). We tested whether the effect of the disease phenotype was random by considering the predictive power of the methylated sites using a machine learning (ML) approach. The resulting models were significantly better than random at distinguishing TB from LTBI (area under the curve [AUC], 0.74; 95% confidence interval [CI], 0.57 to 0.92), with an overall error rate below 30%. Classification between cell types yielded no errors (AUC, 1.00).

As can be expected following PCA, the relative abundances of differential methylation between monocytes and granulocytes within the same individual were much larger (~20% [Fig. S1D]). This confirms a previous study reporting 22% of CpGs to be differentially methylated between these two cell types (19).

To gain insight into the functional role of CpGs with methylation differences between LTBI and TB, we tested enrichment in functional categories of the genes associated with differentially methylated CpGs. BTMs for which the differences between LTBI and TB were significantly enriched resemble a tentative signature/fingerprint with CpGs either hyper- or hypomethylated in LTBI compared to TB within five known functional modules (Fig. 2A). Of particular interest are four CpGs found to be differentially methylated in the “MHC-TLR7-TLR8 cluster,” which all occur in CpG islands (Fig. 2B). While the CpGs associated with HLA-DQB1 (coding for one HLA class II chain) did not reside in a promoter-associated area of the gene, the other three CpGs associated with HLA-F and coding for HLA class I were all located within the promoter. Moreover, these two HLA-related genes to
**FIG 2** Functional association of statistically significant differentially methylated CpGs. (A) Heat map showing differentially methylated CpGs associated with genes from five blood transcriptional modules. Colors are relative as data are scaled row-wise: white indicates lowest methylation for a given CpG, while blue indicates highest methylation for a given CpG. Columns correspond to samples. Multiple CpGs can correspond to a single gene. (B) Dot plots showing changes between LTBI and TB for one CpG associated with the HLA-DQB1 gene and three CpGs associated with the HLA-F gene. Lines connect samples from a single pair of individuals. Blue indicates measurements for samples in LTBI, and red indicates measurements for samples in TB. The q values represent the P values after correction for multiple testing (Benjamini-Hochberg); *, P < 0.05; **, P < 0.01 (Wilcoxon paired test).
which these CpGs have bearing had opposite effects when LTBI and TB were compared: CpGs for HLA-DQB1 were significantly hypomethylated in TB patients compared to their paired LTBI samples, while CpGs at three different loci in the promoter of HLA-F were hypomethylated (Fig. 2).

Hypomethylated HLA-DQB1 in TB and reduced major histocompatibility complex class II (MHC-II) expression, regardless of whether it is a trigger or a result of activation, could benefit M. tuberculosis. This resonates with reports (20, 21) indicating that M. tuberculosis interferes with antigen processing and presentation. Interestingly, an immunogenic peptide from M. tuberculosis is presented by HLA-DQB1, and some HLA-DQB1 alleles associated with TB sensitivity present such peptides resulting in suboptimal antigen-specific interferon gamma (IFN-γ) secretion by CD4 T cells (22). This underlines the relevance of methylation of disease-related MHC-II alleles and warrants further investigation. On the other hand, HLA-F, which has been associated with tumor invasiveness (23) and immune suppression (24), was hypomethylated in TB compared to LTBI-matched individuals. This, along with hypermethylated HLA-DQB1, may result in immune modulation to favor the pathogen.

Global expression of transcripts and miRs. Transcriptomic data from the four experimental groups can be clearly distinguished according to gene expression profiles, with differences between cell types dominating over differences between study groups (see Fig. S2A in the supplemental material). PCA revealed that 33% of the overall variance in the data corresponded to PC1, which was correlated with the two cell types. A further 14% of the variance is explained by PC2 and PC5, which correlate with TB. We have trained ML models to distinguish TB from LTBI and, independently, the two cell types. TB could be distinguished from LTBI with few errors (AUC, 0.99; 95% CI, 0.96 to 1.00), while there were no errors in classification of the cell types.

Second, both cell types showed a number of significantly differentially expressed genes between LTBI and TB (see Table S3 in the supplemental material). Pairwise comparisons between monocytes and granulocytes as well as between the LTBI and TB groups demonstrated a substantial response to TB in both cell types, as well as marked differences between these cell types in both LTBI and TB samples (see Fig. S3B in the supplemental material), as can be seen on the PCA plot (see Fig. S3C). PC1 corresponded to differences between monocytes and granulocytes and was enriched in “mitochondrial and translation-related” genes, and PC2 and PC5 corresponded to differences between LTBI and TB samples.

