A generalisable approach to drug susceptibility prediction for 
*M. Tuberculosis* using machine learning and whole-genome 
sequencing

The CRyPTIC consortium*

* Members of the CRyPTIC consortium are listed at the end.

Abstract

Rapid and up-to-date drug susceptibility testing is urgently needed to address the threat of multidrug resistant tuberculosis. We developed a composite machine learning system to predict susceptibility from whole-genome sequences for 13 anti-tuberculosis drugs. We trained, validated and externally tested the system, and assessed its performance against a previously validated mutation catalogue, existing molecular assays, and World Health Organization Target Product Profiles. 174,492 phenotypes and 26,328 isolates from 15 countries were studied. The sensitivity of the model was greater than 90% for all drugs except ethionamide, clofazimine and linezolid. Specificity was greater than 95% for all drugs except ethambutol, ethionamide, and bedaquiline, delamanid and clofazimine. The machine learning system was more sensitive than the catalogue and assay (all p<0.01), and correctly predicted a pan-susceptible regimen with 98% accuracy in MDR-TB samples. The proposed system can help guide therapy and be updated automatically as new resistance emerges.

Background

In 2019, 10 million individuals fell ill from *Mycobacterium tuberculosis* infection and 1.4 million died. The problem of multidrug resistant tuberculosis (MDR-TB) - defined as resistance to isoniazid and rifampicin – has been described by World Health Organization (WHO) as a global
health crisis\textsuperscript{1}. Despite advances in diagnostics and treatment, MDR-TB remains under-detected and treatment success remains stubbornly below 60% globally\textsuperscript{1,2}. It is expected that the SARS-CoV-2 pandemic will set back progress that has been made by years\textsuperscript{3}.

The WHO has called for universal drug susceptibility testing (DST)\textsuperscript{4}. Culture-based DST is too slow, expensive and technically challenging to offer a realistic solution. Molecular assays can rapidly detect resistance to rifampicin, isoniazid and a subset of second-line drugs, but are limited in the number of resistance-conferring mutations they can detect\textsuperscript{5}, limiting their sensitivity, although more for some drugs than for others. Some countries already rely on whole-genome sequencing (WGS) to identify susceptibility to first-line drugs, but nowhere are routine diagnostic algorithms advanced enough to dispense with culture-based DST where it is available for second-line, new and repurposed drugs\textsuperscript{6}. Indeed, no predictor has yet been demonstrated to meet the WHO Target Product Profile thresholds for clinical application for drugs now recommended for the treatment of MDR-TB\textsuperscript{7–10}.

Artificial intelligence and machine learning algorithms have been suggested as potential solutions where molecular determinants of resistance are either unknown or complex, such as gene-gene interactions, while allowing for real-time updating as new resistant samples are collected\textsuperscript{11–18}. Here, we compare the performance of a previously-validated mutation catalogue with a composite machine learning DST system to priority anti-tuberculosis agents from WGS data. We assess the extent to which machine learning can bridge the gap between the already good performance of catalogue-based predictions for some drugs and what is needed to dispense with routine phenotypic DST for anti-tuberculosis agents in general.

**Results**

**Characteristics of the datasets**

A total of 174,492 phenotypes from 26,328 isolates were studied across two large datasets. The CRyPTIC dataset derived phenotypes from 96-well broth microdilution plates for 10,859 isolates from 10 countries. Lineages 1 to 4, 6 and *Mycobacterium bovis* were represented, with lineage 4
(50%, 5,436/10,859) and lineage 2 (35%, 3,745/10,859) the most common. 28% of samples (3,033/10,859) were MDR. Phenotypes were available for three first-line antibiotics (isoniazid, rifampicin, ethambutol), plus rifabutin, and nine second-line antibiotics used against MDR-TB: two fluoroquinolones (moxifloxacin, levofloxacin), two injectable agents (amikacin, kanamycin), ethionamide, and four new or repurposed drugs (bedaquiline, delamanid, clofazimine, delamanid). Prevalence of resistance ranged from 1% for bedaquiline, to 47% for isoniazid (Table 1). For each new and repurposed drug, a minimum of 69 resistant samples were available. Pyrazinamide was not present on the microdilution plate for technical reasons. Where two drugs of the same class are studied, we report results for the one present in WHO guidelines or the most commonly prescribed in primary results, and for the other in the supplementary appendix, as is the case kanamycin, rifabutin and levofloxacin. A second, independent dataset used MGIT-derived phenotypes and included 15,469 isolates from 9 countries, 21% of which were MDR (3,189/15,469). The independent set included phenotypes for all antibiotics except new-and-repurposed drugs and rifabutin (Table 1).

