

MUBII-TB-DB: a database of mutations associated with antibiotic resistance in *Mycobacterium tuberculosis*.

Jean-Pierre Flandrois, Gérard Lina, Oana Dumitrescu

► **To cite this version:**

Jean-Pierre Flandrois, Gérard Lina, Oana Dumitrescu. MUBII-TB-DB: a database of mutations associated with antibiotic resistance in *Mycobacterium tuberculosis*. BMC Bioinformatics, BioMed Central, 2014, 15 (1), pp.107. <10.1186/1471-2105-15-107>. <inserm-00982132>

HAL Id: inserm-00982132

<http://www.hal.inserm.fr/inserm-00982132>

Submitted on 23 Apr 2014

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

DATABASE

Open Access

MUBII-TB-DB: a database of mutations associated with antibiotic resistance in *Mycobacterium tuberculosis*

Jean-Pierre Flandrois^{1*}, Gérard Lina^{2,3} and Oana Dumitrescu^{2,3}

Abstract

Background: Tuberculosis is an infectious bacterial disease caused by *Mycobacterium tuberculosis*. It remains a major health threat, killing over one million people every year worldwide. An early antibiotic therapy is the basis of the treatment, and the emergence and spread of multidrug and extensively drug-resistant mutant strains raise significant challenges. As these bacteria grow very slowly, drug resistance mutations are currently detected using molecular biology techniques. Resistance mutations are identified by sequencing the resistance-linked genes followed by a comparison with the literature data. The only online database is the TB Drug Resistance Mutation database (TBDRaM database); however, it requires mutation detection before use, and its interrogation is complex due to its loose syntax and grammar.

Description: The MUBII-TB-DB database is a simple, highly structured text-based database that contains a set of *Mycobacterium tuberculosis* mutations (DNA and proteins) occurring at seven loci: *rpoB*, *pncA*, *katG*; *mabA(fabG1)-inhA*, *gyrA*, *gyrB*, and *rrs*. Resistance mutation data were extracted after the systematic review of MEDLINE referenced publications before March 2013. MUBII analyzes the query sequence obtained by PCR-sequencing using two parallel strategies: i) a BLAST search against a set of previously reconstructed mutated sequences and ii) the alignment of the query sequences (DNA and its protein translation) with the wild-type sequences. The post-treatment includes the extraction of the aligned sequences together with their descriptors (position and nature of mutations). The whole procedure is performed using the internet. The results are graphs (alignments) and text (description of the mutation, therapeutic significance). The system is quick and easy to use, even for technicians without bioinformatics training.

Conclusion: MUBII-TB-DB is a structured database of the mutations occurring at seven loci of major therapeutic value in tuberculosis management. Moreover, the system provides interpretation of the mutations in biological and therapeutic terms and can evolve by the addition of newly described mutations. Its goal is to provide easy and comprehensive access through a client-server model over the Web to an up-to-date database of mutations that lead to the resistance of *M. tuberculosis* to antibiotics.

Keywords: Tuberculosis, Antibiotics, Mutation database, Sequence database, Web

Background

Tuberculosis (TB) is an infectious disease caused by a slow-growing bacterium, *Mycobacterium tuberculosis*, which has been linked to humans since the beginning of the early human expansion from east Africa [1]. Tuberculosis was the main cause of deaths in Western Europe

between the seventeenth century and the end of the nineteenth century [2]. It remains a major health threat, killing more than a million individuals every year worldwide. Although the WHO claims that its goal to halt and reverse the TB epidemic by 2015 has already been achieved and that the TB mortality rate has decreased by 41% since 1990, the global burden of TB remains. In 2011, there were an estimated 8.7 million new cases of TB, and 1.4 million people died from TB.

No fully effective vaccination is possible because the Bacille de Calmette et Guérin (BCG)-vaccine protection

* Correspondence: jp.flandrois@univ-lyon1.fr

¹Laboratoire de Biométrie et Biologie Évolutive, UMR CNRS 5558, Université Claude Bernard – Lyon 1, 43 bd. du 11 Novembre 1918, 69622 Villeurbanne Cedex, France

Full list of author information is available at the end of the article

against TB varies among populations [3] and provides only limited protection during childhood [4]. The limitation of the TB expansion is thus based on the improvement and generalization of diagnostic methods [5] and early antibiotic therapy. The WHO recommends the following combined therapy employing first-line antibiotics: rifampicin (or rifabutin), isoniazid, pyrazinamide and ethambutol for two months followed by rifampicin and isoniazid for four months [6].

However, irregular and low-dose treatment has led to the emergence and spread of multidrug-resistant (MDR) and extensively drug-resistant (XDR) *Mycobacterium tuberculosis* complex (MTBC) strains that present significant challenges in disease control [6-8]. The resistance increases in high MDR-TB burden countries, where the number of cases was estimated to be 60,000 worldwide in 2011. In these countries, the proportion of resistant strains varies from 5 to 19% of new TB cases and up to 50% for retreatment cases. This increase in resistance has led to the promotion of laboratory antimicrobial testing for all isolates [6]. In the case of antibiotic resistance, the therapy is conducted using second-line drugs, such as aminoglycosides or non-conventional antibiotics (such as fluoroquinolones).

