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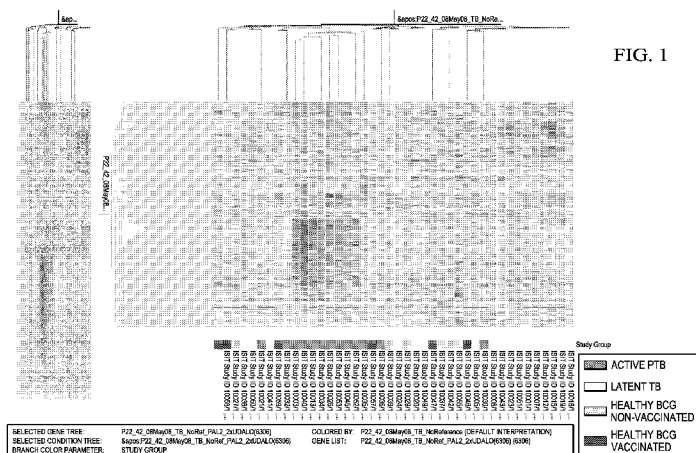


FIG. 1

(57) Abstract: The present invention includes methods, systems and kits for distinguishing between active and latent mycobacterium tuberculosis infection in a patient suspected of being infected with mycobacterium tuberculosis, and distinguishing such patients from uninfected individuals, the method including the steps of obtaining a gene expression dataset from a whole blood obtained sample from the patient and determining the differential expression of one or more transcriptional gene expression modules that distinguish between infected and non-infected patients, wherein the dataset demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent mycobacterium tuberculosis infection.

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BLOOD TRANSCRIPTIONAL SIGNATURE OF *MYCOBACTERIUM TUBERCULOSIS* INFECTION

TECHNICAL FIELD OF THE INVENTION

The present invention relates in general to the field of *Mycobacterium tuberculosis* infection, and more particularly, to a system, method and apparatus for the diagnosis, prognosis and monitoring of latent and active *Mycobacterium tuberculosis* infection and disease progression before, during and after treatment.

LENGTHY TABLE

The patent application contains a lengthy table section. A copy of the table is available in electronic form from the USPTO web site (<http://seqdata.uspto.gov/>). An electronic copy of the table will also be available from the USPTO upon request and payment of the fee set forth in 37 CFR 1.19(b)(3).

BACKGROUND OF THE INVENTION

Without limiting the scope of the invention, its background is described in connection with the identification and treatment of *Mycobacterium tuberculosis* infection.

Pulmonary tuberculosis (PTB) is a major and increasing cause of morbidity and mortality worldwide caused by *Mycobacterium tuberculosis* (*M. tuberculosis*). However, the majority of individuals infected with *M. tuberculosis* remain asymptomatic, retaining the infection in a latent form and it is thought that this latent state is maintained by an active immune response (WHO; Kaufmann, SH & McMichael, AJ., Nat Med, 2005). This is supported by reports showing that treatment of patients with Crohn's Disease or Rheumatoid Arthritis with anti-TNF antibodies, results in improvement of autoimmune symptoms, but on the other hand causes reactivation of TB in patients previously in contact with *M. tuberculosis* (Keane). The immune response to *M. tuberculosis* is multifactorial and includes genetically determined host factors, such as TNF, and IFN- γ and IL-12, of the Th1 axis (Reviewed in Casanova, Ann Rev; Newport). However, immune cells from adult pulmonary TB patients can produce IFN- γ , IL-12 and TNF, and IFN- γ therapy does not help to ameliorate disease (Reviewed in Reljic, 2007, J Interferon & Cyt Res., 27, 353-63), suggesting that a broader number of host immune factors are involved in protection against *M. tuberculosis* and the maintenance of latency. Thus, a knowledge of host factors induced in latent versus active TB may provide information with respect to the immune response which can control infection with *M. tuberculosis*.

The diagnosis of PTB can be difficult and problematic for a number of reasons. Firstly demonstrating the presence of typical *M. tuberculosis* bacilli in the sputum by microscopy examination (smear positive) has a sensitivity of only 50 - 70%, and positive diagnosis requires isolation of *M. tuberculosis* by culture, which can take up to 8 weeks. In addition, some patients are smear negative on sputum or are unable to produce sputum, and thus additional sampling is required by bronchoscopy, an invasive procedure. Due to these

limitations in the diagnosis of PTB, smear negative patients are sometimes tested for tuberculin (PPD) skin reactivity (Mantoux). However, tuberculin (PPD) skin reactivity cannot distinguish between BCG vaccination, latent or active TB. In response to this problem, assays have been developed demonstrating immunoreactivity to specific *M. tuberculosis* antigens, which are absent in BCG. Reactivity to these *M.*
5 *tuberculosis* antigens, as measured by production of IFN- γ by blood cells in Interferon Gamma Release Assays (IGRA), however, does not differentiate latent from active disease. Latent TB is defined in the clinic by a delayed type hypersensitivity reaction when the patient is intradermally challenged with PPD, together with an IGRA positive result, in the absence of clinical symptoms or signs, or radiology suggestive of active disease. The reactivation of latent/dormant tuberculosis (TB) presents a major health hazard with the risk of
10 transmission to other individuals, and thus biomarkers reflecting differences in latent and active TB patients would be of use in disease management, particularly since anti-mycobacterial drug treatment is arduous and can result in serious side-effects.

SUMMARY OF THE INVENTION

The present invention includes methods and kits for the identification of latent versus active tuberculosis (TB) patients, as compared to healthy controls. In one embodiment, microarray analysis of blood of a
15 distinct and reciprocal immune signature is used to determine, diagnose, track and treat latent versus active tuberculosis (TB) patients.

In one embodiment, the present invention includes methods, systems and kits for distinguishing between active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with
20 *Mycobacterium tuberculosis*, the method including the steps of: obtaining a gene expression dataset from a whole blood sample from the patient; determining the differential expression of one or more transcriptional gene expression modules that distinguish between infected patients and non-infected individuals, wherein the dataset demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected individuals, and distinguishing between
25 active and latent *Mycobacterium tuberculosis* (TB) infection based on the one or more transcriptional gene expression modules that differentiate between active and latent infection. In one aspect, the invention may also include the step of using the determined comparative gene product information to formulate a diagnosis.

In another aspect, the method may also include the step of using the determined comparative gene product information to formulate a prognosis or the step of using the determined comparative gene product
30 information to formulate a treatment plan. In one alternative aspect, the method may include the step of distinguishing patients with latent TB from active TB patients. In one aspect, the module may include a dataset of the genes in modules M1.2, M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 to detect active pulmonary infection. In another aspect, the module may include a dataset of the genes in modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 to detect a latent

infection. In yet another aspect, the following genes are down-regulated in active pulmonary infection CD3, CTLA-4, CD28, ZAP-70, IL-7R, CD2, SLAM, CCR7 and GATA-3. In one specific aspect, the expression profile of the modules in Figure 9 is indicative of active pulmonary infection and the expression profile of the modules in Figure 10 is indicative of latent infection. It has been found that the underexpression of genes in modules M3.4, M3.6, M3.7, M3.8 and M3.9 is indicative of active infection. It has also been found that the overexpression of genes in modules M3.1 is indicative of active infection.

In yet another aspect of the present invention, the method may also include the step of distinguishing TB infection from other bacterial infections by determining the gene expression in modules M2.2, M2.3 and M3.5, which are overexpressed by the peripheral blood mononuclear cells or whole blood in infection other than *Mycobacterium*. Alternatively, the method may include the step of distinguishing the differential and reciprocal transcriptional signatures in the blood of latent and active TB patients using two or more of the following modules: M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection. Examples of the genes that are upregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7A, 7D, 7I, 7J and 7K. Further examples of the genes that are downregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7B, 7C, 7E, 7F, 7G, 7H, 7L, 7M, 7N, 7O and 7P. In one specific aspect, the genes that are upregulated in latent TB infection versus a healthy patient may be selected from Table 8B. In another specific aspect, the genes that are downregulated in latent TB infection versus a healthy patient may be selected from Tables 8A, 8C, 8D, 8E and 8F.

Another embodiment of the present invention is a method for distinguishing between active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the method including the steps of: obtaining a first gene expression dataset obtained from a first clinical group with active *Mycobacterium tuberculosis* infection, a second gene expression dataset obtained from a second clinical group with a latent *Mycobacterium tuberculosis* infection patient and a third gene expression dataset obtained from a clinical group of non-infected individuals; generating a gene cluster dataset comprising the differential expression of genes between any two of the first, second and third datasets; and determining a unique pattern of expression/representation that is indicative of latent infection, active infection or being healthy. In one aspect, each clinical group is separated into a unique pattern of expression/representation for each of the 119 genes of Table 6. In another aspect, values for the first and third datasets are compared and the values for the dataset from the third dataset are subtracted therefrom. In another specific aspect, the values for the second and third datasets are compared and the values for the dataset from the third dataset are subtracted therefrom. In one specific embodiment, the method may further include the step of comparing values for two different datasets and subtracting the values for the remaining dataset to distinguish between a patient with a latent infection, a patient with an active infection and a non-

infected individual. In one aspect, the method may further comprise the step of using the determined comparative gene product information to formulate a diagnosis or a prognosis. In yet another aspect, the method includes the step of using the determined comparative gene product information to formulate a treatment plan. The method may also include the step of distinguishing patients with latent TB from active TB patients by analyzing the expression/representation of genes in the gene and patient clusters.

In one specific aspect, the method may further include the step of determining the expression levels of the genes: ST3GAL6, PAD14, TNFRSF12A, VAMP3, BR13, RGS19, PILRA, NCF1, LOC652616, PLAUR(CD87), SIGLEC5, B3GALT7, IBRDC3(NKLAM), ALOX5AP(FLAP), MMP9, ANPEP(APN), NALP12, CSF2RA, IL6R(CD126), RASGRP4, TNFSF14(CD258), NCF4, HK2, ARID3A, PGLYRP1(PGRP), which are underexpressed/underrepresented in the blood of Latent TB patients but not in the blood of Healthy individuals or Active TB patients. In another specific aspect, the method may further include the step of determining the expression levels of the genes: ABCG1, SREBF1, RBP7(CRBP4), C22orf5, FAM101B, S100P, LOC649377, UBTD1, PSTPIP-1, RENBP, PGM2, SULF2, FAM7A1, HOMTES-103, NDUFAF1, CES1, CYP27A1, FLJ33641, GPR177, MID1IP1(MIG-12), PSD4, SF3A1, NOV(CCN3), SGK(SGK1), CDK5R1, LOC642035, which are overexpressed/overrepresented in the blood of Healthy control individuals but were underexpressed/underrepresented in the blood of Latent TB patients, and underexpressed/underrepresented in the blood of Active TB patients. In another specific aspect, the method may further include the step of determining the expression levels of the genes: ARSG, LOC284757, MDM4, CRNKL1, IL8, LOC389541, CD300LB, NIN, PHKG2, HIP1, which are overexpressed/overrepresented in the blood of Healthy individuals, are underexpressed/underrepresented in the blood of both Latent and Active TB patients. In one specific aspect, the method may further include the step of determining the expression levels of the genes: PSMB8(LMP7), APOL6, GBP2, GBP5, GBP4, ATF3, GCH1, VAMP5, WARS, LIMK1, NPC2, IL-15, LMTK2, STX11(FHL4), which are overexpressed/overrepresented in the blood of Active TB, and underexpressed/underrepresented in the blood of Latent TB patients and Healthy control individuals. In one specific aspect, the method may further include the step of determining the expression levels of the genes: FLJ11259(DRAM), JAK2, GSDMDC1(DF5L)(FKSG10), SIPAIL1, [2680400](KIAA1632), ACTA2(ACTSA), KCNMB1(SLO-BETA), which are overexpressed/overrepresented in blood from Active TB patients, and underexpressed/underrepresented in the blood from Latent TB patients and Healthy control individuals. In one specific aspect, the method may further include the step of determining the expression levels of the genes: SPTAN1, KIAAD179(Nnp1)(RRP1), FAM84B(NSE2), SELM, IL27RA, MRPS34, [6940246](IL23A), PRKCA(PKCA), CCDC41, CD52(CDW52), [3890241](ZN404), MCCC1(MCCA/B), SOX8, SYNJ2, FLJ21127, FHIT, which are underexpressed/underrepresented in the blood of Active TB patients but not in the blood of Latent TB patients or Healthy Control individuals. In one specific aspect, the method may further include the step of determining the expression levels of the genes: CDKL1(p42),

MICALCL, MBNL3, RHD, ST7(RAY1), PPR3R1, [360739](PIP5K2A), AMFR, FLJ22471, CRAT(CAT1), PLA2G4C, ACOT7(ACT)(ACH1), RNF182, KLRC3(NKG2E), HLA-DPB1, which are underexpressed/underrepresented in the blood of Healthy Control individuals, overexpressed/overrepresented in the blood of the Latent TB patients, and overexpressed/overrepresented in the blood of Active TB patients.

5 Yet another embodiment of the present invention is a method for distinguishing between active and latent mycobacterium tuberculosis infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the method including the steps of: obtaining a gene expression dataset from a whole blood sample; sorting the gene expression dataset into one or more transcriptional gene expression modules; and mapping the differential expression of the one or more transcriptional gene expression modules that
10 distinguish between active and latent *Mycobacterium tuberculosis* infection, thereby distinguishing between active and latent *Mycobacterium tuberculosis* infection. In one aspect, the dataset includes TRIM genes. In one aspect, the dataset includes TRIM genes, specifically, TRIM 5, 6, 19(PML), 21, 22, 25, 68 are overrepresented/expressed in active pulmonary TB. In one aspect, the dataset of TRIM genes, includes TRIM 28, 32, 51, 52, 68, are underrepresented/expressed in active pulmonary TB.

15 Another embodiment of the present invention is a method of diagnosing a patient with active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with mycobacterium tuberculosis, the method comprising detecting differential expression of one or more transcriptional gene expression modules that distinguish between infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more
20 transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent mycobacterium tuberculosis infection. In another aspect, the method includes one or more of the step of: using the determined comparative gene product information to formulate a diagnosis, the step of using the determined comparative gene product information to formulate a prognosis and the step of using the determined comparative gene product information to formulate a
25 treatment plan. In one alternative aspect, the method may include the step of distinguishing patients with latent TB from active TB patients. In one aspect, the module may include a dataset of the genes in modules M1.2, M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 to detect active pulmonary infection. In another aspect, the module may include a dataset of the genes in modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 to detect a latent infection. In yet another aspect, the
30 following genes are down-regulated in active pulmonary infection CD3, CTLA-4, CD28, ZAP-70, IL-7R, CD2, SLAM, CCR7 and GATA-3. In one specific aspect, the expression profile of the modules in Figure 9 is indicative of active pulmonary infection and the expression profile of the modules in Figure 10 is indicative of latent infection. It has been found that the underexpression of genes in modules M3.4, M3.6, M3.7, M3.8 and M3.9 is indicative of active infection. It has also been found that the overexpression of
35 genes in modules M3.1 is indicative of active infection.

In yet another aspect of the present invention, the method may also include the step of distinguishing TB infection from other bacterial infections by determining the gene expression in modules M2.2, M2.3 and M3.5, which are overexpressed by the peripheral blood mononuclear cells or whole blood in infection other than *Mycobacterium*. Alternatively, the method may include the step of distinguishing the differential and reciprocal transcriptional signatures in the blood of latent and active TB patients using two or more of the following modules: M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection. Examples of the genes that are upregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7A, 7D, 7I, 7J and 7K. Further examples of the genes that are downregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7B, 7C, 7E, 7F, 7G, 7H, 7L, 7M, 7N, 7O and 7P. In one specific aspect, the genes that are upregulated in latent TB infection versus a healthy patient may be selected from Table 8B. In another specific aspect, the genes that are downregulated in latent TB infection versus a healthy patient may be selected from Tables 8A, 8C, 8D, 8E and 8F.

Another embodiment of the present invention is a kit for diagnosing a patient with active and latent mycobacterium tuberculosis infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the kit that includes a gene expression detector for obtaining a gene expression dataset from the patient; and a processor capable of comparing the gene expression to pre-defined gene module dataset that distinguish between infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent *Mycobacterium tuberculosis* infection.

Yet another embodiment includes a system of diagnosing a patient with active and latent *Mycobacterium tuberculosis* infection comprising: a gene expression dataset from the patient; and a processor capable of comparing the gene expression to pre-defined gene module dataset that distinguish between infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent *Mycobacterium tuberculosis* infection, wherein the modules are selected from M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection.

BRIEF DESCRIPTION OF THE DRAWINGS

For a more complete understanding of the features and advantages of the present invention, reference is now made to the detailed description of the invention along with the accompanying figures and in which:

Figure 1 shows the gene array expression results from 42 participants, genes present in at least 2 samples (PAL2), genes 2 folds over or under represented compared with median, clustered by Pearson Correlation comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

5 Figure 2 shows the gene array expression results from PAL2, 2 folds up or down expressed, filtered for statistically significant differences in expression between clinical groups using a non-parametric test (Kruskal-Wallis), $P < 0.01$, with Benjamini-Hochberg correction (1473 genes) and independently clustered using Pearson correlation comparing active PTB, latent TB and healthy controls;

10 Figures 3A – 3D show the gene array expression results from PAL2, 2 folds up or down expressed, filtered for statistically significant differences in expression between clinical groups using a non-parametric test (Kruskal-Wallis), $P < 0.01$, with Benjamini-Hochberg correction, and then filtered for the presence of the gene ontology term for biological process “immune response” in the gene annotation and independently clustered using Pearson correlation (158 genes). These 158 genes are shown separated into 4 figures (3A – 3D) for legibility.

15 Figure 3A shows gene array expression results comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

Figure 3B shows gene array expression results comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

20 Figure 3C shows gene array expression results comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

Figure 3D shows gene array expression results comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

25 Figure 4 shows the gene array expression results from 42 participants, genes present in at least 2 samples (PAL2), genes 2 folds over or under represented compared with median, Genes selected as TRIMs - clustered by Pearson Correlation comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

30 Figure 5A shows detail from the gene array expression results from 42 participants, genes present in at least 2 samples (PAL2), genes 2 folds over or under represented compared with median, clustered by Pearson Correlation comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls, showing that inhibitory immunoregulatory ligands (PDL1/CD274, PDL2/CD273) are overexpressed in active TB patients.

Figure 5B shows the unfiltered gene array expression results that demonstrate that PDL1 is only expressed in active TB patients;

Figure 6 shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, statistically significantly differentially expressed across groups ($P < 0.1$, Kruskal-Wallis non-parametric test with Bonferroni correction) (46 genes) independently clustered using Pearson correlation, comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

Figure 7 shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, statistically significantly differentially expressed across groups ($P < 0.05$, Kruskal-Wallis non-parametric test with Bonferroni correction) (18 genes) independently clustered using Pearson correlation, comparing active PTB, latent TB, healthy BCG non-vaccinated controls and healthy BCG vaccinated controls;

Figure 8A shows that the results of merging different statistical filters applied to the list of genes filtered present in at least 2 samples, 2 folds up or down 'represented' compared to median, discriminates between all three clinical groups. The transcripts shown are statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction) plus the transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction) – 119 genes in total independently clustered using Pearson correlation (clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically); These 119 genes are shown separated into 5 further figures (8B –8F) for legibility and to show that subgroups of these genes may also be used to distinguish between different clinical groups (i.e. between Active, Latent and Healthy).

Figure 8B shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, transcripts statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction) PLUS transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction) (clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically);

Figure 8C shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, transcripts statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction) PLUS transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction);

Figure 8D shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, transcripts statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction)

PLUS transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction) (clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically);

5 Figure 8E shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, transcripts statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction) PLUS transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction) (clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically);

10 Figure 8F shows the gene array expression results filtered for genes present in at least 2 samples, 2 folds up or down 'represented' compared to median, transcripts statistically significantly differentially expressed between Latent and healthy ($P < 0.005$, Wilcoxon-Mann-Whitney non-parametric test with no correction) PLUS transcripts statistically significantly differentially expressed between Active and healthy ($P < 0.5$, Wilcoxon-Mann-Whitney non-parametric test with Bonferroni correction) (clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically);

15 Figure 9 shows the gene array expression results from a gene module analysis of PTB(9) vs Control(6): from 5281 genes, filtered for PAL2, statistically significantly differentially expressed between active PTB and healthy controls by Wilcoxon-Mann-Whitney-test, $p < 0.05$, with no multi-test correction; and

20 Figure 10 shows the gene array expression results from from a gene module analysis of LTB(9) vs Control(6): from - 3137 genes, filtered for PAL2, statistically significantly differentially expressed between active PTB and healthy controls by Wilcoxon-Mann-Whitney-test, $p < 0.05$, with no multi-test correction.

DETAILED DESCRIPTION OF THE INVENTION

25 While the making and using of various embodiments of the present invention are discussed in detail below, it should be appreciated that the present invention provides many applicable inventive concepts that can be embodied in a wide variety of specific contexts. The specific embodiments discussed herein are merely illustrative of specific ways to make and use the invention and do not delimit the scope of the invention.

30 To facilitate the understanding of this invention, a number of terms are defined below. Terms defined herein have meanings as commonly understood by a person of ordinary skill in the areas relevant to the present invention. Terms such as "a", "an" and "the" are not intended to refer to only a singular entity, but include the general class of which a specific example may be used for illustration. The terminology herein is used to describe specific embodiments of the invention, but their usage does not delimit the invention, except as outlined in the claims. Unless defined otherwise, all technical and scientific terms used herein have the meaning commonly understood by a person skilled in the art to which this invention belongs. The following

references provide one of skill with a general definition of many of the terms used in this invention: Singleton et al., Dictionary of Microbiology and Molecular Biology (2d ed. 1994); The Cambridge Dictionary of Science and Technology (Walker ed., 1988); The Glossary of Genetics, 5TH ED., R. Rieger et al. (eds.), Springer Verlag (1991); and Hale & Marham, The Harper Collins Dictionary of Biology (1991).

5 Various biochemical and molecular biology methods are well known in the art. For example, methods of isolation and purification of nucleic acids are described in detail in WO 97/10365; WO 97/27317; Chapter 3 of Laboratory Techniques in Biochemistry and Molecular Biology: Hybridization with Nucleic Acid Probes, Part I. Theory and Nucleic Acid Preparation, (P. Tijssen, ed.) Elsevier, N.Y. (1993); Sambrook, et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press, N.Y., (1989); and Current Protocols in
10 Molecular Biology, (Ausubel, F. M. et al., eds.) John Wiley & Sons, Inc., New York (1987-1999), including supplements.

BIOINFORMATICS DEFINITIONS

As used herein, an “object” refers to any item or information of interest (generally textual, including noun, verb, adjective, adverb, phrase, sentence, symbol, numeric characters, etc.). Therefore, an object is anything
15 that can form a relationship and anything that can be obtained, identified, and/or searched from a source. “Objects” include, but are not limited to, an entity of interest such as gene, protein, disease, phenotype, mechanism, drug, etc. In some aspects, an object may be data, as further described below.

As used herein, a “relationship” refers to the co-occurrence of objects within the same unit (e.g., a phrase, sentence, two or more lines of text, a paragraph, a section of a webpage, a page, a magazine, paper, book,
20 etc.). It may be text, symbols, numbers and combinations, thereof

As used herein, “meta data content” refers to information as to the organization of text in a data source. Meta data can comprise standard metadata such as Dublin Core metadata or can be collection-specific. Examples of metadata formats include, but are not limited to, Machine Readable Catalog (MARC) records used for library catalogs, Resource Description Format (RDF) and the Extensible Markup Language (XML). Meta
25 objects may be generated manually or through automated information extraction algorithms.

As used herein, an “engine” refers to a program that performs a core or essential function for other programs. For example, an engine may be a central program in an operating system or application program that coordinates the overall operation of other programs. The term “engine” may also refer to a program containing an algorithm that can be changed. For example, a knowledge discovery engine may be designed
30 so that its approach to identifying relationships can be changed to reflect new rules of identifying and ranking relationships.

As used herein, “semantic analysis” refers to the identification of relationships between words that represent similar concepts, e.g., through suffix removal or stemming or by employing a thesaurus. “Statistical analysis”

refers to a technique based on counting the number of occurrences of each term (word, word root, word stem, n-gram, phrase, etc.). In collections unrestricted as to subject, the same phrase used in different contexts may represent different concepts. Statistical analysis of phrase co-occurrence can help to resolve word sense ambiguity. “Syntactic analysis” can be used to further decrease ambiguity by part-of-speech analysis. As used herein, one or more of such analyses are referred to more generally as “lexical analysis.” “Artificial intelligence (AI)” refers to methods by which a non-human device, such as a computer, performs tasks that humans would deem noteworthy or “intelligent.” Examples include identifying pictures, understanding spoken words or written text, and solving problems.

Terms such “data”, “dataset” and “information” are often used interchangeably, as are “information” and “knowledge.” As used herein, “data” is the most fundamental unit that is an empirical measurement or set of measurements. Data is compiled to contribute to information, but it is fundamentally independent of it and may be combined into a dataset, that is, a set of data. Information, by contrast, is derived from interests, e.g., data (the unit) may be gathered on ethnicity, gender, height, weight and diet for the purpose of finding variables correlated with risk of cardiovascular disease. However, the same data could be used to develop a formula or to create “information” about dietary preferences, i.e., likelihood that certain products in a supermarket have a higher likelihood of selling.

As used herein, the term “database” refers to repositories for raw or compiled data, even if various informational facets can be found within the data fields. A database may include one or more datasets. A database is typically organized so its contents can be accessed, managed, and updated (e.g., the database is dynamic). The term “database” and “source” are also used interchangeably in the present invention, because primary sources of data and information are databases. However, a “source database” or “source data” refers in general to data, e.g., unstructured text and/or structured data that are input into the system for identifying objects and determining relationships. A source database may or may not be a relational database. However, a system database usually includes a relational database or some equivalent type of database which stores values relating to relationships between objects.

As used herein, a “system database” and “relational database” are used interchangeably and refer to one or more collections of data organized as a set of tables containing data fitted into predefined categories. For example, a database table may comprise one or more categories defined by columns (e.g. attributes), while rows of the database may contain a unique object for the categories defined by the columns. Thus, an object such as the identity of a gene might have columns for its presence, absence and/or level of expression of the gene. A row of a relational database may also be referred to as a “set” and is generally defined by the values of its columns. A “domain” in the context of a relational database is a range of valid values a field such as a column may include.

As used herein, a “domain of knowledge” refers to an area of study over which the system is operative, for example, all biomedical data. It should be pointed out that there is advantage to combining data from several domains, for example, biomedical data and engineering data, for this diverse data can sometimes link things that cannot be put together for a normal person that is only familiar with one area or research/study (one domain). A “distributed database” refers to a database that may be dispersed or replicated among different points in a network.

As used herein, “information” refers to a data set that may include numbers, letters, sets of numbers, sets of letters, or conclusions resulting or derived from a set of data. “Data” is then a measurement or statistic and the fundamental unit of information. “Information” may also include other types of data such as words, symbols, text, such as unstructured free text, code, etc. “Knowledge” is loosely defined as a set of information that gives sufficient understanding of a system to model cause and effect. To extend the previous example, information on demographics, gender and prior purchases may be used to develop a regional marketing strategy for food sales while information on nationality could be used by buyers as a guideline for importation of products. It is important to note that there are no strict boundaries between data, information, and knowledge; the three terms are, at times, considered to be equivalent. In general, data comes from examining, information comes from correlating, and knowledge comes from modeling.

As used herein, “a program” or “computer program” refers generally to a syntactic unit that conforms to the rules of a particular programming language and that is composed of declarations and statements or instructions, divisible into, “code segments” needed to solve or execute a certain function, task, or problem. A programming language is generally an artificial language for expressing programs.

As used herein, a “system” or a “computer system” generally refers to one or more computers, peripheral equipment, and software that perform data processing. A “user” or “system operator” in general includes a person, that uses a computer network accessed through a “user device” (e.g., a computer, a wireless device, etc) for the purpose of data processing and information exchange. A “computer” is generally a functional unit that can perform substantial computations, including numerous arithmetic operations and logic operations without human intervention.

As used herein, “application software” or an “application program” refers generally to software or a program that is specific to the solution of an application problem. An “application problem” is generally a problem submitted by an end user and requiring information processing for its solution.

As used herein, a “natural language” refers to a language whose rules are based on current usage without being specifically prescribed, e.g., English, Spanish or Chinese. As used herein, an “artificial language” refers to a language whose rules are explicitly established prior to its use, e.g., computer-programming languages such as C, C++, Java, BASIC, FORTRAN, or COBOL.

As used herein, “statistical relevance” refers to using one or more of the ranking schemes (O/E ratio, strength, etc.), where a relationship is determined to be statistically relevant if it occurs significantly more frequently than would be expected by random chance.

As used herein, the terms “coordinately regulated genes” or “transcriptional modules” are used interchangeably to refer to grouped, gene expression profiles (e.g., signal values associated with a specific gene sequence) of specific genes. Each transcriptional module correlates two key pieces of data, a literature search portion and actual empirical gene expression value data obtained from a gene microarray. The set of genes that is selected into a transcriptional modules is based on the analysis of gene expression data (module extraction algorithm described above). Additional steps are taught by Chaussabel, D. & Sher, A. Mining microarray expression data by literature profiling. *Genome Biol* 3, RESEARCH0055 (2002), (<http://genomebiology.com/2002/3/10/research/0055>) relevant portions incorporated herein by reference and expression data obtained from a disease or condition of interest, e.g., Systemic Lupus erythematosus, arthritis, lymphoma, carcinoma, melanoma, acute infection, autoimmune disorders, autoinflammatory disorders, etc.).

The Table below lists examples of keywords that were used to develop the literature search portion or contribution to the transcription modules. The skilled artisan will recognize that other terms may easily be selected for other conditions, e.g., specific cancers, specific infectious disease, transplantation, etc. For example, genes and signals for those genes associated with T cell activation are described hereinbelow as Module ID “M 2.8” in which certain keywords (e.g., Lymphoma, T-cell, CD4, CD8, TCR, Thymus, Lymphoid, IL2) were used to identify key T-cell associated genes, e.g., T-cell surface markers (CD5, CD6, CD7, CD26, CD28, CD96); molecules expressed by lymphoid lineage cells (lymphotoxin beta, IL2-inducible T-cell kinase, TCF7; and T-cell differentiation protein mal, GATA3, STAT5B). Next, the complete module is developed by correlating data from a patient population for these genes (regardless of platform, presence/absence and/or up or downregulation) to generate the transcriptional module. In some cases, the gene profile does not match (at this time) any particular clustering of genes for these disease conditions and data, however, certain physiological pathways (e.g., cAMP signaling, zinc-finger proteins, cell surface markers, etc.) are found within the “Underdetermined” modules. In fact, the gene expression data set may be used to extract genes that have coordinated expression prior to matching to the keyword search, i.e., either data set may be correlated prior to cross-referencing with the second data set.

Table 1. Transcriptional Modules

Example Module I.D.	Example Keyword selection	Gene Profile Assessment
M 1.1	Ig, Immunoglobulin, Bone, Marrow, PreB, IgM, Mu.	Plasma cells: Includes genes encoding for Immunoglobulin chains (e.g.IGHM, IGJ, IGLL1, IGKC, IGHD) and the plasma cell marker CD38.
M 1.2	Platelet, Adhesion, Aggregation, Endothelial,	Platelets: Includes genes encoding for platelet glycoproteins (ITGA2B, ITGB3, GP6, GP1A/B), and platelet-derived immune

Example Module I.D.	Example Keyword selection	Gene Profile Assessment
	Vascular	mediators such as PPPB (pro-platelet basic protein) and PF4 (platelet factor 4).
M 1.3	Immunoreceptor, BCR, B-cell, IgG	B-cells: Includes genes encoding for B-cell surface markers (CD72, CD79A/B, CD19, CD22) and other B-cell associated molecules: Early B-cell factor (EBF), B-cell linker (BLNK) and B lymphoid tyrosine kinase (BLK).
M 1.4	Replication, Repression, Repair, CREB, Lymphoid, TNF-alpha	Undetermined. This set includes regulators and targets of cAMP signaling pathway (JUND, ATF4, CREM, PDE4, NR4A2, VIL2), as well as repressors of TNF-alpha mediated NF-KB activation (CYLD, ASK, TNFAIP3).
M 1.5	Monocytes, Dendritic, MHC, Costimulatory, TLR4, MYD88	Myeloid lineage: Includes molecules expressed by cells of the myeloid lineage (CD86, CD163, FCGR2A), some of which being involved in pathogen recognition (CD14, TLR2, MYD88). This set also includes TNF family members (TNFR2, BAFF).
M 1.6	Zinc, Finger, P53, RAS	Undetermined. This set includes genes encoding for signaling molecules, e.g., the zinc finger containing inhibitor of activated STAT (PIAS1 and PIAS2), or the nuclear factor of activated T-cells NFATC3.
M 1.7	Ribosome, Translational, 40S, 60S, HLA	MHC/Ribosomal proteins: Almost exclusively formed by genes encoding MHC class I molecules (HLA-A,B,C,G,E)+ Beta 2-microglobulin (B2M) or Ribosomal proteins (RPLs, RPSs).
M 1.8	Metabolism, Biosynthesis, Replication, Helicase	Undetermined. Includes genes encoding metabolic enzymes (GLS, NSF1, NAT1) and factors involved in DNA replication (PURA, TERF2, EIF2S1).
M 2.1	NK, Killer, Cytolytic, CD8, Cell-mediated, T-cell, CTL, IFN-g	Cytotoxic cells: Includes cytotoxic T-cells and NK-cells surface markers (CD8A, CD2, CD160, NKG7, KLRs), cytolytic molecules (granzyme, perforin, granulysin), chemokines (CCL5, XCL1) and CTL/NK-cell associated molecules (CTSW).
M 2.2	Granulocytes, Neutrophils, Defense, Myeloid, Marrow	Neutrophils: This set includes innate molecules that are found in neutrophil granules (Lactotransferrin: LTF, defensin: DEAF1, Bacterial Permeability Increasing protein: BPI, Cathelicidin antimicrobial protein: CAMP).
M 2.3	Erythrocytes, Red, Anemia, Globin, Hemoglobin	Erythrocytes: Includes hemoglobin genes (HGBs) and other erythrocyte-associated genes (erythrocytic alkaline phosphatase: ANK1, Glycophorin C: GYPC, hydroxymethylbilane synthase: HMBS, erythroid associated factor: ERAF).
M 2.4	Ribonucleoprotein, 60S, nucleolus, Assembly, Elongation	Ribosomal proteins: Including genes encoding ribosomal proteins (RPLs, RPSs), Eukaryotic Translation Elongation factor family members (EEFs) and Nucleolar proteins (NPM1, NOAL2, NAP1L1).
M 2.5	Adenoma, Interstitial, Mesenchyme, Dendrite, Motor	Undetermined. This module includes genes encoding immune-related (CD40, CD80, CXCL12, IFNA5, IL4R) as well as cytoskeleton-related molecules (Myosin, Dedicator of Cytokinesis, Syndecan 2, Plexin C1, Distrobrevin).
M 2.6	Granulocytes, Monocytes, Myeloid, ERK, Necrosis	Myeloid lineage: Related to M 1.5. Includes genes expressed in myeloid lineage cells (IGTB2/CD18, Lymphotoxin beta receptor, Myeloid related proteins 8/14 Formyl peptide receptor 1), such as Monocytes and Neutrophils:
M 2.7	No keywords extracted.	Undetermined. This module is largely composed of transcripts with no known function. Only 20 genes associated with literature, including a member of the chemokine-like factor superfamily (CKLFSF8).
M 2.8	Lymphoma, T-cell, CD4, CD8, TCR, Thymus, Lymphoid, IL2	T-cells: Includes T-cell surface markers (CD5, CD6, CD7, CD26, CD28, CD96) and molecules expressed by lymphoid lineage cells (lymphotoxin beta, IL2-inducible T-cell kinase, TCF7, T-cell differentiation protein mal, GATA3, STAT5B).
M 2.9	ERK, Transactivation,	Undetermined. Includes genes encoding molecules that associate to

Example Module I.D.	Example Keyword selection	Gene Profile Assessment
	Cytoskeletal, MAPK, JNK	the cytoskeleton (Actin related protein 2/3, MAPK1, MAP3K1, RAB5A). Also present are T-cell expressed genes (FAS, ITGA4/CD49D, ZNF1A1).
M 2.10	Myeloid, Macrophage, Dendritic, Inflammatory, Interleukin	Undetermined. Includes genes encoding for Immune-related cell surface molecules (CD36, CD86, LILRB), cytokines (IL15) and molecules involved in signaling pathways (FYB, TICAM2-Toll-like receptor pathway).
M 2.11	Replication, Repress, RAS, Autophosphorylation, Oncogenic	Undetermined. Includes kinases (UHMK1, CSNK1G1, CDK6, WNK1, TAOK1, CALM2, PRKCI, ITPKB, SRPK2, STK17B, DYRK2, PIK3R1, STK4, CLK4, PKN2) and RAS family members (G3BP, RAB14, RASA2, RAP2A, KRAS).
M 3.1	ISRE, Influenza, Antiviral, IFN-gamma, IFN-alpha, Interferon	Interferon-inducible: This set includes interferon-inducible genes: antiviral molecules (OAS1/2/3/L, GBP1, G1P2, EIF2AK2/PKR, MX1, PML), chemokines (CXCL10/IP-10), signaling molecules (STAT1, STAT2, IRF7, ISGF3G).
M 3.2	TGF-beta, TNF, Inflammatory, Apoptotic, Lipopolysaccharide	Inflammation I: Includes genes encoding molecules involved in inflammatory processes (e.g., IL8, ICAM1, C5R1, CD44, PLAUR, IL1A, CXCL16), and regulators of apoptosis (MCL1, FOXO3A, RARA, BCL3/6/2A1, GADD45B).
M 3.3	Granulocyte, Inflammatory, Defense, Oxidize, Lysosomal	Inflammation II: Includes molecules inducing or inducible by Granulocyte-Macrophage CSF (SPI1, IL18, ALOX5, ANPEP), as well as lysosomal enzymes (PPT1, CTSB/S, CES1, NEU1, ASAH1, LAMP2, CAST).
M 3.4	No keyword extracted	Undetermined. Includes protein phosphates (PPP1R12A, PTPRC, PPP1CB, PPM1B) and phosphoinositide 3-kinase (PI3K) family members (PIK3CA, PIK32A, PIP5K3).
M 3.5	No keyword extracted	Undetermined. Composed of only a small number of transcripts. Includes hemoglobin genes (HBA1, HBA2, HBB).
M 3.6	Complement, Host, Oxidative, Cytoskeletal, T-cell	Undetermined. Large set that includes T-cell surface markers (CD101, CD102, CD103) as well as molecules ubiquitously expressed among blood leukocytes (CXCR1: fractalkine receptor, CD47, P-selectin ligand).
M 3.7	Spliceosome, Methylation, Ubiquitin, Beta-catenin	Undetermined. Includes genes encoding proteasome subunits (PSMA2/5, PSMB5/8); ubiquitin protein ligases HIP2, STUB1, as well as components of ubiquitin ligase complexes (SUGT1).
M 3.8	CDC, TCR, CREB, Glycosylase	Undetermined. Includes genes encoding for several enzymes: aminomethyltransferase, arginyltransferase, asparagine synthetase, diacylglycerol kinase, inositol phosphatases, methyltransferases, helicases...
M 3.9	Chromatin, Checkpoint, Replication, Transactivation	Undetermined. Includes genes encoding for protein kinases (PRKPIR, PRKDC, PRKCI) and phosphatases (e.g., PTPLB, PPP1R8/2CB). Also includes RAS oncogene family members and the NK cell receptor 2B4 (CD244).

