Whole-Blood 3-Gene Signature as a Decision Aid for Rifapentine-based Tuberculosis Preventive Therapy

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Background. Systemic drug reaction (SDR) is a major safety concern with weekly rifapentine plus isoniazid for 12 doses (3HP) for latent tuberculosis infection (LTBI). Identifying SDR predictors and at-risk participants before treatment can improve cost-effectiveness of the LTBI program.

Methods. We prospectively recruited 187 cases receiving 3HP (44 SDRs and 143 non-SDRs). A pilot cohort (8 SDRs and 12 non-SDRs) was selected for generating whole-blood transcriptomic data. By incorporating the hierarchical system biology model and therapy–biomarker pathway approach, candidate genes were selected and evaluated using reverse-transcription quantitative polymerase chain reaction (RT-qPCR). Then, interpretable machine learning models presenting as SHapley Additive exPlanations (SHAP) values were applied for SDR risk prediction. Finally, an independent cohort was used to evaluate the performance of these predictive models.

Results. Based on the whole-blood transcriptomic profile of the pilot cohort and the RT-qPCR results of 2 SDR and 3 non-SDR samples in the training cohort, 6 genes were selected. According to SHAP values for model construction and validation, a 3-gene model for SDR risk prediction achieved a sensitivity and specificity of 0.972 and 0.947, respectively, under a universal cutoff value for the joint of the training (28 SDRs and 104 non-SDRs) and testing (8 SDRs and 27 non-SDRs) cohorts. It also worked well across different subgroups.

Conclusions. The prediction model for 3HP-related SDRs serves as a guide for establishing a safe and personalized regimen to foster the implementation of an LTBI program. Additionally, it provides a potential translational value for future studies on drug-related hypersensitivity.

Keywords. interpretable machine learning; latent tuberculosis infection; rifapentine; systemic drug reaction; transcriptome.
facilitate cost-effective implementation of an LTBI program by using the 3HP regimen.

Some studies have proposed gene signatures for LTBI and TB diagnosis [11-13]. However, these studies have not provided absolute thresholds of gene signatures, and the transcriptome-based prediction of SDRs from blood samples before 3HP treatment is lacking. To address these issues, we developed a hierarchical system biology model (HiSBiM) [14] and therapy–biomarker pathway approach to identify a transcriptional signature for 3HP-related SDRs in peripheral blood and then constructed interpretable SDR prediction models with a universal cutoff value by using SHapley Additive exPlanations (SHAP) values [15] to foresee 3HP-related SDRs in LTBI individuals before 3HP treatment.

**METHODS**

**Study Design**

Between January 2017 and January 2020, 447 participants who ever took at least 1 dose of 3HP in 2 medical centers were enrolled. Among them, 44 developed SDRs during 3HP treatment. All of the 44 SDR cases and 143 of the 404 non-SDR cases were selected to identify SDR-related biomarkers. The 8 SDR and 12 non-SDR cases enrolled between January 2017 and June 2019 were grouped as the pilot cohort. The remaining cases, enrolled after July 2019, were randomly divided 80% into the training cohort (28 SDR and 104 non-SDR cases) and 20% into the testing cohort (8 SDR and 27 non-SDR cases).

The study consisted of 4 main parts (Figure 1). First, the pilot cohort was used to identify candidate transcripts that showed a high association with SDR occurrence from peripheral blood transcriptomic profiles through the incorporation of biological similarity score (BS) and HiSBiM (Supplementary Methods). Second, 6 transcripts were selected based on the candidate gene expression profiles by using reverse-transcription quantitative polymerase chain reaction (RT-qPCR; Supplementary Methods, Supplementary Table 1) of pre-3HP samples after pathway-based biomarker selection and validation from 3 SDR and 2 non-SDR samples in the training cohort. Third, random forest (RF) models and SHAP values were applied for SDR predictive model construction by using all samples in the training cohort. Finally, the accuracy of the SDR predictive model was independently evaluated using the testing cohort.

**Participant Selection Criteria**

Individuals were eligible for enrollment if they were aged ≥13 years and received ≥1 dose of 3HP treatment for LTBI, which was diagnosed using QuantiFERON-TB Gold In-tube (Cellestis/Qiagen, Carnegie, Australia). Participants with active TB, a history of LTBI treatment, obesity (BMI >30 kg/m²), which inevitably resulted in a low dosage of isoniazid and rifapentine), malignancy under treatment, living with human immunodeficiency virus, and acute illness with inflammatory symptoms and signs were excluded.

