

Fluoroquinolone resistance and mutational profile of gyrA gene in pulmonary MDR tuberculosis patients

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Research article

Keywords: Mycobacterium tuberculosis; MDR and Pre-XDR TB; fluoroquinolone resistance; MTBDRsl assay.

Posted Date: February 5th, 2020

DOI: <https://doi.org/10.21203/rs.2.15811/v3>

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Version of Record: A version of this preprint was published at BMC Pulmonary Medicine on May 11th, 2020. See the published version at <https://doi.org/10.1186/s12890-020-1172-4>.

Abstract

Background Flouroquinolones (FQs) are the potential drugs that inhibit DNA synthesis and used in the treatment of MDR-TB and anti-TB short term regimens. In recent year's high proportion of flouroquinolone (FQs) resistance in *Mycobacterium tuberculosis* isolates has been observed. The development of FQs resistance among multidrug resistant TB (Pre-XDR TB) negatively impact patient treatment outcome and is a serious threat to control TB. Methods A total of 562 samples were included in the study from patients with pulmonary TB which had been on anti-tuberculosis therapy. MTBDRsl assay was performed for molecular detection of mutations. Sequence analysis was performed for characterization and mutational profiling of FQ resistant isolates. Results FQs resistance was observed in 104 (18.5%) samples and most of them were previously treated and treatment failure cases. A total of 102 isolates had mutations in *gyrA* gene while *gyrB* gene mutations were observed in only two isolates. Mutational analysis showed that the mutations mostly alter protein at codon 94 (D94G) (represents the replacement of aspartic acid with glycine) and 90 (A90V) (substitution of alanine with valine). In MDR and treatment failure cases, the FQs-R was most commonly associated with D94G mutation. Whereas, a high proportion of A90V mutation was observed in MTB isolates which were newly diagnosed. Conclusion The findings suggest that the genotypic studies for FQs resistance should be carried out at the time of initial diagnosis, before starting treatment, to rule out all type of mutations for its potential use in treatment to control the resistance.

Background

Pakistan is among the thirty high Tuberculosis (TB) burden countries where complete elimination of TB is, unfortunately, a distant reality. TB is ranked in ninth number with leading cause of mortality worldwide where, about one third of the world's population is infected with latent infection of TB [1]. According to the World health organization (WHO) 2018 annual report, the incidence rate of TB in Pakistan is 267/100,000 population where mortality rate is reported as 27/100,000 (excludes HIV co-infection cases) [2]. There is significant number of population where TB remains undiagnosed and untreated.

The phenotypic resistance to flouroquinolones (FQs) is associated with mutation in *gyrA* and *gyrB* gene in quinolone resistance-determining region (QRDR), a type II DNA topoisomerase, which target DNA gyrase. Mutations in the DNA *gyr* subunit A confer the high level resistance whereas other confers low level resistance to *gyr* subunit B [3]. FQs have long been using as anti-tuberculosis drugs and their wide spread use has led to the development of resistance in clinical isolates of *Mycobacterium tuberculosis* (*Mtb*). During treatment of TB, MDR patients can develop resistance against flouroquinolones. The development of FQs-R in these patients is a risk factor as additional resistance to this drug can aid in transition of these patients from MDR to pre-XDR or even they can become extensively drug resistant with further resistance to at least one injectable second line drug [4,5].

Geographic differences exist in the frequency of *gyrA* mutations. The understanding of frequency and geographic distribution of the FQs mutation is important in order to maximize its sensitivity and

specificity. Mutations in codon 88-94 of QRDR of *gyrA* gene appear most commonly particularly in codons 88, 90, 91, and 94. Mutations in the region of codon 500 and 538 of *gyrB* gene are most often associated with FQs resistance [6].

The emergence of drug resistance and persistence of infection is serious threat to control TB [7]. This high incidence of resistance severely limits treatment options and requires the use of more toxic and costly treatment regimens [8]. The present study is conducted to detect the mutational profile of FQs-R to help in determining the potential utility and selection of adequate drug regimens.