The traditional phenotypic drug susceptibility test induces serious delays in the detection of resistance due to the extremely slow growth of *M. tuberculosis*, the answer being obtained in most cases two weeks after the isolation of the bacteria. Another drawback of the *in vitro* susceptibility testing is the inadequate detection of resistance to new drugs and to pyrazinamide [9-11]. The rapid diagnosis of drug resistance by molecular methods is essential to initiate effective antibiotic therapies and to prevent the transmission of drug-resistant strains [12].

M. tuberculosis acquires drug resistance primarily through mutations in specific genes [13,14]. The mutations associated with drug resistance occur in *rpoB* for rifampicin (RIF), in *katG* and the *mabA (fabG1)-inhA* operon for isoniazid (INH), in *pncA* for pyrazinamide (PZA), in *rrs* for amikacin (AMK) and in *gyrA* and *gyrB* for fluoroquinolones (FQs) [13,15-17].

Several commercial molecular kits are available to detect *M. tuberculosis* resistance using line-probe assays or real-time PCR, and these kits allow for the prediction of drug resistance in clinical specimens within one working day [9]. They can detect the mutations responsible for resistance to RIF alone (GeneXpert MTB/RIF (Cepheid), INNO-LiPA® Rif. TB (Innogenetics)), to RIF + INH (GenoType® MTBDRplus (Hain LifeScience GmbH)), or to FQs + AMK (GenoType® MTBDRsl (Hain LifeScience GmbH)), but their sensitivity varies, depending on the antibiotic [18]. These kits, unfortunately, detect only the most frequent mutations [18,19]. An alternative approach that allows for the exhaustive detection of

mutations consists of the PCR amplification and sequencing of resistance-linked genes [20,21] and, more experimentally, of complete genome sequencing [22]. It is, however, time consuming to compare the sequences with the reference genome and the mutation identity with the literature data. Recently, Sandgren *et al.* [23] established a comprehensive database that gathers information on the mutations associated with TB drug resistance and the frequency of the most common mutations associated with resistance to specific drugs. The TBDRaM database is a free online resource that allows for the molecular diagnosis of resistant TB after the processing of the amplified sequences by any method. Although very helpful, the TBDRaM database has not been updated since April 2010, and its usage remains time consuming because it demands prior processing of the TB nucleotide sequences to detect mutations and cannot be easily interrogated due to its relaxed grammatical conception.

Here, we describe MUBII-TB-DB, a database of the mutations of *M. tuberculosis* linked to resistance to first-line and second-line antibiotics that can be used to identify the mutations from a submitted sequence. This database and its related software, MUBII, have been developed to satisfy the need of clinical microbiology labs to easily analyze *M. tuberculosis* sequences and to link the results to a potential therapeutic failure. MUBII combines the use of reconstructed mutated gene sequences that can be searched by BLAST, aligned against the wild-type gene sequence, and compared with the mutation database. This concept can be easily adapted to the microbiological identification of other microbial mutations.

Construction and contents

Mutations database

Data

The database was constructed based on a systematic review of the literature, as described below. We focused on publications reporting an association between specific mutations in clinical isolates of *M. tuberculosis* and phenotypic resistance to INH, RIF, PZA, FQs, and AMK. The genes studied were *rpoB* for RIF; *inhA*, *katG*, and the promoter region of the *mabA (FabG1)-inhA* operon for INH; *pncA* for PZA; *gyrA* and *gyrB* for FQs; and *rrs* for AMK. As a starting point, we used the TBDRaM database because it constitutes a comprehensive resource on drug resistance mutations in *M. tuberculosis* based on studies published from January 1966 to December 2009 [23]. Additional publications on *M. tuberculosis* resistance mutations recorded in MEDLINE from December 2009 to March 2013 were included in the analysis (see *infra*). All publications reporting mutations, including the ones recorded in the TBDRaM database, were carefully reviewed for the consistency of the information about the mutated nucleotide and amino acid. We used

the codon numbering given by the annotation of the *M. tuberculosis* whole genome sequence published in [24].

Strategy of the systematic literature review

All studies reporting an association between specific mutations in clinical isolates of *M. tuberculosis* and phenotypic resistance to INH, RIF, PZA, FQs and AMK that were previously selected for the construction of the TBDRaM database have been included again in our survey. Moreover, we searched the MEDLINE database for similar works issued since the TBDRaM database release (Dec 2009 – Mar 2013). We also searched for additional references in the bibliographies of the reports and reviews. Only English language articles were considered. Combinations of the following search terms were used:

Tuberculosis AND (mutation OR mutations) AND (isoniazid OR rifampin OR fluoroquinolones OR amikacin OR pyrazinamide OR katg OR mba OR fabg1 OR inha OR rpoB OR rrs OR pncA OR gyrA OR gyrB).