**Machine learning in the CRyPTIC dataset**

To assess the performance of the machine learning system on the widest possible set of antibiotics, it was initially trained on 75% of the samples in the CRyPTIC data set (8,146 randomly selected isolates). Predictions were made for the remaining 25% (2,713 isolates) (Table 2). For first-line drugs, the sensitivity of the machine learning system was 95% for isoniazid (1,119/1,173), 97% for rifampicin (906/931) and 95% for ethambutol (387/406), with a specificity of 99% (1,195/1,207), 98% (1,287/1,315) and 88% (1,326/1,501) respectively. For second-line drugs against MDR-TB, sensitivity was 96% for moxifloxacin (251/261), 92% for amikacin (146/158) and 88% for ethionamide (271/309), with a specificity of 96% (1,377/1,438), 99% (2,068/2,090) and 89% (1,694/1,901) respectively. Although there were comparatively few phenotypically resistant isolates, sensitivities for bedaquiline, delamanid, clofazimine and linezolid were 94% (15/16), 90% (18/20), 87% (20/23) and 57% (16/28), at the cost of low specificity (71%, 55%, 72% and 96% respectively). Importantly for clinical decisions on whether a drug can be given, the negative predictive value was above 98% for all drugs except isoniazid,
where it was 96%. We note the low prevalence of resistance to new and repurpose drugs (0.7-1%) as a major contributor to high negative predictive value. We assessed whether results were affected by the split of training and validation data, batch effects, or training and testing on genetically-related samples from the same site, by repeating the experiment using a “leave-one-site-out” cross-validation approach, sequentially using each site as the test set, and training the model on the remaining 10 sites (Table S1 in the Supplementary Appendix). Performances were similar across all first- and second-line drugs with the exception of new and repurposed drugs, where sensitivity decreased (67-73%) and specificity increased (73-78%) using the leave-one-site-out approach. Note that a high proportion of resistant samples to these agents were from the same site in South Africa (31/69 resistant to bedaquiline, 55/105 for clofazimine) causing variability when this specific site is used to train or test.

**Machine learning in the independent dataset**

We further assessed the system’s performance in a large independent dataset. As phenotypic DST for the independent dataset was based entirely on MGIT, this would also provide a test of the generalizability of the machine learning system trained on CRyPTIC broth microdilution plates. We re-trained the machine learning system, this time using the entire CRyPTIC dataset, and found that predictions on the independent dataset were similarly accurate to those seen above (Table 3). For first-line drugs, sensitivity was 95% for isoniazid (3,216/3,397), 98% for rifampicin (2,957/3,021) and 94% for ethambutol (1,765/1,877), with specificity of 99% (9,493/9,602), 98% (10,298/10,502) and 92% (10,758/10,502) respectively. For second-line drugs, sensitivity was 93% for moxifloxacin (288/311) and 88% for amikacin (266/302), with specificity of 96% (2,072/2,168) and 95% (2,403/2,535) respectively. Negative predictive value was greater than 98% for all drugs. There were no phenotypes to new or repurposed drugs for which to make predictions for in the independent dataset.