BIOLOGICAL DEFINITIONS

As used herein, the term “array” refers to a solid support or substrate with one or more peptides or nucleic acid probes attached to the support. Arrays typically have one or more different nucleic acid or peptide probes that are coupled to a surface of a substrate in different, known locations. These arrays, also described as “microarrays” or “gene-chips” that may have 10,000; 20,000; 30,000; or 40,000 different identifiable genes based on the known genome, e.g., the human genome. These pan-arrays are used to detect the entire “transcriptome” or transcriptional pool of genes that are expressed or found in a sample, e.g., nucleic acids

that are expressed as RNA, mRNA and the like that may be subjected to RT and/or RT-PCR to make a complementary set of DNA replicons. Arrays may be produced using mechanical synthesis methods, light directed synthesis methods and the like that incorporate a combination of non-lithographic and/or photolithographic methods and solid phase synthesis methods.

5 Various techniques for the synthesis of these nucleic acid arrays have been described, e.g., fabricated on a surface of virtually any shape or even a multiplicity of surfaces. Arrays may be peptides or nucleic acids on beads, gels, polymeric surfaces, fibers such as fiber optics, glass or any other appropriate substrate. Arrays may be packaged in such a manner as to allow for diagnostics or other manipulation of an all inclusive device, see for example, U.S. Pat. No. 6,955,788, relevant portions incorporated herein by reference.

10 As used herein, the term “disease” refers to a physiological state of an organism with any abnormal biological state of a cell. Disease includes, but is not limited to, an interruption, cessation or disorder of cells, tissues, body functions, systems or organs that may be inherent, inherited, caused by an infection, caused by abnormal cell function, abnormal cell division and the like. A disease that leads to a “disease state” is generally detrimental to the biological system, that is, the host of the disease. With respect to the
15 present invention, any biological state, such as an infection (e.g., viral, bacterial, fungal, helminthic, etc.), inflammation, autoinflammation, autoimmunity, anaphylaxis, allergies, premalignancy, malignancy, surgical, transplantation, physiological, and the like that is associated with a disease or disorder is considered to be a disease state. A pathological state is generally the equivalent of a disease state.

Disease states may also be categorized into different levels of disease state. As used herein, the level of a
20 disease or disease state is an arbitrary measure reflecting the progression of a disease or disease state as well as the physiological response upon, during and after treatment. Generally, a disease or disease state will progress through levels or stages, wherein the affects of the disease become increasingly severe. The level of a disease state may be impacted by the physiological state of cells in the sample.

As used herein, the terms “therapy” or “therapeutic regimen” refer to those medical steps taken to alleviate or
25 alter a disease state, e.g., a course of treatment intended to reduce or eliminate the affects or symptoms of a disease using pharmacological, surgical, dietary and/or other techniques. A therapeutic regimen may include a prescribed dosage of one or more drugs or surgery. Therapies will most often be beneficial and reduce the disease state but in many instances the effect of a therapy will have non-desirable or side-effects. The effect of therapy will also be impacted by the physiological state of the host, e.g., age, gender, genetics, weight,
30 other disease conditions, etc.

As used herein, the term “pharmacological state” or “pharmacological status” refers to those samples that will be, are and/or were treated with one or more drugs, surgery and the like that may affect the pharmacological state of one or more nucleic acids in a sample, e.g., newly transcribed, stabilized and/or destabilized as a result of the pharmacological intervention. The pharmacological state of a sample relates to

changes in the biological status before, during and/or after drug treatment and may serve a diagnostic or prognostic function, as taught herein. Some changes following drug treatment or surgery may be relevant to the disease state and/or may be unrelated side-effects of the therapy. Changes in the pharmacological state are the likely results of the duration of therapy, types and doses of drugs prescribed, degree of compliance
5 with a given course of therapy, and/or un-prescribed drugs ingested.

As used herein, the term “biological state” refers to the state of the transcriptome (that is the entire collection of RNA transcripts) of the cellular sample isolated and purified for the analysis of changes in expression. The biological state reflects the physiological state of the cells in the sample by measuring the abundance and/or activity of cellular constituents, characterizing according to morphological phenotype or a
10 combination of the methods for the detection of transcripts.

As used herein, the term “expression profile” refers to the relative abundance of RNA, DNA or protein abundances or activity levels. The expression profile can be a measurement for example of the transcriptional state or the translational state by any number of methods and using any of a number of gene-chips, gene arrays, beads, multiplex PCR, quantitative PCR, run-on assays, Northern blot analysis, Western
15 blot analysis, protein expression, fluorescence activated cell sorting (FACS), enzyme linked immunosorbent assays (ELISA), chemiluminescence studies, enzymatic assays, proliferation studies or any other method, apparatus and system for the determination and/or analysis of gene expression that are readily commercially available.

As used herein, the term “transcriptional state” of a sample includes the identities and relative abundances of
20 the RNA species, especially mRNAs present in the sample. The entire transcriptional state of a sample, that is the combination of identity and abundance of RNA, is also referred to herein as the transcriptome. Generally, a substantial fraction of all the relative constituents of the entire set of RNA species in the sample are measured.

As used herein, the term “modular transcriptional vectors” refers to transcriptional expression data that
25 reflects the “proportion of differentially expressed genes.” For example, for each module the proportion of transcripts differentially expressed between at least two groups (e.g. healthy subjects vs patients). This vector is derived from the comparison of two groups of samples. The first analytical step is used for the selection of disease-specific sets of transcripts within each module. Next, there is the “expression level.” The group comparison for a given disease provides the list of differentially expressed transcripts for each
30 module. It was found that different diseases yield different subsets of modular transcripts. With this expression level it is then possible to calculate vectors for each module(s) for a single sample by averaging expression values of disease-specific subsets of genes identified as being differentially expressed. This approach permits the generation of maps of modular expression vectors for a single sample, e.g., those described in the module maps disclosed herein. These vector module maps represent an averaged expression

level for each module (instead of a proportion of differentially expressed genes) that can be derived for each sample.

Using the present invention it is possible to identify and distinguish diseases not only at the module-level, but also at the gene-level; i.e., two diseases can have the same vector (identical proportion of differentially expressed transcripts, identical “polarity”), but the gene composition of the vector can still be disease-specific. Gene-level expression provides the distinct advantage of greatly increasing the resolution of the analysis. Furthermore, the present invention takes advantage of composite transcriptional markers. As used herein, the term “composite transcriptional markers” refers to the average expression values of multiple genes (subsets of modules) as compared to using individual genes as markers (and the composition of these markers can be disease-specific). The composite transcriptional markers approach is unique because the user can develop multivariate microarray scores to assess disease severity in patients with, e.g., SLE, or to derive expression vectors disclosed herein. Most importantly, it has been found that using the composite modular transcriptional markers of the present invention the results found herein are reproducible across microarray platform, thereby providing greater reliability for regulatory approval.

Gene expression monitoring systems for use with the present invention may include customized gene arrays with a limited and/or basic number of genes that are specific and/or customized for the one or more target diseases. Unlike the general, pan-genome arrays that are in customary use, the present invention provides for not only the use of these general pan-arrays for retrospective gene and genome analysis without the need to use a specific platform, but more importantly, it provides for the development of customized arrays that provide an optimal gene set for analysis without the need for the thousands of other, non-relevant genes. One distinct advantage of the optimized arrays and modules of the present invention over the existing art is a reduction in the financial costs (e.g., cost per assay, materials, equipment, time, personnel, training, etc.), and more importantly, the environmental cost of manufacturing pan-arrays where the vast majority of the data is irrelevant. The modules of the present invention allow for the first time the design of simple, custom arrays that provide optimal data with the least number of probes while maximizing the signal to noise ratio. By eliminating the total number of genes for analysis, it is possible to, e.g., eliminate the need to manufacture thousands of expensive platinum masks for photolithography during the manufacture of pan-genetic chips that provide vast amounts of irrelevant data. Using the present invention it is possible to completely avoid the need for microarrays if the limited probe set(s) of the present invention are used with, e.g., digital optical chemistry arrays, ball bead arrays, beads (e.g., Luminex), multiplex PCR, quantitative PCR, run-on assays, Northern blot analysis, or even, for protein analysis, e.g., Western blot analysis, 2-D and 3-D gel protein expression, MALDI, MALDI-TOF, fluorescence activated cell sorting (FACS) (cell surface or intracellular), enzyme linked immunosorbent assays (ELISA), chemiluminescence studies, enzymatic assays, proliferation studies or any other method, apparatus and system for the determination and/or analysis of gene expression that are readily commercially available.

The “molecular fingerprinting system” of the present invention may be used to facilitate and conduct a comparative analysis of expression in different cells or tissues, different subpopulations of the same cells or tissues, different physiological states of the same cells or tissue, different developmental stages of the same cells or tissue, or different cell populations of the same tissue against other diseases and/or normal cell controls. In some cases, the normal or wild-type expression data may be from samples analyzed at or about the same time or it may be expression data obtained or culled from existing gene array expression databases, e.g., public databases such as the NCBI Gene Expression Omnibus database.

As used herein, the term “differentially expressed” refers to the measurement of a cellular constituent (e.g., nucleic acid, protein, enzymatic activity and the like) that varies in two or more samples, e.g., between a disease sample and a normal sample. The cellular constituent may be on or off (present or absent), upregulated relative to a reference or downregulated relative to the reference. For use with gene-chips or gene-arrays, differential gene expression of nucleic acids, e.g., mRNA or other RNAs (miRNA, siRNA, hnRNA, rRNA, tRNA, etc.) may be used to distinguish between cell types or nucleic acids. Most commonly, the measurement of the transcriptional state of a cell is accomplished by quantitative reverse transcriptase (RT) and/or quantitative reverse transcriptase-polymerase chain reaction (RT-PCR), genomic expression analysis, post-translational analysis, modifications to genomic DNA, translocations, in situ hybridization and the like.

For some disease states it is possible to identify cellular or morphological differences, especially at early levels of the disease state. The present invention avoids the need to identify those specific mutations or one or more genes by looking at modules of genes of the cells themselves or, more importantly, of the cellular RNA expression of genes from immune effector cells that are acting within their regular physiologic context, that is, during immune activation, immune tolerance or even immune anergy. While a genetic mutation may result in a dramatic change in the expression levels of a group of genes, biological systems often compensate for changes by altering the expression of other genes. As a result of these internal compensation responses, many perturbations may have minimal effects on observable phenotypes of the system but profound effects to the composition of cellular constituents. Likewise, the actual copies of a gene transcript may not increase or decrease, however, the longevity or half-life of the transcript may be affected leading to greatly increases protein production. The present invention eliminates the need of detecting the actual message by, in one embodiment, looking at effector cells (e.g., leukocytes, lymphocytes and/or sub-populations thereof) rather than single messages and/or mutations.

The skilled artisan will appreciate readily that samples may be obtained from a variety of sources including, e.g., single cells, a collection of cells, tissue, cell culture and the like. In certain cases, it may even be possible to isolate sufficient RNA from cells found in, e.g., urine, blood, saliva, tissue or biopsy samples and the like. In certain circumstances, enough cells and/or RNA may be obtained from: mucosal secretion, feces, tears, blood plasma, peritoneal fluid, interstitial fluid, intradural, cerebrospinal fluid, sweat or other bodily

fluids. The nucleic acid source, e.g., from tissue or cell sources, may include a tissue biopsy sample, one or more sorted cell populations, cell culture, cell clones, transformed cells, biopies or a single cell. The tissue source may include, e.g., brain, liver, heart, kidney, lung, spleen, retina, bone, neural, lymph node, endocrine gland, reproductive organ, blood, nerve, vascular tissue, and olfactory epithelium.

5 The present invention includes the following basic components, which may be used alone or in combination, namely, one or more data mining algorithms; one or more module-level analytical processes; the characterization of blood leukocyte transcriptional modules; the use of aggregated modular data in multivariate analyses for the molecular diagnostic/prognostic of human diseases; and/or visualization of module-level data and results. Using the present invention it is also possible to develop and analyze
10 composite transcriptional markers, which may be further aggregated into a single multivariate score.

An explosion in data acquisition rates has spurred the development of mining tools and algorithms for the exploitation of microarray data and biomedical knowledge. Approaches aimed at uncovering the modular organization and function of transcriptional systems constitute promising methods for the identification of robust molecular signatures of disease. Indeed, such analyses can transform the perception of large scale
15 transcriptional studies by taking the conceptualization of microarray data past the level of individual genes or lists of genes.

The present inventors have recognized that current microarray-based research is facing significant challenges with the analysis of data that are notoriously “noisy,” that is, data that is difficult to interpret and does not compare well across laboratories and platforms. A widely accepted approach for the analysis of microarray
20 data begins with the identification of subsets of genes differentially expressed between study groups. Next, the users try subsequently to “make sense” out of resulting gene lists using pattern discovery algorithms and existing scientific knowledge.

Rather than deal with the great variability across platforms, the present inventors have developed a strategy that emphasized the selection of biologically relevant genes at an early stage of the analysis. Briefly, the
25 method includes the identification of the transcriptional components characterizing a given biological system for which an improved data mining algorithm was developed to analyze and extract groups of coordinately expressed genes, or transcriptional modules, from large collections of data.

Pulmonary tuberculosis (PTB) is a major and increasing cause of morbidity and mortality worldwide caused by *Mycobacterium tuberculosis* (*M. tuberculosis*). However, the majority of individuals infected with *M.*
30 *tuberculosis* remain asymptomatic, retaining the infection in a latent form and it is thought that this latent state is maintained by an active immune response. Blood is the pipeline of the immune system, and as such is the ideal biologic material from which the health and immune status of an individual can be established. Here, using microarray technology to assess the activity of the entire genome in blood cells, we identified distinct and reciprocal blood transcriptional biomarker signatures in patients with active pulmonary

tuberculosis and latent tuberculosis. These signatures were also distinct from those in control individuals. The signature of latent tuberculosis, which showed an over-representation of immune cytotoxic gene expression in whole blood, may help to determine protective immune factors against *M. tuberculosis* infection, since these patients are infected but most do not develop overt disease. This distinct transcriptional biomarker signature from active and latent TB patients may be also used to diagnose infection, and to monitor response to treatment with anti-mycobacterial drugs. In addition the signature in active tuberculosis patients will help to determine factors involved in immunopathogenesis and possibly lead to strategies for immune therapeutic intervention. This invention relates to a previous application that claimed the use of blood transcriptional biomarkers for the diagnosis of infections. However, this previous application did not disclose the existence of biomarkers for active and latent tuberculosis and focused rather on children with other acute infections (Ramillo, Blood, 2007).

The present identification of a transcriptional signature in blood from latent versus active TB patients can be used to test for patients with suspected *Mycobacterium tuberculosis* infection as well as for health screening/early detection of the disease. The invention also permits the evaluation of the response to treatment with anti-mycobacterial drugs. In this context, a test would also be particularly valuable in the context of drug trials, and particularly to assess drug treatments in Multi-Drug Resistant patients. Furthermore, the present invention may be used to obtain immediate, intermediate and long term data from the immune signature of latent tuberculosis to better define a protective immune response during vaccination trials. Also, the signature in active tuberculosis patients will help to determine factors involved in immunopathogenesis and possibly lead to strategies for immune therapeutic intervention.

Blood represents a reservoir and a migration compartment for cells of the innate and the adaptive immune systems, including either neutrophils, dendritic cells and monocytes, or B and T lymphocytes, respectively, which during infection will have been exposed to infectious agents in the tissue. For this reason whole blood from infected individuals provides an accessible source of clinically relevant material where an unbiased molecular phenotype can be obtained using gene expression microarrays as previously described for the study of cancer in tissues (Alizadeh AA., 2000; Golub, TR., 1999; Bittner, 2000), and autoimmunity (Bennet, 2003; Baechler, EC, 2003; Buczynski, ME, 2005; Chaussabel, D., 2005; Cobb, JP., 2005; Kaizer, EC., 2007; Allantaz, 2005; Allantaz, 2007), and inflammation (Thach, DC., 2005) and infectious disease (Ramillo, Blood, 2007) in blood or tissue (Bleharski, JR et al., 2003). Microarray analyses of gene expression in blood leucocytes have identified diagnostic and prognostic gene expression signatures, which have led to a better understanding of mechanisms of disease onset and responses to treatment (Bennet, L 2003; Rubins, KH., 2004; Baechler, EC, 2003; Pascual, V., 2005; Allantaz, F., 2007; Allantaz, F., 2007). These microarray approaches have been attempted for the study of active and latent TB but as yet have yielded small numbers of differentially expressed genes only (Jacobsen, M., Kaufmann, SH., 2006; Mistry, R, Lukey, PT, 2007),

and in relatively small numbers of patients (Mistry, R., 2007), which may not be robust enough to distinguish between other inflammatory and infectious diseases.

To define an immune signature in TB, the blood of active and latent TB patients and controls were analyzed; patients were selected using very stringent clinical criteria. Patients were recruited from London, UK, where numbers of active TB cases are increasing, and most importantly where the risk of confounding coinfections is minimal, to yield a robust signature that may distinguish latent from active TB. Microarrays were used to analyze the whole genome and subsequent data mining revealed a large number of genes found to be differentially expressed at a statistically significant level across all groups of patients, including active and latent TB patients and healthy controls. Next, a novel approach based on a modular data mining strategy was used, this approach provided a basis for the selection of clinically-relevant transcriptional biomarkers for the analysis of blood microarray transcriptional profiles in SLE and other diseases, and improved our understanding of disease pathogenesis (Chaussabel, 2008, Immunity). The module maps defined in this study provide a means to organize and reduce the dimension of complex data, whilst still retaining the large number of genes expressed in human blood, thus allowing visualization of specific disease fingerprints (Chaussabel, 2008, Immunity). Using this modular approach clearly defined modular transcriptional signatures were obtained that are distinct and reciprocal in the whole blood of active and latent TB patients, and which also differ from healthy controls. The biomarkers described herein improve the diagnosis of PTB, and furthermore will help to define host factors important in the protection against *M. tuberculosis* in latent TB patients, and those involved in the immunopathogenesis of active TB, and thus be used to reduce and manage TB disease.

PATIENTS, MATERIALS AND METHODS.

Participant recruitment and Patient characterization: Participants were recruited from St. Mary's Hospital TB Clinic, Imperial College Healthcare NHS Trust, London, with healthy controls recruited from volunteers at the National Institute for Medical Research (NIMR), Mill Hill, London. The study was approved by the local NHS Research Ethics Committee at St Marys Hospital (LREC), London, UK. All participants (aged 18 and over) gave written informed consent. Strict clinical criteria were satisfied before recruited participants had their provisional study grouping confirmed and were only then allocated to the final group for analysis. The patient and control cohorts were as follows: (i) Active PTB based on clinical diagnosis subsequently confirmed by laboratory isolation of *M. tuberculosis* on mycobacterial culture; (ii) Latent TB - defined by a positive tuberculin skin test (TST, Using 2TU tuberculin (Serum Statens Institute, Copenhagen, Denmark) ≥ 6 mm if BCG unvaccinated, ≥ 15 mm if BCG vaccinated, together with a positive result using an Interferon Gamma Release Assay (IGRA, specifically the Quantiferon-TB Gold In-tube assay, Cellestis, Australia). This IGRA assay measured reactivity to antigens (ESAT-6/CFP-10/TB 7.7 - present in *M. tuberculosis* but not in most environmental mycobacteria or the *M. bovis* BCG vaccine) by IFN- γ release from whole blood.

Latent TB patients also had to have evidence of exposure to infectious TB cases, either through close household or workplace contact, or as recent 'new entrants' from endemic areas; Patients with incidental findings of TST positivity without evidence of exposure to infected persons, were not eligible for inclusion in the study (iii) Healthy volunteer controls (BCG vaccinated and unvaccinated, ≤ 14 mm or ≤ 5 mm by TST
5 respectively; and negative by IGRA). Participants who were pregnant, known to be immunosuppressed, taking immunosuppressive therapies or have diabetes, or autoimmune disease were also ineligible and excluded from this initial study. HIV positive individuals (Only 1% of the TB patients in London present with previously undiagnosed HIV) were excluded from the study. Blood from active and latent PTB patients was collected for the study before any anti-mycobacterial drugs were administered, and then subsequently at
10 set time intervals for the longitudinal part of the study for later study.

Detailed clinical information was collected prospectively for every participant and has been entered into a web-accessible database developed by the present inventors. Using this recorded clinical data, and immune-based assays as described above, 15 out of 58 participants were excluded from the study as they did not meet the standard criteria for the study. This resulted in cohorts of 6 BCG unvaccinated healthy volunteers; 6 BCG
15 vaccinated healthy volunteers, 17 latent TB patients and 14 active PTB patients, all of these samples were then used for RNA isolation. One sample from an active TB patient did not yield sufficient globin reduced RNA after processing to proceed and was therefore excluded from the final analysis.

RNA sampling, extraction, processing for microarray: Whole blood from the above patient cohorts was collected into Tempus tubes (Applied Biosystems, Foster City, CA, USA) and stored between -20°C and $-$
20 80°C before RNA extraction. Total RNA was isolated using the PerfectPure RNA Blood kit (5 PRIME Inc, Gaithersburg, MD, USA). Samples were homogenized with 100% cold ethanol, vortexed, then centrifuged at 4000g for 60 minutes at 0°C , and the supernatant discarded. 300 μl lysis solution was then added to the pellet and vortexed. RNA binding, Dnase treatment, wash and RNA elution steps were then performed according to the manufacturer's instructions. Isolated total RNA was then globin reduced using the GLOBINclear™ 96-
25 well format kit (Ambion, Austin, TX, USA) according to the manufacturer's instructions. Total and globin-reduced RNA integrity was assessed using an Agilent 2100 Bioanalyzer (Agilent, Palo Alto, CA). One sample from an active TB patient did not yield sufficient globin reduced RNA after processing to proceed and was therefore excluded from the final analysis. Biotinylated, amplified RNA targets (cRNA) were then prepared from the globin-reduced RNA using the Illumina CustomPrep RNA amplification kit (Ambion,
30 Austin, TX, USA). Labeled cRNA was hybridized overnight to Sentrix Human-6 V2 BeadChip array (>48,000 probes, Illumina Inc, San Diego, CA, USA), washed, blocked, stained and scanned on an Illumina BeadStation 500 following the manufacturer's protocols. Illumina's BeadStudio version 2 software was used to generate signal intensity values from the scans, subtract background, and scale each microarray to the

median average intensity for all samples (per-chip normalization). This normalized data was used for all subsequent data analysis.

Microarray data analysis: A gene expression analysis software program, Genespring, version 7.1.3 (Agilent), was used to perform statistical analysis and hierarchical clustering of samples. Differentially expressed genes were selected and clustered as described in Results and Figure legends.

RESULTS AND DISCUSSION.

Blood signatures distinguish active and latent TB patients from each other, and from healthy control individuals: To determine whether blood sampled from patients with active and latent TB carry gene expression signatures that allow discrimination between active and latent TB as compared to healthy controls, a step-wise analysis was conducted. After filtering out undetected transcripts and genes with a deviation from the median of less than 2 fold, i.e. with a flat profile, 6269 genes were used for unsupervised clustering analyses by Pearson correlation of the expression profiles obtained from the whole blood RNA samples from active and latent TB and healthy controls (Figure 1). This unsupervised analysis identified distinct signatures, which were found to correspond to distinct clinical phenotypes: in patients with active pulmonary TB (active PTB); and: in individuals with latent tuberculosis (latent TB). The grouping of samples was not perfect (10 of 13 patients with active TB, and 11 of 17 patients with latent TB). Nonetheless, the majority of active PTB and latent TB patients in this group from the training set of patients appeared to have clear and distinct transcriptional signatures. Importantly these signatures appeared to be represented across the broad number of ethnicities collected for the study, including White, Black African, Asian Indian, Asian Bangladeshi, Asian Other, White Irish, Mixed White, Black Caribbean (details of this data are not shown).

This list of 6269 genes was then further analysed using a non-parametric statistical group comparison (Kruskal-Wallis test) to identify genes that were significantly differentially expressed between groups. Using a moderately stringent multiple comparison correction for controlling Type I error (Benjamini-Hochberg correction), 1473 genes were differentially expressed/represented across the active TB and latent TB, and healthy controls ($P < 0.01$) (Figure 2; and listing of 1473 genes in LENGHTY TABLE, filed herewith). These clusters of genes were then correlated with relevant findings in the literature. Filtering of these genes for the ontological term "Immune response" generated a list of 158 such genes (Figures 3A-D; Table 2). This pattern of expression/representation of 158 genes (Figure 3A – 3D) allows discrimination of the group of Active TB patients from the Latent TB patients and from the Healthy control individuals.

Table 2. List of 158 genes annotated with gene ontology term biological process: immune response and found to be significantly differentially expressed ($p < 0.01$) between active TB and other clinical groups.

Gene Symbol	Description
LILRB3	leukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 3
PGLYRP1	peptidoglycan recognition protein 1
FAS	Fas (TNF receptor superfamily, member 6)
IFITM3	interferon induced transmembrane protein 3 (1-8U)
FCGR2A	Fc fragment of IgG, low affinity IIa, receptor (CD32)
FCGR2A	Fc fragment of IgG, low affinity IIa, receptor (CD32)
ST6GAL1	ST6 beta-galactosamide alpha-2,6-sialyltransferase 1
ETS1	v-ets erythroblastosis virus E26 oncogene homolog 1 (avian)
CYBB	cytochrome b-245, beta polypeptide (chronic granulomatous disease)
IFNAR1	interferon (alpha, beta and omega) receptor 1
LY96	lymphocyte antigen 96
TRIM22	tripartite motif-containing 22
GBP2	guanylate binding protein 2, interferon-inducible
DDX58	DEAD (Asp-Glu-Ala-Asp) box polypeptide 58
LAX1	lymphocyte transmembrane adaptor 1
IFI16	interferon, gamma-inducible protein 16
LCK	lymphocyte-specific protein tyrosine kinase
IL32	interleukin 32
CXCL16	chemokine (C-X-C motif) ligand 16
CD40LG	CD40 ligand (TNF superfamily, member 5, hyper-IgM syndrome)
TNFSF13B	tumor necrosis factor (ligand) superfamily, member 13b
IRF2	interferon regulatory factor 2
C5	complement component 5
CD46	CD46 molecule, complement regulatory protein
TNFAIP6	tumor necrosis factor, alpha-induced protein 6
DPP4	dipeptidyl-peptidase 4 (CD26, adenosine deaminase complexing protein 2)
EBI2	Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor)
NFX1	nuclear transcription factor, X-box binding 1
MICB	MHC class I polypeptide-related sequence B
GBP3	guanylate binding protein 3
SLAMF7	SLAM family member 7
CARD12	NLR family, CARD domain containing 4
GBP6	guanylate binding protein family, member 6
IFIT3	interferon-induced protein with tetratricopeptide repeats 3
TAP2	transporter 2, ATP-binding cassette, sub-family B (MDR/TAP)
HLA-DPB1	major histocompatibility complex, class II, DP beta 1
CD3G	CD3g molecule, gamma (CD3-TCR complex)
PRKCQ	protein kinase C, theta
IL7R	interleukin 7 receptor
SLAMF1	signaling lymphocytic activation molecule family member 1
CD274	CD274 molecule
GBP1	guanylate binding protein 1, interferon-inducible, 67kDa
IFITM2	interferon induced transmembrane protein 2 (1-8D)
ITK	IL2-inducible T-cell kinase
APOL2	apolipoprotein L, 2

Gene Symbol	Description
PSME1	proteasome (prosome, macropain) activator subunit 1 (PA28 alpha)
LAT2	linker for activation of T cells family, member 2
IL18RAP	interleukin 18 receptor accessory protein
OSM	oncostatin M
CD6	CD6 molecule
WWP1	WW domain containing E3 ubiquitin protein ligase 1
CD3E	CD3e molecule, epsilon (CD3-TCR complex)
VIPR1	vasoactive intestinal peptide receptor 1
TNFSF10	tumor necrosis factor (ligand) superfamily, member 10
PRKRA	protein kinase, interferon-inducible double stranded RNA dependent activator
TNFRSF1A	tumor necrosis factor receptor superfamily, member 1A
BCL6	B-cell CLL/lymphoma 6 (zinc finger protein 51)
IL8	interleukin 8
OAS3	2'-5'-oligoadenylate synthetase 3, 100kDa
IFIH1	interferon induced with helicase C domain 1
SIGIRR	single immunoglobulin and toll-interleukin 1 receptor (TIR) domain
SIGIRR	single immunoglobulin and toll-interleukin 1 receptor (TIR) domain
SIT1	signaling threshold regulating transmembrane adaptor 1
ITGAM	integrin, alpha M (complement component 3 receptor 3 subunit)
C1QB	complement component 1, q subcomponent, B chain
IL27RA	interleukin 27 receptor, alpha
ALOX5AP	arachidonate 5-lipoxygenase-activating protein
SERPING1	serpin peptidase inhibitor, clade G (C1 inhibitor), member 1, (angioedema, hereditary)
IL1RN	interleukin 1 receptor antagonist
IL1RN	interleukin 1 receptor antagonist
CLEC4D	C-type lectin domain family 4, member D
ICOS	inducible T-cell co-stimulator
OAS1	2',5'-oligoadenylate synthetase 1, 40/46kDa
ZAP70	zeta-chain (TCR) associated protein kinase 70kDa
IL1B	interleukin 1, beta
C4BPA	complement component 4 binding protein, alpha
TNFSF13	tumor necrosis factor (ligand) superfamily, member 13
IFI30	interferon, gamma-inducible protein 30
HPSE	heparanase
CD59	CD59 molecule, complement regulatory protein
CTLA4	cytotoxic T-lymphocyte-associated protein 4
BCL2	B-cell CLL/lymphoma 2
TNFRSF7	CD27 molecule
FPR1	formyl peptide receptor 1
IL2RA	interleukin 2 receptor, alpha
GATA3	GATA binding protein 3
S100A9	S100 calcium binding protein A9
TLR8	toll-like receptor 8
NCF1	neutrophil cytosolic factor 1, (chronic granulomatous disease, autosomal 1)
BCL6	B-cell CLL/lymphoma 6 (zinc finger protein 51)
BST1	bone marrow stromal cell antigen 1
G1P2	ISG15 ubiquitin-like modifier
C1QA	complement component 1, q subcomponent, A chain
TCF7	transcription factor 7 (T-cell specific, HMG-box)

Gene Symbol	Description
IFITM1	interferon induced transmembrane protein 1 (9-27)
TAPBPL	TAP binding protein-like
AIM2	absent in melanoma 2
CCR7	chemokine (C-C motif) receptor 7
LTBR	lymphotoxin beta receptor (TNFR superfamily, member 3)
FYB	FYN binding protein (FYB-120/130)
NFIL3	nuclear factor, interleukin 3 regulated
LAT	linker for activation of T cells
CBLB	Cas-Br-M (murine) ecotropic retroviral transforming sequence b
CD74	CD74 molecule, major histocompatibility complex, class II invariant chain
TAP2	transporter 2, ATP-binding cassette, sub-family B (MDR/TAP)
FLJ14466	transmembrane protein 142A
PSMB9	proteasome (prosome, macropain) subunit, beta type, 9 (large multifunctional peptidase 2)
PSMB8	proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional peptidase 7)
FAIM3	Fas apoptotic inhibitory molecule 3
LTA4H	leukotriene A4 hydrolase
IRF1	interferon regulatory factor 1
OAS2	2'-5'-oligoadenylate synthetase 2, 69/71kDa
RELB	v-rel reticuloendotheliosis viral oncogene homolog B, nuclear factor of kappa light polypeptide gene enhancer in B-cells 3 (avian)
TRA@	T cell receptor alpha locus
LTB4R	leukotriene B4 receptor
PIK3R1	phosphoinositide-3-kinase, regulatory subunit 1 (p85 alpha)
OASL	2'-5'-oligoadenylate synthetase-like
OASL	2'-5'-oligoadenylate synthetase-like
PSME2	proteasome (prosome, macropain) activator subunit 2 (PA28 beta)
CLEC6A	C-type lectin domain family 6, member A
NBN	nibrin
FCGR1A	Fc fragment of IgG, high affinity Ia, receptor (CD64)
SH2D1A	SH2 domain protein 1A, Duncan's disease (lymphoproliferative syndrome)
IL15	interleukin 15
LY9	lymphocyte antigen 9
LILRB1	leukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 1
APOL3	apolipoprotein L, 3
PSMB8	proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional peptidase 7)
CCR6	chemokine (C-C motif) receptor 6
PDCD1LG2	programmed cell death 1 ligand 2
CD96	CD96 molecule
EPHX2	epoxide hydrolase 2, cytoplasmic
BST2	bone marrow stromal cell antigen 2
RIPK2	receptor-interacting serine-threonine kinase 2
SCAP1	src kinase associated phosphoprotein 1
GBP5	guanylate binding protein 5
TRAT1	T cell receptor associated transmembrane adaptor 1
ALOX5	arachidonate 5-lipoxygenase
LY9	lymphocyte antigen 9
TAP1	transporter 1, ATP-binding cassette, sub-family B (MDR/TAP)
RHOH	ras homolog gene family, member H

Gene Symbol	Description
IFI35	interferon-induced protein 35
CD28	CD28 molecule
FYB	FYN binding protein (FYB-120/130)
IFIT2	interferon-induced protein with tetratricopeptide repeats 2
TLR7	toll-like receptor 7
CD2	CD2 molecule
FCER1G	Fc fragment of IgE, high affinity I, receptor for; gamma polypeptide
SMAD3	SMAD family member 3
FCER1A	Fc fragment of IgE, high affinity I, receptor for; alpha polypeptide
SERPINA1	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 1
SERPINA1	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 1
SECTM1	secreted and transmembrane 1
NMI	N-myc (and STAT) interactor
TLR5	toll-like receptor 5
IFIT3	interferon-induced protein with tetratricopeptide repeats 3
IFIT3	interferon-induced protein with tetratricopeptide repeats 3
CD5	CD5 molecule

Genes over-expressed/represented in active TB: Of interest is that a large number of IFN-associated/inducible genes were expressed: for example interferon (IFN)-inducible genes, e.g., SOCS1, STAT1, PML (TRIM19), TRIM22, many guanylate binding proteins, and many other IFN-inducible genes as indicated in Table 2, as expected in active TB, but interestingly these were not evident in latent TB patients, although these patients representation/expression of IFN- γ transcripts in whole blood was in fact higher than the active TB patients. To focus in on this, certain families of genes, some of which are known to be upregulated by IFNs and others not, were further studied, including the TRIM family.

A subset of TRIMS are over-expressed/represented in Active TB: The tripartite motif (TRIM) family of proteins are characterized by a discreet structure (Reymond, A., EMBO J., 2001) and have been shown to have multiple functions, including E3 ubiquitin ligases activity, induction of cellular proliferation, differentiation and apoptosis, immune cell signalling (Meroni, G., Bioessays, 2005). Their involvement has been implicated in protein-protein interactions, autoimmunity and development (Meroni, G., Bioessays, 2005). Furthermore, a number of TRIM proteins have been found to have anti-viral activity and are possibly involved in innate immunity (Nisole, F, 2005, Nat. Rev. Microbiol.; Gack, MU., 2007, Nature). Interestingly, 30 TRIM transcripts (some overlapping probes) were shown to be expressed in active TB, with some also expressed in latent TB and healthy control blood (Figure 4; Table 3). The majority of these TRIMs have been previously shown to be expressed in both human macrophages and mouse macrophages and dendritic cells (Rajsbaum, 2008, EJI; Martinez, FO., J. Imm., 2006) and regulated by IFNs, whereas TRIMs shown to be constitutively expressed in DC or in T cells (Rajsbaum, 2008, EJI) were not detected or were not found to be differentially expressed in active or latent TB versus healthy control blood. Interestingly, it was found that TRIM 5, 6, 19(PML), 21, 22, 25, 68 are overrepresented/expressed; while the

others are underrepresented/expressed: TRIM 28, 32, 51, 52, 68. Of interest a group of TRIMs was highly expressed in active TB, but low to undetectable in latent TB and healthy controls, and four of these (TRIM 5, 6, 21, 22) have been show to cluster on human chromosome 11, and reported to have anti-viral activity (Song, B., 2005, J. Virol.); Li, X, Virology, 2007). A group of TRIMs however, were found to be under-
 5 expressed in the blood of active TB patients versus that of latent TB and healthy controls, including TRIM 28, 32, 51, 52 68, and these have been reported to either not be expressed in human blood-derived macrophages (TRIM 51) or only expressed in undifferentiated monocytes (TRIM-28, 52) or non-activated macrophages or alternately activated macrophages (TRIM-32), or only upregulated to a low level in activated macrophages differentiated from human blood (TRIM-68) (Martinez, FO., J. Imm., 2006).