Inclusion criteria

A study was included in the database under the following conditions: 1) it was a survey of clinical *M. tuberculosis* isolates; 2) drug sensitivity testing was performed on all isolates tested for mutations; and 3) the nucleotide or codon position and the nucleotide or amino acid change were given.

Data retrieval

We extracted and recorded information on the following: 1) the gene in which putative resistance mutations were found and the nucleotide and/or codon position of the mutation and 2) nucleotide and/or amino acid changes. Studies that did not allow the extraction of the above data or showed discrepancies in the wild-type sequences compared with published *M. tuberculosis* genomes sequences were excluded. The workflow of the literature review is summarized in Figure 1. The extracted mutations and original citations are supplied as Additional file 1.

Therapeutic relevance

Mutations with therapeutic relevance are tagged as “high-confidence” mutations in the MUBII-TB-DB database. High-confidence mutations have been previously defined by Sandgren *et al.* [23] as mutations associated with in vitro documented resistance reported by at least 10 publications based on the analysis of different sets of strains.

Reference sequences

The reference sequences of the resistance genes are presented in Table 1.

Database grammar

For each gene, there is a corresponding separate subset of the database, a text file with the description of a given mutation on each line. The line contains both the descriptions of the mutation (nucleotides and amino acids, if applicable), notes on the mutation (for instance, therapeutic relevance) and rules to apply.

The database grammar (examples are presented in Table 2) is as follows:

```
wildNPmutatedN ~ wildXPmutatedX[~Notes[~RULE::  
type[,action][~Remarks]]/~Remarks]
```

where P is the position on the nucleotide or protein chain; N is a nucleotide (or short sequence); and X is an amino-acid or a short peptide sequence. Notes are written freely and may contain alphanumeric characters and punctuation, except the tilde sign. The specific tag “Rules:” identifies the actions to apply to clearly differentiate the rules from the remarks. Rules modify the computed result, for example, to suppress the peptide sequence if a stop codon is created or to correct the result in the case of ambiguous positions of indels that occur within repeated features. Remarks are not used by the programs but are information, such as references and dates, linked to the database entry.

Other data are also recorded in the database: the position of the first codon of the nucleotide chain, the status of the final main product (DNA/protein), and the DNA sequence of the wild strain (the reference sequence). Therefore, each gene-mutation database is a series of flat files containing the descriptions of the mutations and a series of files containing the character sequences in FASTA format as well as the locations of the final gene products and non-coding zones.