**Comparison to the catalogue, molecular assays and target product profiles**

We next used a validated mutation catalogue to make predictions for the independent set to see how it compares to the machine learning system (Table 3). For first-line drugs, sensitivity for the catalogue was 94% for isoniazid (3,177/3,397), 97% for rifampicin (2,936/3,021) and 89% for
ethambutol (1,676/1,877), with specificity of 99% (9,525/9,602), 99% (10,394/10,502) and 96% (11,185/10,502) respectively. These results were consistent with the previously-described performance of this catalogue and that led to its clinical implementation for DST to first-line drugs in a number of countries. Nevertheless, these sensitivity results still fell short of the machine learning system’s performance, which was superior by 1% for isoniazid and rifampin, and by 5% for ethambutol (p<0.001). The improved sensitivity of the machine learning system however came at a small cost in specificity which was 1% less for rifampin and 4% less for ethambutol. The machine learning system proved more sensitive than the catalogue for moxifloxacin (93% vs 86%, p<0.001), amikacin (88% vs 85%, p<0.001) and ethionamide (84% vs 50%, p<0.01), with 1% less specificity for moxifloxacin, 3% for amikacin and 21% for ethionamide.

Greater sensitivities can be generated from the catalogue if predictions are reserved for isolates containing genomic variation that is known to the catalogue. However, returning “indeterminate” predictions where novel candidate gene variation is seen in an isolate does not align well with recent WHO target product profiles that require an optimal indeterminate rate of less than 3% for DST implementation. For the catalogue, this rate would have been 6% for isoniazid, 2% for rifampin, 10% for ethambutol, 9% for moxifloxacin, 14% for amikacin and 36% for ethionamide – all of which were predicted as susceptible in this analysis. The machine learning system has the advantage of providing predictions for all isolates (Table S2). We examined phenotypically resistant samples where an “indeterminate” prediction would have been made by the catalogue. In those, the machine learning system correctly predicted 41/106 isolates phenotypically resistant to isoniazid that were missed by the catalogue, 81/94 for ethambutol, 12/15 for moxifloxacin, 2/11 for amikacin and 9/12 for ethambutol. The specificity of the machine learning system for isolates that would have been called “indeterminate” by the catalogue ranged from 88% for ethambutol to 99.7% for rifampin (Table S3).

As most patients in the world have little, or no access to phenotypic DST, we assessed the performance of our system against the expected performance of Xpert MTB/RIF and Xpert XDR in anticipation of its wider uptake to address the WHO’s call for universal DST. We compared the
performance of the machine learning system to the expected combined performance of the Xpert platforms for the independent set. The sensitivity of the machine learning system was 4% greater than that of Xpert for isoniazid (95% vs 91%, p<0.001) and rifampicin (98% vs 94%, p<0.001), 7% for moxifloxacin (93% vs 86%, p<0.001) and 3% for amikacin (88% vs 85%, p<0.05). Specificity was equal or within 1% for each drug, with the exception of amikacin (95% vs 98%). In other words, if 1,000 isolates were resistant to a second-line quinolone or an injectable drug, the machine learning system would accurately find between 30 and 113 phenotypically resistant isolates predicted as ‘not resistant’ by Xpert, at the cost of calling between 0 and 34 phenotypically susceptible isolates ‘resistant’ (Table 2).

The World Health Organization target product profiles (TPP) for rapid molecular DST assays require a minimum sensitivity of 95% for rifampicin, 90% for isoniazid and fluoroquinolones and 80% for other second-line agents; a specificity of 98% for all drugs; and a minimum indeterminate rate of less than 10% (optimal indeterminate rate of less than 3%). In the CRyPTIC dataset, the machine learning system met the minimum TPP sensitivity threshold for all drugs, with the exception of sensitivity for linezolid (57%). Specificity thresholds were met for isoniazid, rifampicin, levofloxacin and amikacin. They were not met for ethambutol (88%), moxifloxacin (96%), ethionamide (89%) and new and repurposed drugs (55-72%) - although still outperforming the specificity of the catalogue and existing molecular assays for each (Table S4). The machine learning system met the optimal requirement for indeterminate results for all drugs as it provides predictions for all samples.