Methods

Sample collection

Mycobacterium tuberculosis isolates were procured from patients diagnosed with pulmonary TB. The samples were collected from PMDT sites (Programmatic management of drug resistant tuberculosis) of seven different districts (Lahore, Faisalabad, Gujranwala, Sahiwal, Sargodha, Sialkot, and Bahawalpur) from period of May 2018 to March 2019 (Fig. S1). A total of 562 suspected MDR-TB cases were included in the study. GeneExpert and MTBDR plus assay was performed for susceptibility against first line anti-TB drugs. History of anti-tuberculosis treatment was obtained from patients which includes newly diagnosed cases and previously treated cases (treatment failure and treatment default).

This study was approved by the Research Ethics and Biosafety Committee (No.D/650/MMG) of Department of Microbiology and Molecular Genetics, University of the Punjab, Lahore, Pakistan.

Sample processing

Initially, the sputum samples were decontaminated using standard NALC-NaOH (N-acetyl-L-cysteine sodium hydroxide) method [9] and smear positive samples were directly processed for DNA extraction. Smear negative samples were primarily cultured on MGIT Bactec 960® medium. DNA was extracted using Genolyse version 1.0 kit method (Hain Lifescience, Germany). After DNA extraction, supernatant was collected and transferred into fresh tube and stored at -20°C for further processing.

Molecular detection of FQs resistance

The phenotypic resistance to second line drugs including flouroquinolones was determined using GenoType MTBDRs/ version 2.0 kit method. The whole procedure of molecular detection with GenoType MTBDRs/ includes three steps i.e. DNA extraction, amplification with biotinylated primers, and reverse hybridization. The test was considered valid when all control bands appeared correctly.

PCR and sequencing

Primers were designed against QRDR region of *gyrA* gene; forward primer 5-GATGCAGCGCAGCTACATCGAC-3 and Reverse primer 5-GATGCAGCGCAGCTACATCGAC-3. Cyclic

parameters for the amplification reaction were 95°C for 5 min followed by 40 cycles of 95°C for 30 sec, 61°C for 45 sec, 72°C for 50 sec and final elongation for 10 min at 72°C. The PCR products were purified using a Qiagen PCR purification kit and eluted in TE buffer. The sequencing of isolates was performed by 1st Base sequencer, Malaysia.

Results

GeneExpert assay was performed for the confirmation of MDR-TB. The susceptibility of first line drugs was determined against rifampicin (RIF) and isoniazid (INH). The results have shown that 430 samples were MDR (resistant to both isoniazid and rifampicin), 91 samples were mono-resistant (57 resistant to RIF and 34 resistant to INH), and 41 samples were susceptible to these two drugs.

The data analysis showed that out of 562 cases, 313 were the newly diagnosed, 97 were treatment failure cases (who completed the treatment but still positive for MTB), 59 were treatment default cases (previously taking anti-tuberculosis therapy, at least for one month, but did not complete treatment), and for the 93 samples treatment history was unknown.

Detection of FQs resistance

The frequency of FQs-R and its mutational profiling was determined in *gyr* gene. Genotype MTBDRsl assay was performed to determine resistance to second line drugs. Both *gyrA* and *gyrB* were examined for FQs resistance. A total of 104/562 (18.5%) samples were found resistant to FQs.

Among FQs resistant isolates, 8 isolates were rifampicin sensitive, 8 were mono resistant (resistant for rifampicin and sensitive for isoniazid), and 88 were MDR (multi drug resistant). Most of the patients were previously treated and appeared with category 1 failure treatment and some were on category 4 treatment. In total, 102/104 showed resistance for *gyrA* gene while *gyrB* gene mutations were prevalent only in 2/104 cases with mutation in E540V. The resistance was interpreted according to the presence and absence of wild type and mutant probe. When all wild type probes appeared it was interpreted with no detectable mutation. In the absence of any wild type probe, the respective amplicon cannot bind to the corresponding wild type probe resulting in detectable mutation.

Mutational profiling of *gyrA* gene

The mutation probes detect some of the most common resistance-mediating mutations. Table 1 shows all types of mutations in the *gyrA* gene and the pattern of corresponding wild and mutant bands. Failure of *gyrA* WT3 band and the development of MUT3C was the most common pattern of mutation. Mutational analysis showed that these mutations mostly alter the protein at codon 94 and 90 which represent the replacement of aspartic acid with glycine and alanine with valine, respectively. Aspartic acid also replaced with alanine, asparagine and tyrosine in other types of mutations at codon 94. Mutation at codon 91 was also found where serine was replaced with proline.