10 Table 3. TRIM genes differentially expressed in active pulmonary tuberculosis, latent tuberculosis and healthy controls.

Common Name	Gene Symbol	Description
RNF94; STAF50; GPSTAF50	TRIM22	tripartite motif-containing 22
RNF91; SPRING; KIAA0282	TRIM9	tripartite motif-containing 9
MYL; RNF71; PP8675; TRIM19	PML	promyelocytic leukemia
RNF89	TRIM6	tripartite motif-containing 6
TRIM51; MGC10977	TRIM51	SPRY domain containing 5
RNF9; HERF1; RFB30; MGC141979	TRIM10	tripartite motif-containing 10
PML	PML	promyelocytic leukemia; synonyms: MYL, RNF71, PP8675, TRIM19; isoform 7 is encoded by transcript variant 7; promyelocytic leukemia, inducer of; tripartite motif protein TRIM19; promyelocytic leukemia protein; Homo sapiens promyelocytic leukemia (PML), transcript variant 7, mRNA.
RNF88; TRIM5alpha	TRIM5	tripartite motif-containing 5
RNF88; TRIM5alpha	TRIM5	tripartite motif-containing 5
BIA2; DKFZp434C091	TRIM58	tripartite motif-containing 58
Trif; HSD34; RNF36	TRIM69	tripartite motif-containing 69
RNF88; TRIM5alpha	TRIM5	tripartite motif-containing 5
SSA; RO52; SSA1; RNF81	TRIM21	tripartite motif-containing 21
KIAA0129	TRIM14	tripartite motif-containing 14
RNF9; HERF1; RFB30; MGC141979	TRIM10	tripartite motif-containing 10
EFP; Z147; RNF147; ZNF147	TRIM25	tripartite motif-containing 25
HLS5; MAIR; KIAA1098; MGC17233	TRIM35	tripartite motif-containing 35
RNF86; KIAA0517	TRIM2	tripartite motif-containing 2
RNF9; HERF1; RFB30; MGC141979	TRIM10	tripartite motif-containing 10
GNIP; RNF90	TRIM7	tripartite motif-containing 7
KIAA0129	TRIM14	tripartite motif-containing 14
TRIM50B; MGC45477	TRIM50B	tripartite motif-containing 73
4732463G12Rik	TRIM65	tripartite motif-containing 65

Common Name	Gene Symbol	Description
MRF1; TSBF1; RNF104; TRIM57; MGC26631; MGC129860; MGC129861	TRIM59	tripartite motif-containing 59
FMF; MEF; TRIM20; MGC126560; MGC126586	MEFV	Mediterranean fever
TRIM52		Tripartite motif-containing 52
CAR; LEU5; RFP2; DLEU5; RNF77	RFP2	tripartite motif-containing 13
KAP1; TF1B; RNF96; TIF1B; FLJ29029	TRIM28	tripartite motif-containing 28
SS-56; RNF137; FLJ10369; MGC126176	TRIM68	tripartite motif-containing 68
HT2A; BBS11; TATIP; LGMD2H	TRIM32	tripartite motif-containing 32

Selective over-expression/representation of specific immunomodulatory ligands in Active TB Patients: Analysis of the distinct transcriptional profiles revealed that transcripts from the genes CD274 (PDL1) and PCDLG2 (PDL2, CD273) are expressed only in the active TB patients (Figures 5A and B). These molecules have been previously shown to be involved in the regulation of the immune response to both acute and chronic viral infection (A Sharpe, Ann. Rev. Imm.). These molecules act as inhibitory co-stimulatory receptors for the molecule PD1 in interactions between T cells and APCs, and blockade of this pathway has been shown to restore the proliferative and effector functions of antigen specific T cells in HIV, Hepatitis B and C infection.

Genes under-expressed/represented in active TB: Strikingly, a number of genes known to be expressed in T cells (some also on NK and B cells), were found to be profoundly down-regulated/under-represented in the blood of active TB patients (Figure 3D), (but not in latent TB or healthy controls, including, CD3, CTLA-4, CD28, ZAP-70 (T, NK and B cells), IL-7R, CD2 (also on B cells), SLAM (also on NK cells), CCR7, GATA-3 (also in NK cells). This could indicate that gene expression was down-regulated in T, NK and B cells during active PTB, or that the cells had been recruited elsewhere (e.g., the lung) as a result of infection with *M. tuberculosis*. This is currently under investigation using flow cytometric analysis of blood from the different patient groups, as well as by transcriptional analysis of purified populations of T cells from the different patient groups.

Higher Stringency Statistical analysis of transcriptional profiles in latent and active TB patients versus healthy controls. Statistical group comparison was further performed as before by identifying differentially expressed genes between the groups using the non-parametric Kruskal-Wallis test, but now using the most stringent multiple comparison correction for controlling Type I error (Bonferroni correction). With this increased stringency 46 genes ($P < 0.1$) and 18 genes ($P < 0.05$) were identified as differentially expressed between groups (Figures 6 and 7; Tables 4 and 5). Of the 46 genes a large number of IFN-inducible genes, such as STAT-1, GBP and IRF-1 were still observed to be over-expressed/represented in the blood from

active TB patients, and either down-regulated or unchanged in the latent patients or healthy controls. A number of these genes were also found to be over-expressed/represented in the blood of active TB patients, even with the highest stringency analysis which still extracted genes (Bonferroni correction, $P < 0.05$). Only 3 transcripts in active TB were still observed to be down-regulated/under-represented within the 46 gene group, including IL-7R (expressed in T cells), the chemokine receptor CXCR3 (lost at higher statistical stringency) and alpha II-spectrin. The underexpression/representation of CXCR3 is of interest since this chemokine receptor has been shown to be highly expressed in Th1 cells required for protection against mycobacterial infection, which may reflect their suppression or migration out of blood to infected tissue. Table 5 includes 18 genes, with IL7R and SPTAN1 being underrepresented/expressed in active PTB, and all others being overrepresented/expressed and diagnostic for active disease.

Table 4. Genes significantly differentially expressed between active TB and other clinical groups.

Gene Symbol	Description
FAM84B	family with sequence similarity 84, member B
CXCR3	chemokine (C-X-C motif) receptor 3
ETV7	ets variant gene 7 (TEL2 oncogene)
DUSP3	dual specificity phosphatase 3 (vaccinia virus phosphatase VH1-related)
WARS	tryptophanyl-tRNA synthetase
CNIH4	cornichon homolog 4 (Drosophila)
STAT1	signal transducer and activator of transcription 1, 91kDa
IRF1	interferon regulatory factor 1
LILRB1	leukocyte immunoglobulin-like receptor, subfamily B (with TM and ITIM domains), member 1
SIPA1L1	signal-induced proliferation-associated 1 like 1
GSDMDC1	gasdermin domain containing 1
DYNLT1	dynein, light chain, Tctex-type 1
DKFZp761E198	DKFZp761E198 protein
LOC400759	
GBP1	guanylate binding protein 1, interferon-inducible, 67kDa
GBP5	guanylate binding protein 5
FLJ11259	damage-regulated autophagy modulator
LYPLA1	lysophospholipase I
RHBDF2	rhomboid 5 homolog 2 (Drosophila)
PLEK	pleckstrin
ANKRD22	ankyrin repeat domain 22
CASP1	caspase 1, apoptosis-related cysteine peptidase (interleukin 1, beta, convertase)
FLJ39370	chromosome 4 open reading frame 32
FBXO6	F-box protein 6
GCH1	GTP cyclohydrolase 1 (dopa-responsive dystonia)
GBP4	guanylate binding protein 4
IFI30	interferon, gamma-inducible protein 30
VAMP5	vesicle-associated membrane protein 5 (myobrevin)
GBP2	guanylate binding protein 2, interferon-inducible
STX11	syntaxin 11
SPTAN1	spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)
POLB	polymerase (DNA directed), beta

Gene Symbol	Description
IL7R	interleukin 7 receptor
APOL6	apolipoprotein L, 6
ATG3	ATG3 autophagy related 3 homolog (<i>S. cerevisiae</i>)
SQRDL	sulfide quinone reductase-like (yeast)
PSME2	proteasome (prosome, macropain) activator subunit 2 (PA28 beta)
FLJ10379	S1 RNA binding domain 1
WDFY1	WD repeat and FYVE domain containing 1
TAP2	transporter 2, ATP-binding cassette, sub-family B (MDR/TAP)
NPC2	Niemann-Pick disease, type C2
ATF3	activating transcription factor 3
VAMP3	vesicle-associated membrane protein 3 (cellubrevin)
PSMB8	proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional peptidase7)
JAK2	Janus kinase 2 (a protein tyrosine kinase)

Table 5. 18 genes significantly differentially expressed between active TB and other clinical groups.

Gene Symbol	Description
VAMP5	vesicle-associated membrane protein 5 (myobrevin)
GBP2	guanylate binding protein 2, interferon-inducible
STX11	syntaxin 11
SPTAN1	spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)
POLB	polymerase (DNA directed), beta
IL7R	interleukin 7 receptor
APOL6	apolipoprotein L, 6
ATG3	ATG3 autophagy related 3 homolog (<i>S. cerevisiae</i>)
SQRDL	sulfide quinone reductase-like (yeast)
PSME2	proteasome (prosome, macropain) activator subunit 2 (PA28 beta)
FLJ10379	S1 RNA binding domain 1
WDFY1	WD repeat and FYVE domain containing 1
TAP2	transporter 2, ATP-binding cassette, sub-family B (MDR/TAP)
NPC2	Niemann-Pick disease, type C2
ATF3	activating transcription factor 3
VAMP3	vesicle-associated membrane protein 3 (cellubrevin)
PSMB8	proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional peptidase7)
JAK2	Janus kinase 2 (a protein tyrosine kinase)

Improved discrimination between patients with active and latent TB and healthy controls: The approaches described above although able to discriminate active TB from latent TB and healthy controls are less able to discriminate between all three clinical groups. To select discriminating genes the following approach was used. First, genes expressed in blood from healthy individuals were compared versus latent TB patients, using the Wilcoxon-Mann-Whitney test at a $p < 0.005$, which yielded 89 discriminatory genes. Genes expressed in blood from healthy individuals versus active TB patients were then compared, again using the Wilcoxon-Mann-Whitney test but with a $p < 0.5$, and the most stringent Bonferroni correction factor, which yielded a list of 30 discriminatory genes. This list was combined to give a total list of 119 discriminating genes (Table 6). This list of genes was then used to interrogate the dataset of all clinical groups using unsupervised clustering analysis by Pearson correlation. This analysis generated three distinct clusters of clinical groups (Figures 8A to 8F): one cluster is composed of 11 out of 13 of the active TB patients (Figure

8, Cluster C); a second cluster is composed of 16 out of 17 latent TB patients, and 1 active TB patient (Figure 8, Cluster B); a third cluster contains all 12 healthy controls included in the study, plus 1 active TB and 1 latent TB outlier (Figure 8, Cluster A). For each of Figures 8A to 8F, clusters of patients/clinical groups are presented horizontally and clusters of genes are presented vertically. This pattern of expression/representation of the whole list of 119 genes (Figure 8A) now allows discrimination of all three clinical groups from each other: i.e., allows discrimination of Active TB, Latent TB and Healthy individuals from each other, each clinical group exhibiting a unique pattern of expression/representation of these 119 genes or subgroups thereof. The skilled artisan will recognize that 1, 2, 3, 4, 5, 6, 7, 8, 10, 12, 15, 20, 25, 30, 35 or more genes may be placed in a dataset that represents a cluster of genes that may be compared across clusters of clinical groups A (Healthy), B (Latent), C (Active), and that either alone or in combination with other such clusters, each clinical group can exhibit a unique pattern of expression/representation obtained from these 119 genes.

Specifically, Figure 8B demonstrates that the genes ST3GAL6, PAD14, TNFRSF12A, VAMP3, BR13, RGS19, PILRA, NCF1, LOC652616, PLAUR(CD87), SIGLEC5, B3GALT7, IBRDC3(NKLAM), ALOX5AP(FLAP), MMP9, ANPEP(APN), NALP12, CSF2RA, IL6R(CD126), RASGRP4, TNFSF14(CD258), NCF4, HK2, ARID3A, PGLYRP1(PGRP) are underexpressed/underrepresented in the blood of Latent TB patients but not in the blood of Healthy individuals or of Active TB patients.

The genes presented in Figure 8C, ABCG1, SREBF1, RBP7(CRBP4), C22orf5, FAM101B, S100P, LOC649377, UBTD1, PSTPIP-1, RENBP, PGM2, SULF2, FAM7A1, HOM-TES-103, NDUFAF1, CES1, CYP27A1, FLJ33641, GPR177, MID1IP1(MIG-12), PSD4, SF3A1, NOV(CCN3), SGK(SGK1), CDK5R1, LOC642035, are shown to be overexpressed/overrepresented in the blood of Healthy control individuals but were underexpressed/underrepresented in the blood of Latent TB patients, and to a great extent were underexpressed/underrepresented in the blood of Active TB patients.

The pattern of genes in Figure 8D, ARSG, LOC284757, MDM4, CRNKL1, IL8, LOC389541, CD300LB, NIN, PHKG2, HIP1, were shown to be overexpressed/overrepresented in the blood of Healthy individuals but were underexpressed/underrepresented in the blood of both Latent and Active TB patients. Conversely, the genes in Figure 8D, PSMB8(LMP7), APOL6, GBP2, GBP5, GBP4, ATF3, GCH1, VAMP5, WARS, LIMK1, NPC2, IL-15, LMTK2, STX11(FHL4), were shown to be overexpressed/overrepresented in the blood of Active TB, but underexpressed/underrepresented in the blood of Latent TB patients and Healthy control individuals.

The pattern of genes in Figure 8E, of FLJ11259(DRAM), JAK2, GSDMDC1(DF5L)(FKSG10), SIPAIL1, [2680400](KIAA1632), ACTA2(ACTSA), KCNMB1(SLO-BETA), were all overexpressed/overrepresented in blood from Active TB patients but not represented or even underexpressed/underrepresented in the blood from Latent TB patients and Healthy control individuals. Conversely, the genes SPTANI,

KIAAD179(Nnp1)(RRP1), FAM84B(NSE2), SELM, IL27RA, MRPS34, [6940246](IL23A), PRKCA(PKCA), CCDC41, CD52(CDW52), [3890241](ZN404), MCCC1(MCCA/B), SOX8, SYNJ2, FLJ21127, FHIT, were underexpressed/underrepresented in the blood of Active TB patients but not in the blood of Latent TB patients or Healthy Control individuals, where they were overexpressed/overrepresented.

5 Many of the genes (within these 119 genes selected by this method described above) found to be overexpressed/overrepresented in the blood of Active TB patients listed in Figures 8D and 8E, were common to those identified by the alternative method using Higher Stringency Analysis of transcriptional profiles in active, latent TB patients and healthy controls described earlier (genes shown as underlined above from Figures 8D and 8E are contained in list of genes in Figure 7, Table 5, 18 genes $p < 0.05$; genes shown as
10 italicised above from Figures 8D and 8E are contained in list of genes in Figure 6, Table 4, 46 genes $P < 0.1$).

The pattern of genes shown in Figure 8F, CD52(CDW52), [3890241](ZNF404), MCCC1(MCCA/B), SOX8, SYNJ2, FLJ21127, FHIT, were underexpressed/underrepresented in the blood of Active TB patients but not in the blood of Latent TB patients or Healthy Control individuals, where they were if anything overexpressed/overrepresented. This is also presented (overlap) in Figure 8E. Genes CDKL1(p42),
15 MICALCL, MBNL3, RHD, ST7(RAY1), PPR3R1, [360739](PIP5K2A), AMFR, FLJ22471, CRAT(CAT1), PLA2G4C, ACOT7(ACT)(ACH1), RNF182, KLRC3(NKG2E), HLA-DPB1, were underexpressed/underrepresented in the blood of Healthy Control individuals, but were overexpressed/overrepresented in the blood of the Latent TB patients, and overexpressed/overrepresented in the blood of most Active TB patients (Figure 8F). To conclude, the aggregate pattern of expression of the
20 total of 119 genes in Figure 8A (broken down for legibility of genes and specificity between clinical states in Figures 8B - 8F) that distinguishes between infected (Active TB and Latent TB) patients from non-infected patients (Healthy Controls) and additionally, distinguishes between the two groups of infected patients, that is Active and Latent TB patients. Many of the genes overexpressed in the blood of active TB patients via this method were the same genes as those identified using the strictest statistical filtering (shown in Figure 7,
25 Table 6), and many were IFN-inducible and/or involved in endocytic cellular traffic and/or lipid metabolism.

Table 6. Genes found to be significantly differentially expressed between latent and healthy or between active and healthy, which when used in combination differentiate between active, healthy and latent using unsupervised pearson correlation clustering algorithms (119 genes).

Gene Symbol	Description
HMFN0839	lung cancer metastasis-associated protein
LOC653820	
MID1IP1	MID1 interacting protein 1 (gastrulation specific G12 homolog (zebrafish))
SPTAN1	spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)
NALP12	NLR family, pyrin domain containing 12
PSMB8	proteasome (prosome, macropain) subunit, beta type, 8 (large multifunctional peptidase 7)
RNF182	ring finger protein 182

Gene Symbol	Description
KCNMB1	potassium large conductance calcium-activated channel, subfamily M, beta member 1
	Interleukin 23, alpha subunit p19
CDKL1	cyclin-dependent kinase-like 1 (CDC2-related kinase)
IL8	interleukin 8
NOV	nephroblastoma overexpressed gene
APOL6	apolipoprotein L, 6
KLRC3	killer cell lectin-like receptor subfamily C, member 3
SOX8	SRY (sex determining region Y)-box 8
B3GALT7	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminyltransferase 8
GCH1	GTP cyclohydrolase 1 (dopa-responsive dystonia)
IL6R	interleukin 6 receptor
RASGRP4	RAS guanyl releasing protein 4
SGK	serum/glucocorticoid regulated kinase
LOC389541	similar to CG14977-PA
MICALCL	MICAL C-terminal like
VAMP3	vesicle-associated membrane protein 3 (cellubrevin)
NPC2	Niemann-Pick disease, type C2
SYNJ2	synaptojanin 2
NIN	ninein (GSK3B interacting protein)
MBNL3	muscleblind-like 3 (Drosophila)
FLJ11259	damage-regulated autophagy modulator
NALP12	NLR family, pyrin domain containing 12
LIMK1	
ARSG	arylsulfatase G
FLJ33641	chromosome 5 open reading frame 29
PADI4	peptidyl arginine deiminase, type IV
RENBP	renin binding protein
SULF2	sulfatase 2
GSDMDC1	gasdermin domain containing 1
ST7	suppression of tumorigenicity 7
RBP7	retinol binding protein 7, cellular
HK2	hexokinase 2
VAMP5	vesicle-associated membrane protein 5 (myobrevin)
GPR177	G protein-coupled receptor 177
CES1	carboxylesterase 1 (monocyte/macrophage serine esterase 1)
CD52	CD52 molecule
ABCG1	ATP-binding cassette, sub-family G (WHITE), member 1
GBP5	guanylate binding protein 5
MDM4	Mdm4, transformed 3T3 cell double minute 4, p53 binding protein (mouse)
SIGLEC5	sialic acid binding Ig-like lectin 5
ARID3A	AT rich interactive domain 3A (BRIGHT-like)
KIAA0179	ribosomal RNA processing 1 homolog B (S. cerevisiae)
PSD4	pleckstrin and Sec7 domain containing 4
ALOX5AP	arachidonate 5-lipoxygenase-activating protein
CSF2RA	colony stimulating factor 2 receptor, alpha, low-affinity (granulocyte-macrophage)
MMP9	matrix metalloproteinase 9 (gelatinase B, 92kDa gelatinase, 92kDa type IV collagenase)
PGLYRP1	peptidoglycan recognition protein 1
CYP27A1	cytochrome P450, family 27, subfamily A, polypeptide 1
LMTK2	lemur tyrosine kinase 2
BRI3	brain protein I3

Gene Symbol	Description
PILRA	paired immunoglobulin-like type 2 receptor alpha
	Zinc finger protein 404
FLJ21127	tectonic 1
GBP2	guanylate binding protein 2, interferon-inducible
ST3GAL6	ST3 beta-galactoside alpha-2,3-sialyltransferase 6
PLAUR	plasminogen activator, urokinase receptor
NCF4	neutrophil cytosolic factor 4, 40kDa
JAK2	Janus kinase 2 (a protein tyrosine kinase)
SREBF1	sterol regulatory element binding transcription factor 1
SELM	selenoprotein M
PPP3R1	protein phosphatase 3 (formerly 2B), regulatory subunit B, alpha isoform
PRKCA	protein kinase C, alpha
PLA2G4C	phospholipase A2, group IVC (cytosolic, calcium-independent)
GBP4	guanylate binding protein 4
HIP1	huntingtin interacting protein 1
PGM2	phosphoglucomutase 2
	KIAA1632
S100P	S100 calcium binding protein P
IL27RA	interleukin 27 receptor, alpha
IL15	interleukin 15
FHIT	fragile histidine triad gene
FAM84B	family with sequence similarity 84, member B
MCCC1	methylcrotonoyl-Coenzyme A carboxylase 1 (alpha)
ACOT7	acyl-CoA thioesterase 7
TNFRSF12A	tumor necrosis factor receptor superfamily, member 12A
SF3A1	splicing factor 3a, subunit 1, 120kDa
TNFSF14	tumor necrosis factor (ligand) superfamily, member 14
CD300LB	CD300 molecule-like family member b
ANPEP	alanyl (membrane) aminopeptidase (aminopeptidase N, aminopeptidase M, microsomal aminopeptidase, CD13, p150)
FAM7A1	
RHD	Rh blood group, D antigen
HOM-TES-103	hypothetical protein LOC25900
CCDC41	coiled-coil domain containing 41
CRNKL1	crooked neck pre-mRNA splicing factor-like 1 (Drosophila)
NCF1	neutrophil cytosolic factor 1, (chronic granulomatous disease, autosomal 1)
UBTD1	ubiquitin domain containing 1
FLJ22471	coiled-coil domain containing 92
FAM101B	family with sequence similarity 101, member B
LOC284757	
LOC649377	
CDK5R1	cyclin-dependent kinase 5, regulatory subunit 1 (p35)
	Full-length cDNA clone CS0DC025YP03 of Neuroblastoma Cot 25-normalized of Homo sapiens (human)
MBNL3	muscleblind-like 3 (Drosophila)
PSTPIP1	proline-serine-threonine phosphatase interacting protein 1
WARS	tryptophanyl-tRNA synthetase
HLA-DPB1	major histocompatibility complex, class II, DP beta 1
LOC652616	

Gene Symbol	Description
ACTA2	actin, alpha 2, smooth muscle, aorta
IBRDC3	IBR domain containing 3
PHKG2	phosphorylase kinase, gamma 2 (testis)
	Phosphatidylinositol-4-phosphate 5-kinase, type II, alpha
LOC642035	
AMFR	
RGS19	regulator of G-protein signalling 19
C22orf5	chromosome 22 open reading frame 5
ATF3	activating transcription factor 3
SIPA1L1	signal-induced proliferation-associated 1 like 1
MRPS34	mitochondrial ribosomal protein S34
ADAL	adenosine deaminase-like
NDUFAF1	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, assembly factor 1
CRAT	carnitine acetyltransferase
STX11	syntaxin 11

Different and reciprocal immune signatures in active and latent TB are revealed using a modular approach. To yield further information on pathogenesis, the normalised per chip data was then further analyzed using a recently described stable modular analysis framework based on pre-defined clusters of genes transcripts shown to be coordinately expressed across a wide range of diseases, and often representing a cluster of molecules or cells related at a function level (Chaussabel et al., 2008, Immunity).

As the aim of this analysis was to yield functional information about genes contained within the transcriptional signatures for each group, the analysis was focused on subsets of patients found to cluster tightly together in our previous analyses, excluding outliers, reasoning that such groups would be more likely to reveal common pathways and processes involved in the disease process.

10 Nine patients with active TB, six healthy controls and nine patients with latent TB were selected and used in the modular analysis. Each comparison was performed separately, thus nine active TB patients were compared with six healthy controls in one analysis, and then nine latent TB patients were compared with the same six healthy controls in a separate analysis. Transcripts were filtered to exclude any not detected in at least two individuals from either group being compared. Statistical comparisons between patient and healthy control groups were then performed (Non parametric Wilcoxon-Mann-Whitney test, $P < 0.05$), in order to identify genes that were differentially expressed between the patient group and healthy controls. These differentially expressed genes were then separated into those upregulated / overrepresented in disease group compared with control, and those down-regulated/underrepresented in disease group compared with control. These lists are then analysed on a module by module basis. Differentially expressed genes are either predominantly over-expressed or predominantly under-expressed in each module. To ensure validity each module must have >25% of the total genes change in the direction represented and the number of genes changing in a particular direction must be >10. To graphically present the global transcriptional changes, in active TB versus healthy control, or latent TB versus healthy controls, spots are aligned on a grid, with each

position corresponding to a different module based on their original definition Spot intensity indicates proportion of differentially expressed transcripts changing in the direction shown out of the total number of transcripts detected for that module, while spot color indicates the polarity of the change (red: overexpressed/represented, blue: underexpressed/represented). In addition, modules' coordinates can be associated to functional annotations to facilitate data interpretation (Chaussabel, Immunity, 2008; and Figures 9 and 10).

A modular map of active TB compared to healthy control (Figure 9, Table 7A – P; and Table 8) was shown to be distinct to the map of latent TB as compared to healthy controls (Figure 10, Table 7A – F; and Table 9). In fact these independently derived module maps from active TB and latent TB show an inverse pattern of gene expression/representation, in modules which show changes in both disease states when compared with healthy controls. Genes in module M2.1 associated with cytotoxic cells were underexpressed/represented (36% - 18 genes underexpressed/represented out of 50 detected in the module, genes listed in Table 6F) in active TB and yet overexpressed/represented (43% - 22 genes overexpressed/represented out of 51 detected in the module, genes listed in Table 7B) in latent TB. On the other hand, a number of genes in M3.2 and M3.3 (“inflammation”) (genes listed in Tables 6J and 6K) were overexpressed/represented in active TB patients but underexpressed/represented in latent TB patients (genes listed in Table 7E and 7F). Likewise genes in M1.5 (“myeloid lineage”) were overexpressed/represented in active TB (genes listed in Table 6D) whereas they were underexpressed/represented in latent TB (genes listed in Table 7A). Genes in a module M2.10, which did not form a coherent functional module but consisted of an apparently diverse set of genes, were underexpressed/represented in latent TB (genes listed in Table 7D) but not over or underexpressed/represented in active TB as compared to controls. One of these genes is the toll-like receptor adaptor, TRAM, which is downstream of TLR-4 (LPS) and TLR-3 (dsRNA) signalling (Akira, Nat. Rev. Imm.).

For Tables 7A to 7O, relative normalized expression for active TB is given as expression in active patients relative to control. In Tables 8A to 8F, relative normalized expression for latent TB is given as expression in healthy controls relative to latent patients.

Table 7A M1.2 PTB v. Control, Genes Overrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Test_UP_M1.2		
2.447	KX; X1k; XKR1	XK	X-linked Kx blood group (McLeod syndrome)
2.239	CD62; GRMP; PSEL; CD62P; GMP140; PADGEM; FLJ45155	SELP	selectin P (granule membrane protein 140kDa, antigen CD62)
2.161	URG	EGF	epidermal growth factor (beta-urogastrone)
2.133	JAMC; JAM-C; FLJ14529	JAM3	junctional adhesion molecule 3

Relative normalised expression	Common Name	Gene Symbol	Description
2.13	H2B; GL105; H2B.1; H2B/q; H2BFQ; MGC129733; MGC129734	HIST2H2BE	histone cluster 2, H2be
1.889	4.1O; P410; EPB41L4O; MGC20553; RP11-439K3.2	FRMD3	FERM domain containing 3
1.875	CKLFSF5; FLJ37521	CMTM5	CKLF-like MARVEL transmembrane domain containing 5
1.829	ECM; MMRN; GPIa*; EMILIN4	MMRN1	multimerin 1
1.757	PSA; PROS; PS21; PS22; PS23; PS24; PS25; PS 26; Protein S; protein Sa	PROS1	protein S (alpha)
1.752	F13A	F13A1	coagulation factor XIII, A1 polypeptide
1.698	H2B/S; H2BFT; H2BFAiii; MGC131989	HIST1H2BK	histone cluster 1, H2bk
1.638		RTN2	
1.59	TMSA; HTM-alpha; TPM1-alpha; TPM1-kappa	TPM1	tropomyosin 1 (alpha)
1.419		C6orf79	
1.408	BSS; GPIB; CD42B; MGC34595; CD42b-alpha	GPIBA	glycoprotein Ib (platelet), alpha polypeptide
1.338	CD61; GP3A; GPIIIa	ITGB3	integrin, beta 3 (platelet glycoprotein IIIa, antigen CD61)
1.183	CMIP; KIAA1694	CMIP	c-Maf-inducing protein

Table 7B M1.3 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_ PAL2Test_DOWN_M1.3		
0.82	FLJ31738; KIAA1209	PLEKHG1	pleckstrin homology domain containing, family G (with RhoGef domain) member 1
0.778	SPI-B	SPIB	Spi-B transcription factor (Spi-1/PU.1 related)
0.767	EV19; CTIP1; BCL11A-L; BCL11A-S; FLJ10173; FLJ34997; KIAA1809; BCL11A-XL	BCL11A	B-cell CLL/lymphoma 11A (zinc finger protein)
0.715	MGC20446	CYBASC3	cytochrome b, ascorbate dependent 3
0.677	NIDD; MGC42530	ZDHHC23	zinc finger, DHHC-type containing 23
0.629	ESG; ESG1; GRG1	TLE1	transducin-like enhancer of split 1 (E(sp1) homolog, Drosophila)
0.612	B29; IGB	CD79B	CD79b molecule, immunoglobulin-associated beta
0.581	LYB2; CD72b	CD72	CD72 molecule
0.559	KIAA0977	COBLL1	COBL-like 1
0.556	BASH; Ly57; SLP65; BLNK-s; SLP-65; MGC111051	BLNK	B-cell linker
0.543	TCL1	TCL1A	T-cell leukemia/lymphoma 1A
0.518	c-Myc	MYC	v-myc myelocytomatosis viral oncogene homolog (avian)
0.512	BANK; FLJ20706; FLJ34204	BANK1	B-cell scaffold protein with ankyrin repeats 1
0.51	B4; MGC12802	CD19	CD19 molecule
0.496	FCRH1; IFGP1; IRTA5; RP11-	FCRL1	Fc receptor-like 1

Relative normalised expression	Common Name	Gene Symbol	Description
	367J7.7; DKFZp667O1421		
0.487	FLJ00058	GNG7	guanine nucleotide binding protein (G protein), gamma 7
0.482	FLJ21562; FLJ43762	C13orf18	chromosome 13 open reading frame 18
0.477	BRDG1; STAP1	BRDG1	BCR downstream signaling 1
0.471	MGC10442	BLK	B lymphoid tyrosine kinase
0.467	R1; JPO2; RAM2; DKFZp762L0311	CDCA7L	cell division cycle associated 7-like
0.445	ORP10; OSBP9; FLJ20363	OSBPL10	oxysterol binding protein-like 10
0.397	8HS20; N27C7-2	VPREB3	pre-B lymphocyte gene 3
0.361	LAF4; MLLT2-like	AFF3	AF4/FMR2 family, member 3
0.334	FCRL; FREB; FCRLX; FCRLb; FCRLd; FCRLe; FCRLM1; FCRLc1; FCRLc2; MGC4595; RP11-474I16.5	FCRLM1	Fc receptor-like A

Table 7C M1.4 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_DOWN_M1.4		
0.907	FLJ12298; ZKSCAN14	ZNF394	zinc finger protein 394
0.835	JMY; FLJ37870; MGC163496	JMY	junction-mediating and regulatory protein
0.825	C1; C2; HNRNP; SNRPC; hnRNP; MGC104306; MGC105117; MGC117353; MGC131677	HNRNP	heterogeneous nuclear ribonucleoprotein C (C1/C2)
0.78	SON3; BASS1; DBP-5; NREBP; C21orf50; FLJ21099; FLJ33914; KIAA1019	SON	SON DNA binding protein
0.77	HMGE; FLJ25609	GRPEL1	GrpE-like 1, mitochondrial (E. coli)
0.747	HEPP; FLJ20764; MGC19517	CDCA4	cell division cycle associated 4
0.723	RITA; ZNF361; ZNF463; DKFZp686L0787	ZNF331	zinc finger protein 331
0.698	FLJ12670; FLJ20436	C12orf41	chromosome 12 open reading frame 41
0.698	DRBF; MMP4; MPP4; NF90; NFAR; TCP80; DRBP76; NFAR-1; MPHOSPH4; NF-AT-90	ILF3	interleukin enhancer binding factor 3, 90kDa
0.689	TIMAP; ANKRD4; KIAA0823	PPP1R16B	protein phosphatase 1, regulatory (inhibitor) subunit 16B
0.678	PRP21; PRPF21; SAP114; SF3A120	SF3A1	splicing factor 3a, subunit 1, 120kDa
0.667	SDS; SWDS; CGI-97; FLJ10917	SBDS	Shwachman-Bodian-Diamond syndrome
0.665	BL11; HB15	CD83	CD83 molecule
0.645	NOT; RNR1; HZF-3; NURR1; TINUR	NR4A2	nuclear receptor subfamily 4, group A, member 2
0.62	H1RNA	RNASEH1	ribonuclease H1

Table 7D M1.5 PTB v. Control, Genes Overrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_UP_M1.5		
2.384	VHR	DUSP3	dual specificity phosphatase 3 (vaccinia virus phosphatase VH1-related)
2.139	4.1B; DAL1; DAL-1; FLJ37633; KIAA0987	EPB41L3	erythrocyte membrane protein band 4.1-like 3
2.014	HXK3; HKIII	HK3	hexokinase 3 (white cell)
1.972	HL14; MGC75071	LGALS2	lectin, galactoside-binding, soluble, 2
1.844	KYNU	KYNU	kynureninase (L-kynurenine hydrolase)
1.618	BLVR; BVRA	BLVRA	biliverdin reductase A
1.594	RP35; SEMB; SEMAB; CORD10; FLJ12287; RP11-54H19.2	SEMA4A	sema domain, immunoglobulin domain (Ig), transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 4A
1.535		GRN	
1.531	G6S; MGC21274	GNS	glucosamine (N-acetyl)-6-sulfatase (Sanfilippo disease IIID)
1.524	FOAP-10; EMILIN-2; FLJ33200	EMILIN2	elastin microfibril interfacier 2
1.507	cent-b; HSA272195	CENTA2	centaurin, alpha 2
1.449	APPS; CPSB	CTSB	cathepsin B
1.438	ASGPR; CLEC4H1; Hs.12056	ASGR1	asialoglycoprotein receptor 1
1.433	CD32; FCG2; FcGR; CD32A; CDw32; FCGR2; IGFR2; FCGR2A1; MGC23887; MGC30032	FCGR2A	Fc fragment of IgG, low affinity IIa, receptor (CD32)
1.425	TIL4; CD282	TLR2	toll-like receptor 2
1.424	PI; A1A; AAT; PI1; A1AT; MGC9222; PRO2275; MGC23330	SERPINA1	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 1
1.413	TEM7R; FLJ14623	PLXDC2	plexin domain containing 2
1.41	CD14	CD14	CD14 molecule
1.398	Rab22B	RAB31	RAB31, member RAS oncogene family
1.386	FEX1; FEEL-1; FELE-1; STAB-1; CLEVER-1; KIAA0246	STAB1	stabilin 1
1.352	MYD88	MYD88	myeloid differentiation primary response gene (88)
1.349	MLN70; S100C	S100A11	S100 calcium binding protein A11
1.347	FLJ22662	FLJ22662	hypothetical protein FLJ22662
1.346	CLN2; GIG1; LPIC; TPP I; MGC21297	TPP1	tripeptidyl peptidase I
1.251	p75; TBPII; TNFBR; TNFR2; CD120b; TNFR80; TNF-R75; p75TNFR; TNF-R-II	TNFRSF1B	tumor necrosis factor receptor superfamily, member 1B
1.239	JTK9	HCK	hemopoietic cell kinase
1.172	IBA1; AIF-1; IRT-1	AIF1	allograft inflammatory factor 1