In agreement with previous studies, TB patients showed a significant increase in expression of several genes identified as relevant in previous studies (25–27). These include significant upregulation of CD64 (Fc-γ receptor IA), and guanylate-binding proteins (GBP) in TB. In granulocytes, we found a higher expression in TB for several MHC-II-related genes, including the CD74, HLA-DRA, and HLA-DMB genes. While the expression of these genes was significantly lower in granulocytes than in monocytes, no differences between LTBI and TB were apparent in monocytes.

Differential transcription analysis between LTBI and TB followed by functional analysis revealed several BTMs that were statistically significantly enriched for among differentially expressed genes (q < 0.05) (Table 1; Fig. 3A). Notably, the BTM designated “B cell surface signature” was significantly enriched on both the methylation and transcription levels. Likewise, transcriptional differences between monocytes and granulocytes were coherent with the corresponding module enrichments in the methylation data set (see Table S4 in the supplemental material). The responses to TB in monocytes and granulocytes were largely similar in the mRNA data set. The overall Pearson correlation coefficient between the log2-fold changes in monocytes and granulocytes was 0.42 (q ~ 0). In line with this, there were no genes with a statistically significant interaction between cell type and disease state—that is, genes with a different response to TB in monocytes compared to granulocytes.

However, based on visual inspection and uncorrected P values, we marked several candidate genes with an apparent difference between LTBI and TB for 1 cell type only. Employing quantitative reverse transcription-PCR (qRT-PCR), we were able to confirm differential expression in both cell types for GBP5 and signal transducer and activator of transcription 1 (STAT1), while for STAT1 in separate analyses for monocytes and granulocytes, only the monocytes revealed a statistically significant difference (n = 10 per group; P < 0.05, Wilcoxon paired test) (Fig. 3B). Even though statistical analyses fail to point out singular gene products to differ between LTBI and TB after correction for multiple testing, these data were confirmed by a second technique. Moreover, this supports the importance of interrogating cellular subsets independently when investigating the transcriptome for differential markers in TB.

In the miR data set, the largest portion of variation can be explained by differences between cell type and disease phenotype (see Fig. S3 in the supplemental material). Gender played a far less significant role than DNA methylation. The separation between LTBI and TB was less clear in the PCA for mRNA. The random forest ML models showed high performance both for separation of TB from LTBI samples (1 error; AUC, 1.00) and monocytes from granulocytes (no errors; AUC, 1.00).

Several miRs were significantly differently expressed between LTBI and TB in both cell types as well as in one cell type (see Table S5 in the supplemental material). One such example is miR-146a-5p (see Fig. S4A in the supplemental material), for which the largest relative upregulation was in monocytes from LTBI compared to TB. This miR has recently been reported to play a central role in the immune response (28) and to be upregulated in peripheral blood mononuclear cells of control subjects versus TB patients (29). miR-146a is upregulated in response to microbial stimuli and proinflammatory cytokines and has also been negatively correlated with interferon (IFN) type I signaling (30). We have validated this result using a TaqMan miR real-time quantitative PCR (qPCR) assay (see Fig. S4B).

Of note is the relatively large fraction of variance explained by gender in the full data set of DNA methylation as opposed to those for mRNA and miR. Transcripts such as mRNA and miR have a very short half-life. Therefore, changes in transcription rate rapidly affect the number of transcripts, which in turn causes a relatively high baseline variation due to rapid change in regulatory signals. Relative to this high baseline variation in transcripts, the contribution of gender to the total variation will be small. On the other hand, DNA methylation is a biochemically stable modification that results in markedly less baseline variation than that in mRNA and miR transcripts.

DNA methylation at CpGs had been documented repeatedly to be influenced by gender (31–34), not only on X and Y chromosomes but also CpGs on autosomal genes (32, 34). Moreover,
DNA methylation patterns are thought to be established largely during early embryonic development and then stably propagated via mitosis.

Since the samples for this study were collected from adult participants, we can expect to see a relatively small contribution to variation by gender compared to the large baseline variation in transcript data sets but a relatively high contribution of variation by gender against the low baseline variation in the DNA methylation set.

**Phagocyte proteomics during active TB.** To shed light on the relationship between active disease and protein synthesis in professional phagocytes, we measured the relative abundances for 3,047 unique proteins and an additional 429 isoforms. Variance in these data was largely accounted for by differences observed in cell types (Fig. 4A). Yet, in PC3 and PC5, the majority of variance is explained by differences between LTBI and TB, revealing that at the proteomic level in professional phagocytes, LTBI and TB can be discriminated (Fig. 4C). Accordingly, ML models could correctly classify both TB versus LTBI samples (AUC, 0.93; 95% CI, 0.84 to 1.00) and monocytes/granulocytes (no errors; AUC, 1.00).

