MOLECULAR CHARACTERISATION AND DIAGNOSIS

Simultaneous identification and typing of multi-drug-resistant *Mycobacterium tuberculosis* isolates by analysis of *pncA* and *rpoB*

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In *Mycobacterium tuberculosis* there is a strong correlation between in-vitro resistance to rifampicin (RIF) and pyrazinamide (PZA) and mutations in *rpoB* and *pncA*, respectively. Approximately 50 mutations associated with resistance have been reported for *rpoB* and 70 for *pncA*, and, theoretically, many more are possible. Therefore, the identification of *rpoB* and *pncA* mutations in *M. tuberculosis* might be used for the simultaneous determination of resistance and for typing multi-drug-resistant (MDR) strains during possible outbreaks. The present study examined four sensitive and six MDR isolates of *M. tuberculosis* from Turkey and eight isolates from a nosocomial MDR tuberculosis (TB) outbreak in the UK. Gene mutations were identified by the Innogenetics LIPA *rpoB* assay or automated sequencing, or both. All the sensitive isolates had *rpoB* and *pncA* wild-type genotypes, whereas all the RIF- and PZA-resistant isolates had *rpoB* and *pncA* mutations. All four mutations seen in *rpoB*, but none of the six in *pncA*, had been reported previously. The *rpoB* and *pncA* mutations seen in the Turkish isolates defined six distinct genotypes amongst the six MDR isolates, while standard IS6110 typing discriminated only four. All isolates from the single strain MDR-TB outbreak had identical genotypes. Rapid genotyping was performed on the sputum from a patient who presented 2 years after the initial MDR-TB outbreak and this showed *rpoB* and *pncA* genotypes identical to the other outbreak isolates. This result was available within 36 h. The analysis of *rpoB* and *pncA* is a rapid and practical means of simultaneously identifying and typing MDR isolates of *M. tuberculosis*.

Introduction

A combination of four drugs, isoniazid (INH), rifampicin (RIF), pyrazinamide (PZA) and ethambutol (EMB), with or without streptomycin (STR), is recommended for the standard or first-line treatment of tuberculosis (TB) [1]. Antibiotic resistance has emerged to each of these agents, as has multi-drug resistance (MDR-TB) where the isolate is resistant to at least INH and RIF [2]. The rapid identification of MDR-TB is essential for proper patient management [1, 3].

In *Mycobacterium tuberculosis*, resistance to these first-line drugs is usually associated with mutations in specific genes. EMB resistance is associated with mutations in *embB* in 47–69% of isolates [4, 5]. Resistance to INH has been associated with a limited number of mutations in *katG* and *inhA* [6–8] in up to 75% of cases [8]. Mutations associated with resistance to RIF are located within a 69-bp region of *rpoB* [6] in 94–98% of isolates [9–14]. More recently, mutations within *pncA* have been seen in isolates resistant to PZA [15]. *PncA* mutations, which are distributed throughout the gene, have been reported in 72–100% of PZA-resistant isolates [15–18]. The high correlation between mutations in *rpoB* and *pncA* and phenotypic drug resistance makes these genes attractive targets for the rapid determination of antibiotic susceptibilities of *M. tuberculosis*. The detection of mutations within *rpoB* may be used for the presumptive identification of MDR-TB, as MDR strains are usually rifampicin-resistant. The relatively small size of *pncA* and the limited region of interest within *rpoB* make the routine analysis of these loci practical. Furthermore, the wide

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range of reported and potential variations within these genes suggests that rpoB and pncA genotyping might also be used as a rapid method of typing MDR-TB isolates for epidemiological purposes.

To test whether the rapid genetic analysis of rpoB and pncA can be used for the simultaneous detection of resistance and for typing resistant isolates, the present study analysed these genes in 10 epidemiologically unconnected M. tuberculosis isolates from Istanbul, Turkey, where both primary and acquired resistance is common [19], and isolates from a single strain nosocomial outbreak of MDR-TB in London. In-vitro susceptibilities to RIF and PZA were compared with those predicted by sequence mutations, and the genotypes produced by this method were compared with standard IS6110 typing.