**Full drug regimen prediction for multidrug resistant isolates**

While most DST focuses on predicting susceptibility to individual drugs, clinicians are left with the task of assembling a full regimen themselves. This is especially challenging for rifampicin-resistant (RR) and MDR-TB, where new WHO guidelines recommend the inclusion of new and repurposed drugs like bedaquiline and delamanid for which there is no widely-used DST.

We therefore trained the machine learning system to predict an entire treatment regimen. Regimens were designed according to the latest WHO guidance\(^\text{19}\). A total of 50 possible
regimens were considered, including all combinations of group A, group B and group C drugs meeting WHO standards (Figure 1 and Table S5)\textsuperscript{20}. As only the CRyPTIC dataset included phenotypic DST data for new and repurposed drugs, we used the machine learning system trained on 75% of CRyPTIC to predict regimens for the RR isolates in the 25% test set.

Sufficient phenotypic data were available for 768/931 rifampicin resistant isolates to assess at least one potential regimen for treatment of MDR-TB. The machine learning system predicted a susceptible regimen for 482 of these 768 isolates, and was correct in doing so for 472 (98%). In 8 of the 10 remaining regimens, only one drug in each regimen was phenotypically resistant (Table S6). The system predicted some phenotypic resistance in every potential regimen for the 296 other isolates, of which 139 (47%) isolates had a phenotypically susceptible regimen. Prevalence of bedaquiline, linezolid and clofazimine resistance was 1-2% (9, 7 and 14 samples respectively). Considering each drug individually in phenotypically rifampicin-resistant isolates, the sensitivity for moxifloxacin, levofloxacin and amikacin was respectively 98% (229/233), 96% (256/267) and 96% (133/139), and specificity 90% (357/398), 96% (393/408) and 99% (642/652). Sensitivity for bedaquiline and linezolid was 100% (9/9 and 7/7 respectively), and specificity was 78% and 51% (Table S7 in the Supplementary Appendix).

**Discrepancy analysis**

We reviewed individual cases where the machine learning system made an incorrect prediction in the CRyPTIC set. Where a phenotypically resistant isolate was predicted to be susceptible by the system, we interrogated the predictions from the two subcomponents of the machine learning system for evidence of a predicted increase in MIC, albeit still below the cutoff. For isoniazid, 54/1173 phenotypically resistant samples were predicted to be susceptible in the validation set. One or other of the subcomponents of the machine learning system (ML or the algorithm) predicted an MIC above the base line or near the ECOFF for 16/54 of these (30%). For ethambutol, 13/19 (68%) false negatives led to an increase in MIC from one of the two subcomponents, and 7/19 (37%) from both. Increases in MIC in false negative samples were found in rifampicin (19/25), ethionamid (24/38), levofloxacin (1/22), moxifloxacin (1/10) and amikacin (3/12) (Table S8).
Discussion

We assessed the extent to which machine learning can bridge the gap between the good performance of catalogue-based predictions and what is needed to dispense with routine phenotypic DST not only for first-line drugs but for almost all other anti-tuberculosis drugs too. We trained a machine learning system to predict susceptibility to 13 antituberculosis agents using whole genome sequencing data, and tested its performance on a large independent test set. We followed best practice guidance for studies evaluating the accuracy of rapid tuberculosis drug-susceptibility testing (DST)\(^7\). The machine learning system fully met WHO target product profiles (TPP) for three priority drugs in the CRyPTIC dataset - rifampicin, isoniazid, and amikacin - and met sensitivity but not specificity targets for ethambutol, moxifloxacin, ethionamide and new and repurposed drugs, with the exception of linezolid where no targets were met.

For drugs where the WHO-endorsed molecular GeneXpert assay is available (rifampicin for Xpert MTB/RIF, and isoniazid, fluoroquinolones, aminoglycosides and ethionamide for Xpert MTB/XDR), our system significantly increased the sensitivity and negative predictive values on the validation and test sets compared to the expected performance of these assays, at a small cost to specificity. This can be explained by a variety of factors, including that the assays only look at eight genes and promoter regions and exclude rare variants therein\(^5\), while our system is able to explore genome-wide features, leverage interactions between features and assess lineage and genetic background through genome-wide features.