In both cell types, we found significant differences between LTBI and TB (see Table S6 in the supplemental material). Several of these proteins are functionally related, as has been revealed by interrogation for enriched BTMs. Notably, we identified enrichment of the IFN signaling modules (Fig. 5), including GBP1, GBP3, GBP5, STAT1, STAT2, and IFN-induced proteins with tetra-ricopeptide repeats (IFITs).

**Effect of DNA methylation on transcription.** We next interrogated the extent to which differences in DNA methylation may influence transcription in TB. To test whether, in general, distinct methylation sites were correlated with expression of corresponding genes, we calculated row-wise correlation coefficients between CpG methylation sites and the corresponding genes (83,562 CpG gene pairs in total). The distribution of these correlation coefficients was significantly different from those of randomly paired gene pairs in total. The distribution of these correlation coefficients was significantly different from those of randomly paired gene pairs in total. The distribution of these correlation coefficients was significantly different from those of randomly paired gene pairs in total. The distribution of these correlation coefficients was significantly different from those of randomly paired gene pairs in total.

We then focused on genes for which expression was strongly correlated with methylation of related CpG sites by investigating which BTMs contained such genes. In other words, for each functional group (BTM), we interrogated whether the expression of genes included in that functional group was, on average, correlated with methylation of related CpG sites. If so, we have established a relationship between active disease and protein synthesis in professional phagocytes, LTBI and TB can be discriminated (Fig. 4C). Accordingly, ML models could correctly classify both TB versus LTBI samples (AUC, 0.93; 95% CI, 0.84 to 1.00) and monocytes/granulocytes (no errors; AUC, 1.00).

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methylated or expressed for each pair of a CpG and a corresponding gene. Then, for each BTM, we performed a randomization test to determine whether the average correlation coefficient per BTM was significantly different from a random set of correlation coefficients. The existence of a correlation between the absolute methylation values and gene expression does not necessarily mean that a regulation of gene expression by differential methylation is relevant for the TB disease process. Therefore, we further calculated the correlation coefficients of paired differences in methylation/expression between LTBI and TB in monocytes and granulocytes, as well as between monocytes and granulocytes in LTBI and TB. We found that for 39 BTMs, the average correlation coefficients for the genes in that module were significantly different from 0 (at $q < 0.05$) in a randomization test. Several of these modules were related to the immune response, including “regulation of antigen presentation and immune response,” “enriched in neutrophils,” and “immune regulation—monocytes, T and B cells.” Interestingly, 24 annotated modules, including modules related to antigen processing and presentation, showed a significant average correlation coefficient (at $q < 0.05$) of differences between LTBI and TB (Table 2), indicating that disease-specific expression of genes in these modules is connected to differential methylation.

Furthermore, we applied an alternative approach to elucidate whether a functional link between methylation and the transcriptome exists. We calculated the correlation between methylation and gene expression for each pair consisting of a CpG and a matched gene. Genes were ordered by their highest correlation coefficient with any matched CpG, and enrichment in BTMs was calculated. Thirty modules were significantly enriched at $q < 0.05$,
Using this approach, we further directly investigated whether differences in methylation between LTBI and TB within a matched-pair design had an effect on changes in gene expression. For each gene, the highest correlation of differences was calculated between matched samples in LTBI and TB in methylation and gene expression and tested for enrichment in BTMs accordingly. We found enrichment in 18 modules at $q < 0.05$, including “enriched in monocytes,” “enriched in neutrophils,” and “immune activation—generic cluster.”

These results confirm our findings and demonstrate that differences in methylation between LTBI and TB are functionally linked to differences in gene expression relevant to the immune response.

### Effect of DNA methylation on protein synthesis

The effect of changes in gene expression on the abundance of the final protein product is moderated by various factors (35), and any effect of DNA methylation on protein abundance will be exerted via transcriptional changes. Therefore, we interrogated whether observed changes in DNA methylation could be linked to changes in protein abundance.

Applying the same approaches described above, we first calculated the per-BTM average correlation coefficients and identified modules showing a significant average correlation. Second, we determined module enrichment in genes ordered by their correlation with methylation sites.