Table 7E M1.8 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_UP_M1.5		

Relative normalised expression	Common Name	Gene Symbol	Description
	8_PAL2Ttest_DOWN_M1.8		
0.878	DBP2; PRP8; DDX16; PRO2014	DHX16	DEAH (Asp-Glu-Ala-His) box polypeptide 16
0.858	AN11; HAN11	WDR68	WD repeat domain 68
0.843	NDR; NDR1	STK38	serine/threonine kinase 38
0.821	FLJ20097; FLJ23581; KIAA1861	FLJ20097	coiled-coil domain containing 132
0.814	FLJ42526; FLJ45813; MGC71764	RSBN1L	round spermatid basic protein 1-like
0.809	C9orf55; C9orf55B; FLJ20686; bA513M16.3; DKFZp686I09113	DENND4C	DENN/MADD domain containing 4C
0.808	SON3; BASS1; DBP-5; NREBP; C21orf50; FLJ21099; FLJ33914; KIAA1019	SON	SON DNA binding protein
0.807	p150; VPS15; MGC102700	PIK3R4	phosphoinositide-3-kinase, regulatory subunit 4, p150
0.8	4E-T; Clast4; FLJ21601; FLJ26551	EIF4ENIF1	eukaryotic translation initiation factor 4E nuclear import factor 1
0.798	TAF2D; TAFII100	TAF5	TAF5 RNA polymerase II, TATA box binding protein (TBP)-associated factor, 100kDa
0.793	DBR1	DBR1	debranching enzyme homolog 1 (S. cerevisiae)
0.785	SMAP; p120; SMAP2	BRD8	bromodomain containing 8
0.785		CASP2	
0.772	TRF2; TRBF2	TERF2	telomeric repeat binding factor 2
0.772	hNUP133; FLJ10814; MGC21133	NUP133	nucleoporin 133kDa
0.762	MGC4268; FLJ38552	MGC4268	AMME chromosomal region gene 1-like
0.761	PUMH2; PUML2; FLJ36528; KIAA0235; MGC138251; MGC138253	PUM2	pumilio homolog 2 (Drosophila)
0.751	BYE1; DIO1; DATF1; DIDO2; DIDO3; DIO-1; FLJ11265; KIAA0333; MGC16140; C20orf158; dJ885L7.8; DKFZp434P1115	DIDO1	death inducer-obliterator 1
0.738	KOX5; ZNF13	ZNF45	zinc finger protein 45
0.727	FLJ20558	FLJ20558	chromosome 2 open reading frame 42
0.713	FLJ32343	CWF19L2	CWF19-like 2, cell cycle control (S. pombe)
0.709	MGC16770	RAB22A	RAB22A, member RAS oncogene family
0.708	FLJ14431	CBR4	carbonyl reductase 4
0.704	AASDH; NRPS998; NRPS1098	AASDH	2-aminoadipic 6-semialdehyde dehydrogenase
0.698	ZSCAN11	ZNF232	zinc finger protein 232
0.692	NudCL; KIAA1068	NUDCD3	NudC domain containing 3
0.691	CCA1; MtCCA; CGI-47	TRNT1	tRNA nucleotidyl transferase, CCA-adding, 1
0.689	RBM30; RBM4L; ZCRB3B; ZCCHC15; MGC10871	RBM4B	RNA binding motif protein 4B
0.683	CLF; CRN; HCRN; SYF3; MSTP021	CRNKL1	crooked neck pre-mRNA splicing factor-like 1 (Drosophila)

Relative normalised expression	Common Name	Gene Symbol	Description
0.676	ZBU1; HLTF1; RNF80; HIP116; SNF2L3; HIP116A; SMARCA3	SMARCA3	helicase-like transcription factor
0.666	SWAN; KIAA0765; HRIHFB2091	RBM12	RNA binding motif protein 12
0.658	FLJ10287; FLJ11219	CCDC76	coiled-coil domain containing 76
0.654	INT5; KIAA1698	KIAA1698	integrator complex subunit 5
0.652	IAN7; hIAN7; MGC27027	GIMAP7	GTPase, IMAP family member 7
0.651	TTC20; DKFZP586B0923	KIAA1279	KIAA1279
0.65	RAL; MGC48949	RALA	v-ral simian leukemia viral oncogene homolog A (ras related)
0.639	MPRB; LMPB1; C6orf33; FLJ32521; FLJ46206	PAQR8	progesterin and adipoQ receptor family member VIII
0.634	FLJ11171	FLJ11171	hypothetical protein FLJ11171
0.613	LCF; IL-16; prIL-16; FLJ16806; FLJ42735; FLJ44234; HsT19289	IL16	interleukin 16 (lymphocyte chemoattractant factor)
0.611	FLJ33226; 1190004M21Rik	PYGO2	pygopus homolog 2 (Drosophila)
0.577	GLC1G; UTP21; TAWDRP; TA-WDRP; DKFZp686I1650	WDR36	WD repeat domain 36
0.574	FLJ20287; bA208F1.2; RP11-208F1.2	TEX10	testis expressed 10
0.568	KIAA1982	ZNF721	zinc finger protein 721
0.55	FLJ22457; RP5-1180E21.2	DENND2D	DENN/MADD domain containing 2D
0.545	ozrf1; ZFP260	ZFP260	zinc finger protein 260
0.491	GLS1; FLJ10358; KIAA0838; DKFZp686O15119	GLS	glutaminase

Table 7F M2.1 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_DOWN_M2.01		
0.712	PTPMEG; PTPMEG1	PTPN4	protein tyrosine phosphatase, non-receptor type 4 (megakaryocyte)
0.665	FLJ34563; MGC35163	SAMD3	sterile alpha motif domain containing 3
0.643	STAT4	STAT4	signal transducer and activator of transcription 4
0.638	DIL1; DIL-1; Mindin; M-spondin	SPON2	spondin 2, extracellular matrix protein
0.631	SLP2; SGA72M; CHR11SYT; KIAA1597; MGC102768	SYTL2	synaptotagmin-like 2
0.628	DORZ1; DKFZP564O243	ABHD14A	abhydrolase domain containing 14A
0.615	LPAP; CD45-AP; MGC138602; MGC138603	PTPRCAP	protein tyrosine phosphatase, receptor type, C-associated protein
0.595	PKCL; PKC-L; PRKCL; MGC5363; MGC26269; nPKC-eta	PRKCH	protein kinase C, eta
0.581	MGC33870; MGC74858	NCALD	neurocalcin delta
0.566	T11; SRBC	CD2	CD2 molecule
0.554	KLR; CD314; NKG2D; NKG2-	KLRK1	killer cell lectin-like receptor subfamily K,

	D; D12S2489E		member 1
0.546	LAX; FLJ20340	LAX1	lymphocyte transmembrane adaptor 1
0.529	CD122; P70-75	IL2RB	interleukin 2 receptor, beta
0.515	FEZ1	FEZ1	fasciculation and elongation protein zeta 1 (zygin I)
0.509	CHK; CTK; HYL; Lsk; HYLTK; HHYLTK; MGC1708; MGC2101; DKFZp434N1212	MATK	megakaryocyte-associated tyrosine kinase
0.468	CLIC3	CLIC3	chloride intracellular channel 3
0.439	1C7; CD337; LY117; NKp30	NCR3	natural cytotoxicity triggering receptor 3
0.39	TRYP2	GZMK	granzyme K (granzyme 3; tryptase II)

Table 7G M2.4 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_DOWN_M2.04		
0.858	ATPO; OSCP	ATP5O	ATP synthase, H+ transporting, mitochondrial F1 complex, O subunit (oligomycin sensitivity conferring protein)
0.831	M9; eIF3k; ARG134; PTD001; HSPC029; MSTP001; PLAC-24; PRO1474	EIF3S12	eukaryotic translation initiation factor 3, subunit 12
0.822	RPL8	RPL8	ribosomal protein L8
0.811	EF2; EEF-2	EEF2	eukaryotic translation elongation factor 2
0.804	RPB9; hRPB14.5	POLR2I	polymerase (RNA) II (DNA directed) polypeptide I, 14.5kDa
0.801	RP8; ZMYND7; MGC12347	PDCD2	programmed cell death 2
0.788	ARI2; TRIAD1; FLJ10938; FLJ33921	ARIH2	ariadne homolog 2 (Drosophila)
0.776	Erv46; CGI-54; PRO0989; C20orf47; NY-BR-84; SDBCAG84; dJ477O4.2	ERGIC3	ERGIC and golgi 3
0.771	ART-27	UXT	ubiquitously-expressed transcript
0.769	H12.3; HLC-7; PIG21; RACK1; Gnb2-rs1	GNB2L1	guanine nucleotide binding protein (G protein), beta polypeptide 2-like 1
0.766	eIF3h; eIF3-p40; MGC102958; eIF3-gamma	EIF3S3	eukaryotic translation initiation factor 3, subunit 3 gamma, 40kDa
0.759	HCA56	LGTN	ligatin
0.758	2PP2A; IGAAD; I2PP2A; PHAPII; TAF-IBETA	SET	SET translocation (myeloid leukemia-associated)
0.752	ANG2	C11orf2	chromosome 11 open reading frame2
0.74	C6.1B	MTCP1	mature T-cell proliferation 1
0.736	LCP; HCLP-1	KLHDC2	kelch domain containing 2
0.722	DKFZP566B023	RPL36	ribosomal protein L36
0.712	KOX30	ZNF32	zinc finger protein 32
0.71	AMP; MGC125856; MGC125857; MGC129961; DKFZp686D13177	APRT	adenine phosphoribosyltransferase
0.694	GDH; MGC149525; MGC149526; lambda-CRY	CRYL1	crystallin, lambda 1
0.689	FLJ27451; MGC102930	RPS20	ribosomal protein S20

Relative normalised expression	Common Name	Gene Symbol	Description
0.686	INT6; eIF3e; EIF3-P48; eIF3-p46	EIF3S6	eukaryotic translation initiation factor 3, subunit 6 48kDa
0.68	LK4; hCERK; FLJ21430; FLJ23239; KIAA1646; MGC131878; dA59H18.2; dA59H18.3; DKFZp434E0211	CERK	ceramide kinase
0.675	HINT; PKCI-1; PRKCNH1	HINT1	histidine triad nucleotide binding protein 1
0.675	NHP2; NHP2P	NOLA2	nucleolar protein family A, member 2 (H/ACA small nucleolar RNPs)
0.668	AMP; MGC125856; MGC125857; MGC129961; DKFZp686D13177	APRT	adenine phosphoribosyltransferase
0.667	TOM7	TOMM7	translocase of outer mitochondrial membrane 7 homolog (yeast)
0.655	SIVA; CD27BP; Siva-1; Siva-2	SIVA	SIVA1, apoptosis-inducing factor
0.646	PBP; HCNP; PEBP; RKIP	PEBP1	phosphatidylethanolamine binding protein 1
0.628	PRP9; PRPF9; SAP61; SF3a60	SF3A3	splicing factor 3a, subunit 3, 60kDa
0.62	FLJ12525; dJ475B7.2; RP3-475B7.2	LAS1L	LAS1-like (<i>S. cerevisiae</i>)
0.593	EC45; RPL10; RPLY10; RPYL10; FLJ26304; MGC88603	RPL15	ribosomal protein L15
0.567	HNRNP; JKTBP; JKTBP2; laAUF1	HNRPDL	heterogeneous nuclear ribonucleoprotein D-like
0.562	SMD2; SNRPD1	SNRPD2	small nuclear ribonucleoprotein D2 polypeptide 16.5kDa
0.549		PPIA	
0.527	LOC130074; MGC87527	LOC130074	p20
0.524	RDGBB; RDGBB1; RDGB-BETA	PITPNC1	phosphatidylinositol transfer protein, cytoplasmic 1
0.5	HEI10; C14orf18	CCNB1IP1	cyclin B1 interacting protein 1
0.492	EAP; HBP15; HBP15/L22	RPL22	ribosomal protein L22

Table 7H M2.8 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_DOWN_M2.08		
0.871	KPL1; PHR1; PHRET1	PLEKHB1	pleckstrin homology domain containing, family B (evectins) member 1
0.816	MGC132014	INPP4B	inositol polyphosphate-4-phosphatase, type II, 105kDa
0.732	SEP2; SEPT2; KIAA0128; MGC16619; MGC20339; RP5-876A24.2	6-Sep	septin 6
0.711	GIL	AQP3	aquaporin 3 (Gill blood group)
0.691	FLJ36386	LZTFL1	leucine zipper transcription factor-like 1
0.67	p52; p75; PAIP; DFS70; LEDGF; PSIP2; MGC74712	PSIP1	PC4 and SFRS1 interacting protein 1
0.669	GRG; ESP1; GRG5; TLE5; AES-1; AES-2	AES	amino-terminal enhancer of split

Relative normalised expression	Common Name	Gene Symbol	Description
0.668	p33; TNFC; TNFSF3	LTB	lymphotoxin beta (TNF superfamily, member 3)
0.646	KIAA0521; MGC15913	ARHGEF18	rho/rac guanine nucleotide exchange factor (GEF) 18
0.634	TEM3; TEM7; FLJ36270; FLJ45632; DKFZp686F0937	PLXDC1	plexin domain containing 1
0.626	HPIP	PBXIP1	pre-B-cell leukemia homeobox interacting protein 1
0.621	KIAA0495; MGC138189	KIAA0495	KIAA0495
0.615	KUP; ZNF46	ZBTB25	zinc finger and BTB domain containing 25
0.61	FLJ20729; FLJ20760; NY-BR-75; MGC131963	C1orf181	chromosome 1 open reading frame 181
0.609	AAG6; PKCA; PRKACA; MGC129900; MGC129901; PKC-alpha	PRKCA	protein kinase C, alpha
0.604	CGI-25	NOSIP	nitric oxide synthase interacting protein
0.602	FLJ20152; FLJ22155; FLJ22179	FLJ20152	family with sequence similarity 134, member B
0.599	FRA3B; AP3Aase	FHIT	fragile histidine triad gene
0.596	WDR74	WDR74	WD repeat domain 74; synonyms: FLJ10439, FLJ21730; Homo sapiens WD repeat domain 74 (WDR74), mRNA.
0.595	E25A; BRICD2A	ITM2A	integral membrane protein 2A
0.587	HPF2	ZNF84	zinc finger protein 84
0.58	SEK; HEK8; TYRO1	EPHA4	EPH receptor A4
0.578	SID1; SID-1; FLJ20174; B830021E24Rik	SIDT1	SID1 transmembrane family, member 1
0.557	LTBP2; LTBP-3; pp6425; FLJ33431; FLJ39893; FLJ42533; FLJ44138; DKFZP586M2123	LTBP3	latent transforming growth factor beta binding protein 3
0.556	V; RASGRP; hRasGRP1; MGC129998; MGC129999; CALDAG-GEFI; CALDAG-GEFII	RASGRP1	RAS guanyl releasing protein 1 (calcium and DAG-regulated)
0.546	TTF; ARHH	RHOH	ras homolog gene family, member H
0.545	LAT3; LAT-2; y+LAT-2; KIAA0245; DKFZp686K15246	SLC7A6	solute carrier family 7 (cationic amino acid transporter, y+ system), member 6
0.541	TP120	CD6	CD6 molecule
0.537	MGC29816	CHMP7	CHMP family, member 7
0.53	DAGK; DAGK1; MGC12821; MGC42356; DGK-alpha	DGKA	diacylglycerol kinase, alpha 80kDa
0.523	hly9; mLY9; CD229; SLAMF3	LY9	lymphocyte antigen 9
0.52	EMT; LYK; PSCTK2; MGC126257; MGC126258	ITK	IL2-inducible T-cell kinase
0.519	TACTILE; MGC22596; DKFZp667E2122	CD96	CD96 molecule
0.518	SEP2; SEPT2; KIAA0128; MGC16619; MGC20339; RP5-876A24.2	6-Sep	septin 6
0.501	SCAP1; SKAP55	SCAP1	src kinase associated phosphoprotein 1
0.49	FLJ12884; MGC130014; MGC130015	C10orf38	chromosome 10 open reading frame 38

Relative normalised expression	Common Name	Gene Symbol	Description
0.488	T1; LEU1	CD5	CD5 molecule
0.487	MAL	MAL	mal, T-cell differentiation protein
0.484	SATB1	SATB1	SATB homeobox 1
0.48	LDH-H; TRG-5	LDHB	lactate dehydrogenase B
0.473	Ray; FLJ39121; DKFZP586F1318	SH3YL1	SH3 domain containing, Ysc84-like 1 (S. cerevisiae)
0.466	P19; SGRF; IL-23; IL-23A; IL23P19; MGC79388	IL23A	interleukin 23, alpha subunit p19
0.465	KE6; FABG; HKE6; FABGL; RING2; H2-KE6; D6S2245E; dJ1033B10.9	HSD17B8	hydroxysteroid (17-beta) dehydrogenase 8
0.456	ARH; ARH1; ARH2; FHCB1; FHCB2; MGC34705; DKFZp586D0624	LDLRAP1	low density lipoprotein receptor adaptor protein 1
0.453	MGC45416; DKFZp686C03164	OCIAD2	OCIA domain containing 2
0.451	CD172g; SIRPB2; SIRP-B2; bA77C3.1; SIRPgamma	SIRPB2	signal-regulatory protein gamma
0.435	GP40; TP41; Tp40; LEU-9	CD7	CD7 molecule
0.427	MGC15763	MGC15763	oxidoreductase NAD-binding domain containing 1
0.41	AS160; DKFZp779C0666	TBC1D4	TBC1 domain family, member 4
0.404	HMIC; MANIC; MAN1A3; pp6318	MAN1C1	mannosidase, alpha, class 1C, member 1
0.401	Tp44; MGC138290	CD28	CD28 molecule
0.394	FLJ12586	ZNF329	zinc finger protein 329
0.39	TCF-1; MGC47735	TCF7	transcription factor 7 (T-cell specific, HMG-box)
0.385	ABLIM; LIMAB1; LIMATIN; MGC1224; FLJ14564; KIAA0059; DKFZp781D0148	ABLIM1	actin binding LIM protein 1
0.383	NSE2; BCMP101	FAM84B	family with sequence similarity 84, member B
0.377	TOSO	FAIM3	Fas apoptotic inhibitory molecule 3
0.371	EEIG1; C9orf132; MGC50853; bA203J24.7	C9orf132	family with sequence similarity 102, member A
0.36	RIT1; CTIP2; CTIP-2; hRIT1-alpha	BCL11B	B-cell CLL/lymphoma 11B (zinc finger protein)
0.33	CLP24; FLJ20898; MGC111564	C16orf30	chromosome 16 open reading frame 30
0.315	TCF1ALPHA; DKFZp586H0919	LEF1	lymphoid enhancer-binding factor 1
0.29	BLR2; EB11; CD197; CDw197; CMKBR7	CCR7	chemokine (C-C motif) receptor 7
0.244	STK37; PASKIN; KIAA0135; DKFZP434O051; DKFZp686P2031	PASK	PAS domain containing serine/threonine kinase
0.205	NRP2	NELL2	NEL-like 2 (chicken)

Table 7I M3.1 PTB v. Control, Genes Overrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_UP_M3.1		
17.93	MGC22805	ANKRD22	ankyrin repeat domain 22
14.86	C1IN; C1NH; HAE1; HAE2; C1INH	SERPING1	serpin peptidase inhibitor, clade G (C1 inhibitor), member 1, (angioedema, hereditary)
9.425	cig5; vig1; 2510004L01Rik	RSAD2	radical S-adenosyl methionine domain containing 2
8.938	BRES1; MGC29634	EPST11	epithelial stromal interaction 1 (breast)
8.226	GS3686; C1orf29	IFI44L	interferon-induced protein 44-like
7.566	GBP1	GBP1	guanylate binding protein 1, interferon-inducible, 67kDa
5.677	p44; MTAP44	IFI44	interferon-induced protein 44
4.701	LAP; PEPS; LAPEP	LAP3	leucine aminopeptidase 3
4.401	IRG2; IFI60; IFIT4; ISG60; RIG-G; CIG-49; GARG-49	IFIT3	interferon-induced protein with tetratricopeptide repeats 3
4.091	OIAS; IFI-4; OIASI	OAS1	2',5'-oligoadenylate synthetase 1, 40/46kDa
3.947	p100; MGC133260	OAS3	2'-5'-oligoadenylate synthetase 3, 100kDa
3.944	G1P2; UCRP; IFI15	G1P2	ISG15 ubiquitin-like modifier
3.915	UEF1; DRIF2; C7orf6; FLJ39885; KIAA2005	SAMD9L	sterile alpha motif domain containing 9-like
3.909	MMTRA1B	PLSCR1	phospholipid scramblase 1
3.792	XAF1; BIRC4BP; HSXIAPAF1	BIRC4BP	XIAP associated factor-1
3.731	RIGE; SCA2; RIG-E; SCA-2; TSA-1	LY6E	lymphocyte antigen 6 complex, locus E
3.726	C7; IFI10; INP10; IP-10; crg-2; mob-1; SCYB10; gIP-10	CXCL10	chemokine (C-X-C motif) ligand 10
3.668	FBG2; FBS2; FBX6; Fbx6b	FBXO6	F-box protein 6
3.652	RNF94; STAF50; GPSTAF50	TRIM22	tripartite motif-containing 22
3.619	LOC129607	LOC129607	hypothetical protein LOC129607
3.419	ISGF-3; STAT91; DKFZp686B04100	STAT1	signal transducer and activator of transcription 1, 91kDa
3.398	TRIP14; p59OASL	OASL	2'-5'-oligoadenylate synthetase-like
3.284	IFP35; FLJ21753	IFI35	interferon-induced protein 35
3.154	LOC26010; DNAPTP6; DKFZp564A2416	DNAPTP6	viral DNA polymerase-transactivated protein 6
3.076	BAL; BAL1; FLJ26637; FLJ41418; MGC:7868; DKFZp666B0810; DKFZp686M15238	PARP9	poly (ADP-ribose) polymerase family, member 9
3.032	BAL2; KIAA1268	PARP14	poly (ADP-ribose) polymerase family, member 14
2.977	RIG-B; UBCH8; MGC40331	UBE2L6	ubiquitin-conjugating enzyme E2L 6
2.839	APT1; PSF1; ABC17; ABCB2; RING4; TAP1N; D6S114E; FLJ26666; FLJ41500; TAP1*0102N	TAP1	transporter 1, ATP-binding cassette, sub-family B (MDR/TAP)
2.814	MX; MxA; IFI78; IFI-78K	MX1	myxovirus (influenza virus) resistance 1, interferon-inducible protein p78 (mouse)
2.632		IRF7	
2.511	GCH; DYT5; GTPCH1; GTP-	GCH1	GTP cyclohydrolase 1 (dopa-responsive

Relative normalised expression	Common Name	Gene Symbol	Description
	CH-1		dystonia)
2.434	9-27; CD225; IFI17; LEU13	IFITM1	interferon induced transmembrane protein 1 (9-27)
2.415	G10P2; IFI54; ISG54; cig42; IFI-54; GARG-39; ISG-54K	IFIT2	interferon-induced protein with tetratricopeptide repeats 2
2.414	Hlcd; MDA5; MDA-5; IDDM19; MGC133047	IFIH1	interferon induced with helicase C domain 1
2.378	P113; ISGF-3; STAT113; MGC59816	STAT2	signal transducer and activator of transcription 2, 113kDa
2.321	TL2; APO2L; CD253; TRAIL; Apo-2L	TNFSF10	tumor necrosis factor (ligand) superfamily, member 10
2.32	TEL2; TELB; TEL-2	ETV7	ets variant gene 7 (TEL2 oncogene)
2.214	OIAS; IFI-4; OIASI	OAS1	2',5'-oligoadenylate synthetase 1, 40/46kDa
2.206	APT2; PSF2; ABC18; ABCB3; RING11; D6S217E	TAP2	transporter 2, ATP-binding cassette, sub-family B (MDR/TAP)
2.134	MGC78578	OAS2	2'-5'-oligoadenylate synthetase 2, 69/71kDa
2	VRK2	VRK2	vaccinia related kinase 2
1.975	PN-I; PSN1; UMPH; UMPH1; P5'N-1; cN-III; MGC27337; MGC87109; MGC87828	NT5C3	5'-nucleotidase, cytosolic III
1.895	RNF88; TRIM5alpha	TRIM5	tripartite motif-containing 5
1.89	CGI-34; PNAS-2; C9orf83; HSPC177; SNF7DC2	CHMP5	chromatin modifying protein 5
1.863	ZC3H1; PARP-12; ZC3HDC1; FLJ22693	PARP12	poly (ADP-ribose) polymerase family, member 12
1.845	PKR; PRKR; EIF2AK1; MGC126524	EIF2AK2	eukaryotic translation initiation factor 2-alpha kinase 2
1.842	90K; MAC-2-BP	LGALS3BP	lectin, galactoside-binding, soluble, 3 binding protein
1.807	RNF88; TRIM5alpha	TRIM5	tripartite motif-containing 5
1.743	C15; onzin	PLAC8	placenta-specific 8
1.732	p48; IRF9; IRF-9; ISGF3	ISGF3G	interferon-stimulated transcription factor 3, gamma 48kDa
1.713	CD317	BST2	bone marrow stromal cell antigen 2
1.665	ESNA1; ERAP140; FLJ45605; MGC88425; Nbla00052; Nbla10993; dJ187J11.3	NCOA7	nuclear receptor coactivator 7
1.649	FLJ39275; MGC131926	ZNFX1	zinc finger, NFX1-type containing 1
1.628	VODI; IFI41; IFI75; FLJ22835	SP110	SP110 nuclear body protein
1.627	EFP; Z147; RNF147; ZNF147	TRIM25	tripartite motif-containing 25
1.523	NMI	NMI	N-myc (and STAT) interactor
1.505	TRAP; KIAA1529; PCTAIRE2BP; RP11-508D10.1	TDRD7	tudor domain containing 7
1.499	DSH; G1P1; IFI4; p136; ADAR1; DRADA; DSRAD; IFI-4; K88dsRBP	ADAR	adenosine deaminase, RNA-specific
1.494	C1GALT; T-synthase	C1GALT1	core 1 synthase, glycoprotein-N-acetylgalactosamine 3-beta-galactosyltransferase, 1
1.478		PHF11	
1.461	SCOTIN	SCOTIN	scotin

Relative normalised expression	Common Name	Gene Symbol	Description
1.433	FLJ00340; FLJ34579; DKFZp686E07254	SP100	SP100 nuclear antigen
1.415	FLJ45064	AGRN	agrin
1.351	NFTC; OEF1; OEF2; C7orf5; FLJ20073; KIAA2004	SAMD9	sterile alpha motif domain containing 9
1.26	MEL; RAB8	RAB8A	RAB8A, member RAS oncogene family
1.215	6-16; G1P3; FAM14C; IFI616; IFI-6-16	G1P3	interferon, alpha-inducible protein 6

Table 7J M3.2 PTB v. Control, Genes Overrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_UP_M3.2		
2.767	MGC20461	OSM	oncostatin M
2.202	FHL4; HLH4; HPLH4	STX11	syntaxin 11
2.136	LPCAT2; FLJ20481; LysoPAFAT; DKFZp686H22112	AYTL1	acyltransferase like 1
1.987	UP; UPP; UPASE; UDRPASE	UPP1	uridine phosphorylase 1
1.969	IL-1; IL1F2; IL1-BETA	IL1B	interleukin 1, beta
1.886	SAT; DC21; KFSB; SSAT; SSAT-1	SAT	spermidine/spermine N1-acetyltransferase 1
1.862	PFK2; IPFK2	PFKFB3	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 3
1.755	BB2; CD54; P3.58	ICAM1	intercellular adhesion molecule 1 (CD54), human rhinovirus receptor
1.742	BCL4; D19S37	BCL3	B-cell CLL/lymphoma 3
1.695	KRML; MGC43127	MAFB	v-maf musculoaponeurotic fibrosarcoma oncogene homolog B (avian)
1.686	SRPSOX; CXCLG16; SR-PSOX	CXCL16	chemokine (C-X-C motif) ligand 16
1.658	B3GN-T5; beta3Gn-T5	B3GNT5	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminyltransferase 5
1.62	MLA1; ME491; LAMP-3; OMA81H; TSPAN30	CD63	CD63 molecule
1.562	P21; CIP1; SDI1; WAF1; CAP20; CDKN1; MDA-6; p21CIP1	CDKN1A	cyclin-dependent kinase inhibitor 1A (p21, Cip1)
1.548	URAX1; TAIP-3; FAM130B; DKFZp566F164	AXUD1	AXIN1 up-regulated 1
1.542	NHE8; FLJ42500; KIAA0939; MGC138418; DKFZp686C03237	SLC9A8	solute carrier family 9 (sodium/hydrogen exchanger), member 8
1.542	GS; GLNS; PIG43	GLUL	glutamate-ammonia ligase (glutamine synthetase)
1.504	CD87; UPAR; URKR	PLAUR	plasminogen activator, urokinase receptor
1.474	PBEF; NAMPT; MGC117256; DKFZP666B131; 1110035014Rik	PBEF1	pre-B-cell colony enhancing factor 1
1.472	P47; FLJ27168	PLEK	pleckstrin

Relative normalised expression	Common Name	Gene Symbol	Description
1.45	GNA16	GNA15	guanine nucleotide binding protein (G protein), alpha 15 (Gq class)
1.435	FTH; PLIF; FTHL6; PIG15; MGC104426	FTH1	ferritin, heavy polypeptide 1
1.42	MGC14376; MGC149751; DKFZp6886006159	MGC14376	hypothetical protein MGC14376
1.395	NER; UNR; LXRB; LXR-b; NER-I; RIP15	NR1H2	nuclear receptor subfamily 1, group H, member 2
1.39	TTP; GOS24; GOS24; TIS11; NUP475; RNF162A	ZFP36	zinc finger protein 36, C3H type, homolog (mouse)
1.389	E4BP4; IL3BP1; NFIL3A; NF-IL3A	NFIL3	nuclear factor, interleukin 3 regulated
1.328	C8FW; GIG2; SKIP1	TRIB1	tribbles homolog 1 (Drosophila)
1.296	ARI; HARI; HHARI; UBCH7BP	ARIH1	ariadne homolog, ubiquitin-conjugating enzyme E2 binding protein, 1 (Drosophila)
1.272	FRA2; FLJ23306	FOSL2	FOS-like antigen 2
1.269	RIT; RIBB; ROC1; MGC125864; MGC125865	RIT1	Ras-like without CAAX 1
1.25	RBT1	SERTAD3	SERTA domain containing 3
1.227	MAPKAPK2	MAPKAPK2	mitogen-activated protein kinase-activated protein kinase 2
1.217	PPG; PRG; PRG1; MGC9289; FLJ12930	PRG1	serglycin
1.181	SEI1; TRIP-Br1	SERTAD1	SERTA domain containing 1
1.172	CMT2; KIAA0110; MGC11282; RP1-261G23.6	MAD2L1BP	MAD2L1 binding protein
1.169	UBP; SIH003; MGC129878; MGC129879	USP3	ubiquitin specific peptidase 3

Table 7K M3.3 PTB v. Control, Genes Overrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_UP_M3.3		
3.651	MAYP; MGC34175	PSTPIP2	proline-serine-threonine phosphatase interacting protein 2
3.2	Tiff66; MGC116930; MGC116931; MGC116932; MGC116933	VNN1	vanin 1
2.604	Rsc6p; BAF60C; CRACD3; MGC111010	SMARCD3	SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily d, member 3
2.157	FER1L1; LGMD2B; FLJ00175; FLJ90168	DYSF	dysferlin, limb girdle muscular dystrophy 2B (autosomal recessive)
2.091	ASRT5; IRAKM; IRAK-M	IRAK3	interleukin-1 receptor-associated kinase 3
2.082	p6; CAGC; CGRP; MRP6; CAAF1; ENRAGE	S100A12	S100 calcium binding protein A12
1.888	CGI-44	SQRDL	sulfide quinone reductase-like (yeast)
1.819	FAM31A; FLJ38464; KIAA1608; RP11-230L22.3	DENND1A	DENN/MADD domain containing 1A
1.736	APG3; APG3L; PC3-96; FLJ22125; MGC15201;	ATG3	ATG3 autophagy related 3 homolog (S. cerevisiae)

Relative normalised expression	Common Name	Gene Symbol	Description
	DKFZp564M1178		
1.715	CAT1	CRAT	carnitine acetyltransferase
1.703	MGC2654; FLJ12433	MGC2654	chromosome 16 open reading frame 68
1.7	MD-2	LY96	lymphocyte antigen 96
1.695	AD3; VRP; HBLP1	TBC1D8	TBC1 domain family, member 8 (with GRAM domain)
1.663	FLJ20424	C14orf94	chromosome 14 open reading frame 94
1.638	P28; GSTTLp28; DKFZp686H13163	GSTO1	glutathione S-transferase omega 1
1.635	ATRAP; MGC29646	AGTRAP	angiotensin II receptor-associated protein
1.572	FAT; GP4; GP3B; GPIV; CHDS7; PASIV; SCARB3	CD36	CD36 molecule (thrombospondin receptor)
1.547	EI; LEI; PI2; MNEI; M/NEI; ELANH2	SERPINB1	serpin peptidase inhibitor, clade B (ovalbumin), member 1
1.546	RAB32	RAB32	RAB32, member RAS oncogene family
1.541	CR3A; MO1A; CD11B; MAC-1; MAC1A; MGC117044	ITGAM	integrin, alpha M (complement component 3 receptor 3 subunit)
1.481	ALFY; ZFYVE25; KIAA0993; MGC16461	WDFY3	WD repeat and FYVE domain containing 3
1.467	ARHU; WRCH1; hG28K; CDC42L1; FLJ10616; DJ646B12.2; fJ646B12.2	RHOU	ras homolog gene family, member U
1.459	SELR; SELX; MSRB1; HSPC270; MGC3344	SEPX1	selenoprotein X, 1
1.432	LTA4H	LTA4H	leukotriene A4 hydrolase
1.409	VMP1; DKFZP5661133	TMEM49	transmembrane protein 49
1.405	MGC33054	SNX10	sorting nexin 10
1.376	STX3A	STX3A	syntaxin 3
1.369	TTG2; RBTN2; RHOM2; RBTNL1	LMO2	LIM domain only 2 (rhombotin-like 1)
1.368	DBI; IBP; MBR; PBR; BZRP; PKBS; PTBR; mDRC; pk18	BZRP	translocator protein (18kDa)
1.361	CRE-BPA	CREB5	cAMP responsive element binding protein 5
1.344	MAY1; MGC49908; nPKC-delta	PRKCD	protein kinase C, delta
1.341	AAA; AD1; PN2; ABPP; APPI; CVAP; ABETA; CTFgamma	APP	amyloid beta (A4) precursor protein (peptidase nexin-II, Alzheimer disease)
1.333	CRFB4; CRF2-4; D21S58; D21S66; CDW210B; IL-10R2	IL10RB	interleukin 10 receptor, beta
1.31	DCIR; LLIR; DDB27; CLECSF6; HDCGC13P	CLEC4A	C-type lectin domain family 4, member A
1.304	HUFI-2; FLJ20248; FLJ22683; DKFZp434H2035	LRRFIP2	leucine rich repeat (in FLII) interacting protein 2
1.301	C32; CKLF1; CKLF2; CKLF3; CKLF4; UCK-1; HSPC224	CKLF	chemokine-like factor
1.289		ACSS2	
1.265	ESP-2; HED-2	ZYX	zyxin
1.263	SH3BGR; MGC117402	SH3BGRL	SH3 domain binding glutamic acid-rich protein like
1.239	MTX; MTXN	MTX1	metaxin 1
1.237	ASC; TMS1; CARD5;	PYCARD	PYD and CARD domain containing

Relative normalised expression	Common Name	Gene Symbol	Description
	MGC10332		
1.233	a3; Stv1; Vph1; Atp6i; OC116; OPTB1; TIRC7; ATP6N1C; ATP6V0A3; OC-116kDa	TCIRG1	T-cell, immune regulator 1, ATPase, H+ transporting, lysosomal V0 subunit A3
1.223	JTK8; FLJ26625	LYN	v-yes-1 Yamaguchi sarcoma viral related oncogene homolog
1.209	GAIP; RGS GAIP	RGS19	regulator of G-protein signalling 19
1.186	NEU; SIAL1	NEU1	sialidase 1 (lysosomal sialidase)

Table 7L M3.4 PTB v. Control, Genes Underrepresented in Active TB

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_DOWN_M3.4		
0.921	ZZZ4; FLJ10821; FLJ45574; KIAA0399	ZZEF1	zinc finger, ZZ-type with EF-hand domain 1
0.905	TILZ4a; TILZ4b; TILZ4c; KIAA0669	TSC22D2	TSC22 domain family, member 2
0.891	XTP2; BAT2-iso	BAT2D1	BAT2 domain containing 1
0.885	U2AF65	U2AF2	U2 small nuclear RNA auxiliary factor 2
0.878	DKFZp781I24156	PCNP	PEST proteolytic signal containing nuclear protein
0.876	NY-CO-1; FLJ10051	SDCCAG1	serologically defined colon cancer antigen 1
0.868	GCP16; HSPC041; MGC4876; MGC21096; GOLGA3AP1	GOLGA7	golgi autoantigen, golgin subfamily a, 7
0.866	CPR3; DJA2; DNAJ; DNJ3; RDJ2; HIRIP4; PRO3015	DNAJA2	DnaJ (Hsp40) homolog, subfamily A, member 2
0.863	B2-1; SEC7; D17S811E; FLJ34050; FLJ41900; CYTOHESIN-1	PSCD1	pleckstrin homology, Sec7 and coiled-coil domains 1(cytohesin 1)
0.855	SRrp86; SRrp508; MGC133045; DKFZp564B176	SFRS12	splicing factor, arginine/serine-rich 12
0.84	G3BP2	G3BP2	GTPase activating protein (SH3 domain) binding protein 2
0.831	p532; p619	HERC1	hect (homologous to the E6-AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1
0.826	DKFZP564O0523; HSPC304; DKFZp686D1651	DKFZP564O0523	hypothetical protein DKFZp564O0523
0.823	TSPYL	TSPYL1	TSPY-like 1
0.82	KIP1; MEN4; CDKN4; MEN1B; P27KIP1	CDKN1B	cyclin-dependent kinase inhibitor 1B (p27, Kip1)
0.82	SA2; SA-2; FLJ25871; bA517O1.1; DKFZp686P168; DKFZp781H1753	STAG2	stromal antigen 2
0.815	HR21; MCD1; NXP1; SCC1; hHR21; HRAD21; FLJ25655; FLJ40596; KIAA0078	RAD21	RAD21 homolog (S. pombe)
0.808	GCC185; KIAA0336	GCC2	GRIP and coiled-coil domain containing 2
0.806	PIR1	DUSP11	dual specificity phosphatase 11 (RNA/RNP complex 1-interacting)
0.804	AS3; CG008; PDS5B;	APRIN	androgen-induced proliferation inhibitor

