Resistance to anti leprosy drugs in multi-bacillary leprosy: A cross sectional study from a tertiary care centre in eastern Uttar Pradesh, India

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Abstract

Background: WHO MDT is the main drug regimen for treating leprosy and has been used for more than three decades. Many cases of relapse of leprosy have been reported, which points towards the emergence of drug resistance with the antileprotic drugs.

Objectives: To find the resistance with the antileprotic drugs by detecting the mutations in drug resistance determining region of the rpoB, folP1 and gyrA genes of Mycobacterium leprae.

Methods: Leprosy patients with bacterial index ≥2 were included in the study. The slides were further processed to extract genomic DNA, and polymerase chain reactions were performed to amplify the drug resistance determining region (DRDR) of rpoB, folP1 and gyrA genes. The samples in which genes could be amplified were subjected to DNA sequencing to detect mutations.

Results: Out of 78 samples rpoB gene was amplified in 39 (50%), folP1 in 32 (41%) and gyrA in 45 (57.7%). In 20 (25.6%) samples no gene was amplified. Only 32 samples of rpoB, 25 samples of folP1 and 38 samples of gyrA gene were included in the study, rest were excluded due to sequencing error. No mutation was seen in rpoB gene and in folP1 gene. In gyrA gene samples mutations were seen in 8 (21%) samples, and were present at codon 91 GCA → GTA (Alanine → Valine).

Limitations: Small sample size and less efficient method to detect resistance.

Conclusion: Resistance is not a problem with conventional drugs in MDT. It is more common with quinolones.

Key words: Antileprotic drugs, Hansen’s disease, mutations, resistance

Introduction

Leprosy is a chronic infectious disease which mainly affects skin and peripheral nerves, caused by Mycobacterium leprae (M. leprae). More than 81% of the new cases are reported from three countries – India, Brazil, and Indonesia.¹ Although the prevalence of leprosy in India is less than 1/10,000 since 2005,² it still accounts for 62% of the total new cases reported worldwide.¹ Dapsone was introduced for the treatment of leprosy in 1950s and was used worldwide to treat both multibacillary (MB) and paucibacillary (PB) forms of the disease. Long-term monotherapy with dapsone resulted in the emergence of dapsone-resistant strains of M. leprae leading to treatment failure.³,⁴ Between 1960s and 1970s, rifampicin and clofazimine were added in the treatment of leprosy. Rifampicin is a strong bactericidal drug against M. leprae. However, using it alone could result in the emergence of rifampicin resistant strains of M. leprae.⁵ To overcome the threat posed by the worldwide spread of dapsone and rifampicin resistance and to improve the treatment efficacy, World Health
Organization (WHO) recommended multidrug therapy (MDT) for leprosy in 1982. However, drug resistance has been reported since 1964 for dapsone,\(^5\) since 1976 for rifampin,\(^6\) and since 1996 for ofloxacin.\(^7\) Comprehensive data regarding the magnitude of drug resistance is crucial to evaluate the efficacy of MDT. As *M. leprae* cannot be cultured axenically, detection of drug resistance in leprosy is difficult. Shepard developed mouse foot pad assay to determine *M. leprae*’s susceptibility to anti-leprosy drugs in 1962.\(^8\) Since then, it has been the ‘gold standard’ for drug susceptibility testing. While mouse foot pad assay gives definitive information pertaining to the susceptibility of an *M. leprae* isolate to anti-leprosy drugs, it is a laborious and expensive procedure and is carried out only in a few reference centres in the world. The availability of genomic sequence of *M. leprae*\(^9,10\) and an improved understanding of the genetic basis of drug resistance in mycobacteria have led to the development of molecular methods for the detection of mutations associated with dapsone, rifampicin, and fluoroquinolone resistance.\(^11,12\) Recent studies have identified point mutations in the *folP1* gene, which encodes dihydropteroate synthase (DHPs) in dapsone-resistant *M. leprae*.\(^11\) Rifampicin resistance is associated with mutations in the *rpoB* gene that encodes the β subunit of RNA polymerase.\(^14\) Resistance to ofloxacin is known to be associated with mutation in *gyrA* gene encoding the A subunit of DNA gyrase in *M. leprae*.\(^13,16\) No molecular target has been defined for clofazimine. Thus, by performing polymerase chain reaction and DNA sequencing, we can detect mutations in the drug resistance determining regions (DRDR) of *folP1*, *rpoB*, and *gyrA* genes, responsible for resistance to dapsone, rifampicin and ofloxacin, respectively.\(^17\) The aim of the present study was to look for the reported mutations in *folP1*, *rpoB*, and *gyrA* genes.