The first approach (average correlations in a BTM) revealed that for the general correlation, 22 modules had significant correlation coefficients, including “cell cycle and transcription,” “enriched in monocytes,” and “interferon.” Moreover, several modules showed significant average correlations of changes in methylation and protein abundance between LTBI and TB. These modules included “cell cycle and translation,” “enriched in monocytes,” “T cell activation,” and “inflammation.”

In the second approach, we tested the genes ordered by their respective correlation coefficients for enrichment in BTMs. Here, possibly due to low statistical power, we identified “immune activation—generic cluster” for both general correlations and corre-
lations of differences between LTBI and TB and "enriched in monocytes" for general correlation (Table 3). In any case, these results confirm that disease-specific differences in methylation are correlated with protein expression.

**Conclusions.** To date, defining the diagnostic biosignatures of TB has been largely based on gene expression analysis. However, the complexity of gene expression regulation is greatly simplified in approaches where only the transcriptome of a mixture of cells is...
analyzed. We dissected a fragment of regulation of transcriptomic changes in professional phagocytes of TB patients, elucidating the underlying biological mechanisms and paths of gene regulation. For the first time, parallel analysis of the DNA methylome, transcriptome (mRNA and miR), and proteome revealed disease-specific changes permeating these levels of regulation, commencing with methylation marks on the DNA. We demonstrate that methylation events can distinguish TB disease from healthy, infected LTBI in this sample set. These events are functionally related to unique immune-relevant classes and are manifested on both transcriptomic and proteomic levels. Although the low number of samples constrained our investigation to general effects rather than specific regulatory mechanisms, this study paves the way for further detailed investigations that interrogate the DNA methylome and proteome in addition to the transcriptome of TB patients. These future investigations would necessarily entail also the analysis of professional phagocytes in healthy subjects neither latently infected nor with active TB, allowing a comparison and biomarker identification between the two healthy groups. Whether the observed differences from each of the platforms as well as the correlated platforms will be appreciated in a large sample set representative of a population remains to be shown. We cannot ascertain whether the observed changes in methylation causally contribute to risk of TB and whether the altered methylation patterns are a cause or effect of disease. Given that the differences observed in the DNA methylome are related to functional differences observed in both transcriptome and proteome, we suggest that studying the epigenome can bring us closer to defining biomarkers of predisposition to disease, as well as uniquely contribute to our understanding of TB pathogenesis.

MATERIALS AND METHODS

Ethics statement. Blood samples (20 ml) were collected from participants following written consent (ethical approval from Stellenbosch University N10/08/274).

Clinical procedures and sample isolation. Samples from patients diagnosed with active TB (n = 8) and LTBI participants (n = 8) were obtained from an area of high TB endemicity in Cape Town, South Africa (Ravensmead and Uitsig), conforming to International Conference on Harmonisation good clinical practice (ICH-GCP) procedures. TB participants were included following a chest X ray suggestive of active disease in combination with symptoms of active TB and a confirmed positive M. tuberculosis culture result. LTBI participants were recruited to match TB patients’ age, gender, and ethnicity and were confirmed as controls based on a negative chest X-ray, the presence of no signs of active TB, and a negative M. tuberculosis culture. Both TB and LTBI participants were HIV

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**TABLE 2** Blood transcriptional modules with significant average correlation coefficients between the differences in DNA methylation and gene expression in comparison of LTBI and TB

| Module ID | Module title                                      | r   | q value |
|-----------|---------------------------------------------------|-----|--------|
| LL.M200   | Antigen processing and presentation               | 0.25| 3.66E−16|
| LL.M95.0  | Enriched in antigen presentation (II)             | 0.22| 3.43E−14|
| DC.M8.83  | Immune responses                                  | 0.22| 1.6E−10 |
| LL.M71    | Enriched in antigen presentation (I)              | 0.15| 9.94E−08|
| LL.M37.0  | Immune activation—generic cluster                 | -0.07| 5.50E−06|
| LL.M17.3  | Hox cluster IV                                    | -0.25| 9.40E−06|
| LL.S2     | B cell surface signature                          | 0.07| 1.06E−05|
| LL.M146   | MHC-TLR7-TLR8 cluster                             | 0.09| 0.0002 |
| LL.M168   | Enriched in dendritic cells                       | -0.28| 0.001  |
| LL.M5.0   | Regulation of antigen presentation and immune response | 0.09| 0.002  |
| LL.M17.1  | Hox cluster II                                    | -0.03| 0.003  |
| LL.M7.0   | Hox cluster I                                     | -0.19| 0.003  |
| LL.M96    | Hox cluster V                                     | -0.03| 0.004  |
| DC.M5.15  | Neutrophils                                       | -0.11| 0.005  |
| LL.M24    | Cell activation (IL-15, IL-23, TNF)               | -0.18| 0.009  |
| LL.M74    | Transcriptional targets of glucocorticoid receptor | -0.12| 0.02   |
| DC.M4.13  | Inflammation                                      | -0.18| 0.02   |
| LL.M160   | Leukocyte differentiation                         | -0.09| 0.02   |
| LL.M37.1  | Enriched in neutrophils (I)                       | -0.26| 0.03   |
| LL.M75    | Antiviral IFN signature                           | -0.14| 0.03   |
| DC.M4.2   | Inflammation                                      | -0.22| 0.03   |
| LL.M12.0  | Complement activation (I)                         | -0.11| 0.04   |
| DC.M3.2   | Inflammation                                      | -0.18| 0.04   |
| LL.M57    | Immuno-regulation—monocytes, T and B cells        | -0.21| 0.04   |