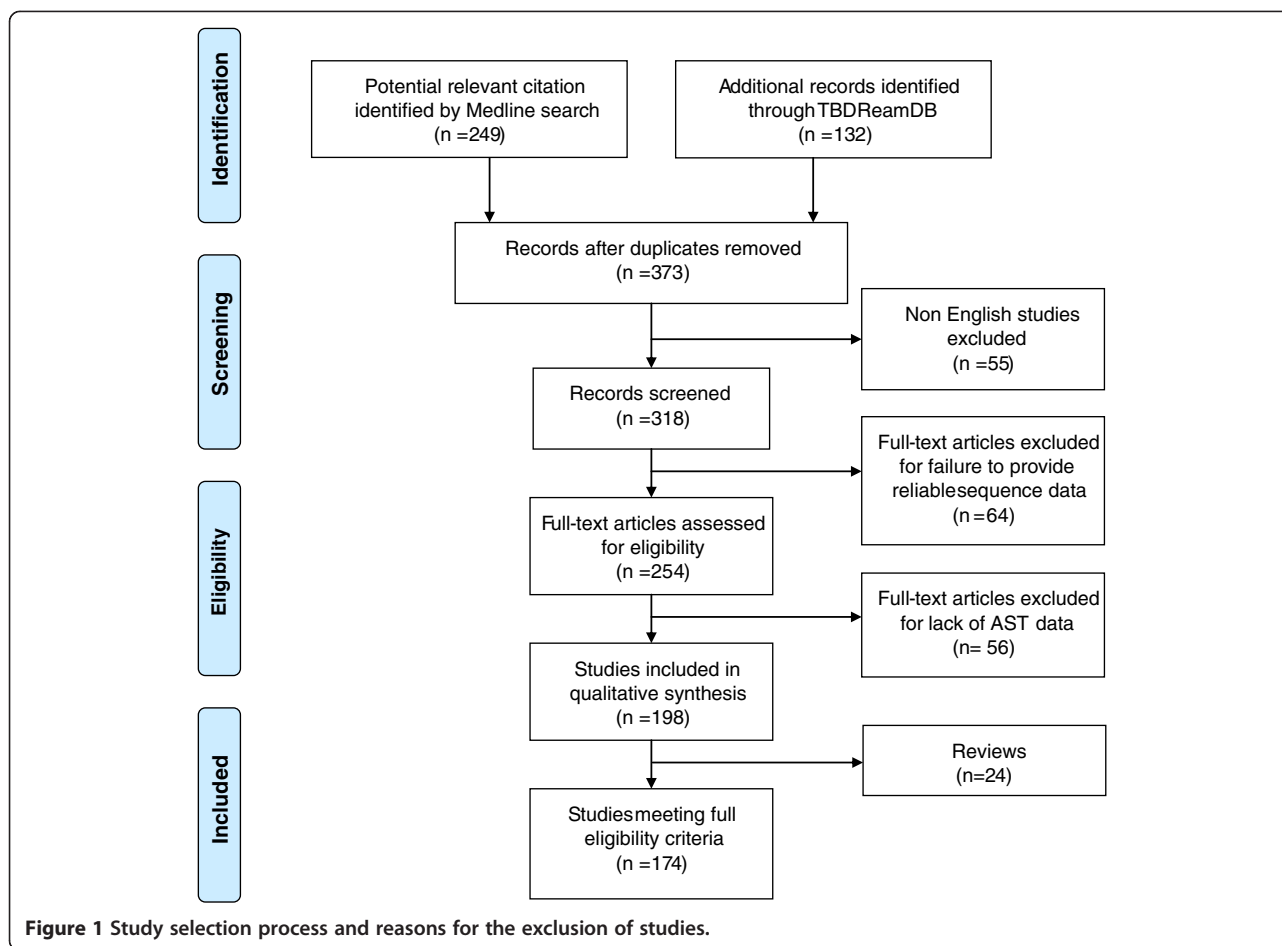
Implementation

MUBII is the analysis and interpretation engine and is entirely written in Python 2.7. The external programs that are called from Python are BLAST2seq [25], BLASTN, BLASTX [26] and the alignment tools MUSCLE [27] and MAFFT [28]. The routines transeq and showalign from EMBOSS [29] are also used. The scheme of the global organization is shown in Figure 2.

The Web interface exclusively uses CGI and HTML. Cascading Style Sheets are used to color the alignments. These decisions were made to achieve simplicity and reliability on every browser. The Web server runs under the APACHE MPM Prefork http server on a LINUX Debian server.

De novo constructed BLASTN database

Only a few mutated sequences are available in GenBank; therefore, the construction of a database containing the sequences of all the described mutations is necessary. A sequence database is built containing a unique mutation



event for each of the sequences. It contains the descriptions of the mutations present in MUBII-TB-DB, the reference sequence and the position of the promoter if required. The mutated sequences are written in the FASTA format at the nucleotide level with the description of the mutation as the descriptor. This database is rebuilt when changes occur within MUBII-TB-DB and is compiled as a BLAST database.

Table 1 List of the reference gene sequences of *Mycobacterium tuberculosis*

Gene	GenBank id.
<i>rpoB</i>	[AL123456] REGION: 759808 to 763325
<i>pncA</i> (including the promoter)	[AL123456] REGION: 2288681 to 2289280
promoter <i>inhA</i>	[AL123456] REGION: 1673325 to 1673439
<i>inhA</i>	[AL123456] REGION: 1674202 to 1675011
<i>katG</i>	[AL123456] REGION: 2153889 to 2156111 identical to [x68081.1]
<i>gyrA</i>	[AL123456] REGION: 7302 to 9818
<i>gyrB</i>	[AL123456] REGION: 5123 to 7267
<i>rrs</i>	[AL123456] REGION: 1471846 to 1473762 identical to [NR_102810]

DNA sequence analysis

The query sequence for a given gene is submitted in the FASTA format to the program through an HTML interface. A preliminary nucleotide BLAST2seq against the gene reference sequence is used to test if the query is given as a + direct string and, if not, computes its complementary strand. BLAST2seq does not allow for the identification of mutations that may occur in the extremities of the sequence.

Alignment to the reference and extraction of the core of the aligned sequences

The query sequence is aligned to the reference sequence by MAFFT. An algorithm identifies the core of the alignment by eliminating the trailing short sequences that are not perfectly aligned at the extremities. The core of the alignment is further used for mutation detection.

The alignment was initially produced using MUSCLE, but as the program tries to align the whole length of the reference sequence to a query that is often a partial sequence, the ends of the query were frequently assigned to positions near the ends of the reference. This problem was observed especially in the low quality-extremities of

Table 2 Example of a mutation database: a part of the *rpoB* mutation database for *Mycobacterium tuberculosis*

Mutation description	Explanation
A1291G ~ S431G ~ ~	Nucleotide chain: A at position 1291 is replaced by G; Protein chain: S at position 431 is replaced by G
+1300TTC ~ +434 F ~ ~	Insertion of TTC at position 1300 (F at position 434 of the protein)
CAG1306- ~ Q436- ~ ~	Deletion of CAG at position 1306 (deletion of Q at position 436 of the protein)
C1331T ~ T4441 ~ High Confidence ~ See mutDB2012	Example with a note and a remark
C1294T ~ Q432- ~ Rare ~ RULE::STOPCODON	Example with a note and a rule
CATGGACCAGAA1299- ~ MDQN434- ~ ~RULE::AMBIGUOUS=> N_ATGGACCAGAAC1300-	Example of a deletion occurring within a repeated zone. The position 1299 deduced from the nucleotide-level sequence alignment is corrected in the results