**Methods**

**Study design**

This was a cross-sectional observational study conducted at a tertiary health care centre during October 2013 to May 2015. The patients included in the present study belonged to the Gangetic belt of the eastern part of north India. The study was approved by the ethics committee of the institute. Written consent was taken from all participants of the study. Patients suffering from leprosy, with bacteriological index (BI) ≥2 were included in the study. The samples (slit-skin smears) were obtained from patients before starting MDT, patients on MDT, defaulters of MDT (defined as who fail to complete treatment within the maximally allowed time frame i.e. six months treatment for PB leprosy must be completed within a maximum period of 9 months, similarly 12 months treatment for MB leprosy must be completed within 18 months), and patients who had relapse (defined as patients who developed new skin lesions after completion of MDT). Diagnosis was primarily on clinical grounds and confirmed by staining for acid fast bacilli (AFB) from samples obtained from ear lobes, forehead, or lesions on other sites of the body, by slit skin smear method, according to the standard procedures recommended by the WHO. Three smears were taken on a sterile glass slide and were air dried. One sample was stained by Ziehl–Neelsen staining and BI was calculated according to the Ridley’s logarithmic scale.\(^18\) Smears on the other two slides were scraped by a sterile blade and suspended in 1 ml sterile saline and stored in 1.5 ml air-tight micro-centrifuge tubes until processed in the department of microbiology.

**Genomic DNA extraction**

Samples were centrifuged at 10,000 revolutions per minute (rpm) for 5 min. Supernatants were discarded and pellets were re-suspended in 250 µl of tris-Cl EDTA-buffer (TE, pH 8.0) and mixed by vortexing. Then, 100 µl of 10% sodium dodecyl sulfate (SDS) and 3 µl of protease-K (20 mg/ml) were added and mixed by inverting the tubes. Cell suspensions were incubated at 37°C overnight. Subsequently, 100 µl of 5 M NaCl and 80 µl of 10% cetyl trimethyl ammonium bromide (C-TAB) were added to each tube and mixed well and incubated at 60°C for 10 min. Equal volume of phenol: chloroform: isooamy alcohol (IAA) (25:24:1) were added to the samples and centrifuged at 12,000 rpm for 10 min at 4°C to separate the aqueous and organic phase. Aqueous phase was collected in new tubes and mixed with equal volume of chloroform: IAA (24:1), and centrifuged at 12,000 rpm for 10 min at 4°C. Aqueous phase was collected again and mixed with equal volume of isopropanol and kept at room temperature for 10 min to precipitate the genomic DNA. Tubes were centrifuged at 12,000 rpm for 10 min at 4°C to precipitate the pellet. Supernatants were discarded and DNA pellets were washed with 70% ethanol by centrifugation at 8,000 rpm for 5 min. Pellets were dried at 37°C in inverted condition and dissolved in 50 µl TE.

**Polymerase chain reaction amplification**

Each genomic DNA sample was submitted for primary and nested polymerase chain reaction (PCR) for amplification of drug resistance determining region of the *rpoB*, *folP1*, and *gyrA* genes with the primers used in earlier studies.\(^19\) Primer sets *folP1*: F1/R1, *rpoB*: F1/R1 and *gyrA*: F1/R1 [Table 1] were used for primary PCR to amplify the corresponding regions of respective genes. All the tubes were kept in thermocycler and standardized programs were used for PCR. For amplification of the target region of the *folP1* gene, cycling conditions used were 95°C for 30 s, 60°C for 30 s, and 72°C for 30 s for 35 cycles. For amplification of target regions of *rpoB* and *gyrA* genes, a programme of 30 s at 95°C, 30 s at 56°C, and 30 s at 72°C for 35 cycles were used. Primer sets\(^17\) *folP1*: F2/R2, *rpoB*: F2/R2, and *gyrA*: F2/R2 [Table 1] were used for the second round of amplification (nested PCR) using similar programs. Amplified products were resolved by agarose gel electrophoresis and visualized by ethidium bromide (EtBr) staining. Before submitting the amplified products for sequencing, every PCR product was purified using gel elution kit (Qiagen) according to the manufacturer’s instructions. DNA sequencing was performed at Eurofin Genomics India Pvt. Ltd. (Bangalore, India).