**TABLE 3** Blood transcriptional modules with significant average correlation coefficients between the differences in DNA methylation and protein abundance in comparison of latent M. tuberculosis infection and TB

| Module ID | Module title                                      | r   | q value |
|-----------|---------------------------------------------------|-----|--------|
| LL.M37.0  | Immune activation—generic cluster                 | -0.27| 1.06E−06|
| LL.M7.4   | T cell activation (III)                           | 0.07| 0.01   |
| LL.M4.0   | Cell cycle and transcription                      | -0.15| 0.03   |
| LL.M11.0  | Enriched in monocytes (II)                        | -0.14| 0.03   |
| DC.M4.2   | Inflammation                                      | -0.28| 0.03   |
| DC.M4.14  | Monocytes                                         | -0.25| 0.04   |

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In this table, **“Module ID”** refers to the original publication, where the prefix “LI” refers to BTMs according to Li et al. (17), and the prefix “DC” refers to BTMs according to Chaussabel et al. (18). Only modules with a functional annotation are shown. **q value** is the average correlation coefficient in the module, and the **q value** is the P value in a randomization test corrected for multiple testing.

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mbio.asm.org September/October 2015 Volume 6 Issue 5 e01187-15
negative. LTBI participants for this study were not followed up longitudinally.

Blood was collected from patients presenting with TB symptoms (n = 8), who were recruited on the day of diagnosis. Subsequently, blood was collected from LTBI participants who were recruited to match the patients’ age and gender. Granulocytes and monocytes were sequentially separated from peripheral blood with magnetic beads by magnetically activated cell sorting (MACS) (Miltenyi Biotec GmbH) (CD15+ and CD14+, respectively) according to the manufacturer’s instructions. Total RNA (rRNA), genomic DNA (gDNA), and protein were isolated using TRizol reagent (Life Technologies Corporation) according to the manufacturer’s instructions. The quality and quantity of nucleic acids were determined by electrophoresis (Agilent 2100, BioAnalyser; Agilent Technologies) and spectrophotometry (NanoDrop 2000c; Thermo Scientific).

**DNA methylation.** To investigate DNA methylation marks on CpGs, genomic DNA (500 ng) was bisulfate converted using the EZ-96 DNA methylation kit (Zymo Research Corporation) and whole-genome amplified, fragmented enzymatically, and then applied to the arrays. After extension, arrays were fluorescently stained and scanned, and the intensities of the nonmethylated and methylated bead types were measured using the Infinium human methylation450K BeadChip (Illumina). DNA methylation values (described as β values and expressed as fractions of the total number of available oligomers on the bead to anneal to) were recorded for each locus in each sample and analyzed using the software GenomeStudio (Illumina, Genomestudio 2011.1, methylation module 1.9.0). The raw data of the microarrays were uploaded to Gene Expression Omnibus.

**Transcription.** mRNA, long intergenic noncoding RNA (lincRNA), and miR abundances were measured using microarrays from Agilent Technologies (human 8-by-60,000 custom layout design 041580 containing the whole human genome), novel content for lincRNAs (from Agilent-028004), and Broad Institute human lincRNA together with Broad Institute TUCP transcripts (from Agilent-039494) and 8-by-60,000 (release 16) unrestricted human miR microarrays (Agilent-031181). Sample labeling and microarray processing were done according to the manufacturer’s instructions, and features were extracted with Agilent Feature Extraction 11.5.1.1 using the recommended protocols and settings. Data were background corrected and normalized using the R package limma version 3.20. To detect differentially expressed genes, we used the linear model (lmFit) using the R package FitDek (Illumina, Genomestudio 2011.1, methylation module 1.9.0). The raw data of the microarrays were uploaded to Gene Expression Omnibus.