Relative normalised expression	Common Name	Gene Symbol	Description
	FLJ23236; KIAA0979; RP1-267P19.1		
0.803		LOC58486	
0.798		SLTM	
0.795	AS; ANCR; E6-AP; HPVE6A; EPVE6AP; FLJ26981	UBE3A	ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome)
0.793	DKFZp686C1054	THUMPD1	THUMP domain containing 1
0.791	SIR2L1	SIRT1	sirtuin (silent mating type information regulation 2 homolog) 1 (<i>S. cerevisiae</i>)
0.79	FLJ40359	TPP2	tripeptidyl peptidase II
0.789	DKFZP564D172	C5orf21	chromosome 5 open reading frame 21
0.788	PALBH; CALPAIN7; FLJ36423	CAPN7	calpain 7
0.775	KIAA1116	RBM16	RNA binding motif protein 16
0.771	FLJ42355; KIAA0276	DCUN1D4	DCN1, defective in cullin neddylation 1, domain containing 4 (<i>S. cerevisiae</i>)
0.768	Rhe; FLJ33619; DKFZp586K0717	FIP1L1	FIP1 like 1 (<i>S. cerevisiae</i>)
0.766	RCP9; RCP; CRCP; CGRP-RCP; MGC111194	RCP9	calcitonin gene-related peptide-receptor component protein
0.764	DIF3; LZK1; DIF-3; LCRG1; ZFP403; FLJ21230; FLJ22561; FLJ42090	ZNF403	zinc finger protein 403
0.76	AD013; CReMM; KISH2; PRIC320	CHD9	chromodomain helicase DNA binding protein 9
0.757	VACM1; VACM-1	CUL5	cullin 5
0.755	MGC13407	NUP54	nucleoporin 54kDa
0.751	ENTH; EPN4; EPNR; CLINT; EPSINR; KIAA0171	ENTH	clathrin interactor 1
0.743	SEC24B	SEC24B	SEC24 related gene family, member B (<i>S. cerevisiae</i>); synonyms: SEC24, MGC48822; isoform a is encoded by transcript variant 1; secretory protein 24; Sec24-related protein B; protein transport protein Sec24B; Homo sapiens SEC24 related gene family, member B (<i>S. cerevisiae</i>) (SEC24B), transcript variant 1, mRNA.
0.742	HAKAI; RNF188; FLJ23109; MGC163401; MGC163403	CBLL1	Cas-Br-M (murine) ecotropic retroviral transforming sequence-like 1
0.738	XE7; 721P; XE7Y; CCDC133; CXYorf3; DXYS155E; MGC39904; MGC125365; MGC125366	DXYS155E	splicing factor, arginine/serine-rich 17A
0.737	NGB; CRFG; FLJ10686; FLJ10690; FLJ39774	GTPBP4	GTP binding protein 4
0.734	VELI3; LIN-7C; MALS-3; LIN-7-C; FLJ11215	LIN7C	lin-7 homolog C (<i>C. elegans</i>)
0.732	JTK5; RYK1; JTK5A; D3S3195	RYK	RYK receptor-like tyrosine kinase
0.731	K10; KPP; CK10	KRT10	keratin 10 (epidermolytic hyperkeratosis; keratosis palmaris et plantaris)
0.728	CYP-M; MGC22229	CYP20A1	cytochrome P450, family 20, subfamily A,

Relative normalised expression	Common Name	Gene Symbol	Description
			polypeptide 1
0.725	CHP1	CHORDC1	cysteine and histidine-rich domain (CHORD)-containing 1
0.724	NET1A; ARHGEF8	NET1	neuroepithelial cell transforming gene 1
0.723	ZF5; ZBTB14; ZNF478; MGC126126	ZFP161	zinc finger protein 161 homolog (mouse)
0.718	JAK1A; JAK1B	JAK1	Janus kinase 1 (a protein tyrosine kinase)
0.717	p5; p6; RRP45; PMSCL1; Rrp45p; PM/Sc1-75	EXOSC9	exosome component 9
0.716	GR; GCR; GRL; GCCR	NR3C1	nuclear receptor subfamily 3, group C, member 1 (glucocorticoid receptor)
0.713	L9mt	MRPL9	mitochondrial ribosomal protein L9
0.705	GRB1; p85-ALPHA	PIK3R1	phosphoinositide-3-kinase, regulatory subunit 1 (p85 alpha)
0.7	MST4; MASK	MASK	serine/threonine protein kinase MST4
0.7	UPF3; HUPF3A; RENT3A	UPF3A	UPF3 regulator of nonsense transcripts homolog A (yeast)
0.698	p17; YBL1; CHRAC17; CHARAC17	POLE3	polymerase (DNA directed), epsilon 3 (p17 subunit)
0.694	PCGF4; RNF51; MGC12685	PCGF4	BMI1 polycomb ring finger oncogene
0.692	MIF2; CENPC; hcp-4; CENP-C	CENPC1	centromere protein C 1
0.686	YAF9; GAS41; NUBI-1; 4930573H17Rik; B230215M10Rik	YEATS4	YEATS domain containing 4
0.679	R3HDM; FLJ23334; KIAA0029	R3HDM1	R3H domain containing 1
0.676	FBX21; FLJ90233; KIAA0875; MGC26682; DKFZp434G058	FBXO21	F-box protein 21
0.665	GRIFE; TULIP1; KIAA0884; DKFZp566D133; DKFZp667F074	GARNL1	GTPase activating Rap/RanGAP domain-like 1
0.663	BRL; BRPF1; BRPF2; DKFZp686F0325	BRD1	bromodomain containing 1
0.651	TIFIA; MGC104238; DKFZp566E104	RRN3	RRN3 RNA polymerase I transcription factor homolog (S. cerevisiae)
0.65	DKFZP586L0724	NOL11	nucleolar protein 11
0.645	FLJ20628; DKFZp564I2178	FLJ20628	hypothetical protein FLJ20628
0.642	FLJ21657; MGC90226; MGC149524	FLJ21657	chromosome 5 open reading frame 28
0.638	NS3TP1; FLJ20752; NBLA00058	ASNSD1	asparagine synthetase domain containing 1
0.636	MEX3C; BM-013; MEX-3C; RNF194; FLJ38871	RKHD2	ring finger and KH domain containing 2
0.628	E6BP; ERC55; ERC-55	RCN2	reticulocalbin 2, EF-hand calcium binding domain
0.613	PHLL1	CRY1	cryptochrome 1 (photolyase-like)
0.612	cdc14; hCDC14; Cdc14A1; Cdc14A2	CDC14A	CDC14 cell division cycle 14 homolog A (S. cerevisiae)
0.576	LCA; LY5; B220; CD45; T200; GP180	PTPRC	protein tyrosine phosphatase, receptor type, C
0.521	PBF; PRF1; HDBP2; PRF-1; Si-1-8-14; DKFZp434K1210	ZNF395	zinc finger protein 395

Table 7M M3.6 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_DOWN_M3.6		
0.898	ABHS; ORF20; TTDN1	C7orf11	chromosome 7 open reading frame 11
0.852	BTF2; TFIIH	GTF2H1	general transcription factor IIH, polypeptide 1, 62kDa
0.845	MGC51029	FUNDC1	FUN14 domain containing 1
0.844	SCOCO; HRIHFB2072	SCOC	short coiled-coil protein
0.839	IF-3mt; IF3(mt)	MTIF3	mitochondrial translational initiation factor 3
0.816	DAB1; MPRP-1; YKR087C; ZMPOMA1; FLJ33782; 2010001009Rik	OMA1	OMA1 homolog, zinc metallopeptidase (<i>S. cerevisiae</i>)
0.815		LOC644560	
0.795	JNKK; MEK4; MKK4; SEK1; JNKK1; SERK1; MAPKK4; PRKMK4	MAP2K4	mitogen-activated protein kinase kinase 4
0.775	REPA2; RPA32	RPA2	replication protein A2, 32kDa
0.765	AMMERC1	AMMECR1	Alport syndrome, mental retardation, midface hypoplasia and elliptocytosis chromosomal region, gene 1
0.741	CBX; M31; MOD1; HP1-BETA; HP1Hs-beta	CBX1	chromobox homolog 1 (HP1 beta homolog <i>Drosophila</i>)
0.739	DLTA; PDCE2; PDC-E2	DLAT	dihydrolipoamide S-acetyltransferase (E2 component of pyruvate dehydrogenase complex)
0.732	p38; AHA1; C14orf3	AHSA1	AHA1, activator of heat shock 90kDa protein ATPase homolog 1 (yeast)
0.731	VEZATIN; DKFZp761C241	VEZT	vezatin, adherens junctions transmembrane protein
0.728	HDPY-30	LOC84661	dpy-30-like protein
0.727	DERP6; MST071; HSPC002; MSTP071	C17orf81	chromosome 17 open reading frame 81
0.723	EFG; GFM; EFG1; EFGM; EGF1; hEFG1; COXPD1; FLJ12662; FLJ13632; FLJ20773	GFM1	G elongation factor, mitochondrial 1
0.721	MGC3232; hAtNOS1; mAtNOS1	C4orf14	chromosome 4 open reading frame 14
0.72	P15RS; FLJ10656; MGC19513	P15RS	hypothetical protein FLJ10656
0.719	MGC9912	C14orf126	chromosome 14 open reading frame 126
0.704	CCR4; KIAA1194	CNOT6	CCR4-NOT transcription complex, subunit 6
0.7	PRED31; HSPC230; FLJ34245; RP11-5919.1	C6orf203	chromosome 6 open reading frame 203
0.696	76P; GCP4	76P	gamma tubulin ring complex protein (76p gene)
0.694	FLJ10422	ELP3	elongation protein 3 homolog (<i>S. cerevisiae</i>)
0.677	MGC13379	MGC13379	HSPC244
0.677	CCTE; KIAA0098; CCT-epsilon; TCP-1-epsilon	CCT5	chaperonin containing TCP1, subunit 5 (epsilon)
0.675		MTMR12	

Relative normalised expression	Common Name	Gene Symbol	Description
0.671	ABRA1; FLJ11520; FLJ12642; FLJ13614	FLJ13614	coiled-coil domain containing 98
0.671	CDG1; CDGS; CDG1a	PMM2	phosphomannomutase 2
0.646	TPA1; FLJ10826; KIAA1612	OGFOD1	2-oxoglutarate and iron-dependent oxygenase domain containing 1
0.641	HV1; MGC15619	MGC15619	hydrogen voltage-gated channel 1
0.639	JJJ3; ZCSL3	ZCSL3	DPH4, JJJ3 homolog (<i>S. cerevisiae</i>)
0.631	GI008; RPMS13; MRP-S13; MRP-S26; NY-BR-87; C20orf193; dJ534B8.3	MRPS26	mitochondrial ribosomal protein S26
0.63	RPMS6; MRP-S6; C21orf101	MRPS6	mitochondrial ribosomal protein S6
0.622	CGI-55; CHD3IP; HABP4L; PAIRBP1; FLJ90489; PAI-RBP1; DKFZp564M2423	SERBP1	SERPINE1 mRNA binding protein 1
0.621	MRP-S14; HSMRPS14; DJ262D12.2	MRPS14	mitochondrial ribosomal protein S14
0.542	LOC153364; MGC46734; DKFZp686P15118	LOC153364	similar to metallo-beta-lactamase superfamily protein

Table 7N M3.7 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_DOWN_M3.7		
0.914	RED; CSA2; MGC59741; IK protein	IK	IK cytokine, down-regulator of HLA II
0.875	IBP	DEF6	differentially expressed in FDCP 6 homolog (mouse)
0.861	NAT3; dJ1002M8.1	NAT5	N-acetyltransferase 5
0.857	OFOX D; OFOX D1; FLJ20308	ALKBH5	alkB, alkylation repair homolog 5 (<i>E. coli</i>)
0.848	H-IDHB; MGC903; FLJ11043	IDH3B	isocitrate dehydrogenase 3 (NAD ⁺) beta
0.846	PGR1; PAM14	MRFAP1	Mof4 family associated protein 1
0.845	B17.2; DAP13	NDUFA12	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 12
0.836	MGC11134	TRPT1	tRNA phosphotransferase 1
0.832	H-l(3)mbt-1	L3MBTL2	l(3)mbt-like 2 (<i>Drosophila</i>)
0.831	HSCARG; FLJ25918	HSCARG	NmrA-like family domain containing 1
0.817	ABC27; ABC50	ABCF1	ATP-binding cassette, sub-family F (GCN20), member 1
0.816	LOC124512	LOC124512	hypothetical protein LOC124512
0.815	HSPC203	C14orf112	chromosome 14 open reading frame 112
0.814	EXOSC1	EXOSC1	exosome component 1; synonyms: p13, CSL4, SKI4, Csl4p, Ski4p, hCsl4p, CGI-108, RP11-452K12.9; homolog of yeast exosomal core protein CSL4; 3'-5' exoribonuclease CSL4 homolog; CSL4 exosomal core protein homolog; Homo sapiens exosome component 1 (EXOSC1), mRNA.
0.81	p14; DOC-1R; FLJ10636	CDK2AP2	CDK2-associated protein 2
0.81	MGC14833; baA6B20.2	C6orf125	chromosome 6 open reading frame 125

Relative normalised expression	Common Name	Gene Symbol	Description
0.809	SRP68	SRP68	signal recognition particle 68kDa
0.805	MGC3320; FLJ14936; RP5-965L7.1	PRPF38A	PRP38 pre-mRNA processing factor 38 (yeast) domain containing A
0.805	DBP-RB; UKVH5d	DDX1	DEAD (Asp-Glu-Ala-Asp) box polypeptide 1
0.804	ACRP; FSA-1; MGC20134	SPAG7	sperm associated antigen 7
0.802	MDHA; MOR2; MDH-s; MGC:1375	MDH1	malate dehydrogenase 1, NAD (soluble)
0.801	MDS016; RPMS21; MRP-S21	MRPS21	mitochondrial ribosomal protein S21
0.8	AIBP; MGC119143; MGC119144; MGC119145	APOA1BP	apolipoprotein A-I binding protein
0.8	ERV29; FLJ22993; MGC102753	SURF4	surfeit 4
0.797	MGC874	CXorf26	chromosome X open reading frame 26
0.795	FLJ22789	C12orf26	chromosome 12 open reading frame 26
0.795	RC68; INT11; RC-68; INTS11; CPSF73L; FLJ13294; FLJ20542	CPSF3L	cleavage and polyadenylation specific factor 3-like
0.793	HSPC196	HSPC196	transmembrane protein 138
0.79	DS-1	ICT1	immature colon carcinoma transcript 1
0.789	SIAHBP1; FIR; PUF60; RoBPI; FLJ31379	SIAHBP1	fuse-binding protein-interacting repressor
0.788	bMRP36a; MGC17989; MGC48892	MRPL43	mitochondrial ribosomal protein L43
0.788	HIT-17	HINT2	histidine triad nucleotide binding protein 2
0.785	MGC2714; FLJ32431	DCUN1D5	DCN1, defective in cullin neddylation 1, domain containing 5 (S. cerevisiae)
0.784	WDC146; FLJ11294	WDR33	WD repeat domain 33
0.775	N27C7-4; MGC70831	C22orf16	chromosome 22 open reading frame 16
0.774		LOC653709	
0.772	CGI-138; HSPC329; MRP-S23	MRPS23	mitochondrial ribosomal protein S23
0.769	P54; NMT55; NRB54; P54NRB	NONO	non-POU domain containing, octamer-binding
0.764	NSE2; MMS21; C8orf36; FLJ32440	C8orf36	non-SMC element 2, MMS21 homolog (S. cerevisiae)
0.764	C8orf40	C8orf40	chromosome 8 open reading frame 40
0.763	FLJ31795	CCDC43	coiled-coil domain containing 43
0.755	NSE1	NSMCE1	non-SMC element 1 homolog (S. cerevisiae)
0.753	MY105; THY28; MDS012; HSPC144; THY28KD; MGC12187	THYN1	thymocyte nuclear protein 1
0.752	YSA1H; hYSAH1	NUDT5	nudix (nucleoside diphosphate linked moiety X)-type motif 5
0.751	TOK-1	BCCIP	BRCA2 and CDKN1A interacting protein
0.747	VARSL; VARS2L; MGC138259; MGC142165	VARSL	valyl-tRNA synthetase 2, mitochondrial (putative)
0.732	FLJ13657; RP11-337A23.1	C9orf82	chromosome 9 open reading frame 82
0.728	GLOD2	MCEE	methylmalonyl CoA epimerase
0.728	C40	C2orf29	chromosome 2 open reading frame 29
0.726	MGC12966	MGC12966	hypothetical protein LOC84792; Homo sapiens hypothetical protein LOC84792 (MGC12966), mRNA.

Relative normalised expression	Common Name	Gene Symbol	Description
0.722	FLJ14803	FLJ14803	hypothetical protein FLJ14803
0.717	HSPC335; MRP-S24	MRPS24	mitochondrial ribosomal protein S24
0.716	RALBP1	REPS1	RALBP1 associated Eps domain containing 1
0.712	CAF1; hCAF-1	CNOT7	CCR4-NOT transcription complex, subunit 7
0.711	A1U; UBIN; C1orf6	UBQLN4	ubiquilin 4
0.71	CGI-118; MGC13323	MRPL48	mitochondrial ribosomal protein L48
0.701	Gm83; HSPC064; MGC126859; MGC138247; DKFZP564O0463	WDSOF1	WD repeats and SOF1 domain containing mitochondrial methionyl-tRNA formyltransferase
0.701	FMT1	MTFMT	formyltransferase
0.697	DKFZp686E10109	NUDCD2	NudC domain containing 2
0.697	MGC11321	MRPL45	mitochondrial ribosomal protein L45
0.691	SDOS; MGC11275	NUDT16L1	nudix (nucleoside diphosphate linked moiety X)-type motif 16-like 1
0.683	FLJ20989	C8orf33	chromosome 8 open reading frame 33
0.681	AK6; FIX; AK3L1; AKL3L; AKL3L1	AK3	adenylate kinase 3
0.671	RIP; HRIP; MGC4189	RIP	RPA interacting protein
0.666	PRP8; RP13; HPRP8; PRPC8	PRPF8	PRP8 pre-mRNA processing factor 8 homolog (<i>S. cerevisiae</i>)
0.664	PCMT; PPMT; PCCMT; HSTE14; MST098; MSTP098; MGC39955	ICMT	isoprenylcysteine carboxyl methyltransferase
0.66	YTM1; FLJ10881; FLJ12719; FLJ12720	WDR12	WD repeat domain 12
0.646	GAB1; CDC91L1; MGC40420	CDC91L1	phosphatidylinositol glycan anchor biosynthesis, class U
0.613	MGC4248	C10orf58	chromosome 10 open reading frame 58
0.613	sen15	C1orf19	chromosome 1 open reading frame 19
0.599	MGC2404	ACBD6	acyl-Coenz A binding domain containing 6

Table 70 M3.8 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTBvCSelect_09May08_PAL2Ttest_DOWN_M3.8		
0.841	MAP; RUSC3; SGSM3; DKFZp761D051	RUTBC3	RUN and TBC1 domain containing 3
0.84	FLJ13848	FLJ13848	N-acetyltransferase 11
0.827	HEL308; MGC20604	HEL308	DNA helicase HEL308
0.826	dgkd-2; DGKdelta; KIAA0145	DGKD	diacylglycerol kinase, delta 130kDa
0.814	DKFZp779L2418	SFRS14	splicing factor, arginine/serine-rich 14
0.814	HMMH; MUTM; OGH1; HOGG1	OGG1	8-oxoguanine DNA glycosylase
0.808	PRO9856; LAVS3040; DKFZp434D0711; DKFZp686L0539	BRD9	bromodomain containing 9
0.807	HCDI	C14orf124	chromosome 14 open reading frame 124

Relative normalised expression	Common Name	Gene Symbol	Description
0.798	GTF2D; SCA17; TFIIID; GTF2D1; MGC117320; MGC126054; MGC126055	TBP	TATA box binding protein
0.772	ZIS; ZIS1; ZIS2; ZNF265; FLJ41119; DKFZp686J1831; DKFZp686N09117	ZNF265	zinc finger, RAN-binding domain containing 2
0.764		OGT	
0.762	MTMR8; C8orf9; LIP-STYX; MGC126672; DKFZp434K171	MTMR9	myotubularin related protein 9
0.76	TDP-43	TARDBP	TAR DNA binding protein
0.754	FPM315; ZKSCAN12	ZNF263	zinc finger protein 263
0.754	C42; CGI-05; HSPC167; C20orf34; CDK5RAP1.3; CDK5RAP1.4	CDK5RAP1	CDK5 regulatory subunit associated protein 1
0.747	P50; P85; PAK3; PIXB; COOL1; P50BP; P85SPR; BETA-PIX; KIAA0142; KIAA0412; P85COOL1; Nbla10314; DKFZp761K1021	ARHGEF7	Rho guanine nucleotide exchange factor (GEF) 7
0.745	NAC; CARD7; NALP1; SLEV1; DEFCAP; PP1044; VAMAS1; CLR17.1; KIAA0926; DEFCAP-L/S; DKFZp586O1822	NALP1	NLR family, pyrin domain containing 1
0.744	KIAA0388	EZH1	enhancer of zeste homolog 1 (Drosophila)
0.741	MGC19570; dJ34B21.3	C6orf130	chromosome 6 open reading frame 130
0.737	RP11-336K24.1	KIAA0907	KIAA0907
0.732	LAM; TSC; KIAA0243; MGC86987	TSC1	tuberous sclerosis 1
0.725	LRS; LEUS; LARS1; LEURS; PIG44; RNTLS; HSPC192; hr025Cl; FLJ10595; FLJ21788; KIAA1352	LARS	leucyl-tRNA synthetase
0.724	HZF1	ZNF266	zinc finger protein 266
0.72	FAC1; FALZ; NURF301	FALZ	bromodomain PHD finger transcription factor
0.72	FLJ12892; FLJ41065; DKFZp434L1050	CCDC14	coiled-coil domain containing 14
0.708	TIR8; MGC110992	SIGIRR	single immunoglobulin and toll-interleukin 1 receptor (TIR) domain
0.7	FLJ21007; RP11-459E2.1	TDRD3	tudor domain containing 3
0.691	CGI75; mtTFB; CGI-75	TFB1M	transcription factor B1, mitochondrial
0.689	FP977; FLJ12270; MGC11230	WDR59	WD repeat domain 59
0.684	TS11	ASNS	asparagine synthetase
0.677	MGC111199	NIT2	nitrilase family, member 2
0.675		ASB1	
0.663	MCAF2; FLJ12668	ATF7IP2	activating transcription factor 7 interacting protein 2
0.648	SIN; RPC5	POLR3E	polymerase (RNA) III (DNA directed) polypeptide E (80kD)
0.646	BMS1L; KIAA0187	BMS1L	BMS1 homolog, ribosome assembly protein (yeast)
0.636	CBX7	CBX7	chromobox homolog 7

Relative normalised expression	Common Name	Gene Symbol	Description
0.63	PAN2; hPAN2; FLJ39360; KIAA0710	USP52	ubiquitin specific peptidase 52
0.623	MSK1; RLPK; MSPK1; MGC1911	RPS6KA5	ribosomal protein S6 kinase, 90kDa, polypeptide 5
0.612	SYB1; VAMP-1; DKFZp686H12131	VAMP1	vesicle-associated membrane protein 1 (synaptobrevin 1)
0.601	ALC1; CHDL; FLJ22530	CHD1L	chromodomain helicase DNA binding protein 1-like
0.587	KIAA0355	KIAA0355	KIAA0355
0.557	KIAA1615	ZNF529	zinc finger protein 529
0.554	MGC2146	IL11RA	interleukin 11 receptor, alpha
0.552	RNF84; MGC:39780	TRAF5	TNF receptor-associated factor 5
0.551	FLJ11795; MGC126013; MGC126014	FLJ11795	ankyrin repeat domain 55
0.548	DKFZp686O1788	MTX3	metaxin 3
0.544	DABP	DBP	D site of albumin promoter (albumin D-box) binding protein
0.541	FISH; SH3MD1	SH3PXD2A	SH3 and PX domains 2A
0.524	CLAX; LLT1; OCIL	CLEC2D	C-type lectin domain family 2, member D
0.518	HPF1; FLJ11015; FLJ14876; FLJ90585; MGC33853	ZNF83	zinc finger protein 83
0.514	ZCW4; ZCWCC2; FLJ11565; dJ75H8.2	MORC4	MORC family CW-type zinc finger 4
0.512	RTS; TYMSAS; RTS beta; HSRTSBETA; RTS alpha	ENOSF1	enolase superfamily member 1
0.483	C7orf32; ATP6V0E2L	ATP6V0E2L	ATPase, H ⁺ transporting V0 subunit e2
0.458	PLC1; PLC-II; PLC148; PLCgamma1	PLCG1	phospholipase C, gamma 1
0.428	RLK; TKL; BTKL; PTK4; PSCTK5; MGC22473	TXK	TXK tyrosine kinase
0.367	T14; S152; Tp55; TNFRSF7; MGC20393	TNFRSF7	CD27 molecule

Table 7P M3.9 PTB v. Control, Genes Underrepresented in Active TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_PTbvCSelect_09May08_PAL2Ttest_DOWN_M3.9		
0.869	ABC43; PMP70; PXMP1	ABCD3	ATP-binding cassette, sub-family D (ALD), member 3
0.86	SPG8; MGC111053	KIAA0196	KIAA0196
0.859	PUMH; HSPUM; PUMH1; PUM1; KIAA0099	PUM1	pumilio homolog 1 (Drosophila)
0.856	ASF; SF2; SF2p33; SRp30a; MGC5228	SFRS1	splicing factor, arginine/serine-rich 1 (splicing factor 2, alternate splicing factor)
0.848	DKFZp779N2044	KIAA0528	KIAA0528
0.843	ALG6	ALG6	asparagine-linked glycosylation 6 homolog (S. cerevisiae, alpha-1,3-glucosyltransferase)
0.829	MGC111579; DKFZp781B11202	DARS	aspartyl-tRNA synthetase

Relative normalised expression	Common Name	Gene Symbol	Description
0.829	ADDL	ADD3	adducin 3 (gamma)
0.829	KOX18; ZNF36; PHZ-37; ZNF139; MGC138429; 9130423L19Rik	ZKSCAN1	zinc finger with KRAB and SCAN domains 1
0.826	RPD3; YAF1	HDAC2	histone deacetylase 2
0.825	FLJ21634; MGC71630	GALNT11	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 11 (GalNAc-T11)
0.816	POLZ; REV3	REV3L	REV3-like, catalytic subunit of DNA polymerase zeta (yeast)
0.812	Ki; PA28G; REG-GAMMA; PA28-gamma	PSME3	proteasome (prosome, macropain) activator subunit 3 (PA28 gamma; Ki)
0.811	BRM; SNF2; SWI2; hBRM; Sth1p; BAF190; SNF2L2; SNF2LA; hSNF2a; FLJ36757; MGC74511	SMARCA2	SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 2
0.807	ZNT5; ZTL1; ZNTL1; ZnT-5; MGC5499; FLJ12496; FLJ12756	SLC30A5	solute carrier family 30 (zinc transporter), member 5
0.802	RAB7L; DKFZp686P1051	RAB7L1	RAB7, member RAS oncogene family-like 1
0.796	ASCIZ; KIAA0431; DKFZp779K1455	ASCIZ	ATM/ATR-Substrate Chk2-Interacting Zn2+-finger protein
0.796	TAF2B; CIF150; TAFIII150	TAF2	TAF2 RNA polymerase II, TATA box binding protein (TBP)-associated factor, 150kDa
0.786	N4WBP5; MGC10924	NDFIP1	Nedd4 family interacting protein 1
0.782	PAP41; MGC117304; MGC126719; MGC126721	PRPSAP2	phosphoribosyl pyrophosphate synthetase-associated protein 2
0.779	FLJ22584	TTC13	tetratricopeptide repeat domain 13
0.775	CLCI; ICln; CLNS1B	CLNS1A	chloride channel, nucleotide-sensitive, 1A
0.772	LRRC5; FLJ10470; FLJ20403	LRRC8D	leucine rich repeat containing 8 family, member D
0.77	CCT6; Cctz; HTR3; TCPZ; TCP20; MoDP-2; TTCP20; CCT-zeta; MGC126214; MGC126215; CCT-zeta-1; TCP-1-zeta	CCT6A	chaperonin containing TCP1, subunit 6A (zeta 1)
0.765	TOK-1	BCCIP	BRCA2 and CDKN1A interacting protein
0.764	G3BP; HDH-VIII; MGC111040	G3BP	GTPase activating protein (SH3 domain) binding protein 1
0.763	FACT; CDC68; FACTP140; FLJ10857; FLJ14010; FLJ34357; SPT16/CDC68	SUPT16H	suppressor of Ty 16 homolog (S. cerevisiae)
0.757	FBP2; FLJ12799; FLJ38170	C14orf135	chromosome 14 open reading frame 135
0.753	GCP3; SPBC98; Spc98p	TUBGCP3	tubulin, gamma complex associated protein 3
0.752	FLJ13576; DKFZp564C012	FLJ13576	transmembrane protein 168
0.751	SRP72	SRP72	signal recognition particle 72kDa
0.75	CIA1; WDR39	WDR39	cytosolic iron-sulfur protein assembly 1 homolog (S. cerevisiae)
0.738	HPT; MRS2; MGC78523	MRS2L	MRS2-like, magnesium homeostasis factor

Relative normalised expression	Common Name	Gene Symbol	Description
			(S. cerevisiae)
0.729	CED-4; FLASH; RIP25; FLJ11208; KIAA1315	CASP8AP2	CASP8 associated protein 2
0.728	PTPLB	PTPLB	protein tyrosine phosphatase-like (proline instead of catalytic arginine), member b
0.724	CHAC; FLJ42030; KIAA0986	VPS13A	vacuolar protein sorting 13 homolog A (S. cerevisiae)
0.724	REC14	WDR61	WD repeat domain 61
0.719	EB9; PDAF; RCAS1	EBAG9	estrogen receptor binding site associated, antigen, 9
0.712	SNX4	SNX4	sorting nexin 4
0.704	TOPIIB; top2beta	TOP2B	topoisomerase (DNA) II beta 180kDa
0.704	CGI-12; FLJ10939	MTERFD1	MTERF domain containing 1
0.703	CBC2; NIP1; CBP20; PIG55	NCBP2	nuclear cap binding protein subunit 2, 20kDa
0.702	HAD; HHF4; HADH1; SCHAD; HADHSC; M/SCHAD; MGC8392	HADHSC	hydroxyacyl-Coenzyme A dehydrogenase
0.701	p56; HSD8; FLJ11088; DKFZP779L1558; DKFZp779L1558	DKFZP779L1558	coiled-coil domain containing 91
0.701	CREB; MGC9284	CREB1	cAMP responsive element binding protein 1
0.7	AIP5; Tiu11; hSDRP1; DKFZp434D2111	WWP1	WW domain containing E3 ubiquitin protein ligase 1
0.681	TAT-SF1; dJ196E23.2	HTATSF1	HIV-1 Tat specific factor 1
0.674	LDLC	COG2	component of oligomeric golgi complex 2
0.671	HC71; CGI-150; C17orf25	C17orf25	glyoxalase domain containing 4
0.67	GABAT; NPD009; GABA-AT	ABAT	4-aminobutyrate aminotransferase
0.668	AKAP18	AKAP7	A kinase (PRKA) anchor protein 7
0.661	LSFC; GP130; LRP130; CLONE-23970	LRPPRC	leucine-rich PPR-motif containing
0.644	SCC-112; PIG54; FLJ41012; KIAA0648; MGC131948; MGC161503; DKFZp686B19246	SCC-112	SCC-112 protein
0.643	GDE	AGL	amylo-1, 6-glucosidase, 4-alpha-glucanotransferase (glycogen debranching enzyme, glycogen storage disease type III)
0.643	NIP3	BNIP3	BCL2/adenovirus E1B 19kDa interacting protein 3
0.64	HSSB; RF-A; RP-A; REPA1; RPA70	RPA1	replication protein A1, 70kDa
0.63	TAF2C; TAF4A; TAF2C1; FLJ41943; TAFII130; TAFII135	TAF4	TAF4 RNA polymerase II, TATA box binding protein (TBP)-associated factor, 135kDa
0.626	TMP21; S31I125; Tmp-21-I; S31III125; P24(Delta)	TMED10	transmembrane emp24-like trafficking protein 10 (yeast)
0.617	FLJ20397; FLJ25564; FLJ31671; FLJ39381	FLJ20397	HEAT repeat containing 2
0.612	CHA; Fig1b; E2BP-1; MGC46135	TCFL5	transcription factor-like 5 (basic helix-loop-helix)
0.588	SRB; Cctd; MGC126164; MGC126165	CCT4	chaperonin containing TCP1, subunit 4 (delta)

Relative normalised expression	Common Name	Gene Symbol	Description
0.582	Seh1; SEH1A; SEH1B; SEC13L	SEH1L	SEH1-like (<i>S. cerevisiae</i>)
0.527	HSU79274	C12orf24	chromosome 12 open reading frame 24

Table 8A M1.5 LTB v. Control, Genes Underrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May 08_PAL2Ttest_DOWN_M1.5		
2.007	STF1; STFA	CSTA	cystatin A (stefin A)
1.915	LSH; NRAMP; NRAMP1	SLC11A1	solute carrier family 11 (proton-coupled divalent metal ion transporters), member 1
1.903	EZI; Zfp467	ZNF467	zinc finger protein 467
1.813	TIL4; CD282	TLR2	toll-like receptor 2
1.811	HSULF-2; FLJ90554; KIAA1247; MGC126411; DKFZp313E091	SULF2	sulfatase 2
1.716	FLJ22662	FLJ22662	hypothetical protein FLJ22662
1.691	FDF03	PILRA	paired immunoglobulin-like type 2 receptor alpha
1.686	HET; ITM; BWR1A; IMPT1; TSSC5; ORCTL2; BWSCR1A; SLC22A1L; p45-BWR1A; DKFZp667A184	SLC22A18	solute carrier family 22 (organic cation transporter), member 18
1.682	ILT1; LIR7; CD85H; LIR-7	LILRA2	leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 2
1.657	C1QR1; C1qRP; CDw93; MXRA4; C1qR(P); dJ737E23.1	C1QR1	CD93 molecule
1.636	NCF; MGC3810; P40PHOX; SH3PXD4	NCF4	neutrophil cytosolic factor 4, 40kDa
1.623	NOXA2; p67phox; P67-PHOX	NCF2	neutrophil cytosolic factor 2 (65kDa, chronic granulomatous disease, autosomal 2)
1.542	FLJ10357; SOLO	FLJ10357	hypothetical protein FLJ10357
1.525	JTK9	HCK	hemopoietic cell kinase
1.521	FEM-2; POPX2; hFEM-2; CaMKPase; KIAA0015	PPM1F	protein phosphatase 1F (PP2C domain containing)
1.498	CD32; FCG2; FcGR; CD32A; CDw32; FCGR2; IGFR2; FCGR2A1; MGC23887; MGC30032	FCGR2A	Fc fragment of IgG, low affinity IIa, receptor (CD32)
1.493	DHRS8; PAN1B; RETSDR2; 17-BETA-HSD11; 17-BETA- HSDXI	DHRS8	hydroxysteroid (17-beta) dehydrogenase 11
1.482	FLJ11151; CSTP1	FLJ11151	hypothetical protein FLJ11151
1.478	CD31; PECAM-1	PECAM1	platelet/endothelial cell adhesion molecule (CD31 antigen)
1.469	DORA	IGSF6	immunoglobulin superfamily, member 6
1.452	GP; G1RZFP; GOLIATH; MGC99542; MGC117241; MGC138647	RNF130	ring finger protein 130

Relative normalised expression	Common Name	Gene Symbol	Description
1.45	MLN70; S100C	S100A11	S100 calcium binding protein A11
1.449	MGC3886	CTSS	cathepsin S
1.425	APPH; APPL2; CDEBP	APLP2	amyloid beta (A4) precursor-like protein 2
1.41	IMPD; RP10; IMPD1; LCA11; sWSS2608; DKFZp781N0678	IMPDH1	IMP (inosine monophosphate) dehydrogenase 1
1.406	FCNM	FCN1	ficolin (collagen/fibrinogen domain containing) 1
1.376	MYD88	MYD88	myeloid differentiation primary response gene (88)
1.371	B144; LST-1; D6S49E; MGC119006; MGC119007	LST1	leukocyte specific transcript 1
1.348	OS9	OS9	amplified in osteosarcoma
1.334	TEM7R; FLJ14623	PLXDC2	plexin domain containing 2
1.334	Rab22B	RAB31	RAB31, member RAS oncogene family
1.301	TS; TXS; CYP5; THAS; TXAS; CYP5A1	TBXAS1	thromboxane A synthase 1 (platelet, cytochrome P450, family 5, subfamily A)
1.292	HXK3; HKIII	HK3	hexokinase 3 (white cell)
1.292	RISC; HSCP1	SCPEP1	serine carboxypeptidase 1
1.283	IBA1; AIF-1; IRT-1	AIF1	allograft inflammatory factor 1
1.283	CD14	CD14	CD14 molecule
1.27	PI; A1A; AAT; PI1; A1AT; MGC9222; PRO2275; MGC23330	SERPINA1	serpin peptidase inhibitor, clade A (alpha-1 antiprotease, antitrypsin), member 1
1.261	LIR6; CD85I; LIR-6; MGC126563	LILRA1	leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1
1.221	CAP102; FLJ36832	CTNNA1	catenin (cadherin-associated protein), alpha 1, 102kDa
1.192	BCKDK	BCKDK	branched chain ketoacid dehydrogenase kinase
1.137	p75; TBPII; TNFBR; TNFR2; CD120b; TNFR80; TNF-R75; p75TNFR; TNF-R-II	TNFRSF1B	tumor necrosis factor receptor superfamily, member 1B