The WHO guidelines for the management of MDR-TB recommend giving all patients on long MDR-TB regimens bedaquiline, linezolid and clofazimine\(^20\). Sensitivity and specificity for these three drugs fall below WHO TPP requirements. Sensitivity of 93%, 90% and 87% in the CRyPTIC set for bedaquiline, delamanid and clofazimine reflect the very low prevalence of resistance (15,18 and 20 resistant samples respectively) - while also explaining the high negative predictive values of >99% for all three drugs. As more resistant isolates are collected, we expect the sensitivity and specificity of the machine learning system to increase, and NPV to decrease, as has been the case for other drugs. Nevertheless, a test with a NPV of >99% and sensitivity of
70% would provide value to clinicians who currently have no other test for these new and repurposed drugs and hence treat their patients empirically in the absence of reliable, rapid and robust molecular or genotypic DST, playing a key role in preventing the amplification and dissemination of resistance.

A key benefit of our approach over molecular DST is the ability to update and train automatically as new resistant samples are added. This is critical as resistance to existing and new agents like bedaquiline emerge, avoiding the expensive multi-phase multi-year development times of molecular assays, or the need to update catalogues through expert review. The U.S. Food and Drug Administration (FDA) recently released a regulatory framework for ‘live’ modifications to artificial intelligence and machine learning-based software as a medical device and has recently provided clearance or approval for several such diagnostic devices, paving the way for clinical implementation and dissemination.

We note a number of further novelties and benefits of our systems approach. First, by combining machine learning with an algorithmic catalogue generation we leverage existing knowledge, including known genes associated with resistance, avoiding a common complaint against pure machine learning systems. Second, a prediction can be made for all isolates in a given set, while previous published catalogue-based methods that met clinical thresholds required the exclusion of 4-10% of samples with unknown mutations in candidate genes. Third, using kmers from sequencing reads allows for genome-wide analysis while being robust to potential errors or variability in genotype mapping or variant calling, known to affect prediction of transmission inferences and resistance prediction. The method also naturally uses wild-type sequences (the explicit "presence of normal") as prediction features for the machine learning model - analogous to the stretch of 81 nucleotides in the \( rpoB \) gene probed by Xpert - rather than a list of mutations described by a vcf file, where the "absence of abnormal" is inferred from the absence of mutations. Fourth, the model predicts MIC as an intermediate step. Although we have focussed on predicting binary DST results so that we can perform external validation on MGIT data, our system predicts MICs which would allow treatment to be individualized both in terms of drugs selection and dosing. MIC predictions could also be used...
to assess confidence in a susceptibility prediction and mitigate future errors, with isolates without any predicted elevation less likely to be resistant than isolates with a sub-resistant increase in MIC. Fifth, by using an interpretable supervised machine learning algorithm, we provide a list of useful features used for prediction, which in turn can be used as hypotheses for potential causal mutations, when combined with protein analysis.

A limitation of this study is the use of a previously-published literature-derived catalogue, rather than the more cutting-edge, recently-published WHO-endorsed catalogue\textsuperscript{19}. This was impossible as the WHO catalogue was developed using samples from both the CRyPTC and independent sets. Second, we were unable to access an external dataset with sufficient resistance to perform independent validation of predictions to bedaquiline, linezolid, delamanid and clofazimine. As a result, accuracy metrics for the CRyPTIC dataset which does contain DST for these compounds in large numbers are reported using both a left-out validation set and cross-validation of models tested on each site and trained on all other sites. Third, we report the performance of GeneXpert \textit{in silico}, but clinical performance of the actual method might of course be different.

This study demonstrates that WGS can now be used to provide clinically actionable susceptibility prediction for drugs recommended for the treatment of susceptible and of MDR-TB, using an composite machine-learning system. This system can be used to guide therapy, and can be straightforwardly updated as resistant samples to new and repurposed drugs arise and are collected.