| Genes | Primer Sequence (5’-3’)|
|-------|------------------------|
| *folP1* gene  |          |
| Outer primers | folp1-F1  | CTTGATCCTGACGATGCTGT  |
|            | folp1-R1  | CCACCAAGACATCGTGGAC  |
| Inner primers | folp1-F2  | GATCCTGACGATGCTGCCAG  |
|            | folp1-R2  | ACATCGTTGACGATCGTG  |
| *rpoB* gene  |          |
| Outer primers | rpoB-F1  | ACCTGATCAAATTACCTGTC  |
|            | rpoB-R1  | GTATCTGACGTCGTGCTGA  |
| Inner primers | rpoB-F2  | CTGATCAATTCGCTCCGTT  |
|            | rpoB-R2  | CGACATGAACAGCATCAG  |
| *gyrA* gene  |          |
| Outer primers | gyrA-F1  | ATGACTGATACACCGTCGCA  |
|            | gyrA-R1  | AATAACGCTACGGCCTGGTG  |
| Inner primers | gyrA-F2  | GATGCTCTCAAACCGTGACAT  |
|            | gyrA-R2  | ACCCGGCGAAATTTG  |

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**Table 1: Primers used in this study**
using chain-termination (Sanger’s) method. DNA sequence analysis was done by pair-wise and multiple-sequence alignments using CLUSTAL W2 tool.

**Results**

Ninety-two patients were enrolled but only 78 (66 males and 12 females, mean age 35.73 ± 10.89 years) among them were included in the study. Remaining 14 patients (5 before MDT, 6 on MDT, 2 defaulters and 1 relapse) smears failed to demonstrate *M. leprae*. The most common clinical type of leprosy was borderline lepromatous (41%) followed by lepromatous (38.5%), mid-borderline (15.4%), and histoid (5.1%). Primary amplification yielded the product of 254 base pair (bp), 345 bp, and 390 bp for folP1, rpoB, and gyrA genes, respectively. The secondary round of nested PCR yielded products of 255 bp, 242 bp, and 225 bp for rpoB, folP1, and gyrA, respectively. In 22 (28.2%) patients all the three genes could be amplified. In 39 (50%) cases rpoB, 32 (41.02%) folP1 and 45 (57.7%) cases gyrA were amplified. In 20 (25.6%) patients no gene could be amplified, in seven (8.9%) only rpoB, in four (5.1%) only folP1, and in 11 (14.1%) only gyrA were amplified. In 24 (30.8%) cases combination of rpoB and folP1, in 30 (38.5%) cases rpoB and gyrA and in 26 (33.3%) cases folP1 and gyrA were amplified by PCR. Forty five cases were new, 20 were on MDT, six were defaulters, and seven were relapsed cases. Figures 1–3 show PCR amplified products on agarose gel electrophoresis and visualized by ethidium bromide (EtBr) staining.

DNA sequencing of 32 samples of rpoB, 25 of folP1 and 38 of gyrA genes were successfully done. In the remaining samples, there were sequencing errors and therefore those cases were excluded from this analysis. Sequencing of rpoB gene was done on 22 new cases, six patients on MDT, two defaulters, and two cases of relapse. The expected mutation was not found in any of the samples. Sequencing of folP1 gene was done on 17 new cases, six patients on MDT and two defaulters. No mutation was found in any of them. Sequencing for gyrA gene was done on 24 new cases, seven patients on MDT, four defaulters and three cases of relapse. The expected mutations could be seen in only eight cases at 91 codon (GCA → GTA), of which three were new cases, two were on MDT, and three were defaulers. Details of patient characteristics with drug resistance are provided in Table 2.

**Discussion**

At present it is not technically possible to provide direct evidence for the mechanisms of resistance of *M. leprae* to most antileprosy drugs. Current understanding regarding this aspect is based on our knowledge about drug resistance of *M. tuberculosis*. According to several previous studies, drug resistance in *M. leprae* may be primarily attributed to mutations in genes encoding drug targets. For effective treatment and containment of drug resistant strains, it is mandatory to have local and global data on the drug resistance pattern of the bacterium. In the present study, we have examined the mutations possibly associated with drug resistance in the target genes by PCR-based amplification and sequencing. It is really heartening to note that none of the randomly selected 32 amplicons subjected for sequencing showed nucleotide mutation in rpoB, indicating