Different hybridization patterns were observed for FQ resistance in *gyrA* gene and 19 isolates from all possible patterns were selected for further mutational analysis in the hot spot regions of *gyrA* gene loci using H37RV (AGH06049.1) as a reference strain. In these isolates, amino acid changes were determined due to these mutations. We found variation in four isolates with respect to wild type and mutant probe where amino acid changes vary from the most known mutation reported by LPA testing. One isolate substituted from asp→ala, two isolates had ala→val and one had thr→ala mutation in addition to ser/ala→X (an undetermined mutation) (Fig. 1).

Discussion

Fluoroquinolones have long been widely used for several infectious diseases and easily accessible in certain regions even without prescription. Such misuse of FQs has highly contributed to their efficacy in the treatment of TB and emergence of FQ-resistance [10].

In the current cross-sectional study, presumptive multi drug resistant isolates of MTB were included. A high proportion of rifampicin (Rif) and isoniazid (INH) resistance was observed. Of the total, 92% (521/562 i.e. 430 MDR and 91 monoresistant) isolates were resistant to first line drugs (INH and Rif) either both or alone. Among FQs resistant isolates, 8 isolates were rifampicin sensitive, 8 were mono resistant, and 88 were MDR. The isolates having resistance against rifampicin and isoniazid are termed as MDR-TB and if they develop additional resistance against FQs then known as Pre-XDR TB [11].

MTB develop resistance against FQs, mostly, by developing mutations against drug targeted proteins. The detection of gyrase mutations can help in predicting FQs resistance as well as estimating the levels of resistance to various fluoroquinolones [12]. GenoType MTBDRs/ assay can detect mutations in the QRDR region of the gyrase genes (*gyrA* and *gyrB*). GenoType MTBDRs/ assay was used to determine the frequency of FQs resistance of our isolates. A total of 104/562 (18.5%) isolates were found resistant to FQs. The high prevalence of FQs resistance was also reported in other provinces of Pakistan [10,13] and neighboring countries India [14,15], China [16,17] and Bangladesh [18,19].

Short treatment regimens are used to reduce emergence of antimicrobial resistance in MTB. According to National guidelines for control of TB in Pakistan 2019 (adopted by WHO), the anti-TB short regimens include third or fourth generation fluoroquinolones (levofloxacin and moxifloxacin respectively) for 4 months for drug susceptible cases. They are also given in isoniazid resistant and previously treated cases in initial phase of therapy (2 months). The high proportion of FQ-R indicates the patient ineligibility for shorter regimens.

The resistance to FQs occurs due to point mutations in conserved QRDR region of *gyrA* and *gyrB* gene. The mutation in QRDR region changes the structure of drug binding pocket (QBP) of quinolones and results in cross resistance to all FQs. The frequency of *gyrA* mutations was much higher than *gyrB*. It was found in hybridization pattern that most of the isolates had mutation in *gyrA* gene loci with substitution at amino acid 94 and 90 where D94G and A90V were more prevalent. These A90V and D94G mutations are associated with high level of resistance to fluoroquinolone antibiotics (Fig. 2). A90V mutation detects

resistance for levofloxacin but a higher generation of FQ i.e. moxifloxacin can be used at higher dose. But if there is mutation at D94G both levofloxacin and moxifloxacin are ineffective [20] (GLI guideline Line probe assay). S91P, D94A, and D94N/Y were other mutations found in our isolates. With the first two mutations the isolates could be susceptible to moxifloxacin at higher doses but resistant to levofloxacin. However, D94N/Y detects resistance for both levofloxacin and moxifloxacin. These findings correlate with most of the previous studies [6,21,22].

The patient's characteristics and type of mutation are assessed in two ways. The first approach was to determine frequency of particular mutation according to their categorization of resistance to first line drugs and the second approach was by their categorization according to treatment history. In MDR and RRD cases, the FQs-R was most commonly associated with D94G mutation. This mutation shows high level resistance to all fluoroquinolones even to fourth generation moxifloxacin. Monoresistance INH-R showed S91P and D94N/Y. D94N/Y mutation was also observed in RRD. When mutation pattern of newly diagnosed TB isolates with FQs resistance was observed, it showed high proportion of A90V mutation where moxifloxacin still remains the drug of choice at higher doses. In contrast, the mutation pattern of D94G was commonly found in treatment failure and relapse cases.