Table 8B M2.1 LTB v. Control, Genes Overrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May08_PAL2Ttest_UP_M2.01		
0.801	LIME; LP8067; FLJ20406; dJ583P15.4; RP4-583P15.5	LIME1	Lck interacting transmembrane adaptor 1
0.769	FLJ34563; MGC35163	SAMD3	sterile alpha motif domain containing 3
0.763	SISd; SCYA5; RANTES; TCP228; D17S136E; MGC17164	CCL5	chemokine (C-C motif) ligand 5
0.758	ORP7; MGC71150	OSBPL7	oxysterol binding protein-like 7
0.757		LOC387882	
0.736	SLP2; SGA72M; CHR11SYT; KIAA1597; MGC102768	SYTL2	synaptotagmin-like 2
0.735	DORZ1; DKFZP564O243	ABHD14A	abhydrolase domain containing 14A
0.727	MGC33870; MGC74858	NCALD	neurocalcin delta

0.691	LPAP; CD45-AP; MGC138602; MGC138603	PTPRCAP	protein tyrosine phosphatase, receptor type, C-associated protein
0.686	T11; SRBC	CD2	CD2 molecule
0.671	CD8; MAL; p32; Leu2	CD8A	CD8a molecule
0.656	HOP; OB1; LAGY; Toto; Cameo; NECC1; SMAP31; MGC20820	HOP	homeodomain-only protein
0.651	2F1; MAFA; MAFA-L; CLEC15A; MAFA-2F1; MGC13600	KLRG1	killer cell lectin-like receptor subfamily G, member 1
0.65		LOC197135	
0.643	GIG1	NKG7	natural killer cell group 7 sequence
0.638	TSAd; F2771	SH2D2A	SH2 domain protein 2A
0.634	FEOM; CFEOM; FEOM1; CFEOM1; FLJ20052; KIAA1708; DKFZp779C159	KIF21A	kinesin family member 21A
0.627	KIAA0442; MGC13140	AUTS2	autism susceptibility candidate 2
0.583	BFPP; TM7LN4; TM7XN1; DKFZp781L1398	GPR56	G protein-coupled receptor 56
0.572	TARP; CD3G; TCRG; TCRGC1; TCRGC2	TARP	TCR gamma alternate reading frame protein
0.502	519; LAG2; NKG5; LAG-2; D2S69E; TLA519	GNLY	granulysin
0.303	CCP-X; CGL-2; CSP-C; CTLA1; CTSGL2	GZMH	granzyme H (cathepsin G-like 2, protein h-CCPX)

Table 8C M2.6 LTB v. Control, Genes Underrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May 08_PAL2Ttest_DOWN_M2.06		
	Module 2.06, myeloid, fold change is healthy relative to LTB, ie DOWN in LTB		
2.409	HsT287	ZNF516	zinc finger protein 516
2.286	CRISP11; LCRISP2; MGC74865; DKFZP434B044	CRISPLD2	cysteine-rich secretory protein LCCL domain containing 2
2.177	MAG1; GPAT3; AGPAT8; MGC11324	HMFN0839	lung cancer metastasis-associated protein
2.095	CDD	CDA	cytidine deaminase
2.094	CRBP4; CRBP4V; MGC70641	RBP7	retinol binding protein 7, cellular
1.917	SSC1; HsT17287	AQP9	aquaporin 9
1.916	GMR; CD116; CSF2R; CDw116; CSF2RX; CSF2RY; GMCSFR; CSF2RAX; CSF2RAY; MGC3848; MGC4838; GM-CSF-R-alpha	CSF2RA	colony stimulating factor 2 receptor, alpha, low-affinity (granulocyte-macrophage)
1.853	G0S8	RGS2	regulator of G-protein signalling 2, 24kDa
1.734	HKII; HXK2; DKFZp686M1669	HK2	hexokinase 2
1.734	BB1	LENG4	leukocyte receptor cluster (LRC) member 4
1.701	UB1; CEP3; BORG2; FLJ46903	CDC42EP3	CDC42 effector protein (Rho GTPase binding) 3
1.671	SPAL2; FLJ23126; FLJ23632;	SIPA1L2	signal-induced proliferation-associated 1

	KIAA1389		like 2
1.669	ST1; SYCL; MDA-9; TACIP18	SDCBP	syndecan binding protein (syntenin)
1.669	CAN; CAIN; N214; D9S46E; MGC104525	NUP214	nucleoporin 214kDa
1.651		SLC19A1	
1.65	LPB3; S1P3; EDG-3; S1PR3; FLJ37523; MGC71696	EDG3	endothelial differentiation, sphingolipid G-protein-coupled receptor, 3
1.642	FPR; FMLP	FPR1	formyl peptide receptor 1
1.61	GPCR1; GPR86; GPR94; P2Y13; SP174; FKSG77	P2RY13	purinergic receptor P2Y, G-protein coupled, 13
1.606	WDR80; FLJ00012	ATG16L2	ATG16 autophagy related 16-like 2 (S. cerevisiae)
1.601	LENG5; SEN34; SEN34L	TSEN34	tRNA splicing endonuclease 34 homolog (S. cerevisiae)
1.575	FPF; p55; p60; TBP1; TNF-R; TNFAR; TNFR1; p55-R; CD120a; TNFR55; TNFR60; TNF-R-I; TNF-R55; MGC19588	TNFRSF1A	tumor necrosis factor receptor superfamily, member 1A
1.572	PEL12	PEL12	pellino homolog 2 (Drosophila)
1.562	FLJ13052; FLJ37724; dJ283E3.1; RP1-283E3.6	NADK	NAD kinase
1.558	5-LO; 5LPG; LOG5; MGC163204	ALOX5	arachidonate 5-lipoxygenase
1.534	TMPIT	TMPIT	transmembrane protein induced by tumor necrosis factor alpha
1.517	FLJ31978	GLT1D1	glycosyltransferase 1 domain containing 1
1.517	PFKFB4	PFKFB4	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 4
1.516	FLJ22470; KIAA1993; MGC24652; RP11-106H5.1	ZBTB34	zinc finger and BTB domain containing 34
1.482	P39; VATX; VMA6; ATP6D; ATP6DV; VPATPD	ATP6V0D1	ATPase, H ⁺ transporting, lysosomal 38kDa, V0 subunit d1
1.473	PRAM-1; MGC39864	PRAM1	PML-RARA regulated adaptor molecule 1
1.471	BIT; MFR; P84; SIRP; MYD-1; SHPS1; CD172A; PTPNS1; SHPS-1; SIRPalpha; SIRPalpha2; SIRP-ALPHA-1	PTPNS1	signal-regulatory protein alpha
1.463	M130; MM130	CD163	CD163 molecule
1.434	AF-1; IFGR2; IFNGT1	IFNGR2	interferon gamma receptor 2 (interferon gamma transducer 1)
1.405	RALB	RALB	v-ral simian leukemia viral oncogene homolog B (ras related; GTP binding protein)
1.405	SLCO3A1	SLCO3A1	solute carrier organic anion transporter family, member 3A1; synonyms: OATP-D, OATP3A1, FLJ40478, SLC21A11; solute carrier family 21 (organic anion transporter), member 11; Homo sapiens solute carrier organic anion transporter family, member 3A1 (SLCO3A1), mRNA.
1.397	PTPE; HPTPE; DKFZp313F1310; R-PTP- EPSILON	PTPRE	protein tyrosine phosphatase, receptor type, E
1.397	RCC4; FLJ14784	DIRC2	disrupted in renal carcinoma 2
1.396	DAPI2; KARAP; PLOSL	TYROBP	TYRO protein tyrosine kinase binding

			protein
1.371	B144; LST-1; D6S49E; MGC119006; MGC119007	LST1	leukocyte specific transcript 1
1.359	BFD; PFC; PFD; PROPERDIN	PFC	complement factor properdin
1.31	CAG4A; ERDA5; PRAT4A	TNRC5	trinucleotide repeat containing 5
1.307	CD18; TNFCR; D12S370; TNFR-RP; TNFRSF3; TNFR2- RP; LT-BETA-R; TNF-R-III	LTBR	lymphotoxin beta receptor (TNFR superfamily, member 3)
1.305	CEB	VAMP3	vesicle-associated membrane protein 3 (cellubrevin)
1.304	CSC-21K	TIMP2	TIMP metalloproteinase inhibitor 2
1.301	BPOZ; EF1ABP; PP2259; MGC20585	ABTB1	ankyrin repeat and BTB (POZ) domain containing 1
1.294	C6orf209; FLJ11240; bA810I22.1; RP11-810I22.1	LMBRD1	LMBR1 domain containing 1
1.266	PBF; C21orf1; C21orf3	PTTG1IP	pituitary tumor-transforming 1 interacting protein
1.235	ZFYVE10; FLJ32333; KIAA0371; FYVE-DSP1	MTMR3	myotubularin related protein 3
1.216	CFP1; CBCP1; C10orf9	C10orf9	cyclin Y
1.2	SPT4H; SUPT4H	SUPT4H1	suppressor of Ty 4 homolog 1 (S. cerevisiae)

Table 8D M2.10 LTB v. Control, Genes Underrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May 08_PAL2Ttest_DOWN_M2.10		
	Undefined module M2.10, fold change healthy relative to LTB, ie DOWN in LTB		
1.608	JAML; AMICA; Gm638; CREA7-1; CREA7-4; FLJ37080; MGC118814; MGC118815	AMICA1	adhesion molecule, interacts with CXADR antigen 1
1.537	MPEG1; MGC132657; MGC138435	MPEG1	macrophage expressed gene 1
1.514	L13; MGC13061	RNF135	ring finger protein 135
1.507	PAKalpha; MGC130000; MGC130001	PAK1	p21/Cdc42/Rac1-activated kinase 1 (STE20 homolog, yeast)
1.471	T49; pT49	FGL2	fibrinogen-like 2
1.405	KIAA0513	KIAA0513	KIAA0513
1.396	NCKX4; SLC24A2; FLJ38852	SLC24A4	solute carrier family 24 (sodium/potassium/calcium exchanger), member 4
1.358	FLJ34389	MLKL	mixed lineage kinase domain-like
1.348	ETO2; MTG16; MTGR2; ZMYND4	CBFA2T3	core-binding factor, runt domain, alpha subunit 2; translocated to, 3
1.331	IRC1; IRC2; IRp60; IGSF12; CMRF35H; CMRF-35H; CMRF35H9; CMRF-35-H9	CD300A	CD300a molecule
1.3	GLIPR; RTVP1; CRISP7	GLIPR1	GLI pathogenesis-related 1 (glioma)
1.229	ENC-1AS	HEXB	hexosaminidase B (beta polypeptide)
1.222	TIRP; TRAM; TIRAP3;	TICAM2	toll-like receptor adaptor molecule 2

Relative normalised expression	Common Name	Gene Symbol	Description
	TICAM-2; MGC129876; MGC129877		
1.175	FLJ31265	NUDT16	nudix (nucleoside diphosphate linked moiety X)-type motif 16
1.17	FKBP133; KIAA0674	KIAA0674	FK506 binding protein 15, 133kDa

Table 8E M3.2 LTB v. Control, Genes Underrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May 08_PAL2Ttest_DOWN_M3.2		
	Inflammation 3.2 fold change is healthy relative to LTB, ie DOWN in LTB		
4.289	K60; NAF; GCP1; LECT; LUCT; NAP1; 3-10C; CXCL8; GCP-1; LYNAIP; MDNCF; MONAP; NAP-1; SCYB8; TSG-1; AMCF-I; b-ENAP	IL8	interleukin 8
2.068	CD87; UPAR; URKR	PLAUR	plasminogen activator, urokinase receptor
2.009	PBEF; NAMPT; MGC117256; DKFZP666B131; 1110035O14Rik	PBEF1	pre-B-cell colony enhancing factor 1
1.9		IER3	
1.87	TREM-1	TREM1	triggering receptor expressed on myeloid cells 1
1.79	E4BP4; IL3BP1; NFIL3A; NF- IL3A	NFIL3	nuclear factor, interleukin 3 regulated
1.739	KIAA1145	TMCC3	transmembrane and coiled-coil domain family 3
1.728	PINH; FLJ21759; FLJ23500; C20orf110; dJ1181N3.1; DKFZp434B2411; DKFZp434O0827	TP53INP2	tumor protein p53 inducible nuclear protein 2
1.705	MAD; MAD1; MGC104659	MXD1	MAX dimerization protein 1
1.657	SGK1	SGK	serum/glucocorticoid regulated kinase
1.654	SLCO3A1	SLCO3A1	solute carrier organic anion transporter family, member 3A1; synonyms: OATP-D, OATP3A1, FLJ40478, SLC21A11; solute carrier family 21 (organic anion transporter), member 11; Homo sapiens solute carrier organic anion transporter family, member 3A1 (SLCO3A1), mRNA.
1.637	C5orf6	FAM53C	family with sequence similarity 53, member C
1.632	PDLIM7	PDLIM7	PDZ and LIM domain 7 (enigma)
1.591	NIN1; NINJURIN	NINJ1	ninjurin 1
1.572	RIT; RIBB; ROC1; MGC125864; MGC125865	RIT1	Ras-like without CAAX 1
1.567	SB135	MYADM	myeloid-associated differentiation marker
1.54	RCP; NOEL1A; FLJ22524; FLJ22622; MGC78448; rab11-	RAB11FIP1	RAB11 family interacting protein 1 (class I)

Relative normalised expression	Common Name	Gene Symbol	Description
	FIP1; DKFZp686E2214		
1.526	DANGER; bA127L20; bA127L20.2; RP11-127L20.4	KIAA1754	KIAA1754
1.515		SPAG9	
1.499	HSS; JLP; HLC4; PHET; PIG6; FLJ13450; FLJ14006; FLJ26141; FLJ34602; KIAA0516; MGC14967; MGC74461; MGC117291	SPAG9	sperm associated antigen 9
1.496	MGC20461	OSM	oncostatin M
1.444	KIAA1673	CPEB4	cytoplasmic polyadenylation element binding protein 4
1.433	IL-1; IL1F2; IL1-BETA	IL1B	interleukin 1, beta
1.413	TRIP8; FLJ14374; KIAA1380; RP11-10C13.2; DKFZp761F0118	JMJD1C	jumonji domain containing 1C
1.41	FLJ11080; FLJ33961; DKFZP566A1524	FAM49A	family with sequence similarity 49, member A
1.4	EOPA; NUDEL; MITAP1; DKFZp451M0318	NDEL1	nudE nuclear distribution gene E homolog (A. nidulans)-like 1
1.384	NHE8; FLJ42500; KIAA0939; MGC138418; DKFZp686C03237	SLC9A8	solute carrier family 9 (sodium/hydrogen exchanger), member 8
1.379	FLJ14744	PPP1R15B	protein phosphatase 1, regulatory (inhibitor) subunit 15B
1.356	PPG; PRG; PRG1; MGC9289; FLJ12930	PRG1	serglycin
1.348	ATG8; GEC1; APG8L	GABARAPL1	GABA(A) receptor-associated protein like 1
1.332	TTP; GOS24; GOS24; TIS11; NUP475; RNF162A	ZFP36	zinc finger protein 36, C3H type, homolog (mouse)
1.329	PFK2; IPFK2	PFKFB3	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 3
1.31	DKFZp547M072	MIDN	midnolin
1.301	FLJ13448	COQ10B	coenzyme Q10 homolog B (S. cerevisiae)
1.285	C8FW; GIG2; SKIP1	TRIB1	tribbles homolog 1 (Drosophila)
1.284	FLJ13725; KIAA1930	FAM65A	family with sequence similarity 65, member A
1.272	FLJ46337; MGC117209; DKFZP434H132	C15orf39	chromosome 15 open reading frame 39
1.258	AII; AVP; FCU; MWS; FCAS; CIAS1; NALP3; C1orf7; CLR1.1; PYPAF1; AII/AVP; AGTAVPRL	CIAS1	NLR family, pyrin domain containing 3
1.252	BRF1; ERF1; cMG1; ERF-1; Berg36; TIS11B; RNF162B	ZFP36L1	zinc finger protein 36, C3H type-like 1
1.249	FRA2; FLJ23306	FOSL2	FOS-like antigen 2
1.235	GADD34	PPP1R15A	protein phosphatase 1, regulatory (inhibitor) subunit 15A
1.235	p33; p47; p33ING1; p24ING1c; p33ING1b; p47ING1a	ING1	inhibitor of growth family, member 1
1.231	P47; FLJ27168	PLEK	pleckstrin
1.218	UBP; SIH003; MGC129878; MGC129879	USP3	ubiquitin specific peptidase 3

Relative normalised expression	Common Name	Gene Symbol	Description
1.208	Sei-2; TRIP-Br2; MGC126688; MGC126690	SERTAD2	SERTA domain containing 2
1.204	DCTN4	DCTN4	dynactin 4 (p62)
1.192	ROX; MAD6; MXD6	MNT	MAX binding protein
1.165	RBT1	SERTAD3	SERTA domain containing 3
1.157	WIPI3; WIPI-3	WDR45L	WDR45-like
1.156	ERF; RF1; ERF1; TB3-1; D5S1995; SUP45L1; MGC111066	ETF1	eukaryotic translation termination factor 1
1.156	KIAA0118	RAB21	RAB21, member RAS oncogene family
1.098	MAPKAPK2	MAPKAPK2	mitogen-activated protein kinase-activated protein kinase 2

Table 8F M3.3 LTB v. Control, Genes Underrepresented in Latent TB.

Relative normalised expression	Common Name	Gene Symbol	Description
	P22_15_LTBvCSelect_09May08_PAL2Ttest_DOWN_M3.3		
	Inflammation 3.2 fold change is healthy relative to LTB, ie DOWN in LTB		
2.716	QC; GCT	QPCT	glutaminy-peptide cyclotransferase (glutaminy cyclase)
2.579	CRE-BPA	CREB5	cAMP responsive element binding protein 5
2.468	APN; CD13; LAP1; PEPN; gp150	ANPEP	alanyl (membrane) aminopeptidase (aminopeptidase N, aminopeptidase M, microsomal aminopeptidase, CD13, p150)
2.426	PAD; PDI4; PDI5; PADI5	PADI4	peptidyl arginine deiminase, type IV
2.245	MRP; WLS; C1orf139; FLJ23091; MGC14878; MGC131760	GPR177	G protein-coupled receptor 177
2	HIS; HSTD; histidase	HAL	histidine ammonia-lyase
1.963	PYGL	PYGL	phosphorylase, glycogen; liver (Hers disease, glycogen storage disease type VI)
1.948		EGFL5	
1.935	L-H2; ASGP-R; CLEC4H2; Hs.1259	ASGR2	asialoglycoprotein receptor 2
1.892	CD114; GCSFR	CSF3R	colony stimulating factor 3 receptor (granulocyte)
1.882	LAMPB; CD107b; LAMP-2C	LAMP2	lysosomal-associated membrane protein 2
1.813	ALFY; ZFYVE25; KIAA0993; MGC16461	WDFY3	WD repeat and FYVE domain containing 3
1.8	STX3A	STX3A	syntaxin 3
1.771	CR1	CR1	complement component (3b/4b) receptor 1 (Knops blood group); synonyms: KN, C3BR, CD35; isoform F precursor is encoded by transcript variant F; C3-binding protein; CD35 antigen; complement component receptor 1; C3b/C4b receptor; Knops blood group antigen; Homo sapiens complement component (3b/4b) receptor 1

Relative normalised expression	Common Name	Gene Symbol	Description
			(Knops blood group) (CR1), transcript variant F, mRNA.
1.764	DCL-1; BIMLEC; CLEC13A; KIAA0022	CD302	CD302 molecule
1.758	FER1L1; LGMD2B; FLJ00175; FLJ90168	DYSF	dysferlin, limb girdle muscular dystrophy 2B (autosomal recessive)
1.733	TM6SF1	TM6SF1	transmembrane 6 superfamily member 1
1.721	MYO1F	MYO1F	myosin IF
1.691	CPR8; KIAA1254	CCPG1	cell cycle progression 1
1.688	LAB; NTAL; WSCR5; WBSR5; HSPC046; WBSR15	LAT2	linker for activation of T cells family, member 2
1.687	CNAIP; FLJ40652; bK126B4.4	NFAM1	NFAT activating protein with ITAM motif 1
1.659	FVL; PCCF; factor V	F5	coagulation factor V (proaccelerin, labile factor)
1.655	FLJ20273; DKFZp686F02235	FLJ20273	RNA-binding protein
1.647	NR4; CD213A1; IL-13Ra	IL13RA1	interleukin 13 receptor, alpha 1
1.636	NCF; MGC3810; P40PHOX; SH3PXD4	NCF4	neutrophil cytosolic factor 4, 40kDa
1.635	p63; CLIMP-63; ERGIC-63; MGC99554	CKAP4	cytoskeleton-associated protein 4
1.611	SELR; SELX; MSRB1; HSPC270; MGC3344	SEPX1	selenoprotein X, 1
1.6	MD-2	LY96	lymphocyte antigen 96
1.599	NPL1; c112; C1orf13; MGC61869; MGC149582	NPL	N-acetylneuraminic pyruvate lyase (dihydrodipicolinate synthase)
1.59	HAP; ASYIP; NSPL2; NSPLII; RTN3-A1	RTN3	reticulon 3
1.581	VMP1; DKFZP566I133	TMEM49	transmembrane protein 49
1.567	HBP; HEBP	HEBP1	heme binding protein 1
1.562	LAMPB; CD107b; LAMP-2C	LAMP2	lysosomal-associated membrane protein 2
1.559	C32; CKLF1; CKLF2; CKLF3; CKLF4; UCK-1; HSPC224	CKLF	chemokine-like factor
1.538		RASSF2	
1.532	SemE; SEMAE	SEMA3C	sema domain, immunoglobulin domain (Ig), short basic domain, secreted, (semaphorin) 3C
1.53	ARAP3; DRAG1; FLJ21065	CENTD3	centaurin, delta 3
1.516	HIG-1; C14orf75; FLJ36164; MGC135025; DKFZp434N0820	TDRD9	tudor domain containing 9
1.51	CAMKK; CAMKKB; KIAA0787; MGC15254	CAMKK2	calcium/calmodulin-dependent protein kinase kinase 2, beta
1.503	MEKK3; MAPKKK3	MAP3K3	mitogen-activated protein kinase kinase kinase 3
1.488	AC; PHP; ASAH; PHP32; FLJ21558; FLJ22079	ASAH1	N-acylsphingosine amidohydrolase (acid ceramidase) 1
1.484	FCRN; alpha-chain	FCGRT	Fc fragment of IgG, receptor, transporter, alpha
1.479	MGC33054	SNX10	sorting nexin 10
1.474	HO68; VA68; VPP2; Vma1; ATP6A1; ATP6V1A1	ATP6V1A	ATPase, H ⁺ transporting, lysosomal 70kDa, V1 subunit A

Relative normalised expression	Common Name	Gene Symbol	Description
1.466	MGST; GST12; MGST-I; MGC14525	MGST1	microsomal glutathione S-transferase 1
1.466	GAIP; RGS GAIP	RGS19	regulator of G-protein signalling 19
1.461	TKT1; FLJ34765	TKT	transketolase (Wernicke-Korsakoff syndrome)
1.449	S171	NUMB	numb homolog (Drosophila)
1.448	FCHO2	FCHO2	FCH domain only 2
1.444	LOC339745	LOC339745	hypothetical protein LOC339745
1.443	CR3A; MO1A; CD11B; MAC-1; MAC1A; MGC117044	ITGAM	integrin, alpha M (complement component 3 receptor 3 subunit)
1.442	D54; hD54; DKFZp686A1765	TPD52L2	tumor protein D52-like 2
1.432	MY014; KIAA0488; MGC20471; MGC126871; MGC126873	SNX27	sorting nexin family member 27
1.429	QK; Hqk; QK3; DKFZp586I0923	QKI	quaking homolog, KH domain RNA binding (mouse)
1.424	EVDB; D17S376	EVI2B	ecotropic viral integration site 2B
1.424	PPT; CLN1; INCL	PPT1	palmitoyl-protein thioesterase 1 (ceroid-lipofuscinosis, neuronal 1, infantile)
1.405	AOAH	AOAH	acyloxyacyl hydrolase (neutrophil)
1.404	MAY1; MGC49908; nPKC-delta	PRKCD	protein kinase C, delta
1.39	IMPA2	IMPA2	inositol(myo)-1(or 4)-monophosphatase 2
1.382	ZYG11; FLJ13456	ZYG11B	zyg-11 homolog B (C. elegans)
1.366	a3; Stv1; Vph1; Atp6i; OC116; OPTB1; TIRC7; ATP6N1C; ATP6V0A3; OC-116kDa	TCIRG1	T-cell, immune regulator 1, ATPase, H+ transporting, lysosomal V0 subunit A3
1.364	PGCP	PGCP	plasma glutamate carboxypeptidase
1.362	NNA1; KIAA1035; DKFZp686M20191	AGTPBP1	ATP/GTP binding protein 1
1.355	TTG2; RBTN2; RHOM2; RBTNL1	LMO2	LIM domain only 2 (rhombotin-like 1)
1.344	CIP1; FLJ46905	SLC12A9	solute carrier family 12 (potassium/chloride transporters), member 9
1.34	ASRT5; IRAKM; IRAK-M	IRAK3	interleukin-1 receptor-associated kinase 3
1.34	NEU; SIAL1	NEU1	sialidase 1 (lysosomal sialidase)
1.332	CRFB4; CRF2-4; D21S58; D21S66; CDW210B; IL-10R2	IL10RB	interleukin 10 receptor, beta
1.321	ASC; TMS1; CARD5; MGC10332	PYCARD	PYD and CARD domain containing kelch repeat and BTB (POZ) domain containing 11
1.31	KLHDC7C; KIAA0711	KBTBD11	containing 11
1.308	LTA4H	LTA4H	leukotriene A4 hydrolase
1.307	NR2B1; FLJ16020; FLJ16733; MGC102720	RXRA	retinoid X receptor, alpha
1.303	JAM; KAT; JAM1; JAMA; JCAM; CD321; JAM-1; JAM-A; PAM-1	F11R	F11 receptor
1.298	LH; LLH; PLOD	PLOD1	procollagen-lysine 1, 2-oxoglutarate 5-dioxygenase 1
1.285	JTK8; FLJ26625	LYN	v-yes-1 Yamaguchi sarcoma viral related oncogene homolog

Relative normalised expression	Common Name	Gene Symbol	Description
1.281	MTX; MTXN	MTX1	metaxin 1
1.28	CGI-44	SQRDL	sulfide quinone reductase-like (yeast)
1.267	FLJ20424	C14orf94	chromosome 14 open reading frame 94
1.248	DCIR; LLIR; DDB27; CLECSF6; HDCGC13P	CLEC4A	C-type lectin domain family 4, member A
1.238	EI; LEI; PI2; MNEI; M/NEI; ELANH2	SERPINB1	serpin peptidase inhibitor, clade B (ovalbumin), member 1
1.234	3PK; MAPKAP3	MAPKAPK3	mitogen-activated protein kinase-activated protein kinase 3
1.227		ACSS2	
1.217	H2A.y; H2A/y; H2AFJ; mH2A1; H2AF12M; MACROH2A1.1; macroH2A1.2	H2AFY	H2A histone family, member Y
1.213	PP3856	NAPRT1	nicotinate phosphoribosyltransferase domain containing 1
1.212	ESP-2; HED-2	ZYX	zyxin
1.179	SPC18; SPCS4A; SEC11L1; sid2895; 1810012E07Rik	SEC11L1	SEC11 homolog A (<i>S. cerevisiae</i>)
1.173	hEDTP; C3orf29; FLJ22405; FLJ90311	C3orf29	myotubularin related protein 14
1.129	TGN38; TGN46; TGN48; TGN51; TTGN2; MGC14722	TGOLN2	trans-golgi network protein 2

The active TB group showed 5281 genes to be differentially expressed as compared to healthy controls, as compared to the latent group, which showed only differential expression of 3137 genes as compared to controls, possibly reflective of a more subdued, although clearly active immune response as shown by overexpression/representation of genes in the cytotoxic module. As an explanation, and not a limitation of the present invention, these results probably explain the observation that changes in additional modules were seen in active TB patients as compared to controls, but not in latent TB as compared to controls. These included overexpressed/represented genes in M1.2 (platelets, genes listed in Table 7A), and underexpressed/represented genes in M1.3 (B cells, genes listed in Table 7B), and M2.8 (T cells, genes listed in Table 7H), the latter perhaps being expected since in the T cells response to *M. tuberculosis* infection, it is possible that T cells are recruited to the site of infection and/or are suppressed during chronic infection. Genes in module M2.4, under-expressed/represented (genes listed in Table 7G) included transcripts encoding ribosomal protein family members whose expression is altered in acute infection and sepsis (Calvano, 2005; Thach, 2005), and genes in this module have also been shown to be underexpressed in SLE, liver transplant patients and those infected with *Streptococcus (S). pneumoniae* (Chaussabel, Immunity, 2005). The largest set of overexpressed genes (66 genes out of 90 detected, Table 7I) in active TB was observed in module, M3.1, (IFN-inducible), and is in keeping with a role of IFN- γ in protection, however genes in this module were not differentially expressed in latent TB patients, who control the infection, as compared to controls. In

active TB genes were underexpressed in a number of modules (M3.4, M3.6, M3.7, M3.8 and M3.9, genes listed in Tables 7L – 7P) containing genes, which did not present a coherent functional module but consisted of an apparently diverse set of genes, and had also been observed to be underexpressed in liver transplant recipients (Chaussabel., 2008, Immunity).

5 Based on transcriptional analysis of whole blood and using this modular map approach active TB patients could be distinguished from latent TB patients. Furthermore, comparison of the modular map obtained for active TB in this study with other modular maps created for different diseases, it is clear that active TB patients have a distinct global transcriptional profile (Figure 9), than observed in patients with SLE, transplant, melanoma or *S. pneumoniae* patients (Chaussabel, 2008, Immunity). Certain modules may be
10 common to a number of diseases such as M2.4, included transcripts encoding ribosomal protein family members, which is underexpressed in active TB, SLE, liver transplant patients and those infected with *S. pneumoniae*. However, genes in other modules are less widely affected, such as M3.1 (IFN-inducible), which although overexpressed in active TB (Figure 9) and SLE (Chaussabel, 2008, Immunity), but not other diseases, particularly *S. pneumoniae*, which shows no differential gene expression in M3.1 as compared to
15 controls. Transcriptional profiles in SLE differ from active TB with respect to over or underexpression of genes in a number of other modules. Likewise, although overexpression of genes in modules M3.2 and M3.3 (“inflammatory”), M1.2 (platelets) and M1.5 (“myeloid”), and underexpression of genes in M3.4, 5, 6, 7, 8 and 9 (non-functionally coherent modules) is observed in active TB and *S. pneumoniae* these diseases can still be distinguished by this method since genes in modules M2.2 (neutrophils), M2.3 (erythrocytes), M3.5
20 (non-functionally coherent module) are overexpressed in *S. pneumoniae* as compared to controls but not differentially affected in active TB. Thus by retaining the complexity and magnitude of the data, yet organizing and reducing the dimension of the complex data, it is possible to distinguish different infectious and inflammatory diseases by transcriptional profiles of blood (Chaussabel, 2008, Immunity).

The present invention identifies a discreet differential and reciprocal dataset of transcriptional signatures in
25 the blood of latent and active TB patients. Specifically, active TB patients showed an over-expression/representation of genes in functional IFN-inducible, inflammatory and myeloid modules, which on the other hand were down-regulated/under-represented in latent TB. Active TB patients showed and increased expression/over-representation of immunomodulatory genes PDL-1 and PDL-2, which may contribute to the immunopathogenesis in TB. Blood from latent TB patients showed an over-
30 expression/representation of genes within a cytotoxic module, which may contribute to the protective response that contains the infection with *M. tuberculosis* in these patients and could provide biomarkers for testing efficacy of vaccinations in clinical trials. We believe the success of our preliminary study is achieved by the strict clinical criteria we have employed, accompanying immune reactivity studies to support attribution of latency, improved quality of RNA collection and isolation, advanced high throughput whole
35 genome microarray platform, and sophisticated data mining tools to retain the magnitude of the gene

expression but with an accessible format (Chaussabel et al., submitted). Such findings will be of value as diagnostics of latent and active TB, may yield insights into the potential mechanisms of immune protection (Latent TB) versus immune pathogenesis (Active TB), underlying these transcriptional differences, and the design of novel therapies for protection or in the design of immune therapeutics in active TB to achieve more rapid cure with anti-mycobacterial drugs.

It is contemplated that any embodiment discussed in this specification can be implemented with respect to any method, kit, reagent, or composition of the invention, and vice versa. Furthermore, compositions of the invention can be used to achieve methods of the invention.

It will be understood that particular embodiments described herein are shown by way of illustration and not as limitations of the invention. The principal features of this invention can be employed in various embodiments without departing from the scope of the invention. Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, numerous equivalents to the specific procedures described herein. Such equivalents are considered to be within the scope of this invention and are covered by the claims.

All publications and patent applications mentioned in the specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications and patent applications are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

The use of the word “a” or “an” when used in conjunction with the term “comprising” in the claims and/or the specification may mean “one,” but it is also consistent with the meaning of “one or more,” “at least one,” and “one or more than one.” The use of the term “or” in the claims is used to mean “and/or” unless explicitly indicated to refer to alternatives only or the alternatives are mutually exclusive, although the disclosure supports a definition that refers to only alternatives and “and/or.” Throughout this application, the term “about” is used to indicate that a value includes the inherent variation of error for the device, the method being employed to determine the value, or the variation that exists among the study subjects.

As used in this specification and claim(s), the words “comprising” (and any form of comprising, such as “comprise” and “comprises”), “having” (and any form of having, such as “have” and “has”), “including” (and any form of including, such as “includes” and “include”) or “containing” (and any form of containing, such as “contains” and “contain”) are inclusive or open-ended and do not exclude additional, unrecited elements or method steps.

The term “or combinations thereof” as used herein refers to all permutations and combinations of the listed items preceding the term. For example, “A, B, C, or combinations thereof” is intended to include at least one of: A, B, C, AB, AC, BC, or ABC, and if order is important in a particular context, also BA, CA, CB, CBA, BCA, ACB, BAC, or CAB. Continuing with this example, expressly included are combinations that contain

repeats of one or more item or term, such as BB, AAA, MB, BBC, AAABCCCC, CBBAAA, CABABB, and so forth. The skilled artisan will understand that typically there is no limit on the number of items or terms in any combination, unless otherwise apparent from the context.

5 All of the compositions and/or methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations may be applied to the compositions and/or methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the
10 spirit, scope and concept of the invention as defined by the appended claims.

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What is claimed is:

1. A method for distinguishing between active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the method comprising:
obtaining a gene expression dataset from a whole blood sample from the patient;
- 5 determining the differential expression of one or more transcriptional gene expression modules that distinguish between infected patients and non-infected individuals, wherein the dataset demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected individuals, and
distinguishing between active and latent *Mycobacterium tuberculosis* (TB) infection based on the one or
10 more transcriptional gene expression modules that differentiate between active and latent infection.
2. The method of claim 1, further comprising the step of using the determined comparative gene product information to formulate a diagnosis.
3. The method of claim 1, further comprising the step of using the determined comparative gene product information to formulate a prognosis.
- 15 4. The method of claim 1, further comprising the step of using the determined comparative gene product information to formulate a treatment plan.
5. The method of claim 1, further comprising the step of distinguishing patients with latent TB from active TB patients.
6. The method of claim 1, wherein the module comprises a dataset of the genes in modules M1.2, M1.3,
20 M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 to detect active pulmonary infection.
7. The method of claim 1, wherein the module comprises a dataset of the genes in modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 to detect a latent infection.
8. The method of claim 1, wherein the following genes are down-regulated in active pulmonary
25 infection CD3, CTLA-4, CD28, ZAP-70, IL-7R, CD2, SLAM, CCR7 and GATA-3.
9. The method of claim 1, wherein the expression profile of Figure 9 is indicative of active pulmonary infection.
10. The method of claim 1, wherein the expression profile of Figure 10 is indicative of latent infection.
11. The method of claim 1, wherein the underexpression of genes in modules M3.4, M3.6, M3.7, M3.8
30 and M3.9 is indicative of active infection.

12. The method of claim 1, wherein the overexpression of genes in modules M3.1 is indicative of active infection.

13. The method of claim 1, further comprising the step of distinguishing TB infection from other bacterial infections by determining the gene expression in modules M2.2, M2.3 and M3.5, which are overexpressed by the peripheral blood mononuclear cells or whole blood in infection other than *Mycobacterium*.

14. The method of claim 1, further comprising the step of distinguishing the differential and reciprocal transcriptional signatures in the blood of latent and active TB patients using two or more of the following modules: M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection.

15. The method of claim 1, wherein the genes that are upregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7A, 7D, 7I, 7J and 7K.