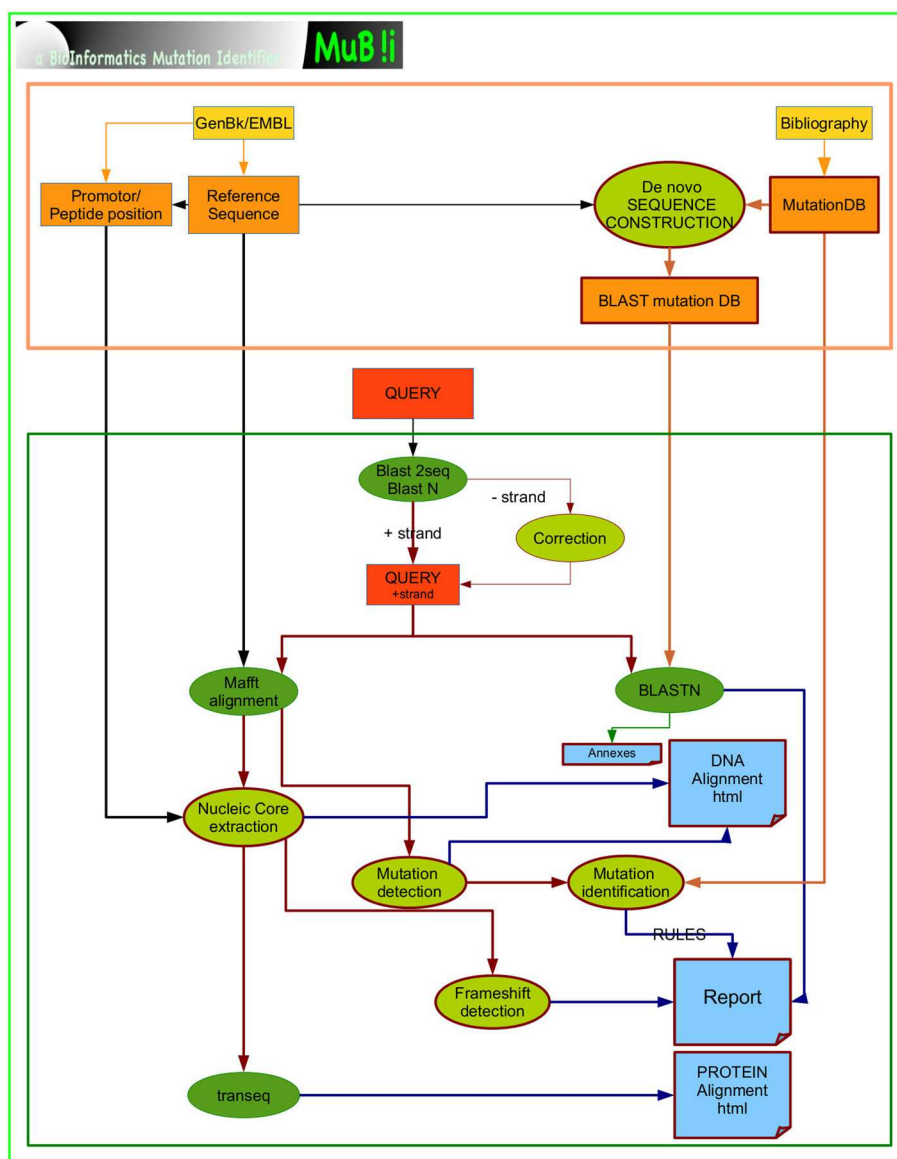


Figure 2 MUBli general organization. The MUBli-TB-DB database and the *de novo* constructed mutation database (orange frame) are built before the query session. The mutation database is used for the results interpretation and for the construction of the BLAST mutation database. Ovals indicate the use of external software (dark green) or of MUBli routines (pale green). The query analysis process (green frame) combines the BLAST result and the expertise of the alignment using the mutation database. Outputs are in blue.

the query and has been found to occur less frequently when using MAFFT.

Extraction of the mutations from the alignment and identification

For the core section of the alignment, a program extracts the non-matching sections and constructs a Python dictionary of the results. Each entry of this dictionary is compared with the entries of a mutation dictionary for the given gene constructed from the MUBII-TB-DB database. The resulting table contains a description of each point mutation, insertion or deletion and indicates its presence or absence in the database. In the case of deletion or insertion, the possibility of frameshift creation is also checked, and the result is added in an “alert” section. Moreover, if the mutation is known and modifies the encoded protein, this information is included. In the case of mutations usually identified by their positions in the *Escherichia coli* gene (some sections of the *rpoB* gene), the *E. coli* gene position is also computed. Finally, if the identified mutation requires interpretation, the result is corrected to fulfill the RULE:: indications. All results are saved in files in a format ready for inclusion in the HTML page.