Since Genotype MTBDRsl assay target only small region of gene with limited number of well-known mutations and sometimes, the interpretations are indistinct for cross-resistance to FQs which occur due to particular *gyrA* mutations. The sequence analysis was performed for understanding of resistance at the genotypic level [23]. There were some isolates (4/102) which showed co-existence of mutations in hybridization pattern. The combination of mutation was D94A with D94H, S91P with D94G, D94G with D94N/Y, and A90V with D94G. Co-existence of mutations was also observed in sequence analysis of *gyrA* gene where S91T mutation was detected in 95% isolates. But this type of mutation is not related with fluoroquinolone resistance. It could be present even in sensitive isolates as determined by other studies [24,25]. The results of all our mutations are in line with a study conducted in Pakistan where they detect mutations in extensively drug resistant strains [26].

Interestingly, we had found some other hot spot mutations in hybridization pattern of co-existence cases where all wild type present with mutant probe. In our two isolates, when MUT3C and MUT2 appeared in the presence of its all wild type probe, substitution of Ala into Val was observed, but when MUT2 probe appeared in the absence of WT3 the mutational change of Asp into Ala occurred. Four isolates selected for our most common pattern i.e. absence WT3 and presence of corresponding mutant probe MUT3C, but, one isolate had shown mutation of Thr into Ala additionally with G/M/R into undetermined amino acid X. For most of the isolates amino acids changed from the pattern of well-known mutations but mostly occurred in the same region i.e. codon 94 and 91. Overall, these findings suggest that the mutation pattern can differ according to the hybridization pattern of wild type and a mutant probe.

The study represents the burden of fluoroquinolone resistance in MDR-TB patients regardless of FQ antibiotic therapy. However, the relevance of genotypic and phenotypic resistance is important to predict accurately FQ-R. Even though we found FQ-R in MTB susceptible isolates but it does not reflect its true

prevalence in these patients. The resistance might develop due to the prior use of FQ antibiotic and the results were not included in the study.

Conclusions

In conclusion, the emergence of flouoroquinolone resistance in clinical isolates is alarming. We found high proportion of FQs resistance in MDR cases and even in mono-resistant and all drugs sensitive isolates. Previously treated and failure TB treatment cases being the most important group for developing resistance. Our findings suggest that the implementation of FQs in these patients should be carefully administered and genotypic studies should be carried out, preferably at the time of initial diagnosis, to rule out all type of mutations for effective treatment and particularly to control its resistance.

Abbreviations

FQ: Flouoroquinolone; FQ-R: Flouoroquinolone Resistant; LPA: Line Probe Assay; MDR: Multidrug Resistant; MGIT: Mycobacteria Growth Indicator Tube; Mtb: *Mycobacterium tuberculosis*; MUT: Mutant; NALC-NaOH: N-acetyl-L-Cysteine Sodium Hydroxide; QRDR: Quinolone Resistance Determining Region; TB: Tuberculosis; WT: Wild Type; XDR: Extensively Drug Resistant.

Declarations

Ethical approval and consent to participate

All study participants were informed verbally (approved by Ethics and Biosafety Committee) and consent to participate was collected from the patients directly or from parents. No study participants were included in the present study involving guardians. The study was approved by Departmental Research Ethics and Biosafety Committee.

Consent for publication

“Not Applicable”.

Availability of data and materials

The datasets generated and/ or analyzed during the current study are part of PhD thesis of the first author and not publicly available. The datasets are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

Funding

No funding was obtained for the current study.

Authors' contributions

SK performed experiments, analyzed the data and wrote the manuscript. conceived the research idea. ZT provided sources. NM helped in experiments and data analysis. Both MS had responsibility for data collection. AR conceived the research idea and helped in manuscript preparation. All authors contributed substantially to the interpretation of the results. All authors approved the final manuscript.