**Methods**

**Study design**

We performed a training, validation and external testing study of a mutation catalogue and a machine learning system to predict susceptibility to 13 anti-tuberculosis antibiotics using whole-genome sequencing (WGS). We trained and validated the system on 10,859 isolates from 11 laboratories in 10 countries collected by the CRyPTIC consortium. Phenotypes were
determined using the UKMYC broth microdilution system\textsuperscript{26}. We then assessed how this system, trained on UKMYC-derived phenotypes, would perform against a commonly used DST method in independent samples. For this we made predictions for an external set of isolates used to derive the WHO catalogue of drug resistant mutations\textsuperscript{26}. We selected only those samples that had been phenotypically characterized by Mycobacteria Growth Indicator Tube (MGIT), namely 15,239 \textit{M. tuberculosis complex} isolates from 9 countries (Table 1 for an overview and Table S9 for a detailed description of each dataset). Approval for the CRyPTIC study was obtained\textsuperscript{26}.

**Whole-genome sequencing**

All isolates were whole-genome sequenced using Illumina next-generation sequencing machines, with sequencing protocols varying between sites as previously described\textsuperscript{26}. Sequencing reads were trimmed and mapped to the reference genome H37Rv, and variants called using Clockwork (v0.8.3) a bespoke processing pipeline built for CRyPTIC and optimized to detect both single nucleotide polymorphisms (SNPs) and insertions and deletions (indels). Prior to mapping and calling, raw nucleotide kmers from sequencing reads were set aside for training the machine learning predictor.

**Phenotypic drug-susceptibility testing**

Phenotypic drug-susceptibility testing (DST) for the CRyPTIC training and validation set was performed across all sites using a standard protocol described elsewhere\textsuperscript{26}. Briefly, samples were subcultured and inoculated into 96-well broth microdilution plates containing 13 drugs and designed by the CRyPTIC consortium and manufactured by Thermo Fisher Inc., U.K.. Between 5-10 doubling dilutions were used for each drug, and minimum inhibitory concentrations (MIC) for each were read after 14 days using three methods for quality assurance. MICs were converted to predictions of resistance or susceptibility using epidemiological cutoffs\textsuperscript{26}. As the plate design was modified during the study, the intersect of both plates was used as the MIC phenotype, and concentrations outside both were right-censored or left-censored as appropriate (Table S10). Phenotypic DST for the external test set used the BACTEC MGIT 960 system.
Susceptibility prediction

DST for each sample was predicted using two methods: a mutation catalogue previously tested and validated in CRyPTIC, and a machine learning system. Although the catalogue had previously been tested on 1st line drugs, here we used targets assayed by commercial molecular assays to expanded the catalogue to cover some second line drugs. The machine learning system was itself a composite of two complementary predictors (Figure 1). The first predictor was a kmer-based, hypothesis-free, genome-wide supervised machine learning algorithm. Raw nucleotide kmers \((k=31)\) from sequencing reads (i.e. prior to mapping or assembly) were used as features. A total of \(1.9 \times 10^9\) individual kmers were considered. Where \(<5\) kmers were identified for an isolate these were considered sequencing errors (Figure S1 in the Supplementary Appendix). We merged features across patterns\(^27\), applied feature selection using the F-test applied to MICs, and trained an optimized tree-based extreme gradient boosting method to allow for rapid training, testing and feature interpretation. After training, the top features relevant to each prediction were mapped to H37Rv using bowtie2 for detailed feature analysis (Figure S2 & Table S11). The second predictor was an algorithmic approach that associates mutations with phenotypic resistance modelled on previously described approaches\(^28\). It focussed on the same pre-determined list of candidate genes and promotor sequences as used for the generation of the WHO \(M.\) \(tuberculosis\) drug-resistance mutations catalogue\(^26\) (Table S12). After the masking of neutral mutations using the same process as described\(^26\), the remaining genetic variation across candidate genes relevant to a drug was taken as a unique genetic signature. After the masking of neutral mutations the remaining genetic variation across candidate genes relevant to a drug was taken as a unique genetic signature. This included the absence of any remaining variation, and where there was just a single remaining mutation. The mode MIC from all isolates sharing that unique genetic signature was then taken to predict MICs in test set isolates that shared the same unique signature. If no exact match was made to a combination of variants, the highest mode MIC associated with any individual mutations was used to predict the MIC. Where no match could be made to any genetic signature described in the training set, the test set phenotype prediction was left as ‘U’ (unknown). Both methods’ outcomes were combined into a final joint
prediction system using an “or” logic gate, in order to optimize sensitivity and negative predictive value. Youden’s J statistic was applied to derive the operating threshold of the system. Performance on the independent test set was generated by training the system on the entire CRyPTIC dataset. Performance on the validation set is reported by training the system on CRyPTIC samples not included in it. P-values were calculated using McNemar chi-square test. To better assess the generalizability of the approach within the CRyPTIC dataset and minimize the risk of training and testing on genomically-related isolates, we compared main validation results to those from a leave-one-site-out approach, where each of the 11 sites is left out in turn for training, but correspondingly used for testing, with performance taken as the mean weighted by resistance prevalence. We benchmarked the performance of the catalogue and machine learning system against the expected performance of Xpert® MDR/RIF and Xpert® XDR (Cepheid, Sunnyvale, U.S.), based on the targets they probe (Table S13).
Ethics