BLASTN on a reconstructed mutated sequence database

A BLASTN of the query against the constructed mutated sequence database is performed, and the descriptors of the best matching sequences are added to the results. The whole BLASTN result is also saved.

DNA translation to protein

The protein translation of the core of the nucleotide alignment is obtained using transeq from the EMBOSS library.

Output page construction

An HTML output file is computed for the whole alignment (DNA and protein levels) that highlights mutations using colored tags. The information gathered in the various results files is returned (detected mutations and frameshifts, mutation identification, BLAST result on the reconstructed sequence database, position within the *M. tuberculosis* wild-type gene, and, if possible, position within the *E. coli* gene). The results, alerts and alignment are placed in iframes that enable horizontal browsing and searching to inspect sequences for mutations. A specific output adapted for printing is constructed. It provides information about the detected mutations along with alignments using the showalign EMBOSS routine.

Quality control

We have used the *in silico* mutation generation routine of MUBII to generate a mutated version of the *M. tuberculosis* H37Rv sequence. For every gene and every

mutation described in MUBII-TB-DB, we have constructed a sequence containing the required modification (e.g., base change, deletion, insertion). The mutated sequence thus obtained was then submitted to MUBII to verify the accuracy of the answer. All described mutations as well as hundreds of random mutations have been checked, and the MUBII results have been carefully analyzed. This procedure has identified uncertainties in mutation identification when an indel occurs near or within repeated features, especially in the *rpoB* gene. An interpretation RULE has been added to indicate such a situation and modify the answer accordingly. Following this extended quality control, both the MUBII-TB-DB database and the MUBII process have been validated for use.

Utility and discussion

Data input

The use of MUBII is straightforward, as the submission of the FASTA formatted sequence and the name of the corresponding gene are performed using a standard Web browser. It is also possible to use a test sequence for demonstration purposes or to verify the system.

Data analysis

The result appears in a new html page embedding the presentation of the DNA and protein alignments along with the sequence of the wild-type strain. The first section of the result pages concerns the nucleotide sequence (Figure 3). This section shows the DNA alignment of the query against the wild-type sequence and highlights any mutations. When mutations are detected, alerts are printed to note the position and the type of the mutation, its status (already described or unknown, therapeutic relevance), and situations such as frameshifts. At the end of this section, the name of the best matching sequence in the BLAST of the query against the reconstructed database of mutated sequences is shown. Access to the whole BLAST result is also possible. Because only a few strains of *M. tuberculosis* carry double mutations, the database contains only singly mutated sequences. The second section concerns the protein corresponding to the core nucleotide sequence. This section shows the peptide-level alignment of the query against the wild-type sequence and the position of mutations. The alignment is deduced from the nucleotide alignment and highlights the mutations and observed changes in the case of frameshift. In this last case, as the gaps are not introduced, the picture shows the actual changes observed in the query. This presentation is more informative, for the biologist, of the real changes that occur (Figure 4). When a stop-codon mutation is created, a specific alert is highlighted, and the shortened version of the protein is shown.



Figure 4 Protein alignment and detected mutations. A shows the detection of a mutation. B shows frameshift creation. C shows stop-codon creation. The whole alignment can be observed by horizontally scrolling the alignment window. These representations emphasize the effect of the mutation on the protein chain to provide solid understanding in terms of establishing therapy and monitoring.

Conclusions

The growing incidence and spread of antibiotic-resistant *M. tuberculosis* has led to efforts to identify the mutations in clinical isolates. The mutations linked to therapy failure are a particularly serious concern for physicians managing TB patients. The reference method is to sequence the gene of interest and then to detect and identify possible mutations. This process is time consuming, as there is no mutation database directly coupled to an analysis tool that allows for the submission of a sequence, its comparison to the database and the immediate analysis of the results. MUBII-TB-DB is a structured database collecting the description and information on modifications to seven loci of major therapeutic value in tuberculosis. The simple structure and grammar of MUBII-TB-DB contribute to the evolving capacity of the database, which can be easily updated with new entries (both mutations and resistance genes). MUBII-TB-DB is adapted to MUBII, a query and interpretation engine that is interrogated via a website. The sequence of a gene implicated in TB antibiotic resistance is analyzed simultaneously as a DNA string by BLAST against a *de novo* constructed sequence database and by an alignment process for the DNA and protein sequences. The results are graphs (alignments) and text (description of the mutation, previous description, therapeutic signification). MUBII-TB-DB is fast and easy to use, even by technicians without bioinformatics training. The system provides access to an interpretation in biological and therapeutic terms. It enables the automation of repetitive and somewhat technically challenging tasks in the tuberculosis laboratory. Its interest is to provide easy and comprehensive access to an up-to-date antibiotic resistance database concerning *M. tuberculosis*. Moreover, our algorithm is applicable to any mutation database and to any microorganism, making MUBII a useful tool for the surveillance and control of multidrug-resistant microorganisms.