Materials and methods

Organisms

Five PZA-resistant and five PZA-sensitive distinct clinical isolates of M. tuberculosis complex were obtained from an Istanbul laboratory. Another seven isolates were collected from an outbreak of MDR-TB [20]; four serial isolates were collected over an 18-month period from one of these patients and one further isolate was from an epidemiologically related patient who presented 24 months later. A sputum specimen from this final patient was analysed directly after being concentrated [21] and the pellet washed. Organisms were identified by standard methods [22]. INH, RIF and PZA susceptibilities were determined by the Bectec radiometric-based culture method (Becton-Dickinson, Oxford) according to the manufacturer’s standard protocols. PZA susceptibility was also determined by detection of pyrazinamide activity [23].

Extraction of DNA

Cells cultured on Lowenstein-Jensen slopes were suspended in a buffer containing 10 mM Tris, 1 mM EDTA and Triton X100 1% at pH 8.0. Suspensions were heated to 100°C for 30 min, vortex mixed and centrifuged briefly at 13 500 rpm.

PCR and sequence analysis of pncA

A PCR product of 599 bp was amplified with primers pncA1u (5'-GCCAGCATTGTAAAAACGACGGCAGTGGAATCCAGATCAGCGGCGCCGGATCGTATGATCATCGTCCG) and pncA2r (5'-GACCAGGAAAACAGCTATGACCCGAAATTCTGTTCACGGAGGCTGCAAACAACTCG), which were designed by using a published pncA sequence (GenBank MTU59967) and adding universal and reverse primer sequences to the 5’ ends to facilitate sequencing. The amplification product was sequenced with 5'-fluorescein-labelled universal and reverse primers (Pharmacia, St Albans), a Thermo-sequenase cycle sequencing kit (Amersham, Little Chalfont) and an ALF DNA sequencer (Pharmacia). The sequence of each isolate was compared with that of the PZA-sensitive wild-type (WT) sequence (GenBank MTU59967). One specimen in this study was analysed by a nested PCR, in which primers P1 (GTCGCTATGCTGCGGACG) and P6 (GCT TTGCGGAGGCGCTTCCA) [15] were used in the additional outer PCR.

Analysis of rpoB

An 80-nucleotide base region of rpoB was analysed and compared with the WT genotype (GenBank L27989) with the Inno-LiPA Rif TB kit (Innogenetics, Zwijnaarde, Belgium) following the manufacturer’s protocol. Briefly, this consisted of a nested PCR with biotin-labelled primers; the resultant ampiclon was hybridised to an array of oligonucleotides immobilised on a nitrocellulose membrane. Hybrids containing biotin were visualised colorimetrically. Hybridisation with the five Innogenetics WT oligonucleotides (S1–S5) indicated that the rpoB was wild-type and the organism was sensitive to RIF; failure to hybridise with one or more of these oligonucleotides indicated a mutant rpoB and, therefore, resistance to RIF was deduced. Four commonly seen specific mutations were identified by hybridisation to one of four mutant oligonucleotides, R1–R4. Other mutants were identified by amplification of a 260-bp region of rpoB with the primers TR1 (5’-TACCGGCGGAGTCTGATCC) and TR2 (5’-TACGGCGGATCGGAAACC) [14] and sequencing the product with 5'-fluorescein-labelled TR1 and TR2 in a Thermo-sequenase cycle sequencing kit and an ALF DNA sequencer, as before. The sequence of each isolate was compared with the RIF-sensitive WT sequence (GenBank L27989).

DNA typing of M. tuberculosis strains

DNA was extracted for restriction fragment-length polymorphism (RFLP) analysis with proteinase K (Boehringer-Mannheim, Lewes) and lysozyme (Sigma), followed by phenol/chloroform extraction. Restriction digests were prepared according to standardised methodology [24] with the restriction endonuclease PvuII. Fragments were separated in an agarose 1.5% gel and capillary-blotted on to nylon membrane. A digoxigenin-labelled IS6110 probe was generated by PCR with 0.4 µm primer 41 and primer 43 [25], 40 µM dNTP and 8 µM DIG-DUTP. Hybridisation was detected as described previously [26].

Results

pncA

The five phenotypically PZA-sensitive Turkish isolates were all genotypically WT and the five phenotypically PZA-resistant isolates were all genotypically mutant.
All five resistant isolates had different \textit{pncA} mutations, as shown in Table 1. Isolate 985040 contained two mutations, one resulting in an amino-acid change when translated and a silent mutation. All other mutations in these Turkish isolates resulted in amino-acid changes when translated. All eight isolates of the London PZA-resistant outbreak strain and the four serial isolates had the same \textit{pncA} mutation, which was different from those seen in the Turkish isolates (Table 1).