16. The method of claim 1, wherein the genes that are downregulated in active pulmonary TB infection versus a healthy patient are selected from Tables 7B, 7C, 7E, 7F, 7G, 7H, 7L, 7M, 7N, 7O and 7P.

17. The method of claim 1, wherein the genes that are upregulated in latent TB infection versus a healthy patient are selected from Table 8B.

18. The method of claim 1, wherein the genes that are downregulated in latent TB infection versus a healthy patient are selected from Tables 8A, 8C, 8D, 8E and 8F.

19. A method for distinguishing between active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the method comprising:

obtaining a first gene expression dataset obtained from a first clinical group with active *Mycobacterium tuberculosis* infection, a second gene expression dataset obtained from a second clinical group with a latent *Mycobacterium tuberculosis* infection patient and a third gene expression dataset obtained from a clinical group of non-infected individuals;

generating a gene cluster dataset comprising the differential expression of genes between any two of the first, second and third datasets; and

determining a unique pattern of expression/representation that is indicative of latent infection, active infection or being healthy.

20. The method of claim 19, wherein each clinical group is separated into a unique pattern of expression/representation for each of the 119 genes of Table 6.

21. The method of claim 19, wherein values for the first and third datasets are compared and the values for the dataset from the third dataset are subtracted therefrom.
22. The method of claim 19, wherein values for the second and third datasets are compared and the values for the dataset from the third dataset are subtracted therefrom.
- 5 23. The method of claim 19, further comprising the step of comparing values for two different datasets and subtracting the values for the remaining dataset to distinguish between a patient with a latent infection, a patient with an active infection and a non-infected individual.
24. The method of claim 19, further comprising the step of using the determined comparative gene product information to formulate a diagnosis or a prognosis.
- 10 25. The method of claim 19, further comprising the step of using the determined comparative gene product information to formulate a treatment plan.
26. The method of claim 19, further comprising the step of distinguishing patients with latent TB from active TB patients.
27. The method of claim 19, further comprising of determining the expression levels of the genes:
15 ST3GAL6, PAD14, TNFRSF12A, VAMP3, BR13, RGS19, PILRA, NCF1, LOC652616, PLAUR(CD87), SIGLEC5, B3GALT7, IBRDC3(NKLAM), ALOX5AP(FLAP), MMP9, ANPEP(APN), NALP12, CSF2RA, IL6R(CD126), RASGRP4, TNFSF14(CD258), NCF4, HK2, ARID3A, PGLYRP1(PGRP), which are underexpressed/underrepresented in the blood of Latent TB patients but not in the blood of Healthy individuals or Active TB patients.
- 20 28. The method of claim 19, further comprising of determining the expression levels of the genes: ABCG1, SREBF1, RBP7(CRBP4), C22orf5, FAM101B, S100P, LOC649377, UBTD1, PSTPIP-1, RENBP, PGM2, SULF2, FAM7A1, HOM-TES-103, NDUFAF1, CES1, CYP27A1, FLJ33641, GPR177, MID1IP1(MIG-12), PSD4, SF3A1, NOV(CCN3), SGK(SGK1), CDK5R1, LOC642035, which are
25 underexpressed/underrepresented in the blood of Healthy control individuals but were underexpressed/underrepresented in the blood of Latent TB patients, and underexpressed/underrepresented in the blood of Active TB patients.
29. The method of claim 19, further comprising of determining the expression levels of the genes: ARSG, LOC284757, MDM4, CRNKL1, IL8, LOC389541, CD300LB, NIN, PHKG2, HIP1, which are
30 overexpressed/overrepresented in the blood of Healthy individuals, are underexpressed/underrepresented in the blood of both Latent and Active TB patients.
30. The method of claim 19, further comprising of determining the expression levels of the genes: PSMB8(LMP7), APOL6, GBP2, GBP5, GBP4, ATF3, GCH1, VAMP5, WARS, LIMK1, NPC2, IL-15,

LMTK2, STX11(FHL4), which are overexpressed/overrepresented in the blood of Active TB, and underexpressed/underrepresented in the blood of Latent TB patients and Healthy control individuals.

31. The method of claim 19, further comprising of determining the expression levels of the genes: FLJ11259(DRAM), JAK2, GSDMDC1(DF5L)(FKSG10), SIPAIL1, [2680400](KIAA1632),
5 ACTA2(ACTSA), KCNMB1(SLO-BETA), which are overexpressed/overrepresented in blood from Active TB patients, and underexpressed/underrepresented in the blood from Latent TB patients and Healthy control individuals.

32. The method of claim 19, further comprising of determining the expression levels of the genes: SPTANI, KIAAD179(Nnp1)(RRP1), FAM84B(NSE2), SELM, IL27RA, MRPS34, [6940246](IL23A),
10 PRKCA(PKCA), CCDC41, CD52(CDW52), [3890241](ZN404), MCCC1(MCCA/B), SOX8, SYNJ2, FLJ21127, FHIT, which are underexpressed/underrepresented in the blood of Active TB patients but not in the blood of Latent TB patients or Healthy Control individuals.

33. The method of claim 19, further comprising of determining the expression levels of the genes: CDKL1(p42), MICALCL, MBNL3, RHD, ST7(RAY1), PPR3R1, [360739](PIP5K2A), AMFR, FLJ22471,
15 CRAT(CAT1), PLA2G4C, ACOT7(ACT)(ACH1), RNF182, KLRC3(NKG2E), HLA-DPB1, which are underexpressed/underrepresented in the blood of Healthy Control individuals, overexpressed/overrepresented in the blood of the Latent TB patients, and overexpressed/overrepresented in the blood of Active TB patients.

34. A method for distinguishing between active and latent mycobacterium tuberculosis infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the method comprising:

20 obtaining a gene expression dataset from a whole blood sample;

sorting the gene expression dataset into one or more transcriptional gene expression modules; and

mapping the differential expression of the one or more transcriptional gene expression modules that distinguish between active and latent *Mycobacterium tuberculosis* infection, thereby distinguishing between active and latent *Mycobacterium tuberculosis* infection.

25 35. The method of claim 34, wherein the dataset comprises TRIM genes.

36. The method of claim 34, wherein the dataset comprises TRIM genes, and TRIM 5, 6, 19(PML), 21, 22, 25, 68 are overrepresented/expressed in active pulmonary TB.

37. The method of claim 34, wherein the dataset comprises TRIM genes, and TRIM 28, 32, 51, 52, 68, are underrepresented/expressed in active pulmonary TB.

30 38. A method of diagnosing a patient with active and latent *Mycobacterium tuberculosis* infection in a patient suspected of being infected with mycobacterium tuberculosis, the method comprising detecting differential expression of one or more transcriptional gene expression modules that distinguish between

infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent mycobacterium tuberculosis infection.

- 5 39. The method of claim 38, further comprising the step of using the determined comparative gene product information to formulate a diagnosis.
40. The method of claim 38, further comprising the step of using the determined comparative gene product information to formulate a prognosis.
41. The method of claim 38, further comprising the step of using the determined comparative gene
10 product information to formulate a treatment plan.
42. The method of claim 38, wherein the module comprises a dataset of the genes in modules M1.2, M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 to detect active pulmonary infection.
43. The method of claim 38, wherein the module comprises a dataset of the genes in modules M1.5,
15 M2.1, M2.6, M2.10, M3.2 or M3.3 to detect a latent infection.
44. The method of claim 38, wherein the following genes are down-regulated in active pulmonary infection CD3, CTLA-4, CD28, ZAP-70, IL-7R, CD2, SLAM, CCR7 and GATA-3.
45. The method of claim 38, wherein the expression profile of modules of Figure 9 is diagnostic of active pulmonary infection.
- 20 46. The method of claim 38, wherein the expression profile of modules of Figure 10 is diagnostic of latent infection.
47. The method of claim 38, wherein the underexpression of genes in modules M3.4, M3.6, M3.7, M3.8 and M3.9 is indicative of active infection.
48. The method of claim 38, wherein the overexpression of genes in modules M3.1 is indicative of active
25 infection.
49. The method of claim 38, further comprising the step of distinguishing TB infection from other bacterial infections by determining the gene expression in modules M2.2, M2.3 and M3.5, which are overexpressed by the peripheral blood mononuclear cells or whole blood in infection other than Mycobacterium.
- 30 50. The method of claim 38, further comprising the step of distinguishing the differential and reciprocal transcriptional signatures in the blood of latent and active TB patients using two or more of the following

modules: M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection.

51. A kit for diagnosing a patient with active and latent mycobacterium tuberculosis infection in a patient suspected of being infected with *Mycobacterium tuberculosis*, the kit comprising:

5 a gene expression detector for obtaining a gene expression dataset from the patient; and
a processor capable of comparing the gene expression to pre-defined gene module dataset that distinguish between infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent
10 *Mycobacterium tuberculosis* infection.

52. A system of diagnosing a patient with active and latent *Mycobacterium tuberculosis* infection comprising:

a gene expression dataset from the patient; and
a processor capable of comparing the gene expression to pre-defined gene module dataset that distinguish
15 between infected and non-infected patients obtained from whole blood, wherein whole blood demonstrates an aggregate change in the levels of polynucleotides in the one or more transcriptional gene expression modules as compared to matched non-infected patients, thereby distinguishing between active and latent
Mycobacterium tuberculosis infection, wherein the modules are selected from M1.3, M1.4, M1.5, M1.8, M2.1, M2.4, M2.8, M3.1, M3.2, M3.3, M3.4, M3.6, M3.7, M3.8 or M3.9 for active pulmonary infection and
20 modules M1.5, M2.1, M2.6, M2.10, M3.2 or M3.3 for a latent infection.

FIG. 1

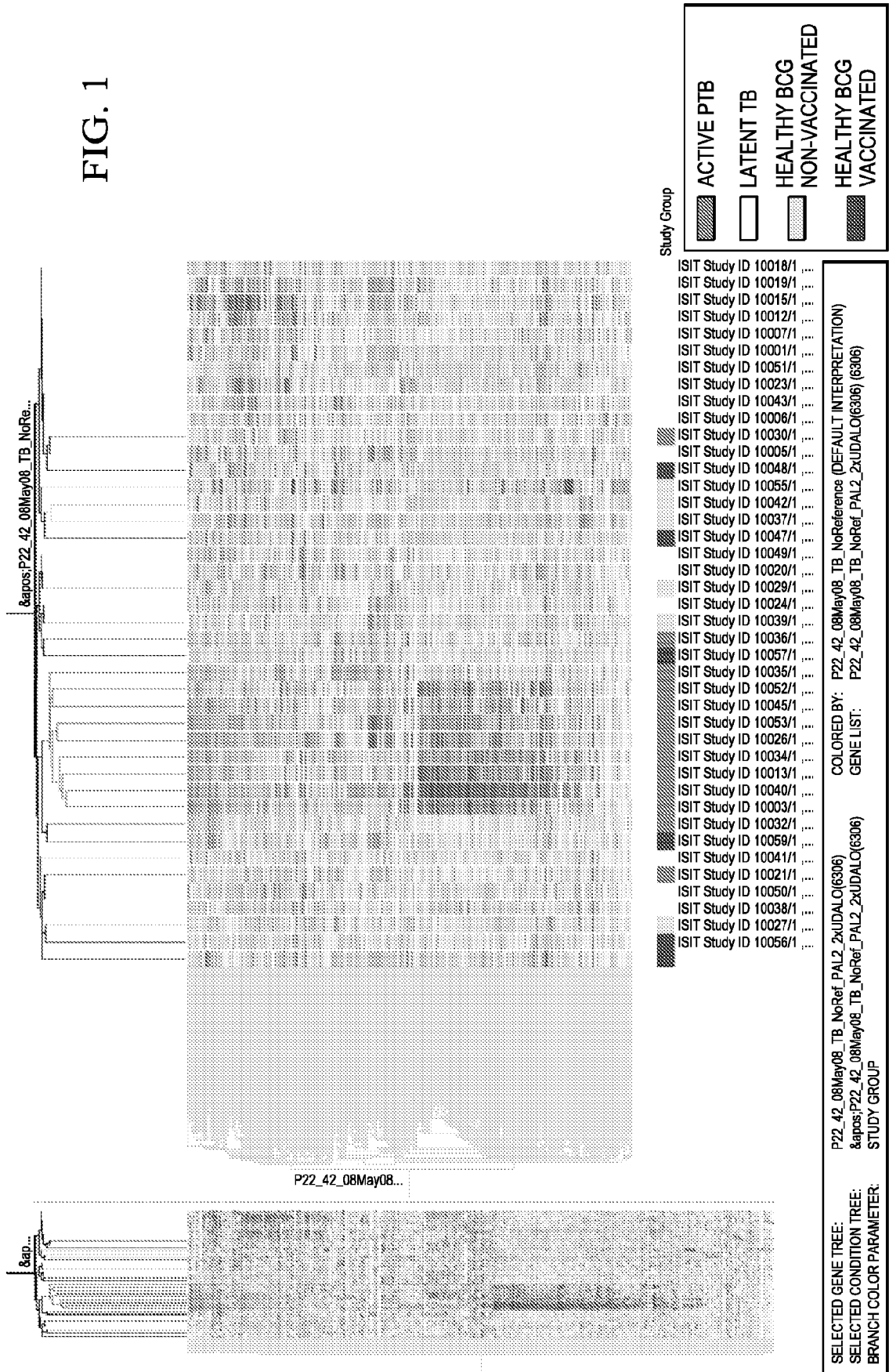
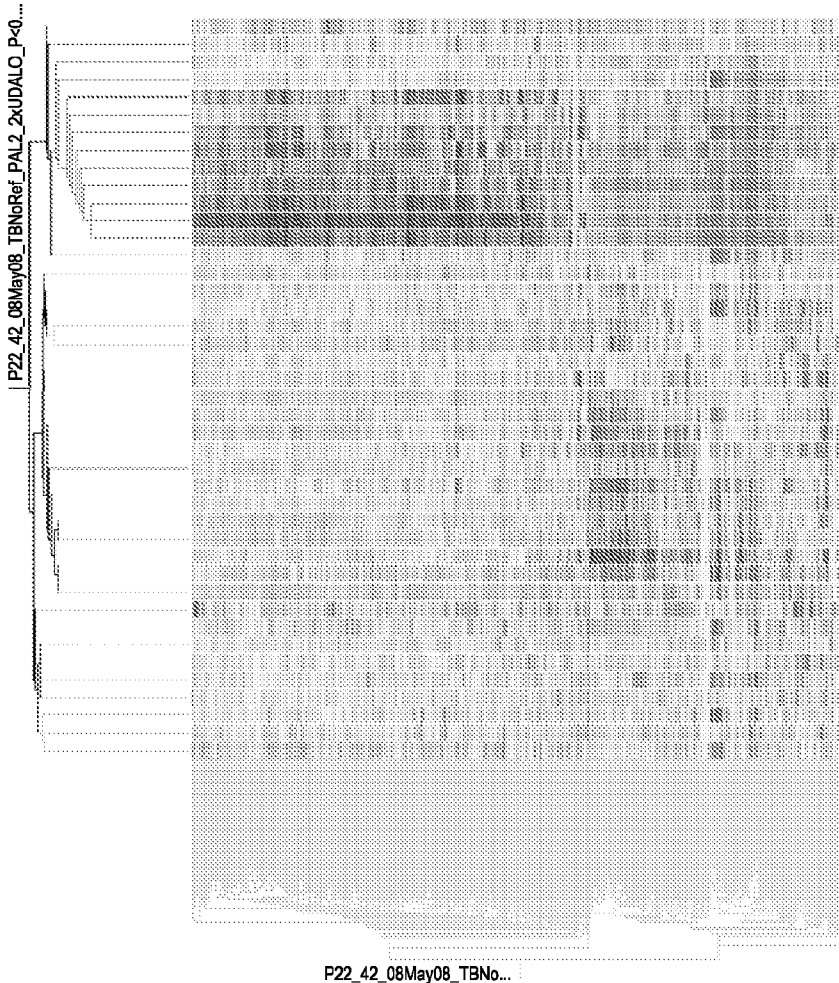


FIG. 2

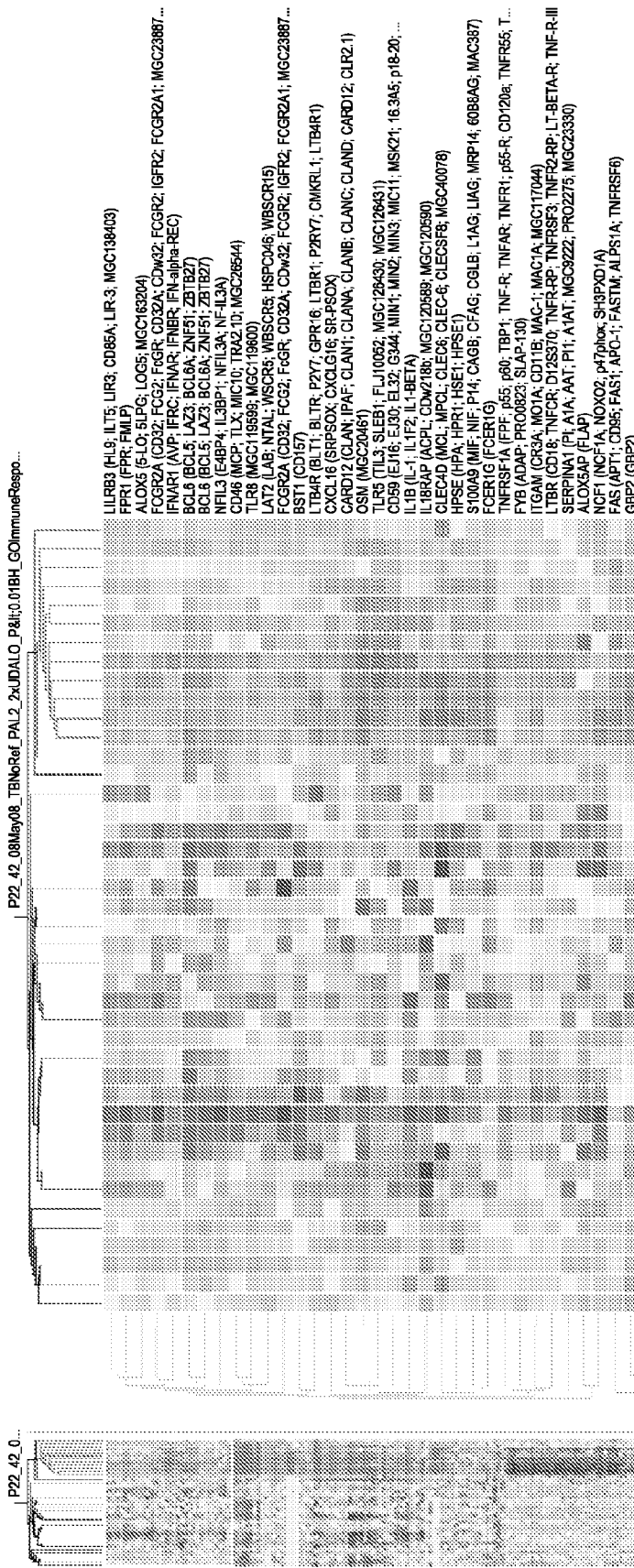


Study Group2

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 SELECTED CONDITION TREE: P22_42_08May08_TBNofRef_PAL2_2xUDALO_P<0.018-H(1473)
 BRANCH COLOR PARAMETER: STUDY GROUP2
 COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_08May08_TBNofRef_PAL2_2xUDALO_P<0.018-H(1473) (1473)

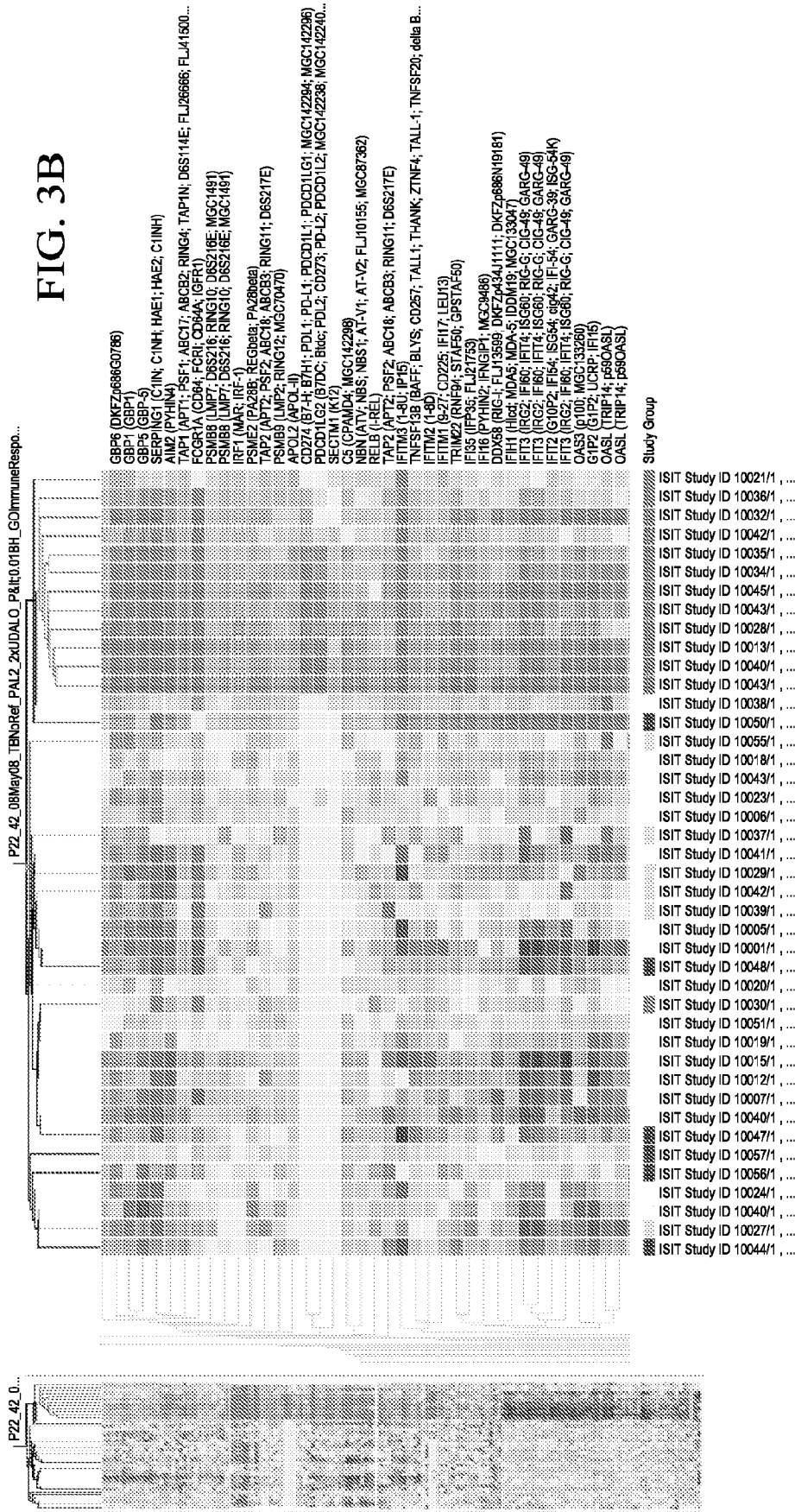
FIG. 3A



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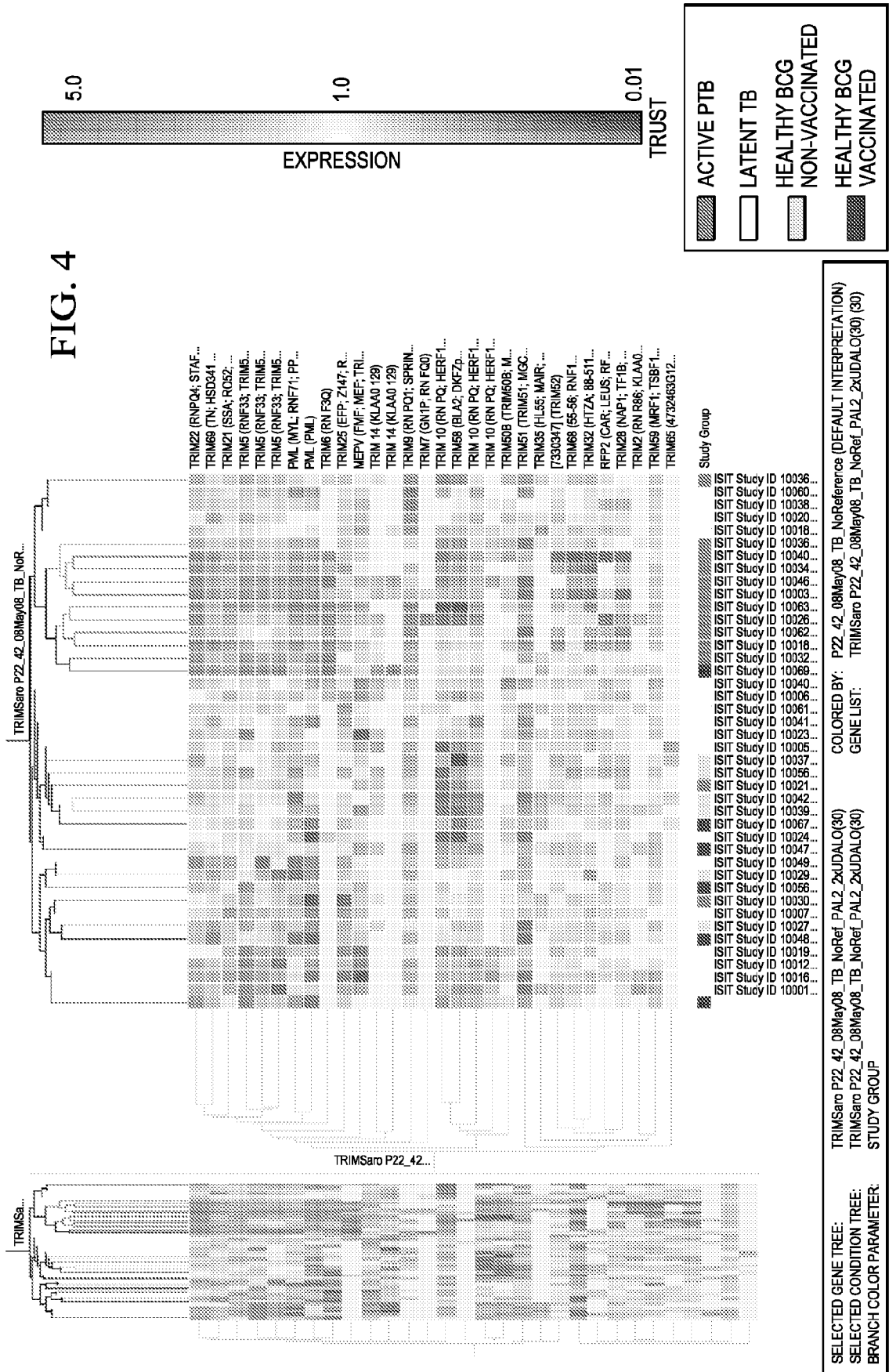
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 SELECTED CONDITION TREE: P22_42_08May08_TBNoRef_PAL2_2xUDALO_P<0.01BH_GOImmuneResponse(158)
 BRANCH COLOR PARAMETER: STUDY GROUP
 COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_08May08_TBNoRef_PAL2_2xUDALO_P<0.01BH_GOImmuneResponse(158) (158)

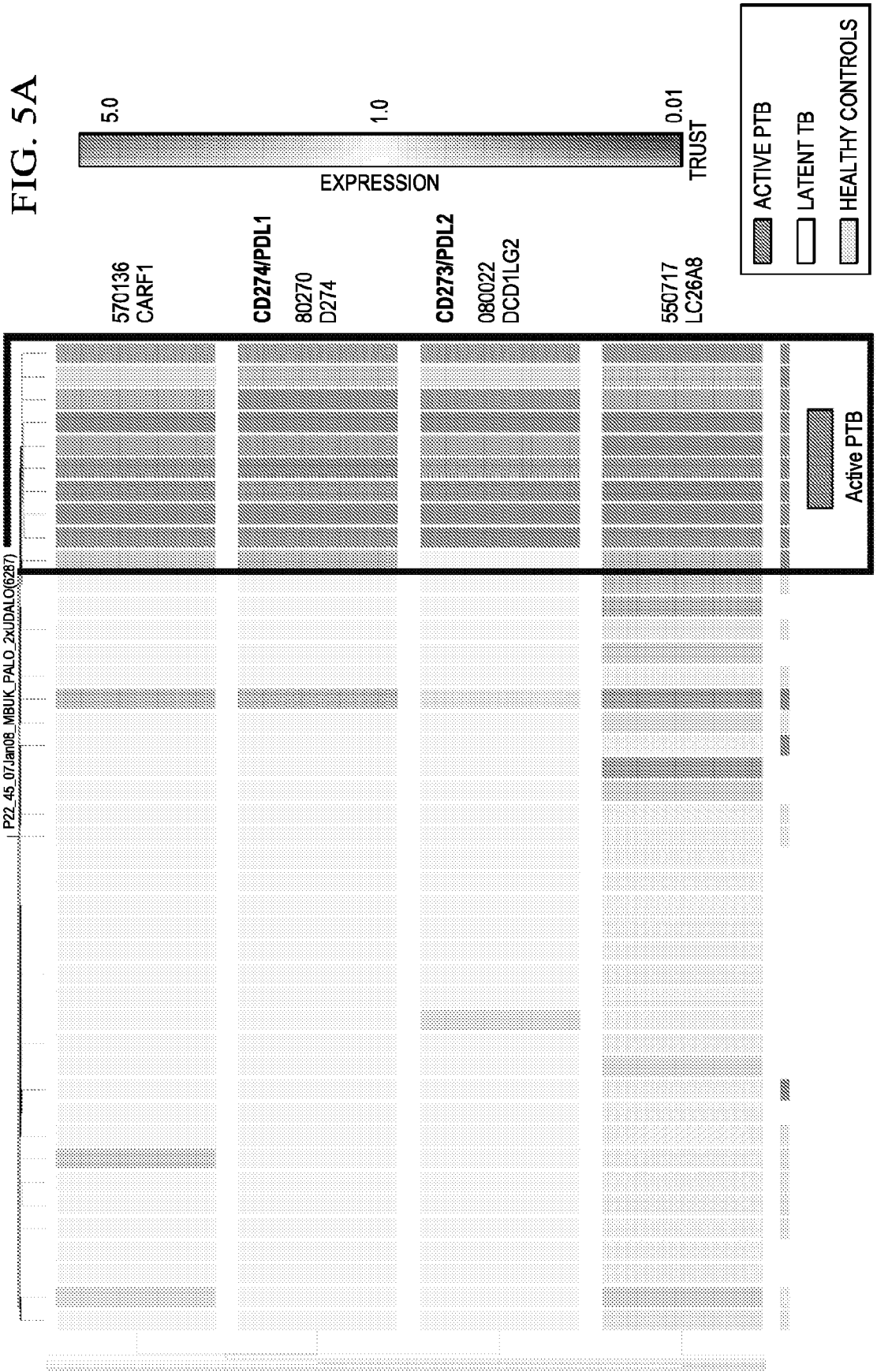
FIG. 3B



SELECTED GENE TREE: P22_42_08May08_TBNoRef_PAL2_2xUDALO_P<0.01BH_GOImmuneResponse(158)
 SELECTED CONDITION TREE: P22_42_08May08_TBNoRef_PAL2_2xUDALO_P<0.01BH_GOImmuneResponse(158) (158)
 BRANCH COLOR PARAMETER: STUDY GROUP

ACTIVE PTB
 LATENT TB
 HEALTHY BCG NON-VACCINATED
 HEALTHY BCG VACCINATED





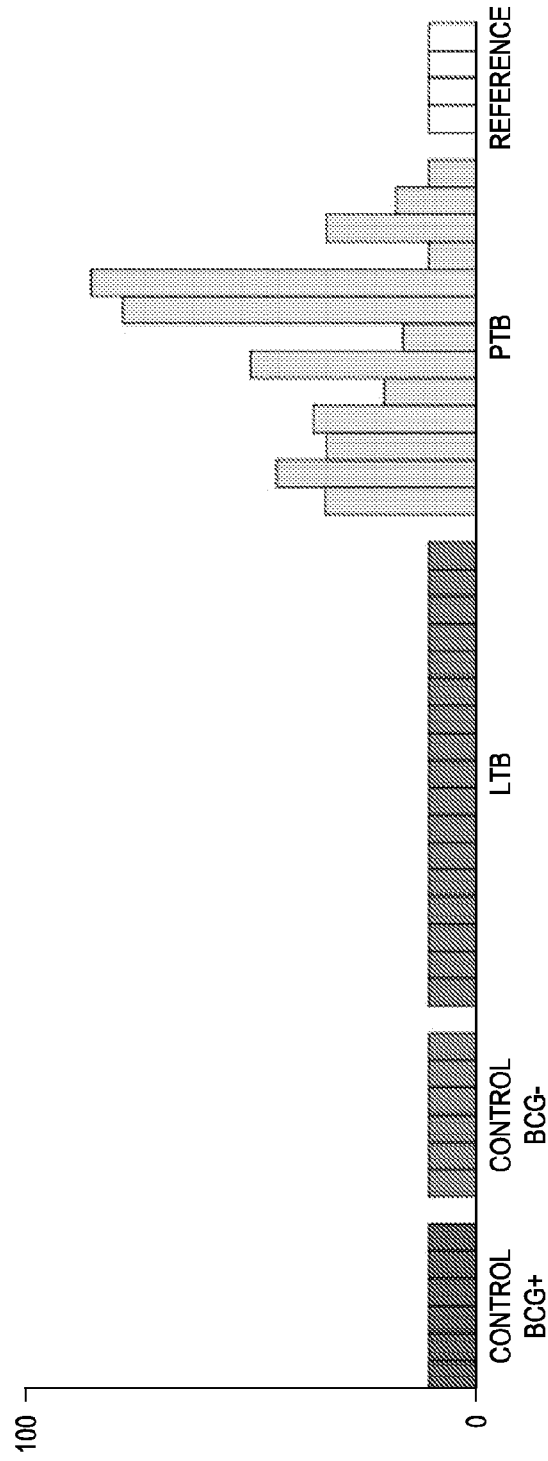


FIG. 5B

FIG. 6

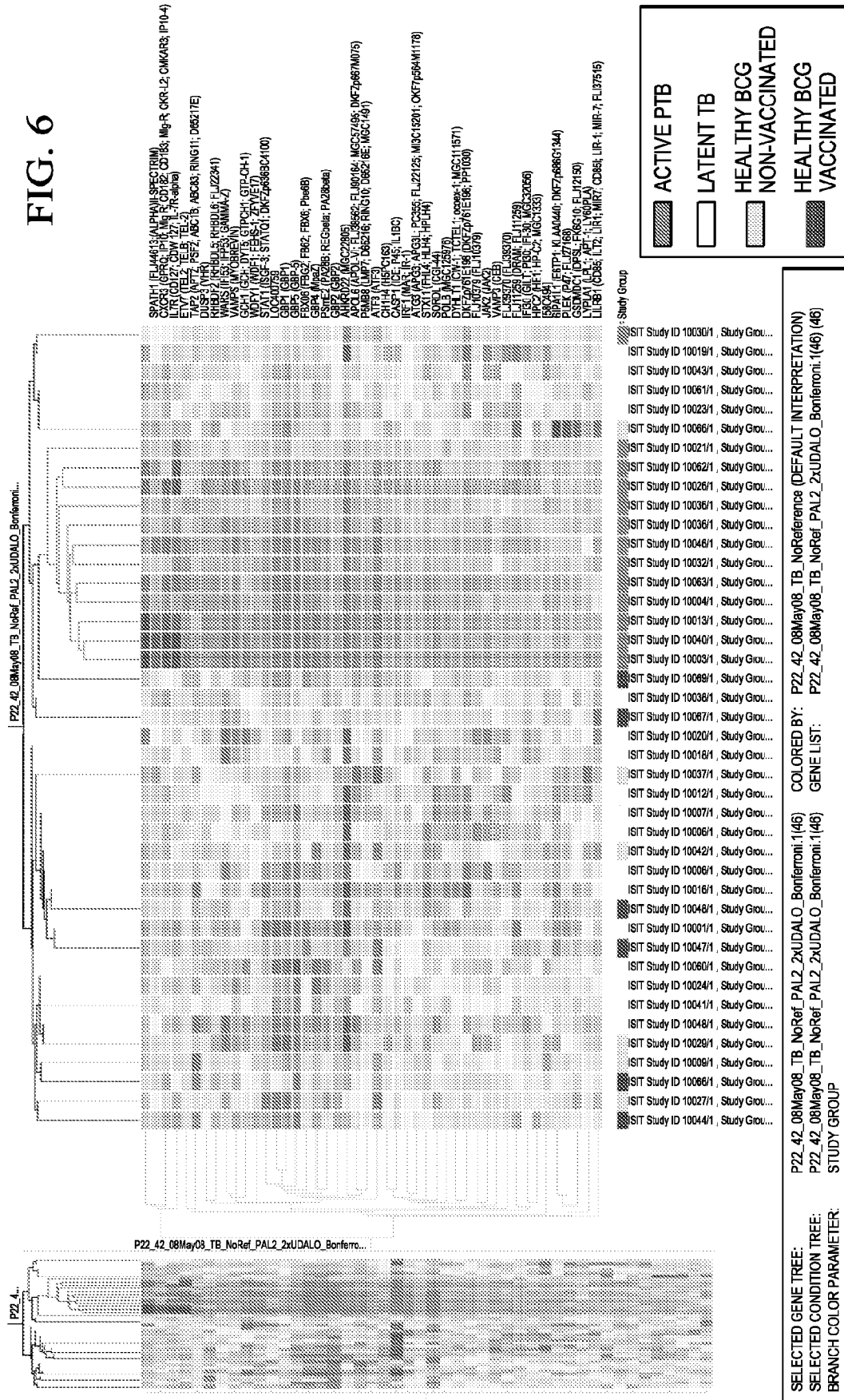
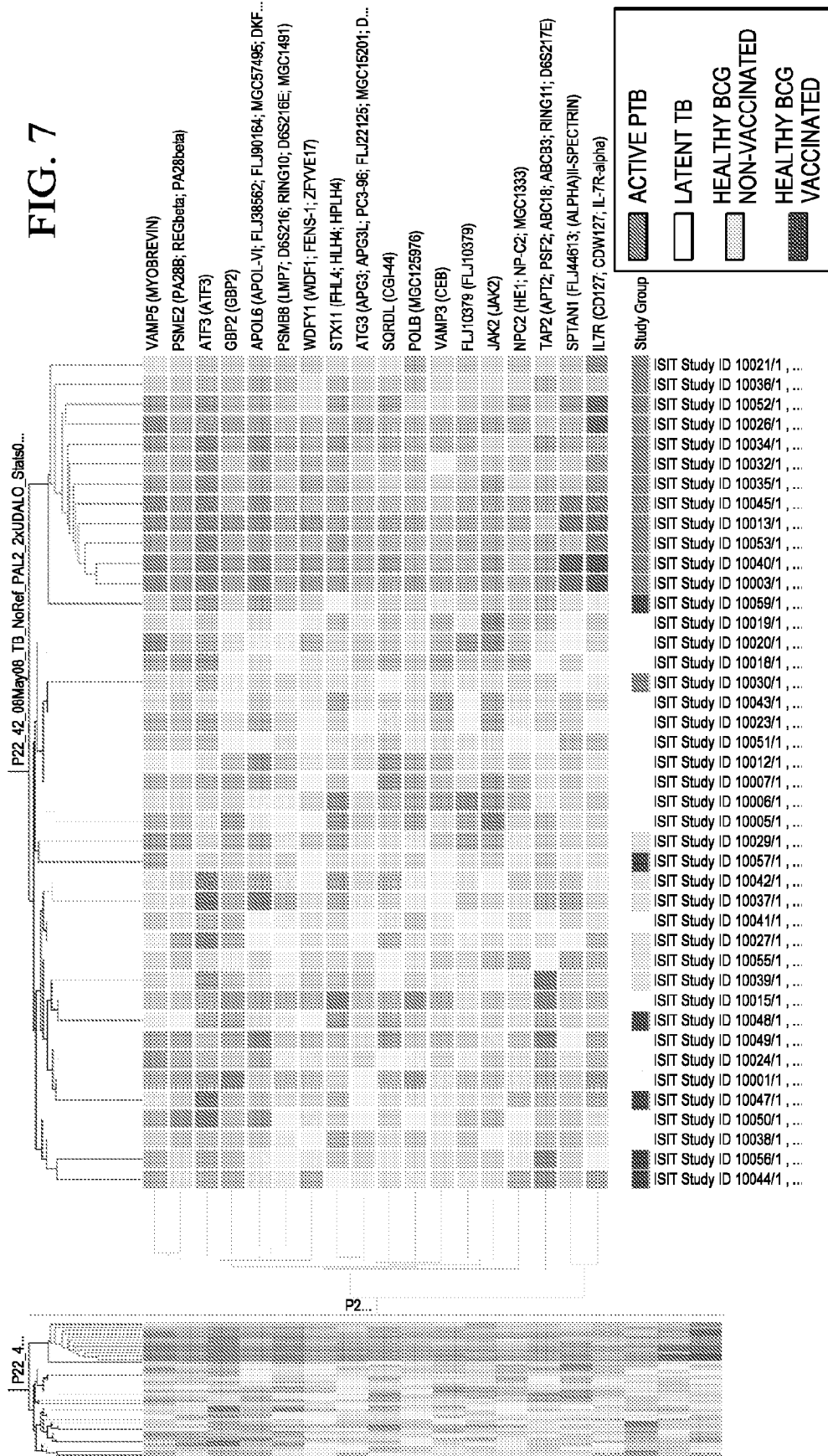