Approval for CRyPTIC study was obtained by Taiwan Centers for Disease Control IRB No. 106209, University of KwaZulu Natal Biomedical Research Ethics Committee (UKZN BREC) (reference BE022/13) and University of Liverpool Central University Research Ethics Committees (reference 2286), Institutional Research Ethics Committee (IREC) of The Foundation for Medical Research, Mumbai (Ref nos. FMR/IEC/TB/01a/2015 and FMR/IEC/TB/01b/2015), Institutional Review Board of P.D. Hinduja Hospital and Medical Research Centre, Mumbai (Ref no. 915-15-CR [MRC]), scientific committee of the Adolfo Lutz Institute (CTC-IAL 47-J / 2017) and in the Ethics Committee (CAAE: 81452517.1.0000.0059) and Ethics Committee review by Universidad Peruana Cayetano Heredia (Lima, Peru) and LSHTM (London, UK).

Members of the CRyPTIC consortium (in alphabetical order)

Correspondence to: Alexander S Lachapelle (alexander.lachapelle@eng.ox.ac.uk)


Institutions

1 IRCCS San Raffaele Scientific Institute, Milan, Italy
2 Oswaldo Cruz Foundation, Rio de Janeiro, Brazil
3 Institute Adolfo Lutz, São Paulo, Brazil
4 University of Oxford, Oxford, UK
5 Stanford University School of Medicine, Stanford, USA
6 Scottish Mycobacteria Reference Laboratory, Edinburgh, UK
7 Yale School of Public Health, Yale, USA
8 Universidad Peruana Cayetano Heredia, Lima, Perú
9 Wadsworth Center, New York State Department of Health, Albany, USA
10 Chinese Center for Disease Control and Prevention, Beijing, China
11 Bill & Melinda Gates Foundation, Seattle, USA
12 UK Health Security Agency, London, UK
13 Vita-Salute San Raffaele University, Milan, Italy
14 University of New South Wales, Sydney, Australia
15 The University of British Columbia, Vancouver, Canada
16 Public Health Ontario, Toronto, Canada
17 SYNLAB Gauting, Munich, Germany
18 Institute of Microbiology and Laboratory Medicine, IMLred, WHO-SRL Gauting, Germany
19 EMBL-EBI, Hinxton, UK
20 National Institute for Communicable Diseases, Johannesburg, South Africa
21 Public Health England, Birmingham, UK
22 Taiwan Centers for Disease Control, Taipei, Taiwan
23 Hinduja Hospital, Mumbai, India
24 University of Cape Town, Cape Town, South Africa
25 University of Surrey, Guildford, UK
26 Imperial College, London, UK
27 Université de Montréal, Canada
28 The Foundation for Medical Research, Mumbai, India
29 Research Center Borstel, Borstel, Germany
30 Africa Health Research Institute, Durban, South Africa
31 London School of Hygiene and Tropical Medicine, London, UK
32 Oxford University Clinical Research Unit, Ho Chi Minh City, Viet Nam
33 University College London, London, UK
34 National University of Singapore, Singapore
35 Instituto Nacional de Salud, Lima, Perú
Additional authors contributing to the CRyPTIC consortium (in alphabetical order)