**Study Protocols and SDR Assessment**

Eligible participants received 12 doses of weight-adjusted weekly high-dose rifapentine plus isoniazid under supervision (Supplementary Methods). Within 2 days after each dose of 3HP, the manifestations [4] and severity [16] of any adverse drug reaction (ADR) were assessed and recorded. Blood samples were collected before 3HP treatment, monthly after 3HP treatment, and after SDR development (Supplementary Methods).

**Definition of SDR**

SDR phenotypes included hypotension (systolic blood pressure <90 mm Hg), urticaria, angioedema, acute bronchospasm, or conjunctivitis and more than 4 of the following symptoms occurring concurrently (more than 1 of which had to be grade ≥2): weakness, fatigue, nausea, vomiting, headache, fever, aches, sweating, dizziness, shortness of breath, flushing, or chills [4]. The causative association of 3HP was determined using the Naranjo algorithm [17].

**Bioinformatics Analysis and Candidate Gene Selection**

For RNA-seq data (GSE174552) obtained from the pilot cohort [18], we used the HisBiM and therapy-biomarker pathway approach to identify transcriptomic signatures (Figure 2). A fold change of ≥1.5 and P < .05 (t test) were set to identify 1243 differentially expressed genes (DEGs) before and after 3HP treatment (Figure 1). To investigate the composition of cell types of each sample, we used CIBERSORT to analyze the signatures of the DEGs [19].

To infer potential biomarkers, we first analyzed enrichment pathways using hypergeometric distribution of these 1243 DEGs. Then, we proposed an integrated gene score (Si) to calculate the importance of each DEG (Supplementary Methods). Third, these DEGs were clustered into 60 groups based on BS by using hierarchical clustering analysis (Supplementary Methods) [20]. We then identified 19 potential biomarkers by integrating BS and Si scores and applied RT-qPCR for further validation (Figure 2A). Finally, 6 genes were determined using the therapy–biomarker pathway approach and domain knowledge (Figure 2B).

**Interpretable Prediction Model**

To build a robust prediction model with a universal cutoff value, the training cohort was used to develop interpretable RF models with SHAP values (Supplementary Methods). First, to avoid overfitting in the prediction of 3HP-related SDRs, we systematically tested parameters (eg, trees, features, samples, and leaf nodes) to investigate the characteristics of RF models through the scikit-learn Python library (sklearn.ensemble.RandomForestClassifier) [21].
Next, the SHAP method was applied to these models to reveal the impact of each gene on each sample. The default values were then used to build simple and robust models. The sum of SHAP values of a sample’s input features was used to compare universal cutoff values for predicting 3HP-related SDRs. Finally, the independent testing cohort was used to evaluate the performance of the models based on sensitivity (sen), specificity (spe), and their geometric mean (G-mean, calculated as $\sqrt{\text{sen} \times \text{spe}}$) [22].

**Sensitivity and Statistical Analyses**

We assessed the performance of the selected SDR predictive models in the joint population of training and testing cohorts as well as various subpopulations, stratified by age, sex, BMI, presence of systemic comorbidity, and renal function (using estimated glomerular filtration rate [23] as a surrogate).

The demographic data, comorbidity status, laboratory data, treatment course, and all ADRs were programmatically collected. The intergroup difference was analyzed using the
Figure 2. Gene expression signature and therapy–biomarker pathway for predicting SDR in participants with latent tuberculosis infection before treatment with 3HP. A, Heat map and hierarchical clustering of gene expression (left) and pathway (right) for 19 potential biomarkers in SA, SB, NA, and NB samples. Of the 19 genes, 4 are significantly upregulated after 3HP treatment (dark green), whereas the other 15 genes are not (light green). Among these genes, the 6 selected potential biomarkers are marked in red. B, Therapy–biomarker pathways for illustrating potential genes associated with SDR development under 3HP treatment. For better visualization, the 19 potential biomarkers are underlined with the 6 selected genes marked in red. C, Bar chart for the expression of ATP5PF, GABARAPL2, ATP6V0E1, PIGX, SPCS1, and DDT in 3 SDR (red) and 2 non-SDR (blue) samples collected before 3HP treatment. The expression levels were validated through reverse-transcription quantitative polymerase chain reaction. Abbreviations: 3HP, weekly rifapentine plus isoniazid for 12 doses; MHC, major histocompatibility complex; NA, samples from the non-SDR group after 3HP treatment; NB, samples from the non-SDR group before 3HP treatment; NAD, nicotinamide adenine dinucleotide; SDR, systemic drug reaction; SA, samples from the SDR group after 3HP treatment; SB, samples from the SDR group before 3HP treatment.
Mann–Whitney U test for continuous variables and the χ² test for categorical variables. Statistical significance was set at 2-sided P < .05. The exact binominal method was applied to calculate 95% confidence intervals (CIs) of sensitivity and specificity. Standard normal distribution was used to calculate the 95% CI of the area under the receiver operating characteristic curve.

