MOLECULAR DIAGNOSTICS

DNA-PCR and RT-PCR for the 18-kDa gene of *Mycobacterium leprae* to assess the efficacy of multi-drug therapy for leprosy

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DNA-PCR and reverse transcription (RT)-PCR for the 18-kDa protein of *Mycobacterium leprae* were used to examine the efficacy of multi-drug therapy (MDT) in leprosy. MDT was administered for 0–24 months. Fourteen (63.6%) of 22 patients showed positive PCR results after treatment for 12 months and the positive results decreased to 30% after 24 months of MDT. These results did not correlate with the bacterial index (BI) or the IgM antibody titre for the phenolic glycolipid (PGL)-1. One-dimensional densitometric analysis of agarose gels from PCR from the longitudinal study showed a gradual reduction of the 360-bp band after 12–24 months of MDT. RT-PCR for mRNA of the 18-kDa protein successfully tracked bacterial RNA changes in the biopsies and confirmed a decrease in the RNA of *M. leprae* in patients after MDT for 12 months. Thus, DNA- and RT-PCR for the 18-kDa protein of *M. leprae* are effective in assessing the efficacy of MDT for leprosy.

Introduction

Leprosy is a chronic infectious disease caused by *Mycobacterium leprae*. Although multi-drug therapy (MDT) has been very successful in reducing the prevalence of the disease, leprosy is still a major health problem globally. A sensitive and specific method for the detection of *M. leprae* would add an unbiased criterion to the available means of diagnosis and it might allow diagnosis of leprosy at a very early stage before the appearance of clinical signs. The lack of methods to identify *M. leprae* quickly and specifically has hampered research into the distribution and spread of the bacillus.

The assessment of chemotherapy efficacy in *M. leprae* infection is difficult, as the only reliable method for determining viability of the organism depends on its growth in mouse footpads. This technique is extremely expensive and time-consuming. It takes at least 10 months for organisms to grow and >5 years for low-grade drug resistance to be proven.

A widely used indirect measure of *M. leprae* viability is the morphological index (MI) [1]. Because *M. leprae* cannot be cultivated in vitro, the MI has become the standard laboratory assay for monitoring the short-term bactericidal activity of new anti-leprosy drugs. In this technique, the integrity of stained bacilli is judged microscopically and the proportion of morphologically intact bacilli is calculated. Although the idea of the technique is simple and it can be used in the field, in practice it requires skilled staff and precise staining conditions. Because of the lack of adequate standardisation and subjective interpretation, the MI is difficult to apply. Mycobacteria can be visualised by staining followed by microscopy, but this is not specific for *M. leprae*. As acid-fast staining requires at least $10^4$ organisms/g of tissue for reliable detection, sensitivity is low, particularly for patients in the tuberculoid leprosy spectrum when acid-fast bacilli (AFB) are rare or absent [2].

Current available serological tests show a relatively low sensitivity, which, in combination with the low prevalence of leprosy, limits their application to early case finding and detection of infection [3]. There are several reports on the measurement of phenolic glycolipid (PGL)-1 antigen titres in serum as a rapid marker for the efficacy of chemotherapy in patients...
with leprosy [4, 5], although the precise relationship of circulating antigen to bacterial viability is not clear.

Recently, several investigators have used the PCR to amplify various genomic sequences of *M. leprae* to improve detection when low numbers of bacteria are present [6–9]. DNA probes offer a route to the more sensitive detection and identification of bacterial DNA, e.g., *M. leprae* DNA in clinical samples, through the application of the PCR with *M. leprae* specific primers. PCR for identification of the DNA, which encodes *M. leprae* proteins of 65 kDa [6], 36 kDa [7] and 18 kDa [8] and the repetitive sequences [9] of *M. leprae*, was introduced as a more sensitive and specific method than the empirical AFB examination.

In this study PCR was used as a measure of *M. leprae* viability on the premise that it detects nucleic acid, which must be present in the living organisms but which on cell death is more susceptible to degradation than other cell components. In this way, the efficacy of MDT was assessed in a longitudinal study.