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**Table 2: Characteristics of patients having drug resistance**

| Age/sex   | Patient status | Duration of illness in months | Bacterial index | Reactional state | Genes amplified with PCR | Mutation detected |
|-----------|----------------|-------------------------------|-----------------|------------------|--------------------------|------------------|
| 28/male   | New            | 12                            | 4               | Nil              | rpoB, folP1 and gyrA     | Mutation in gyrA GCA → GTA at codon 91 |
| 26/male   | On MDT×3 months| 24                            | 5               | Type 2           | rpoB, folP1 and gyrA     | Mutation in gyrA GCA → GTA at codon 91 |
| 32/male   | New            | 24                            | 3               | Type 2           | rpoB, folP1 and gyrA     | Mutation in gyrA GCA → GTA at codon 91 |
| 38/male   | New            | 5                             | 3               | Nil              | gyrA                     | Mutation in gyrA GCA → GTA at codon 91 |
| 21/male   | Relapse        | 36                            | 3               | Type 2           | gyrA                     | Mutation in gyrA GCA → GTA at codon 91 |
| 40/male   | Relapse        | 60                            | 2               | Type 1           | folP1, gyrA              | Mutation in gyrA GCA → GTA at codon 91 |
| 26/male   | On MDT for 10 months | 18                           | 3               | Nil              | gyrA                     | Mutation in gyrA GCA → GTA at codon 91 |
| 45/male   | Defaulter      | 12                            | 3               | Nil              | gyrA                     | Mutation in gyrA GCA → GTA at codon 91 |

MDT: Multidrug therapy, PCR: Polymerase chain reaction
that the *M. leprae* strains on eastern part of North India are 100% sensitive to rifampicin. Our finding is in agreement with another study carried out in South India in 2011 by Sekar et al. On the contrary, one study conducted on relapse cases of leprosy, collected from leprosy hospitals scattered wide across regions endemic for leprosy in India, during 2009 and 2013, showed mutations in rpoB gene of the *M. leprae* in 3.6% (4/111) at codon 439 (Phe → Leu), at 442 (Gln → His), 433 (Thr → Ile), and at 441 (Asp → Tyr). Another study carried out in east India showed 4% (2/50) mutation at codon 442 (Glu → His) in relapse cases. The latter mutation (Glu → His) was common with other Indian studies. There was no sharing with those 4 mutations reported from east and southeast Asian countries in strains of relapsed cases.

The folP1 gene, a target for dapsone, showed no mutation, indicating that dapsone is likely to be very effective in treating patients with leprosy in this region. However, previous studies from other parts of India had shown that the frequency of mutations of *M. leprae* among relapsed cases ranged from 8.1% to 15%. While looking for mutations in the target gene gyrA for quinolone, we found that 21% (8/38) of the strains had only one point mutation i.e. at 91 codon, from GCA → GTA leading to change in amino acid from alanine → valine. These eight cases consisted of three new cases, two patients on MDT, one defaulter, and two cases of relapse. The finding of this mutation in three new cases in our study was surprising. A possible reason is rampant prescription of quinolones to treat other infections. A study carried out on patients attached to Leprosy Mission hospitals in India showed mutations in gyrA gene in 8.1% (9/111) of the participants. Three different mutations were noted in this study; two at codon 91 (Ala → Thr); 91 (Ala → Val), and one at codon 92 (Ser → Ala). However, Sekar et al. from south India did not find any mutation in this gene. A study conducted in east and southeast Asian countries showed mutations in gyrA gene among 6.8% of the participants. A study from Japan found only one strain with mutation at codon 91 (Ala → Val). Surprisingly, several studies carried out in different parts of the world (India, USA, Japan, Vietnam, etc.) reported no mutation in the gyrA gene of *M. leprae*. Our region is endemic for enteric fever also and quinolones were most commonly used to treat these infections. Quinolones were misused to treat many other types of infections and febrile conditions which may lead to drug resistance with these drugs. A high rate of mutation in gyrA gene of *M. leprae* from this region indicates the weak prospects of quinolone as a second line drug.

Small number of cases and not focussing on relapsed cases are possible limitations of this study. Our findings indicate that resistance to first line drugs against leprosy is not a major problem in east Uttar Pradesh. We must counsel the patients about adherence to treatment and follow-up the patients adequately to minimize the number of defaulters. However, it must also be noted that mutations of the three genes rpoB, folP1, and gyrA must not be taken as a definite indicator of drug resistance. The culprit mutation must be verified by mouse foot pad assay meticulously. Further, such mutations must also be examined in strains cured later by standard therapy.

To conclude: Antileprotic drugs present in MDT are still effective against *Mycobacterium leprae*. Quinolones may not be a good choice as an antileprotic agent in our region.

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**Conflicts of interest**

There are no conflicts of interest.

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