Acknowledgements

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Table

Table 1: The frequency and the mutations conferring resistance to FQs.

gyrA Mutation probe	Missing wild type probe	Phenotypic Susceptibility	Mutation	Frequency (n=102)
gyr A MUT1	gyr A WT2	Resistant	A90V	25
gyr A MUT2	gyr A WT2	Resistant	S91P	7
	gyr A WT3			
gyr A MUT3A	gyr A WT3	Resistant	D94A	7
gyr A MUT3B	gyr A WT3	Resistant	D94N/Y	9
gyr A MUT3C	gyr A WT3	Resistant	D94G	45
None	any one wild type probe	Resistant	A90V D94A D94G	9

Figures

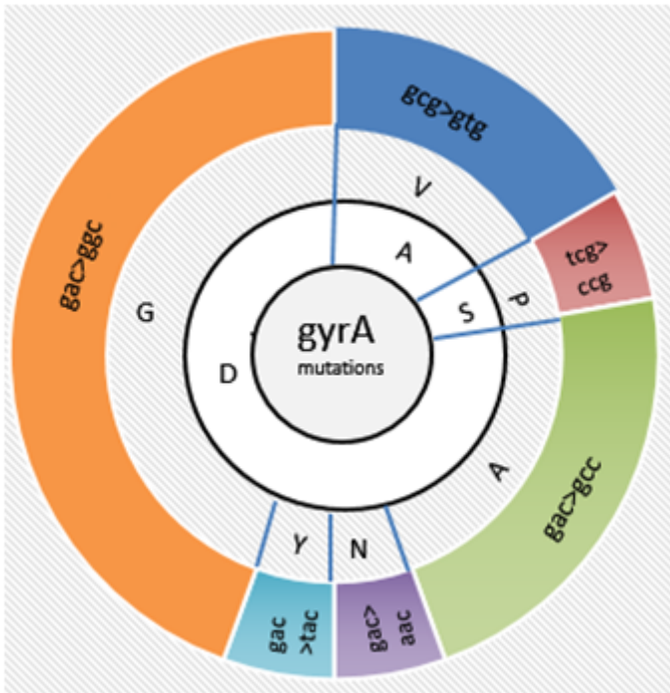


Figure 1

Frequency amino acid and nucleotide change in FQs isolates. In general, amino acid change had found in three amino acids alanine (A into V), serine (S into P) and aspartic acid (D either into A, N, Y or G). The nucleotide change corresponding to these amino acids change is also represented.

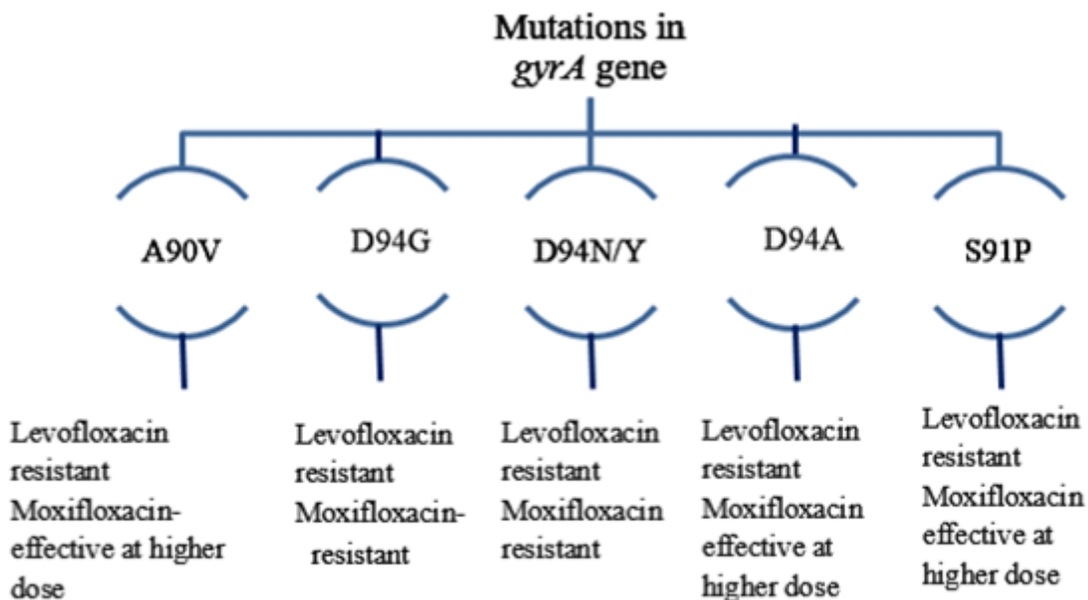


Figure 2

The association of *gyrA* gene mutation with levofloxacin and moxifloxacin resistance.

Supplementary Files

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