An alternative detection method was also developed, which targets the abundant mRNA for the 18-kDa protein of *M. leprae*. Detection of RNA should give increased sensitivity over assays based on the detection of a single copy or even multiple copies of genomic sequences, as each cell contains $10^3–10^4$ copies of mRNA. An RNA-based detection method would be expected to better reflect the number of viable organisms because RNA is generally degraded within a few minutes of cell death. Thus, an RNA-based detection system might be useful for confirmation of the diagnosis in patients for whom a diagnosis is difficult to make, for assessing the efficacy of chemotherapy, in distinguishing relapse from late reaction in previously treated patients and for epidemiological studies.

To examine the efficacy of 18-kDa PCR and RT-PCR for the assessment of MDT, this study compared the changes in the DNA of *M. leprae* with bacterial index (BI) and IgM antibody titre for phenolic glycolipid (PGL)-1 after MDT, determined the band thickness of PCR signals by the one-dimensional gel analysis method from representative biopsies for longitudinal study up to 12 and 24 months after MDT, and evaluated changes in the expression of mRNA for the 18-kDa protein in *M. leprae* during MDT by RT-PCR.

**Materials and methods**

**Specimen collection and bacterial index (BI)**

Punch biopsies, slit smears and sera from the peripheral blood of 31 multi-bacillary case (MB) patients, with a positive BI of >3 at the time of diagnosis were collected from the Institute of Hansen’s Disease in Seoul, the Affiliated Hospital of the Korean Leprosy Control Association in Euiwang City, the Catholic Skin Clinic in Taegu City and the Jesus Clinic in Taegu City. Biopsies were performed on the most representative lesions with patient’s consent at 0, 6, 9, 12, 18 and 24 months during the course of MDT and stored in a cryovial. Biopsies for DNA-PCR were sent to the Institute of Hansen’s Disease in ice, and specimens for RT-PCR were stored in liquid nitrogen and sent to the Institute of Hansen’s Disease in a portable liquid nitrogen carrier.

The highest BI score in lesions was used to describe a patient’s BI, because the average BI from lesions and routine sites not showing visible signs could reduce the BI more than warranted. BI scores were determined at the individual clinics. IgM antibody titre for PGL-1 was assessed at the Affiliated Hospital of the Korean Leprosy Control Association.

**Separation of *M. leprae* DNA from biopsies and infected footpads of nude mice**

After finely cutting half of the biopsy with no. 10 and 15 disposable scalpels in a Petri dish, 300 μl of Dulbecco’s phosphate-buffered saline (DPBS; pH 7.4) were added and the suspension was placed in a microcentrifuge tube. The suspension was homogenised by vortex mixing with three glass beads (3 mm in diameter) for 3 min. The supernate was recovered by centrifugation at 125 g for 10 min at 4°C. A 50-μl sample was subjected to freezing and thawing five times for 1 min each; 10 μl were then used as a DNA template for 18-kDa PCR. The rest of the DNA sample was stored at −70°C until required.

Footpad granuloma from *M. leprae*-infected nude mice was dissected, soaked in iodine 1% solution and chopped finely with no. 10 and 15 disposable scalpels. The sample was then homogenised in 2 ml of DPBS with 25–30 glass beads in a Mickle homogeniser (Mickle Laboratory Engineering, Surrey, UK). Ten μl of supernate were used as a DNA template and part of the supernate was stained with Ziehl-Neelsen’s stain for acid-fast bacilli which were enumerated by the procedure of Shepard and McRae [10].

**18-kDa PCR**

A modification of a previously reported method [11] was used with the same primers used to amplify the DNA encoding the 18-kDa protein and hot start PCR was accomplished. A reaction mixture of 10 μl of template DNA and 100 pmole of primers was overlaid with mineral oil in a reaction tube, then denatured at 94°C for 5 min in a PCR 9600 heating cycler (Perkin Elmer, Branchburg, NJ, USA). A reaction mixture of 125 μM dNTPs, Taq DNA polymerase (Boehringer Mannheim, Mannheim, Germany) 0.5 U, 50 mM KCl, 10 mM Tris-HCl and 1.5 mM MgCl₂ was added to make...
a total volume of 10 µl. The reaction consisted of 40 cycles of 30 s at 94°C, 30 s at 60°C, 1.5 min at 72°C, followed by an extension at 72°C for 10 min. Amplification product was confirmed as a 300-bp band in agarose gel electrophoresis at 80 V. The DNA-PCR for the 18-kDa protein was sensitive enough to detect as few as 10 *M. leprae* bacilli.