Irena Arandjelovic¹, Anna Barbova², Maxine Caws³, Iñaki Comas⁴, Roland Diel⁵, Carla Duncan⁶, Sarah Dunstan⁷, Maha Farha⁸, Margaret Fitzgibbon⁹, Vicky Furio¹⁰, Jennifer Gardy¹¹, Jennifer Guthrie⁶, Dang Thi Minh Ha¹², Kathryn Holt¹³, Michael Inouye¹⁴, Frances Jamieson⁶, Mostofa Kama¹⁵, Julianne Kus⁶, Vanessa Mathys¹⁶, Rick Twee Hee Ong¹⁷, Youwen Qin¹⁸, Tom Rogers¹⁹, Gian Maria Rossolini²⁰, Emma Roycroft⁹, Vitali Sintchenko²¹, Alena Skrahina²², Yik Ying Teo¹⁷, Phan Vuong Khac Thai¹², Dick van Soolingen²³, Mark Wilcox²⁴, Matteo Zignol²⁵

Institutions

¹ University of Belgrade, Belgrade, Serbia
² Ministry of Health of Ukraine, Kyiv, Ukraine
³ Liverpool School of Tropical Medicine, United Kingdom
⁴ Biomedicine Institute of Valencia IBV-CSIC, Spain
⁵ University Medical Hospital Schleswig-Holstein, Germany
⁶ Public Health Ontario, Toronto, Canada
⁷ University of Melbourne, Australia
⁸ Harvard Medical School, Boston, USA
⁹ Irish Mycobacteria Reference Laboratory, Dublin, Ireland
¹⁰ Universitat de València, Spain
¹¹ Bill & Melinda Gates Foundation, Seattle, USA
¹² Pham Ngoc Thach hospital, Ho Chi Minh City, Vietnam
¹³ Monash University, Melbourne, Australia
¹⁴ Baker Institute, Melbourne, Australia and Cambridge University
¹⁵ National Institute of Diseases of the Chest and Hospital, Dhaka, Bangladesh
¹⁶ Sciensano, Belgian reference laboratory for M. tuberculosis
¹⁷ National University of Singapore, Singapore
¹⁸ University of Melbourne and Baker Institute, Melbourne, Australia
¹⁹ Trinity College Dublin, Ireland, and Irish Mycobacteria Reference Laboratory, Dublin, Ireland
²⁰ Careggi University Hospital, Florence, Italy
²¹ The University of Sydney, Australia
²² Republican Scientific and Practical Centre for Pulmonology and TB, Minsk, Belarus
²³ National Institute for Public Health and the Environment, Bilthoven, The Netherlands
²⁴ Leeds Teaching Hospital NHS Trust, Leeds, United Kingdom
²⁵ World Health Organization, Geneva
Author contributions
DAC, DMC, DWC, HH, SJH, NAI, NM, DM, SN, TEAP, CR, GS, PS, GT, ASW, TMW, DJW, ZY contributed to high-level CRyPTIC study design. ASL, TMW, DWC, TEAP, ASW, PWF, DAC designed the specifics of this study. MH, JK, ZI and PWF retrieved and processed genotypic data including kmers. PWF retrieved and processed phenotypic data. ASL, DAC, TMW, SK, YY, PWF, developed the machine learning system. All other authors contributed to the generation of data. ASL and TMW performed all the analysis. ASL and TMW wrote the manuscript with all authors offering feedback.

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**Competing Interest**

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