**Proteomics.** Protein pellets were resuspended in lysis buffer containing 8 M urea, RapiGest (Waters), and ammonium bicarbonate. Proteins were reduced and alkylated, followed by a tryptic digest. The peptide solution was desalted by C18 reverse-phase chromatography, vacuum dried, and resolubilized to a final concentration of 1 mg/ml. Each peptide sample was analyzed on a Thermo Easy-nLC 1000 high-performance liquid chromatography (HPLC) system connected to an Orbitrap Elite mass spectrometer, which was equipped with a nanoelectrospray ion source (Thermo Scientific). Peptides were separated on a 15-cm Acclaim PepMap rapid separation liquid chromatography (RSLC) column (75-μm inner diameter, 2-μm particle size; Thermo, Fisher Scientific) at a flow rate of 300 nL/min⁻¹. Mass spectrometry (MS) spectra were acquired in the Orbitrap with a resolution of 120,000, and tandem MS (MS/MS) spectra were acquired in the linear ion trap at normal scan speed following collision-induced dissociation of the 10 most abundant precursors per cycle (normalized collision energy, 35%). We performed label-free quantification (LFQ) using Progenesis 4.0 (Nonlinear Dynamics) by automatic alignment of total ion chromatograms of raw files, using imported pepXML files from Sequest searches against the human UniprotKB/Swiss-Prot protein database. The search identifications were statistically scored using PeptideProphet (37) within the TPP (38), and all peptides with an iProbability score above 0.9 were considered resulting in a protein false discovery rate (FDR) of 1%. After manually improving the alignment, quantified peaks were filtered for identification by sequence search, and overall protein abundances were calculated thereof. The mass spectrometry discovery peptidomics data have been deposited into the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository.

**Statistical analysis and systems biological approaches.** Unless otherwise stated, statistical analyses were performed as follows: data were tested in each platform’s quality control pipeline. These were then tested for normality, described in general, followed by differential analyses employing Wilcoxon tests (DNA methylation) and limma (gene expression, miR, and proteomics), with correction for multiple testing according to Benjamini-Hochberg (39). For each platform, we performed a PCA and estimated the contribution of controlled variables in explaining the variance of the PCA components by applying PCA regression. For machine learning, random forests were applied as implemented in the R package randomForest version 4.10 (40). Unless otherwise stated, for statistical tests, we used the significance threshold of q ≤ 0.05, where q is the P-value corrected for the family-wise error rate using the Benjamini-Hochberg method (39).

To validate the predictive power of various platforms to differentiate between TB and LTBI, we have applied the random forest ML method (R package randomForest version 4.6.40), cross-validated with a modified leave-one-out (LOO) scheme. Here, at each iteration of the LOO, we have removed from the training a set of matching samples (monocytes and granulocytes) from one TB patient and the matching LTBI control. We have used the remaining samples as the training set and applied the trained model to the four test samples. The variable set used to train the model was the full set, except for the methylation platform, in which we used the filtered variable set. The results are reported as AUC and 95% CIs.

To functionally annotate results of statistical tests, we used the R software package tmod version 0.19 (available from CRAN; http://cran.r-project.org/web/packages/tmod/index.html), with BTMs as described by Li et al. (17) and Chassaing et al. (41). Depending on context, we either used a hypergeometric test for enrichment of modules in a set of differentially regulated genes compared to the genetic background or a summed rank statistics for enrichment in modules in an ordered list of genes. All procedures and R scripts required for replication of results are available upon request.

**Microarray data accession number.** The raw data from the microarrays have been uploaded to the Gene Expression Omnibus (GEO) under SuperSeries accession no. GSE70478. The mass spectrometry discovery peptidomics data have been deposited into the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository with the data set identifier PXD001960.

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01187-15/-/DCSupplemental.

- Figure S1, PDF file, 0.1 MB.
- Figure S2, PDF file, 0.1 MB.
- Figure S3, PDF file, 0.1 MB.
- Figure S4, PDF file, 0.02 MB.
- Table S1, DOC file, 0.1 MB.
- Table S2, DOC file, 0.1 MB.
- Table S3, DOC file, 0.5 MB.
- Table S4, DOC file, 0.3 MB.
- Table S5, DOC file, 0.8 MB.
- Table S6, DOC file, 0.2 MB.

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