Ethic Approval
The study was approved by the institutional ethics committees of National Taiwan University Hospital and Kaohsiung Medical University Hospital. Each participant provided informed consent before enrollment.

RESULTS
Selection of Study Participants
Among the 44 SDR cases and 143 non-SDR cases (Figure 1), some had been reported in previous studies (Supplementary Results). Of the 44 SDR cases, 38 (86.3%) presented as flu-like syndrome (Supplementary Table 2), and 93.2% had ≥1 flu-like symptoms. Most SDRs occurred 4–5 hours after the third dose of 3HP and persisted for a median duration of 18–29 hours. The occurrence of SDRs resulted in permanent discontinuation of 3HP in 13 (30%) SDR cases.

The baseline characteristics of the SDR and non-SDR participants were similar, except that the SDR group in the training cohort had a lower prevalence of hypertension and diabetes mellitus and a lower level of serum alanine transaminase (Table 1; Supplementary Table 3).

Selection of Candidate Transcripts
Because of unqualified RNA, 4 non-SDR samples collected before treatment were excluded from mRNA sequencing (Figure 1). On the basis of SΦ and BS scores, we identified 19 candidate genes with significantly higher expression in the 16 SDR samples (orange) than in the 20 non-SDR samples (purple; Figure 2A, left). The accuracy in the discrimination of SDR and non-SDR populations was 94% (34 of 36).

By analyzing the hypergeometric distribution, 4 genes (ie, ATP5PF, ATF4, NDUFB11, and TUBA1C) were involved in the neural-related pathways in Kyoto Encyclopedia of Genes and

Table 1. Clinical Characteristics of Participants

| Characteristic | Pilot Cohort | Training Cohort | Testing Cohort |
|---------------|-------------|----------------|---------------|
|               | SDR (n = 8) | non-SDR (n = 12) | SDR (n = 28) | non-SDR (n = 104) | SDR (n = 8) | non-SDR (n = 27) |
| Age, y        | 51.2 ± 10.8 | 48.8 ± 10.8     | 47.7 ± 14.1  | 50.8 ± 19.8     | 48.1 ± 14.0 | 55.1 ± 21.3      |
| <35           | 1 (12.5%)  | 1 (8.3%)        | 6 (21.4%)   | 30 (28.8%)      | 2 (25.0%)  | 5 (18.5%)        |
| ≥35           | 7 (87.5%)  | 11 (91.7%)      | 22 (78.6%)  | 74 (71.2%)      | 6 (75.0%)  | 22 (81.5%)       |
| Female sex    | 6 (75.0%)  | 10 (83.3%)      | 17 (60.7%)  | 49 (47.1%)      | 4 (50.0%)  | 12 (44.4%)       |
| Body mass index, kg/m² | 22.6 ± 2.7 | 23.3 ± 2.5     | 23.7 ± 3.3  | 24.7 ± 3.9      | 24.0 ± 3.6 | 25.3 ± 4.6       |
| Diabetes mellitus | 1 (12.5%) | 2 (16.7%)       | 2 (7.1%)    | 25 (24.0%)      | 1 (12.5%)  | 8 (29.6%)        |
| Hypertension  | 1 (12.5%)  | 1 (8.3%)        | 4 (14.3%)   | 36 (34.6%)      | 1 (12.5%)  | 11 (40.7%)       |
| Autoimmunea  | 1 (12.5%)  | 0               | 1 (3.6%)    | 2 (1.9%)        | 0          | 0               |
| Isoniazid/Rifampentine dose, mg/kg | 15.1 ± 1.8 | 15.2 ± 1.5     | 14.4 ± 2.6  | 13.6 ± 2.2      | 14.8 ± 2.3 | 13.4 ± 2.2       |
| Hemoglobin, g/dL | 13.2 ± 1.3 | 13.9 ± 1.1     | 14.0 ± 1.6  | 14.1 ± 1.8      | 13.8 ± 1.4 | 14.1 ± 1.7       |
| Leukocyte, K/µL | 6.6 ± 1.2  | 7.2 ± 1.3       | 6.5 ± 1.3   | 7.2 ± 1.8a      | 6.9 ± 1.4  | 7.9 ± 2.5        |
| Platelet, K/µL | 270 ± 36   | 281 ± 60        | 286 ± 61    | 267 ± 62        | 276 ± 76   | 259 ± 72         |
| Alanine transaminase, U/L | 22.0 ± 12.2 | 18.0 ± 11.3   | 19.6 ± 8.2  | 24.2 ± 14.8a   | 24.4 ± 172 | 20.0 ± 5.8       |
| Estimated glomerular filtration rate, mL/min/1.73m²² | 105 ± 30 | 112 ± 30        | 102 ± 37    | 95 ± 25         | 99 ± 18    | 86 ± 31          |
| QuantiFERON response, IU/mL³ | 2.1 ± 2.1 | 1.9 ± 2.5       | 1.8 ± 1.6   | 2.4 ± 25        | 2.3 ± 3.2  | 3.6 ± 2.8        |
| Any adverse event due to weekly rifampentine plus isoniazid for 12 dosesa | 8 (100.0%) | 8 (66.7%)       | 28 (100.0%) | 63 (60.6%)      | 8 (100.0%) | 12 (44.4%)       |
| Grade 3       | 2 (25.0%)  | 1 (8.3%)        | 3 (10.7%)   | 1 (1.0%)        | 2 (25.0%)  | 0               |
| Grade 2       | 6 (75.0%)  | 1 (8.3%)        | 25 (89.3%)  | 22 (21.2%)      | 6 (75.0%)  | 3 (11.1%)        |
| Grade 1       | 0          | 6 (50.0%)       | 40 (38.5%)  | 0               | 9 (33.3%)  |                |