**Longitudinal comparison of signals from DNA-PCR for the 18-kDa protein by one dimensional densitometry**

*M. leprae* DNA samples from serial biopsies stored after MDT for 0, 6, 12 and 24 months were used to prepare DNA for one-dimensional analysis of bands on agarose gel. Samples were centrifuged at 125 g at 4°C and the supernate was centrifuged at 15 000 g for 20 min. The pellet was overlaid with 50 µl of lysis buffer (proteinase K 10 µl, 1 M Tris-HCl 40 µl, 50 µl of Tween 80 0.5%, DNAase- and RNase-free water 400 µl) and incubated at 60°C for 2 h, and then heat inactivated at 95°C for 10 min. To purify the *M. leprae* DNA, an equal volume of phenol:chloroform:isoamyl alcohol (25:24:1) was added and agitated vigorously. After centrifuging at 15 000 g for 10 min, an equal volume of chloroform:isoamyl alcohol (24:1) was added to the supernate and the sample was re-centrifuged. One-tenth of the volume of 5 M NaCl and three volumes of absolute ethanol were added to the supernate and the tubes were stored at −70°C overnight. Sample were then centrifuged at 15 000 g for 20 min, the precipitate was dried and rehydrated with 20 µl of distilled water and heated at 65°C for 5 min. Ten µl of the precipitate were used as template DNA.

The thickness of bands on agarose gel electrophoresis was compared semi-quantitatively for samples obtained 0, 6, 12 and 24 months after MDT with Bio-1D V96 software for Windows 95 (Vilber Lourmat, Marne La Vallee, France).

**RT-PCR for mRNA of the 18-kDa protein**

Total RNA of the *M. leprae* of infected nude mice and wedge or punch biopsy specimens were separated with Trizol (GibcoBRL Life Technologies, Grand Island, NY, USA) by the manufacturer’s protocol. RNA contents were measured at 260 nm with a UV/VIS spectrophotometer 8450 A (Hewlett Packard, Palo Alto, CA, USA). To decontaminate genomic DNA, DNase I (Boehringer Mannheim, Mannheim, Germany) and RNase inhibitor (Promega, Madison, WI, USA) were added to 1 µg of RNA and incubated at 37°C for 1 h. The pure RNA fraction was extracted with phenol:chloroform:isoamyl alcohol and precipitated with ethanol.

The cDNA was synthesised from 1–5 µg of RNA with 250 ng of random primer (Sigma) for 2 min and cooled rapidly in ice. The reaction mixture, with 0.5 mM dNTPs, 10 mM DTT, RNase inhibitor (Promega) 100 units and H-superscript II reverse transcriptase (GibcoBRL) 200 units was added and held at 42°C for 1 h and then heated to 94°C for 5 min. The 18-kDa PCR confirmed that the cDNA came from the RNA of *M. leprae*. The single strand of the DNA was synthesised by the addition of 50 mM Tris-HCl, 5 mM MgCl₂, 10 mM (NH₄)₂SO₄, 10 mM DTT, 1.5 mM dNTPs, 7 U of RNase H and 1 U of DNA polymerase, and incubation at 16°C for 4 h.

To validate the effectiveness of RT-PCR for 18-kDa mRNA of *M. leprae*, it was compared with the results of RT-PCR for the 16S rRNA [12] with serial dilution of *M. leprae* from infected nude mice.

**IgM antibody titre for PGL-1**

IgM antibody titre for PGL-1 antigen was determined by the ELISA method described previously [4] with sera from patients on MDT. The absorbance was read at 492 nm with a Thermonax plate reader (Molecular Device, Sunnyvale, CA, USA). Values >0.200 were considered positive.