\textbf{rpoB}

The four phenotypically RIF-sensitive Turkish isolates had WT \textit{rpoB} genes. Mutations were detected in \textit{rpoB} in all six Turkish Rif-resistant isolates. Isolate 1028448 was initially designated phenotypically sensitive to RIF but was shown to contain a mutation. On re-testing, this isolate was found to be phenotypically resistant. Four different \textit{rpoB} mutations were seen amongst these six isolates as shown in Table 1. Mutations in isolates 1032166, 985040, 1017379 and 1015307 were detected with the LiPA assay by hybridisation to R5, R2, R5 and R2 respectively. Mutations in isolates 1028448 and 992048 were determined by sequencing. Isolate 1015307 was shown to contain a mixed population of \textit{M. tuberculosis}: a WT and a mutant. All eight of the London Rif-resistant outbreak isolates and the serial isolates contained the same mutation (Table 1).

\textbf{IS6110 typing}

The IS6110-probed restriction patterns for the Turkish isolates investigated in this study are shown in Fig. 1. The four fully sensitive Turkish isolates each had different IS6110 restriction patterns, demonstrating that these WT strains were not linked. Two of the Turkish MDR-TB isolates, 992048 and 1032166, had similar IS6110 typing patterns and resembled that of the WT isolate 1010968. The more intense signal seen in the profile of isolate 1032166 showed an additional fragment not visible in isolates 1010968 and 992048. The other three Turkish MDR-TB isolates had different IS6110 restriction patterns. All the outbreak isolates were indistinguishable by IS6110 typing as previously reported [20] and gave a five band pattern.

\textbf{rpoB/\textit{pncA} genotyping}

The nucleotide variations seen within the \textit{rpoB} and \textit{pncA} sequences of the Rif- and PZA-resistant isolates could be used to genotype them. In this study, all the five Turkish MDR-TB isolates had different \textit{rpoB} or \textit{pncA} sequences, or both, including the two MDR-TB isolates with similar IS6110 patterns. Thus \textit{rpoB}/\textit{pncA} genotyping discriminated five genotypes amongst the five MDR-TB isolates compared with the four seen with IS6110 typing (Fig. 1). All eight of the London MDR-TB outbreak isolates contained the same

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|}
\hline
Isolate no. & \textbf{rpoB} genotype & \textbf{pncA} genotype & Wt/Aux. \textbf{RIF} & Amo acid. \textbf{HIN} \\
\hline
985040 & 2406→T & 2406→T & 2406→T & 2406→T \\
1017379 & 2406→T & 2406→T & 2406→T & 2406→T \\
1015307 & 2406→T & 2406→T & 2406→T & 2406→T \\
1010968 & 2406→T & 2406→T & 2406→T & 2406→T \\
992048 & 2406→T & 2406→T & 2406→T & 2406→T \\
1028448 & 2406→T & 2406→T & 2406→T & 2406→T \\
93041 & 2406→T & 2406→T & 2406→T & 2406→T \\
1032166 & 2406→T & 2406→T & 2406→T & 2406→T \\
\hline
\end{tabular}
\caption{Phenotype and genotype of \textit{rpoB} and \textit{pncA} from 10 Turkish \textit{M. tuberculosis} isolates (1-10) and a representative isolate from a London outbreak of MDR-TB (STI1-8). (All eight London outbreak isolates gave identical results.)}
\end{table}
**roB/pncA** mutation combination, as did the serial isolates.

### Discussion

In this study, the Turkish and UK MDR *M. tuberculosis* isolates contained mutations in *roB* and *pncA* that were associated with phenotypic resistance to Rif and PZA, respectively, whereas none of the RIF- or PZA-sensitive isolates examined contained mutations. These results confirm those of other workers reported from other geographical locations [9–18].

The determination of genotype to predict phenotypic resistance to Rif and PZA is much more rapid than traditional culture methods, taking hours rather than weeks. Standard methods of pyrazinamide susceptibility testing can be unreliable [27,28] and are usefully supplemented by testing for pyrazinamidase production [29]. However, *pncA* genotyping can be performed several weeks earlier than pyrazinamidase production, as it can be done directly from a primary isolation slope and is probably more reliable.