FIG. 7



SELECTED GENE TREE: P22_42_08May08_TB_NoRef_PAL2_2xUDALO_Stats0... (18)
 SELECTED CONDITION TREE: P22_42_08May08_TB_NoRef_PAL2_2xUDALO_Stats0.05Bonferromi(18)
 BRANCH COLOR PARAMETER: STUDY GROUP

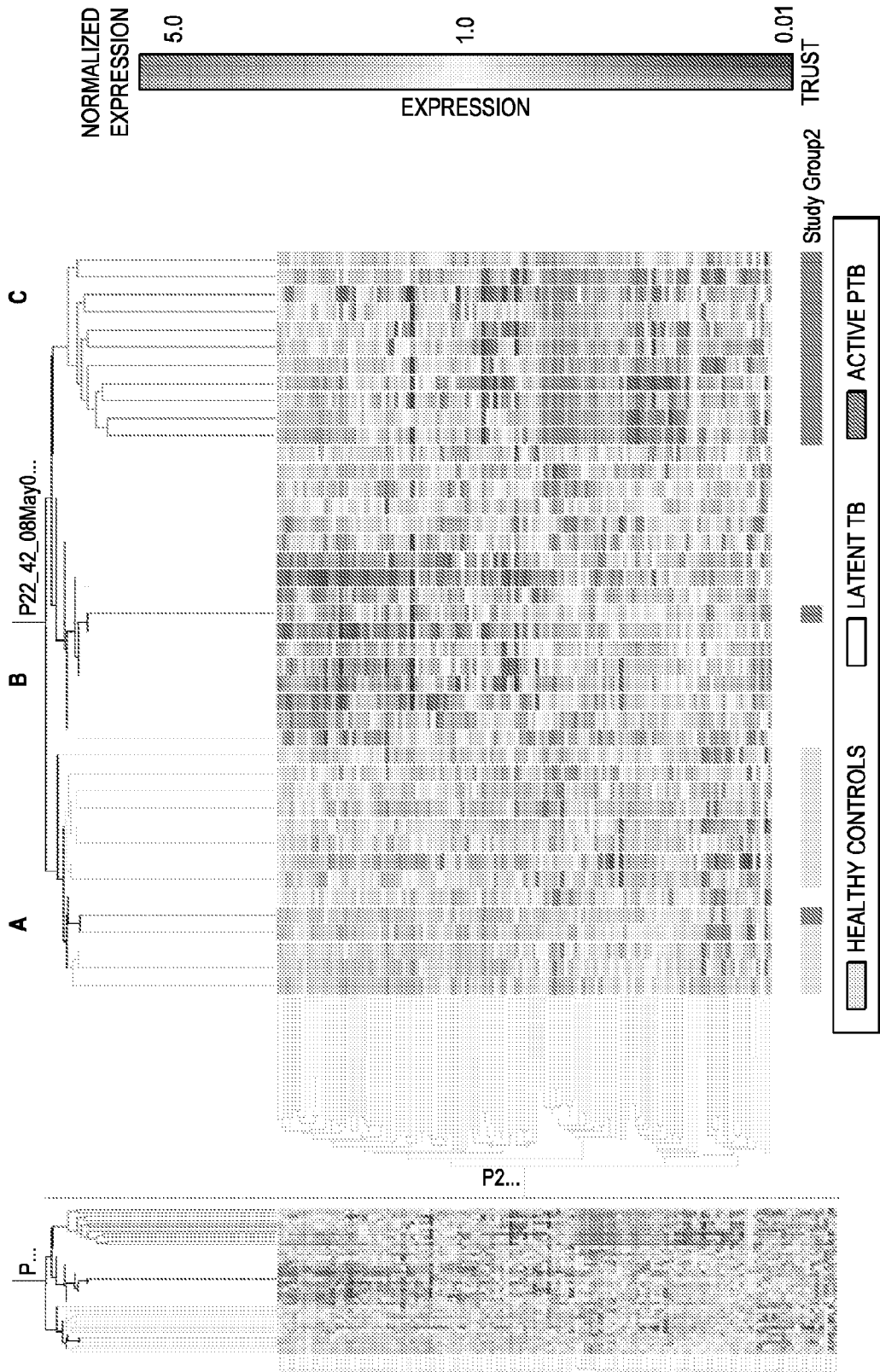
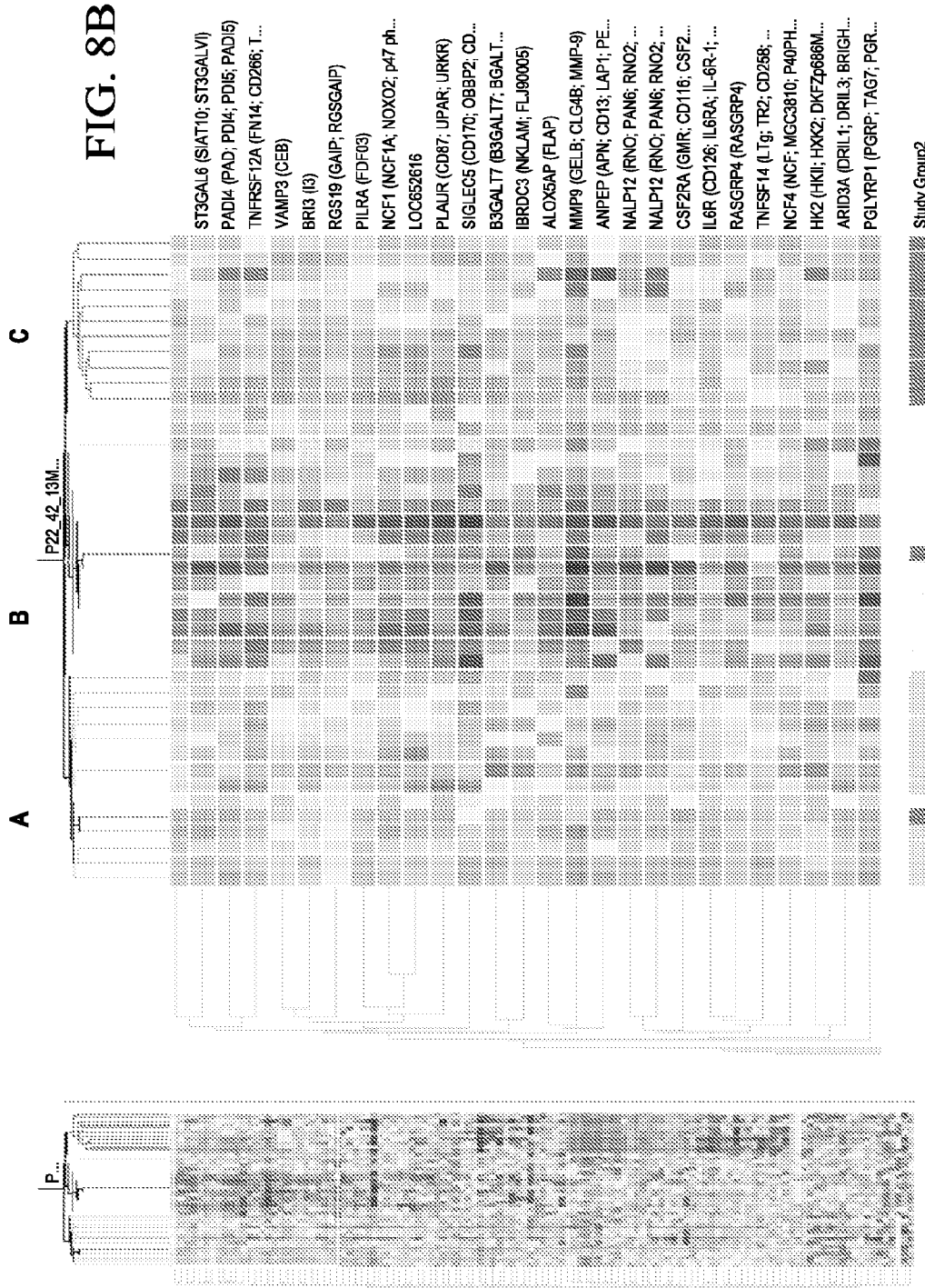
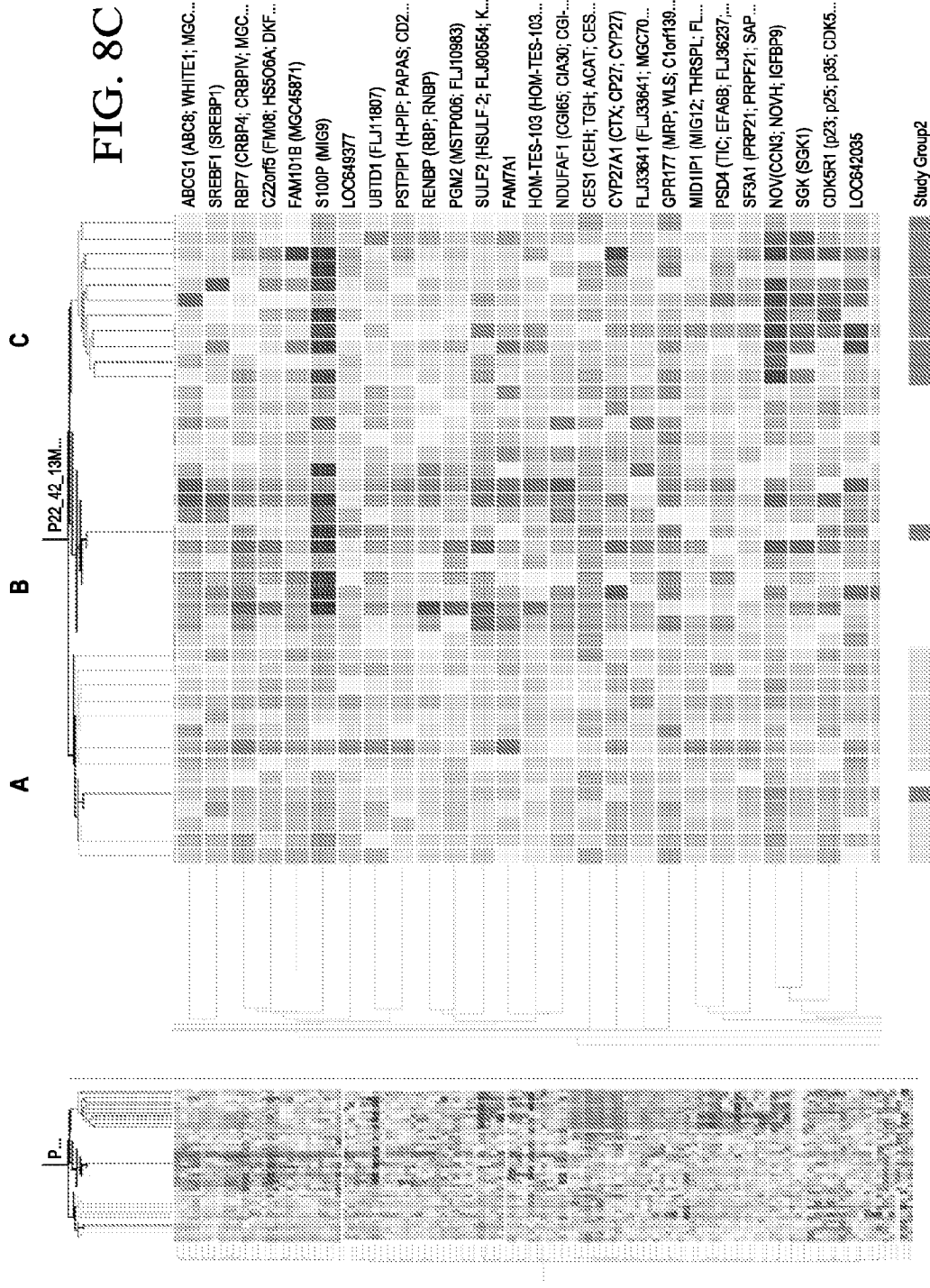


FIG. 8A



SELECTED GENE TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbVLTbVC(119)
 SELECTED CONDITION TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbVLTbVC(119)
 BRANCH COLOR PARAMETER: STUDY GROUP2

COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbVLTbVC(119) (119)



SELECTED GENE TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119)
 SELECTED CONDITION TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119)
 BRANCH COLOR PARAMETER: STUDY GROUP2
 COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119) (119)

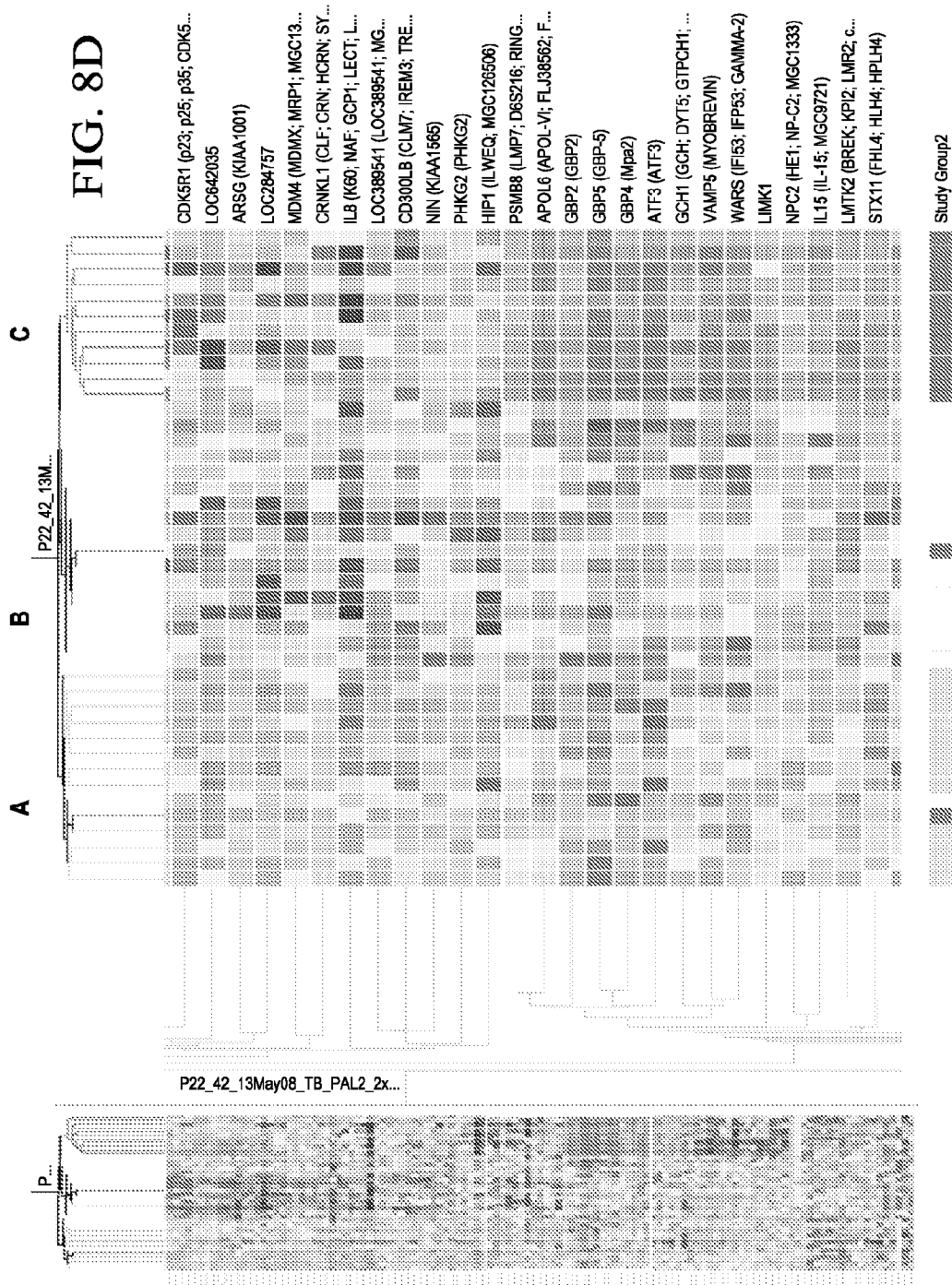
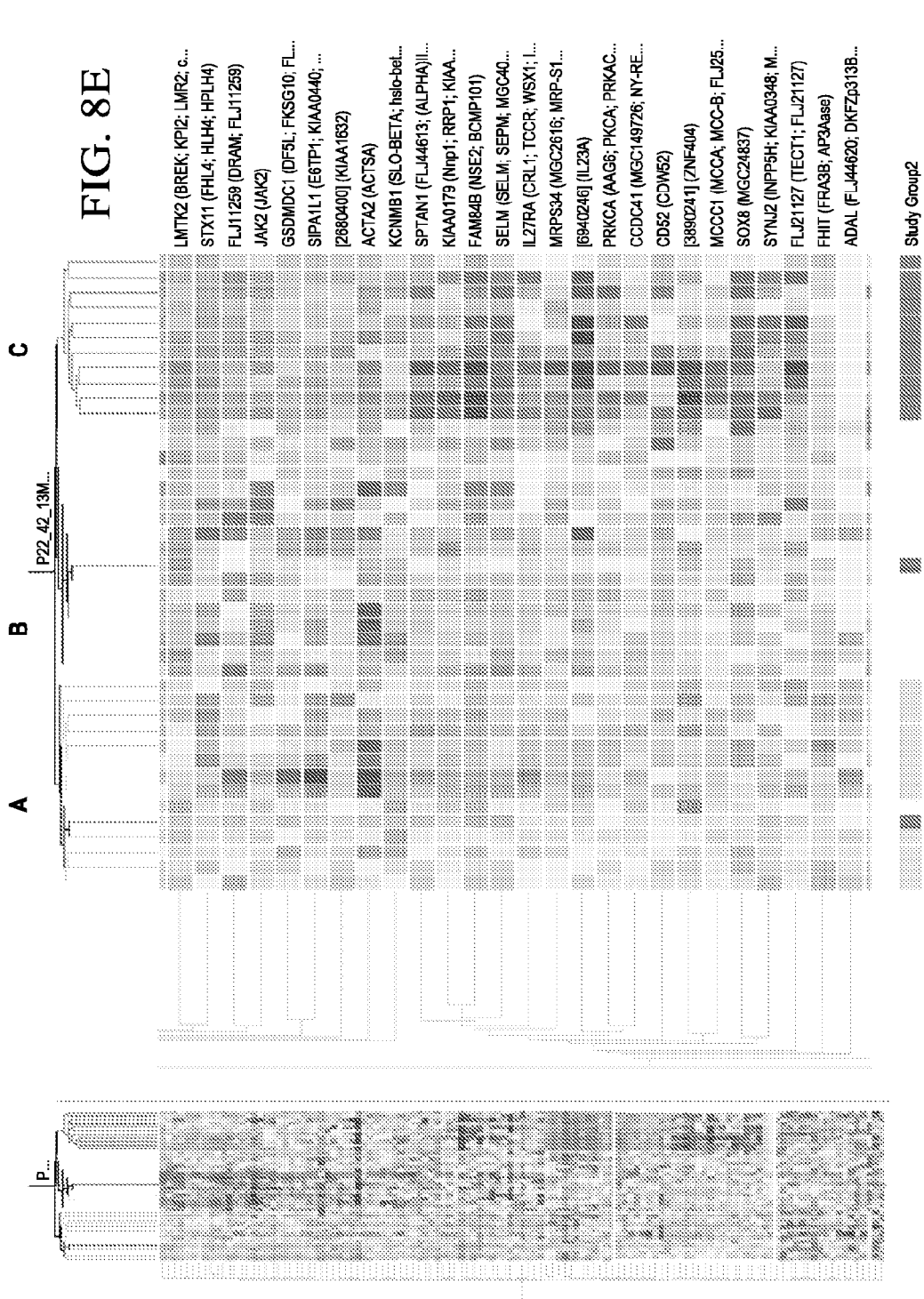


FIG. 8D

SELECTED GENE TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl.TbxC(119)
 SELECTED CONDITION TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl.TbxC(119)
 BRANCH COLOR PARAMETER: STUDY GROUP2

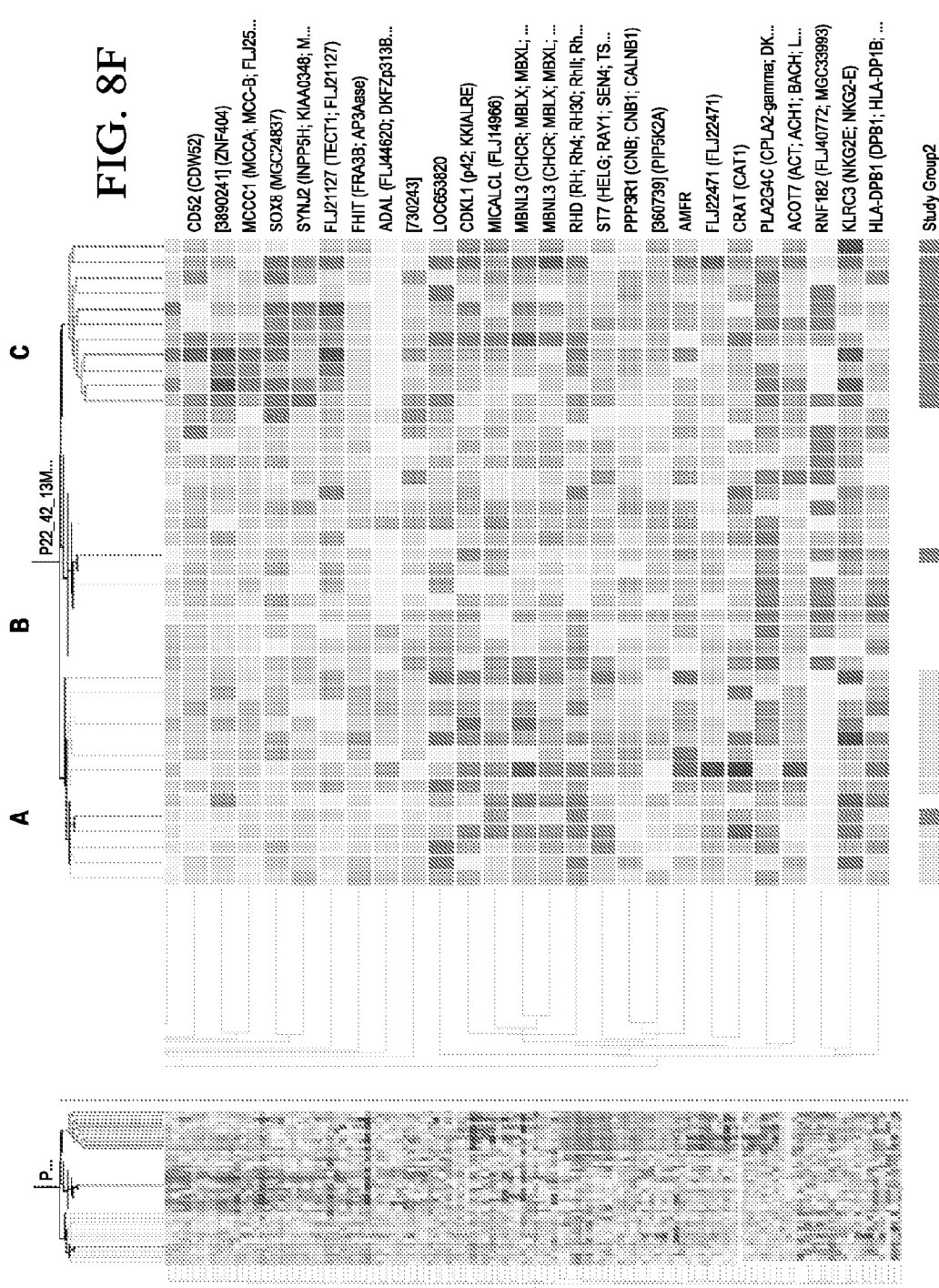
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FIG. 8E



SELECTED GENE TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119)
 SELECTED CONDITION TREE: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119)
 BRANCH COLOR PARAMETER: STUDY GROUP2

COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_13May08_TB_PAL2_2xUDALO_StatsDiff_PTbMl_TbVc(119) (119)



SELECTED GENE TREE: P22_42_13May08_TB_PAL2_2XUDALO_StatsDiff_PTbMl_TbVc(119)
 SELECTED CONDITION TREE: P22_42_13May08_TB_PAL2_2XUDALO_StatsDiff_PTbMl_TbVc(119)
 BRANCH COLOR PARAMETER: STUDY GROUP2

COLORED BY: P22_42_08May08_TB_NoReference (DEFAULT INTERPRETATION)
 GENE LIST: P22_42_13May08_TB_PAL2_2XUDALO_StatsDiff_PTbMl_TbVc(119) (119)

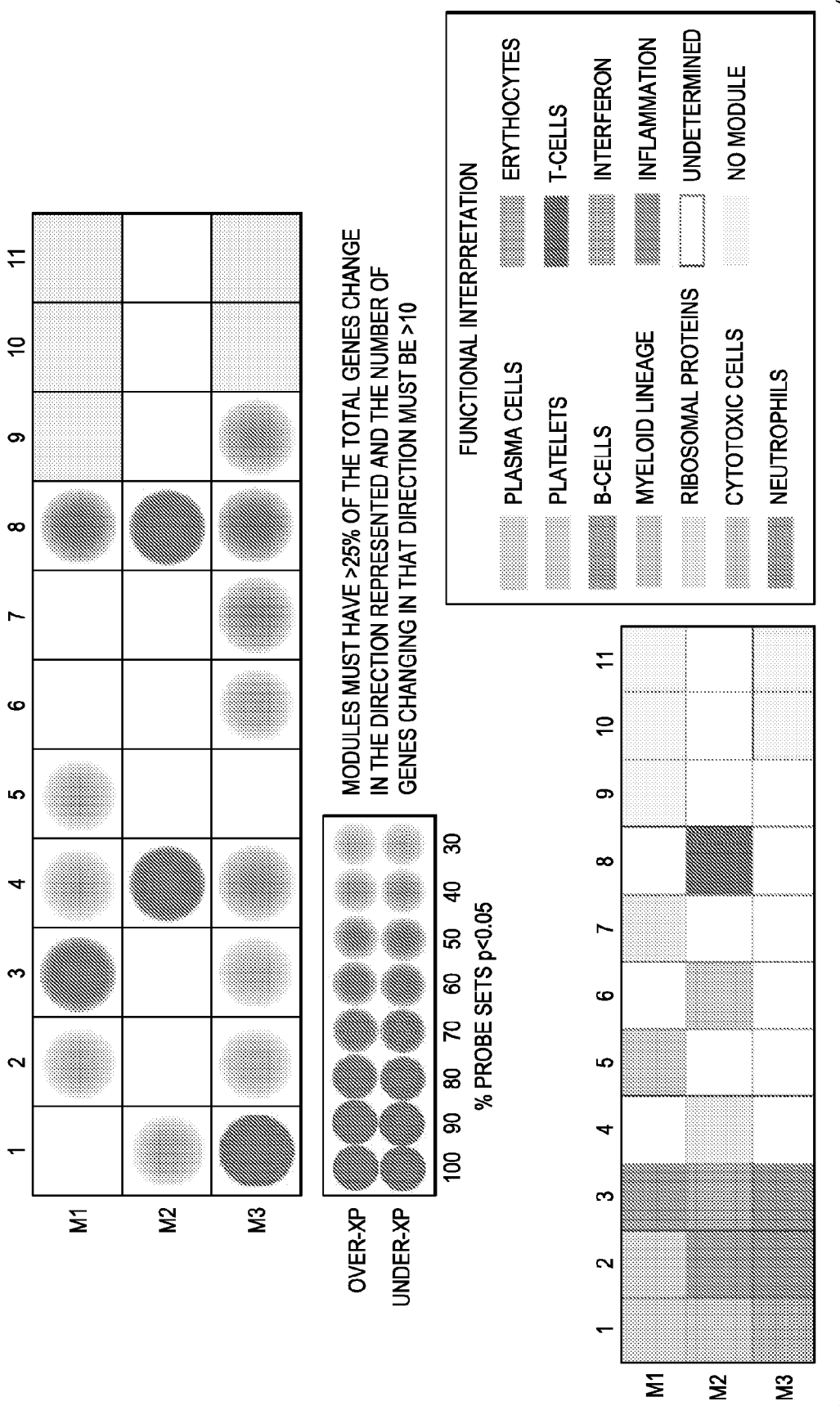


FIG. 9

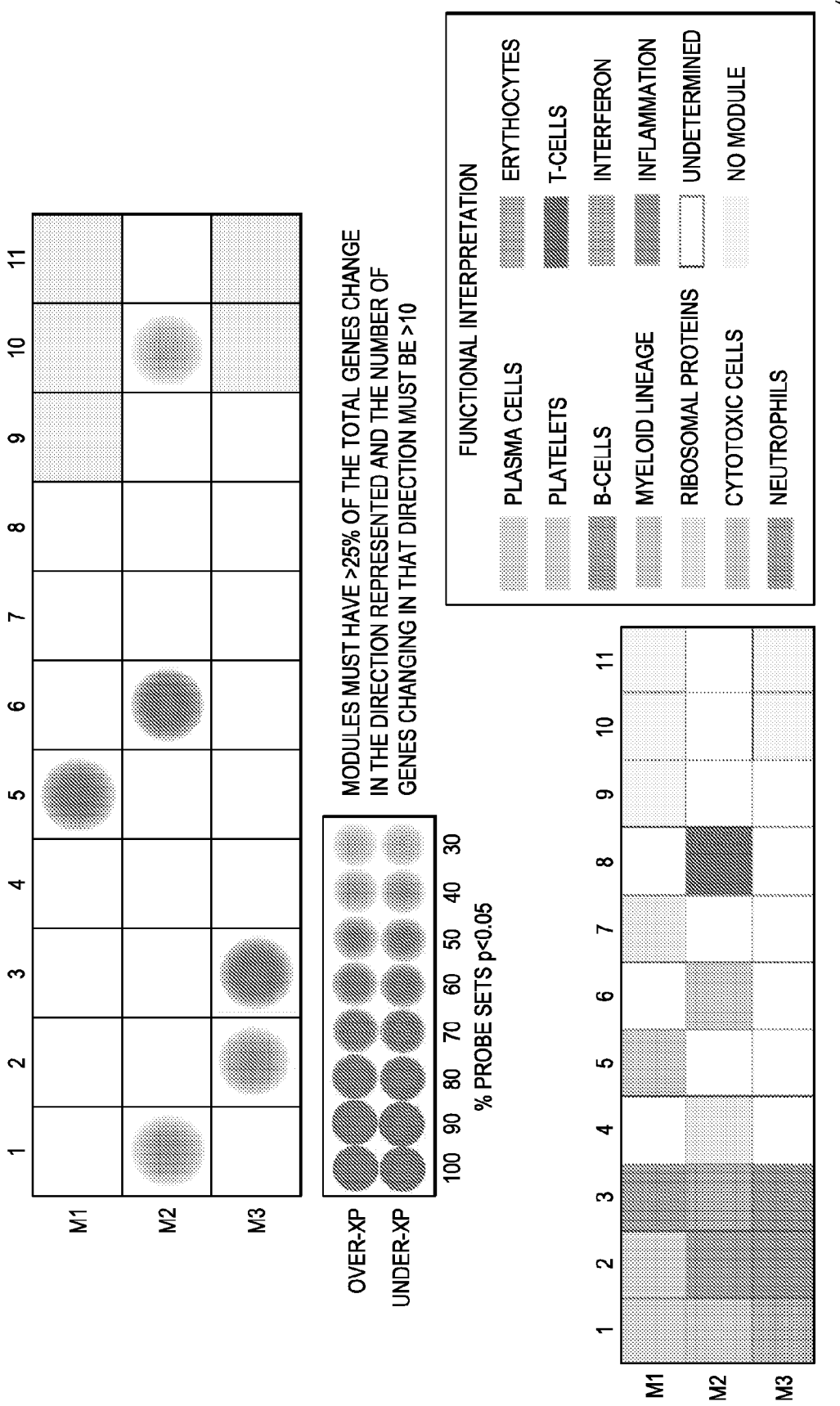


FIG. 10

专利名称(译)	结核分枝杆菌感染的血液转录特征		
公开(公告)号	EP2300823A4	公开(公告)日	2012-03-14
申请号	EP2009771053	申请日	2009-06-25
[标]申请(专利权)人(译)	贝勒研究协会 在NAT INST FOR医疗资源		
申请(专利权)人(译)	BAYLOR研究所 全国学院为医学研究 帝国理工学院医疗保健NHS信托		
当前申请(专利权)人(译)	BAYLOR研究所 全国学院为医学研究 帝国理工学院医疗保健NHS信托		
[标]发明人	BANCHEREAU JACQUES F CHAUSSABEL DAMIEN OGARRA ANNE BERRY MATTHEW KON ONN MIN		
发明人	BANCHEREAU, JACQUES, F. CHAUSSABEL, DAMIEN O'GARRA, ANNE BERRY, MATTHEW KON, ONN, MIN		
IPC分类号	G01N33/50 G01N33/53 C12Q1/00		
CPC分类号	C12Q1/6883 C12Q2600/112 C12Q2600/158 G01N33/5695 G01N2800/60 C12R1/32 G16B99/00		
优先权	61/075728 2008-06-25 US		
其他公开文献	EP2300823A2		
外部链接	Espacenet		

摘要(译)

本发明包括用于区分怀疑感染结核分枝杆菌的患者中活动性和潜伏性结核分枝杆菌感染的方法，系统和试剂盒，并将这些患者与未感染的个体区分开，该方法包括从中获得基因表达数据集的步骤。全血从患者获得样品并确定区分感染和未感染患者的一个或多个转录基因表达模块的差异表达，其中数据集显示一个或多个转录基因表达中多核苷酸水平的总体变化与匹配的非感染患者相比，模块，从而区分活动和潜伏的结核分枝杆菌感染。