**Results**

A longitudinal study of the changes in *M. leprae* in biopsy specimens was conducted by DNA-PCR and RT-PCR for the 18-kDa protein. The DNA-PCR had a detection limit of 10 *M. leprae* bacilli (Fig. 1). Twenty-two patients who had a positive BI of >3 at the start of MDT gave positive 18-kDa PCR results. The PCR result remained positive at 0 month (22/22 patients), 6 months (21/22), 12 months (14/22) and 24 months (6/20) after MDT (Table 1). There were no positive PCR results in patients with a BI score of ≤1 during treatment (Table 2). In the case that showed a negative PCR result but a BI of 4, the PGL-1 titre supported the high score (Table 2, patient no. 19). Thus, there was not a close correlation between BI and PCR positivity.

However, when one-dimensional densitometry of the bands from serial biopsies was performed, it demonstrated a clear and gradual decrease after MDT for 12 and 24 months (Table 2). When biopsy specimens with the highest BI from each patient were chosen for analysis, four of 11 patients had a highest BI of ≤2.0 for the specimens. Of the specimens from the four patients, one was PCR positive and the others were PCR negative (Table 3). There is a slight correlation between BI and PCR results.

The RT-PCR method based on the 18-kDa mRNA showed successful RNA separation and synthesis of cDNA as confirmed by the amplification of the gene for the 18-kDa protein. The results of the RT-PCR
showed a detection limit of $\geq 10^2$ M. leprae bacilli, which was more effective than the results from RT-PCR, that targets 16S rRNA with serial dilution of M. leprae [17]. RT-PCR for the 18-kDa gene showed gradual reductions in band intensity during MDT over 0, 3, 6 and 12 months (Fig. 2). There were marked changes in band thickness in RT-PCR of patient nos. 16 and 18, but no change in the intensity of bands was found in patient no. 31 after MDT for 0, 3 and 12 months (Fig. 2).

Discussion

The present study shows that DNA-PCR for the 18-kDa gene detected M. leprae DNA for up to 24 months qualitatively and serial decreases of band density from biopsies semi-quantitatively. The introduction of PCR improved the diagnosis and follow-up of MDT. PCR has an advantage over BI and IgM antibody titre for PGL-1 because it provides proof of the changes in the numbers and the viability of M. leprae [13]. Others have reported a steep decrease in positive PCR results in 8 of 13 patients at 6 months and in 5 of 11 patients at 24 months [14]. Tissue fluid from a slit smear was often used as an alternative to skin biopsy, but the PCR results were observed to fall faster than when biopsy specimens were used [15, 16].

The differences noted above could depend on the method used to separate the DNA and the target DNA to be amplified. While other studies separated DNA from frozen sections, the present study used biopsy specimens; and with respect to the amplified gene, the gene for the 18-kDa protein was targeted. However, others have used the pra gene for the 36-kDa protein. Recently, Donoghue et al. reported that PCR based on the amplification of the gene for the 18-kDa antigen rather than the 36-kDa antigen has an advantage in terms of its lower detection limit for M. leprae [17]. For routine 18-kDa PCR, the present study adopted physical methods (freezing-thawing) to extract the M. leprae DNA, but to compare the thickness of bands from serial biopsies a proteinase K and extraction methods were used, which gave more favourable results than the physical method.

DNA-PCR identified only the presence of M. leprae, but it did not determine the viability of the organisms. Recently, RT-PCR for 16S rRNA of M. leprae, which evaluates the viability of M. leprae in the lesion after MDT, was developed [12, 18]. RT-PCR for mRNA of the 18-kDa gene gave clearer band changes than 18-kDa PCR during MDT. RT-PCR showed a gradual decrease in the intensity of PCR bands over 0, 3, 6 and 12 months of MDT (Fig. 2). This means that 18-kDa mRNA is one of the well-conserved RNAs like 16S rRNA. The results showed a step-wise reduction in the strength of bands after MDT for 1 year. The other method, NASBA, which targets 16S rRNA, resulted in a steep decrease in 5 (19.2%) of 26 patients at 6 months and in 1 (16.7%) of 6 patients (16.7%) at 12 months [19]. The cause of differences in results seems to be associated with the frozen section specimen preparation used.