Previous studies have described 71 different mutations in *pncA* associated with PZA resistance [15–18]. These consisted of 55 substitutions, 12 deletions and six insertions. Two previously described *pncA* mutations, consisting of a base change at the −11 position [18] and a deletion of 1–11 [15], would not be detected by the methodology described in this report, which uses primers based on the first and last 20 bases of the *pncA* coding region. None of the six *pncA* missense mutations seen in the present study, consisting of five substitutions and one deletion, have been described previously. One of the resistant isolates with a missense mutation also contained a silent *pncA* mutation. A silent mutation in *roB* has been seen before [30], but has not been reported previously in *pncA*. The existence of silent mutations demonstrates the value of identifying all nucleotide base changes in order to predict amino-acid sequence and hence antibiotic susceptibility.

The five Turkish PZA-resistant isolates in the present study had five different *pncA* genotypes, but only four IS6110 types, suggesting that, for resistant isolates, *pncA* typing may be more useful for discriminating potentially related strains than IS6110 typing. Only four of the approximately 70 *pncA* genotypes reported so far have been seen in more than one geographical location, and each of these occurred in two locations only [15–18]. There are 1235 possible single base mutations in *pncA* that can result in an amino-acid change or a stop codon. There are even more possible variants if the less common mutations are included, such as multiple base changes, deletions, insertions and mutations outside the *pncA* coding region. If these mutations are stable, the characterisation of *pncA* could provide a powerful tool for studying the epidemiology of MDR-TB.

Fifty mutations have been reported in *roB* in isolates from various geographical locations [9–14]. Some mutations occur much more frequently than others. For example S53L, which was seen in two isolates in the present study, is common [9–14], whereas N518H has been reported only once [11]. The reason for this variation in frequency of mutant genotypes is not clear. All four mutations in *roB* seen in this study have been reported previously. One of the cultures yielded two *roB* genotypes: the mutant 2386A→T and a WT. It is thought that resistant *M. tuberculosis* cells may appear in WT populations by spontaneous mutation; in the presence of Rif these cells are selected, shifting the population from predominantly WT to a predominantly resistant phenotype. This mixed culture may have been from a patient where this change was taking place.

The six Turkish Rif-resistant isolates in the present study could be separated into four *roB* genotypes and five IS6110 types. Although, in this sample, IS6110
typing gave a greater discrimination than the rpoB genotypes, the two isolates with similar IS6110 types had different rpoB genotypes.

When both pncA and rpoB sequence analysis were used to type the six Turkish MDR-TB isolates, six genotypes were discriminated, compared with only five by IS6110 typing. Although the present study examined only a small number of resistant strains, the ability of rpoB/pncA typing to distinguish all of them is striking. Furthermore, the large number of reported and theoretical mutations associated with resistance suggests that analysis of these genes would be useful for the epidemiology of resistant tuberculosis.

The rpoB/pncA genotype was identical in isolates from the eight epidemiologically linked patients [20], as were serial isolates obtained from one patient over an 18-month period. Also, one of the cases was identified as being positive for MDR-TB 2 years after the initial outbreak. It has been shown that IS6110 probing is a stable typing tool [31]. These results suggest that the genotyping used in the present study is also stable. This is shown during patient-to-patient transmission (the outbreak isolates), during infection (the isolate cultured subsequent to the main outbreak) and during active disease (the serial isolates).

The use of these gene mutations for the simultaneous detection of antimicrobial resistance and for rapid typing is particularly useful for determining whether two or more hospital isolates of MDR-TB represent an outbreak, as described above. In this setting, this genotyping technique may be more useful than IS6110 typing, because it is rapid, the interpretation of the results is simple, the method is highly discriminatory and it can be performed directly from specimens without waiting for culture. Of course, rpoB/pncA typing does not have the universal application of IS6110 typing, as it can be applied only to resistant isolates of M. tuberculosis, different strains may develop the same rpoB/pncA mutations, and the technique cannot identify relatedness between sensitive and resistant isolates of the same strain.

The characterisation of rpoB and pncA provides a practical and rapid prediction of M. tuberculosis susceptibility to RIF and PZA, and can be used simultaneously for outbreak typing of resistant strains. An obvious limitation of this method is that automated sequencing is required for the analysis of pncA and for mutations in rpoB for which there are no LIPa probes. However, for laboratories with the necessary equipment and expertise, analysis of rpoB and pncA can be done routinely and rapidly on selected specimens and isolates. Simpler methods for detecting genotype variations in isolates or directly in clinical specimens would be powerful clinical tools for the diagnosis and management of MDR-TB.

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References

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