Data are presented as number (percentage) or mean ± standard deviation. Abbreviation: SDR, systemic drug reaction.

aP < .05 between SDR and non-SDR groups.
bThe autoimmune disease was rheumatoid arthritis for the pilot cohort and 2 non-SDR cases in the training cohort. The autoimmune disease was Sjögren syndrome for the SDR case in the training cohort.

³QuantiFERON response was defined as the difference between the interferon-gamma levels of Antigen and Nil tubes as obtained using the QuantiFERON-TB Gold in-Tube test (Cellestis/Qiagen, Carnegie, Australia).

²For detailed information on each adverse event, please see Supplementary Table 3.
Genomes ($P < .05$; Figure 2A, right; Supplementary Figure 1). Other genes were involved in immune, infection, and metabolism-related pathways, implying the hyperinflammatory status. Furthermore, the therapy--biomarker pathway was constructed based on above 19 genes to illustrate and classified into 6 biological functions (Figure 2B; Supplementary Results).

The RT-qPCR results of 3 SDR and 2 non-SDR samples (Supplementary Table 4) in the training cohort revealed that 11 of the 19 genes are consistently highly expressed in SDR samples (Figure 2C; Supplementary Figure 2). According to the therapy--biomarker pathway approach and RT-qPCR results of these 5 samples, 6 potential biomarkers, namely, $ATP5PF$, $ATP6V0E1$, $PIGX$, $SPCS1$, $GABARAPL2$, and $DDT$, were further selected (Supplementary Table 5).

According to CIBERSORT analysis, a high proportion of activated dendritic cells (11.2% vs 2.6%, $P = .006$) and a
low proportion of monocytes (3.7% vs 14.3%, \(P = .009\)) before 3HP treatment were significantly associated with SDRs (Supplementary Figure 3).

**SDR Prediction Models**

To establish a robust prediction model with a universal cutoff value, we developed interpretable RF models with SHAP. Comparisons between RF and SHAP models can be summarized as follows: first, the SHAP model (yellow) has similar performance, with a G-mean of approximately 0.7 for unbalanced and balanced datasets (Figure 3A). Conversely, the G-mean of the RF model for the unbalanced dataset (green) was significantly lower than the G-means of the RF models for balanced datasets and the G-means of the SHAP models for unbalanced datasets \((P < .001)\). Second, SHAP models significantly outperformed RF models in various parameter settings, including default, 500 trees, 6 genes, 5 samples required to be at a leaf node, and 5 samples required to split an internal node \((P < .001)\); Figure 3B). On the basis of these results, we used the SHAP model to provide a universal cutoff value and foresee 3HP-related SDRs before preventive therapy.