Availability

The database is available at <http://umr5558-bibiserv.univ-lyon1.fr/mubii/mubii-select.cgi>.

Additional file

Additional file 1: Mutation database references spreadsheet.

Abbreviations

TB: Tuberculosis; WHO: World Health Organization; MDR: Multidrug Resistant; XDR: Extensively Drug Resistant; BCG: The vaccine using the attenuated "Bacille de Calmette et Guérin"; MTBC: *Mycobacterium tuberculosis* complex; RIF: Rifampicin; INH: Isoniazid; PZA: Pyrazinamide; AMK: Amikacin; FQs: Fluoroquinolones.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

JPF and GL conceived the project; JPF designed the concept and implemented the system and website; OD implemented the data extraction and the resulting database; GL controlled the database; GL, OD, and JPF discussed the system during its development; OD supervised the quality assurance and the tests; GL and OD supervised the tests; JPF, OD and GL wrote the manuscript. All authors have read and approved the final manuscript.

Acknowledgements

We thank Manolo Gouy for useful discussions during the elaboration of the manuscript.

Author details

¹Laboratoire de Biométrie et Biologie Évolutive, UMR CNRS 5558, Université Claude Bernard – Lyon 1, 43 bd. du 11 Novembre 1918, 69622 Villeurbanne Cedex, France. ²CIRI, International Center for Infectiology Research, LabEx Ecofect, Université Lyon1; Inserm, U1111, Ecole Normale Supérieure de Lyon; CNRS, UMR5308, 69000 Lyon, France. ³Hospices Civils de Lyon, 69002 Lyon, France.

Received: 9 December 2013 Accepted: 7 April 2014

Published: 14 April 2014

References

- Comas I, Coscolla M, Luo T, Borrell S, Holt KE, Kato-Maeda M, Parkhill J, Malla B, Berg S, Thwaites G, Yeboah-Manu D, Bothamley G, Mei J, Wei L, Bentley S, Harris SR, Niemann S, Diel R, Aseffa A, Gao Q, Young D, Gagneux S: **Out-of-Africa migration and Neolithic coexpansion of *Mycobacterium tuberculosis* with modern humans.** *Nat Genet* 2013, **45**(10):1176–1182.
- Wilson LG: **Commentary: Medicine, population, and tuberculosis.** *Int J Epidemiol* 2005, **34**:521–524.
- Abubakar I, Pimpin L, Ariti C, Beynon R, Mangtani P, Sterne J, Fine P, Smith P, Lipman M, Elliman D, Watson J, Drumright L, Whiting P, Wynnycky E, Rodrigues L: **Systematic review and meta-analysis of the current evidence on the duration of protection by bacillus Calmette-Guérin vaccination against tuberculosis.** *Health Technol Assess* 2013, **17**:1–372.
- Colditz GA: **Efficacy of BCG Vaccine in the Prevention of Tuberculosis Meta-analysis of the Published Literature.** *JAMA* 1994, **271**:698.