The MDT regimen used was based on rifampin, clofazimine and DDS (dapsone). However, rifampin, ofloxacin and minocycline (ROM) were prescribed for patients who refused to take clofazimine fearing skin discoloration.

No change in the intensity of bands was found in patient no. 31 after 0, 3 and 12 months of MDT (Fig. 2). The patient was subjected to treatment with

Table 1. Results of DNA-PCR for the 18-kDa gene after MDT for 0–24 months

<table>
<thead>
<tr>
<th>Length of MDT (months)</th>
<th>Number of patients tested</th>
<th>Number (%) of patients with positive PCR result</th>
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<tbody>
<tr>
<td>0</td>
<td>22</td>
<td>22 (100)</td>
</tr>
<tr>
<td>6</td>
<td>22</td>
<td>21 (95.5)</td>
</tr>
<tr>
<td>12</td>
<td>22</td>
<td>14 (63.6)</td>
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<tr>
<td>24</td>
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<td>6 (30)</td>
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ofloxacin and was found to have a mutation in the gyrA of *M. leprae* in single-stranded conformational polymorphism analysis (data not shown). The gyrA gene is known as a target gene of the quinolones. Therefore, it is likely that the mutation in the *gyrA* induced the resistance to ofloxacin, one of the quinolones, and that no change was found in the intensity of bands.

It usually takes 6 or 12 months [16] for a patient to become PCR-negative after MDT and sometimes the patients show higher BI scores of up to 4–6 positive during the treatment periods, which represents a large discrepancy between the results of BI and PCR. PCR results seemed to be better correlated with the morphological index (MI) than the BI [13]. PCR titre with 36-kDa PCR which amplifies the DNA of the *pra* gene has demonstrated a rapid and steep decrease to zero between 12 months and 24 months after the initiation of MDT [13, 15]. These results are not consistent with the fact that the BI usually decreases after MDT to a level of $\log_{10} 1.7–1.9$ in MB cases within 12–24 months, from an average BI of 3.2–3.7 at the commencement of MDT [20].

Although the BI indicates the presence of *M. leprae* and its DNA fragments in the lesions, some PCR methods could not identify *M. leprae* in the biopsies. This could limit the use of PCR for assessment of MDT. However, the PCR methods used may not reflect the state of the DNA of *M. leprae* during the MDT. It is possible that certain DNA of *M. leprae* is more vulnerable to the drugs that were used in MDT. Application of RT-PCR promises to be a more useful tool in assessing the efficacy of MDT. Comparison of PCR results among longitudinal biopsies could show more dynamic changes in the numbers of *M. leprae* in the lesions.

It is well known that the removal of *M. leprae* and its DNA depends on macrophages in the leprosy granuloma. It usually takes 5–6 years to abolish all traces of *M. leprae* in MB cases. This conclusion is supported by the fact that the IgM antibody for PGL-1 persisted for several years after the cessation of MDT. We could not find any report about changes in the RNA of *M. leprae* after MDT, but it is plausible that some of the RNA can last for >1 year. This proposition is supported by the results of the present study, which showed a clear band of RT-PCR after MDT, and by the observations of Jamil *et al.*, who noted that the MI of patients was 1–3% at 12 and 24 months after MDT was started [14]. Thus, the results of the present study showed that 18-kDa PCR and RT-PCR for mRNA of the 18-kDa gene are more effective tools for following the efficacy of MDT.
than the BI or IgM antibody titre for PGL-1 in MB cases.

This work was supported by a grant (#HMP-96-M-2-0025) from the Good Health R Project, Ministry of Health and Welfare, ROK and the Catholic Medical Center Research Foundation in the programme year of 1999. We thank Mr Se-Kon Kim, Mr Pil-Soo Kang and Ms Hee-Suk Lee for their technical support.

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