The expression profiles by RT-qPCR of all 6 genes were significantly higher in the SDR group compared with the non-SDR group in the training cohort \((P < .005)\); Supplementary Figure 6. The performance of the 4 selected SDR prediction models for the testing samples is shown in Supplementary Table 7. The area under the curve of the 2 best prediction models, that is, ATP6V0E1-PIGX-SPCS1 (yellow) and ATP6V0E1-PIGX-SPCS1-DDT (green), were 0.921 and 0.894, respectively (Figure 4A). The SHAP values of SDR samples were significantly higher than those of non-SDR samples \((P < .001)\); Figure 4B). The interpretation models provide the SHAP output value (Figure 4C), and PIGX is often the most influential in prediction.

**SDR Prediction Model in Subpopulations**

The best SDR prediction models ATP6V0E1-PIGX-SPCS1 and ATP6V0E1-PIGX-SPCS1-DDT for the joint population from training and testing cohorts had G-means of 0.959 (sen = 0.972, spec = 0.921).

Furthermore, SPCSI and GABARAPL2 had a similar expression level (Supplementary Figure 4). On the basis of the findings of redundancy, we selected 4 models with a high G-mean, shown in Figure 3D (details in Supplementary Table 6) for further analysis of SDR prediction. The box plots show significant differences in the SHAP output value between SDR and non-SDR samples \((P < .001)\); Figure 3D). Under a universal cutoff, the SHAP model provided an output value for each sample in these 4 selected models (Supplementary Figure 5).

**Testing of SDR Prediction Models**

The expression profiles based on RT-qPCR of all 6 selected genes were significantly higher in the SDR group compared with the non-SDR group in the testing cohort \((P < .005)\); Supplementary Figure 6. The performance of the 4 selected SDR prediction models for the testing samples is shown in Supplementary Table 7. The area under the curve of the 2 best prediction models, that is, ATP6V0E1-PIGX-SPCS1 (yellow) and ATP6V0E1-PIGX-SPCS1-DDT (green), were 0.921 and 0.894, respectively (Figure 4A). The SHAP values of SDR samples were significantly higher than those of non-SDR samples \((P < .001)\); Figure 4B). The interpretation models provide the SHAP output value (Figure 4C), and PIGX is often the most influential in prediction.

**Figure 4.** SHAP models of the 2 selected models to discriminate pretreatment samples collected from participants with and without SDR in the testing cohort. A, Receiver operating characteristic curve and AUC of the 3-gene (ATP6V0E1-PIGX-SPCS1) and 4-gene (ATP6V0E1-PIGX-SPCS1-DDT) models in the testing cohort. B, Box plot of the SHAP output values of 2 models for the 8 SDR (orange) and 27 non-SDR (steel blue) testing samples. Boxes indicate median and interquartile range, whereas bars and colored dots indicate the range and outliers, respectively. Data were analyzed using the Mann–Whitney U test. C, Interpretation of SDR predictive models with universal cutoffs for SDR (orange) and non-SDR (blue) testing samples. The SHAP output value in each sample is a red diamond, and the universal cutoff is 0 (deep red line). G-mean represents the geometric mean of sensitivity and specificity. Abbreviations: AUC, area under the curve; SDR, systemic drug reaction; SHAP, SHapley Additive exPlanations.
spe = 0.947; Figure 5A) and 0.955 (sen = 0.972, spe = 0.939; Figure 5B), respectively, by using 0 as the cutoff value for SHAP output values. The performance of these SDR prediction models was similar across different subpopulations (sen = 0.933–1.000, spe = 0.900–0.986).

DISCUSSION

This is the first study to provide an accurate prediction model with a universal cutoff value through the integration of clinical samples, bioinformatic techniques, and explainable machine learning to foresee SDR occurrence before 3HP treatment for LTBI. In the management of 3HP-related SDRs, in addition to treatment and supportive care, we herein propose a practical module that consists of the expression profiles of 3–4 genes by using peripheral blood samples to provide a decision aid for safely selecting people with LTBI to undergo 3HP, the most convenient regimen for TB prevention. The SDR prediction model represents a major step forward for precision medicine in TB preventive therapy. As the ancient Greek physician Hippocrates said, “the first and most important oath for being a doctor is do no harm.” This is especially true in the field of preventive medicine. Using the SDR prediction model invented in the current study, almost 97.2% of 3HP-related SDRs can be foreseen and prevented. The intervention program for LTBI treatment could therefore be successfully, safely, and cost-effectively rolled out.