5. Boehme CC, Saacks S, O'Brien RJ: **The changing landscape of diagnostic services for tuberculosis.** *Semin Respir Crit Care Med* 2013, **34**:17–31.
6. World Health Organization: *WHO Global tuberculosis report 2012.* Geneva, Switzerland: World Health Organization; 2012.
7. Sotgiu G, Centis R, D'Ambrosio L, Tadolini M, Castiglia P, Migliori GB: **Do we need a new Fleming époque: The nightmare of drug-resistant tuberculosis.** *Int J Mycobacteriology* 2013, **2**:123–125.
8. Lynch JB: **Multidrug-resistant Tuberculosis.** *Med Clin North Am* 2013, **97**(4):553–579. ix–x.
9. Wilson ML: **Rapid diagnosis of *Mycobacterium tuberculosis* infection and drug susceptibility testing.** *Arch Pathol Lab Med* 2013, **137**:812–819.
10. Horne DJ, Pinto LM, Arentz M, Lin S-YG, Desmond E, Flores LL, Steingart KR, Minion J: **Diagnostic accuracy and reproducibility of WHO-endorsed phenotypic drug susceptibility testing methods for first-line and second-line antituberculosis drugs.** *J Clin Microbiol* 2013, **51**:393–401.
11. Piersimoni C, Mustazzolu A, Giannoni F, Bornigia S, Gherardi G, Fattorini L: **Prevention of false resistance results obtained in testing the susceptibility of *Mycobacterium tuberculosis* to pyrazinamide with the Bactec MGIT 960 system using a reduced inoculum.** *J Clin Microbiol* 2013, **51**:291–294.
12. Falzon D, Jaramillo E, Schünemann HJ, Arentz M, Bauer M, Bayona J, Blanc L, Caminero JA, Daley CL, Duncombe C, Fitzpatrick C, Gebhard A, Getahun H, Henkens M, Holtz TH, Keravec J, Keshavjee S, Khan AJ, Kulier R, Leimane V, Lienhardt C, Lu C, Mariandyshev A, Migliori GB, Mirzayev F, Mitnick CD, Nunn P, Nwagboniwe G, Oxlade O, Palmero D, et al: **WHO guidelines for the programmatic management of drug-resistant tuberculosis: 2011 update.** *Eur Respir J* 2011, **38**:516–528.
13. Heym B, Cole ST: **Multidrug resistance in *Mycobacterium*.** *Int J Antimicrob Agents* 1997, **8**:61–70.
14. Warner DF, Mizrahi V: **Complex genetics of drug resistance in *Mycobacterium tuberculosis*.** *Nat Genet* 2013, **45**:1107–1108.
15. Musser JM: **Antimicrobial agent resistance in mycobacteria: molecular genetic insights . Antimicrobial Agent Resistance in Mycobacteria: Molecular Genetic Insights.** *Clin Microbiol* 1995, **8**:496–514.
16. Almeida Da Silva PEA, Palomino JC: **Molecular basis and mechanisms of drug resistance in *Mycobacterium tuberculosis*: classical and new drugs.** *J Antimicrob Chemother* 2011, **66**:1417–1430.
17. Zhang Y, Yew WW: **STATE OF THE ART Mechanisms of drug resistance in *Mycobacterium tuberculosis*.** *Int J Tuberc Lung Dis* 2009, **13**:1320–1330.
18. Kontsevaya I, Ignatyeva O, Nikolayevskyy V, Balabanova Y, Kovalyov A, Kritsky A, Matskevich O, Drobniowski F: **Diagnostic accuracy of the genotype MTBDRsl assay for rapid diagnosis of extensively drug-resistant tuberculosis in HIV-coinfected patients.** *J Clin Microbiol* 2013, **51**:243–248.
19. Alonso M, Palacios JJ, Herranz M, Penedo A, Menéndez A, Bouza E, García de Viedma D: **Isolation of *Mycobacterium tuberculosis* strains with a silent mutation in *rpoB* leading to potential misassignment of resistance category.** *J Clin Microbiol* 2011, **49**:2688–2690.
20. Campbell PJ, Morlock GP, Sikes RD, Dalton TL, Metchock B, Starks AM, Hooks DP, Cowan LS, Plikaytis BB, Posey JE: **Molecular detection of mutations associated with first- and second-line drug resistance compared with conventional drug susceptibility testing of *Mycobacterium tuberculosis*.** *Antimicrob Agents Chemother* 2011, **55**:2032–2041.
21. Feuerriegel S, Oberhauser B, George AG, Dafaie F, Richter E, Rüscher-Gerdes S, Niemann S: **Sequence analysis for detection of first-line drug resistance in *Mycobacterium tuberculosis* strains from a high-incidence setting.** *BMC Microbiol* 2012, **12**:90.
22. Zhang H, Li D, Zhao L, Fleming J, Lin N, Wang T, Liu Z, Li C, Galwey N, Deng J, Zhou Y, Zhu Y, Gao Y, Wang T, Wang S, Huang Y, Wang M, Zhong Q, Zhou L, Chen T, Zhou J, Yang R, Zhu G, Hang H, Zhang J, Li F, Wan K, Wang J, Zhang X-E, Bi L: **Genome sequencing of 161 *Mycobacterium tuberculosis* isolates from China identifies genes and intergenic regions associated with drug resistance.** *Nat Genet* 2013, **45**:1255–1260.
23. Sandgren A, Strong M, Muthukrishnan P, Weiner BK, Church GM, Murray MB: **Tuberculosis drug resistance mutation database.** *PLoS Med* 2009, **6**:e2.
24. Cole S, Brosch R, Parkhill J, Garnier T: **Deciphering the biology of *Mycobacterium tuberculosis* from the complete genome sequence.** *Nature* 1998, **393**:537–544.
25. Tatusova TA, Madden TL: **BLAST 2 Sequences, a new tool for comparing protein and nucleotide sequences.** *FEMS Microbiol Lett* 1999, **174**:247–250.
26. Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ: **Basic local alignment search tool.** *J Mol Biol* 1990, **215**:403–410.
27. Edgar RC: **MUSCLE: multiple sequence alignment with high accuracy and high throughput.** *Nucleic Acids Res* 2004, **32**:1792–1797.
28. Katoh K, Toh H: **Recent developments in the MAFFT multiple sequence alignment program.** *Brief Bioinform* 2008, **9**:286–298.
29. Rice P, Longden I, Bleasby A: **EMBOSS: The European Molecular Biology Open Software Suite.** *Trends Genet* 2000, **16**:2–3.

doi:10.1186/1471-2105-15-107

Cite this article as: Flandrois et al.: MUBII-TB-DB: a database of mutations associated with antibiotic resistance in *Mycobacterium tuberculosis*. *BMC Bioinformatics* 2014 **15**:107.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at
www.biomedcentral.com/submit