An SDR usually occurred 4–5 hours after the third 3HP dose, most likely due to immune response to certain drug acting as an allergen to elicit endogenous proteins or peptides [24]. Results of CIBERSORT analysis suggest that the difference in immune cell constitution before treatment may be an important driver for T cell–mediated hypersensitivity reaction against 3HP. The proliferation of circulating activated dendritic cells at...
baseline may result in the augmentation of immune response with reexposure to the same antigen [25].

SDR prediction models are based on interpretable SHAP models rather than RF models because of 3 advantages (Figure 3A and 3B). First, the SHAP model is suitable for unbalanced data without the need for adjusting any model parameters. Second, under various machine learning parameters, the SHAP model is more stable and accurate than the RF model. Last, the SHAP model can provide a universal cutoff value based on the comprehensive consideration of the contribution of every feature in each sample.

The underlying pathogenetic mechanism of the 3HP-related hypersensitivity reactions remains unclear. PIGX, a type I transmembrane protein in the endoplasmic reticulum, is required for the complete activation of the regulatory and effector functions of T lymphocytes [26]. GABARAPL2 [27, 28] and ATP6V0E1 [29] both encode proteins essential for autophagy in macrophages. DDT (D-dopachrome tautomerase) regulates a diverse range of physiological functions related to innate immunity and inflammation [30]. SPCSI (signal peptidase complex subunit 1) is involved in the post-translational processing of the structural proteins of Japanese encephalitis virus, Zika virus, and hepatitis C virus [31, 32]. ATP5PF encodes ATP synthase-coupling factor 6 and enhances oxidative phosphorylation in mitochondria during inflammation [33]. However, the exact physiological functions and molecular interactions of the aforementioned genes on 3HP-related SDRs remain undetermined.

The current study has 3 unique characteristics. First, we used a rather large number of clinical samples with clear clinical phenotypes [3]. Second, we proposed a therapy–biomarker pathway approach to identify candidate signatures and developed interpretable machine learning models with SHAP values to provide a universal cutoff value for SDR prediction. Third, the accuracy of the SDR prediction model constructed in the training cohort was confirmed to be high by using an independent testing cohort under a standardized framework of validation. We believe that the multidisciplinary collaboration and the way the clinical problem was addressed as well as the way the SDR prediction model was constructed and tested could be a research model for other severe ADRs in the future.

The current study has some limitations. First, SDRs are likely to be heterogeneous. The exact mechanisms that surround both the predictive transcriptomes and the intercellular processing of host immunity remain unexplained. Second, we cannot identify the offending drug for SDRs. However, the finding that 4 of the 19 potential genes identified in the pilot cohort were involved in the neural-related pathways and the well-documented neurotoxicity of isoniazid (INH) [34] suggest that further investigation of the potential contribution of INH to SDR is necessary. Third, this study only recruited Asians. External validation using non-Asian populations is necessary to confirm the performance of the current SDR prediction models. Fourth, the best signatures depended on the PIGX transcript, yet it was present in tiny amounts. However, it may not be a critical limitation in clinical application given that the average cycle threshold value was 30.3 ± 2.8 and 31.2 ± 1.9 in SDR and non-SDR groups, respectively, suggesting that the low expression level of PIGX in most clinical samples should remain detectable by RT-qPCR.

Last, the current predictive tool would likely be unaffordable or difficult to logistically implement in resource-limited countries, which have the highest TB burden. Further optimization and simplification of the technology or expanding our evaluation to assess its utility in whole blood, as others have similarly explored, might make this tool more feasible to implement.

CONCLUSIONS
By using RNA sequencing to create a global picture of cellular function across all expressed genes, bioinformatic tools to select candidate genes, and machine learning to interpret RT-qPCR results, we provided SDR prediction models with a universal cutoff value to foresee 3HP-related SDRs and provide an aid for treatment decisions before preventive therapy. The SDR prediction model represents a major step forward for precision medicine in TB preventive therapy and may speed up the global uptake of public health programs against LTBI for TB elimination.

Supplementary Data
Supplementary materials are available at Clinical Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copylefted and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes
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