Long Lasting Disease: Leprosy

Pramoda Earla

Department of Microbiology, Aditya Degree College [PG Courses], Andhra University, India

*Corresponding author: Pramoda Earla, Department of Microbiology, Aditya Degree College [PG Courses], Affiliated to Andhra University, Kakinada, East Godavari District, Andhra Pradesh, India, Tel: +91-7416948660; E-mail: pramodaearla@gmail.com

Rec date: Mar 20, 2015, Acc date: Mar 25, 2015, Pub date: Mar 27, 2015

Copyright: © 2015 Earla P. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Abstract

There are several infectious diseases from ancient times to which people got affected, suffered and died too. Leprosy can be regarded as the most infectious, transmittable and long lasting disease among all infectious diseases. The other name of leprosy is Hansen’s disease which was named after the physician Gerhard Armauer Hansen. The first causative agent of leprosy disease in humans is Mycobacterium leprae (M. Leprae) which has identified by microscopy technique. It is a rod-shaped gram-positive acid fast bacterium. The phenolic glycolipid 1 is the glycolipid material present in cell wall of this bacterium which generally shows immunological specificity in M. leprae. Survival of this acid fast bacterium in the host cell depends on the cell wall structure. Mycobacterium lepromatosis is a newly emerged leprosy-causing organism. It is emerging day by day as one of the major infectious diseases all over the world including developing countries. It is estimated that approximately 90% of the population develop protective immunity towards this disease, and, therefore, do not get sick after getting effected with this leprosy. Genetic and environmental factors are playing vital role in leprosy infection. The main symptoms are skin sores, bumps or lumps that will never go away after several weeks or months and which will become permanent if untreated foe a long time. It will mainly affect skin region. We cannot treat leprosy as a highly infectious disease. It is probably transmitting through droplets from the mouth and nose during close and frequent contacts with untreated patients. These bacteria mainly infect skin macrophages and Schwann cells in peripheral nerves. The involvement of autonomic fibers causes alteration in glandular functions. It will lead to dry mucous membrane and dry skin and which is responsible for the loss of tactile, thermal and pain sensibility. Incubation period of leprosy is usually two to four years with major manifestations. The Semmes-Weinstein technique is a widely used technique to evaluate plantar sensibility. Multi drug therapy (MDT) and early diagnosis are the key elements in eliminating the leprosy disease as a concern of public health. The ultimate aim is the development of a prophylactic vaccine, to protect against both drug-resistant and drug-susceptible strains. However, immunoprophylaxis for the leprosy disease continues to be largely speculative. The research in the area of leprosy remains an active area of scientific research.

Keywords: Leprosy; Mycobacterium leprae, Mycobacterium lepromatosis; Acid fast bacterium; Microscopy; Clinical susceptibility; Pedigree analysis; Genetic marker analysis; Twin studies; Linkage studies; Association studies; Skin macrophages; Schwann cells; Peripheral nerves; Muscle atrophy; Anti-leprosy agents; Early diagnosis; Azathioprine; Cyclosporine A; Corticosteroids

Abbreviations

DLL: Diffuse Lepromatous Leprosy; MDT: Multi Drug Therapy; SNP: Single Nucleotide Polymorphism; T-lep: Tuberculoid leprosy; L-Lep: Lepromatous Leprosy; CMI: Cell Mediated Immunity; SW Test: Semmes-Weinstein Test

Introduction

There are several infectious diseases from ancient times to which people got affected, suffered and died too [1]. Leprosy can be regarded as the most infectious, transmittable and long lasting disease among all infectious diseases. It is a chronic and granulomatous disease, mainly caused by two bacteria named Mycobacterium leprae and Mycobacterium lepromatosis. The other name of leprosy is Hansen’s disease [2-11] which was named after the physician Gerhard Armauer Hansen.
M. leprae is having a cell envelope made up of plasma membrane and cell wall made up of lipid-rich outer layer when examined under electron microscope. The phenolic glycolipid 1 is the glycolipid material present in cell wall of this bacterium which generally shows immunological specificity in M. leprae. Survival of this acid fast bacterium in the host cell depends on the cell wall structure [3].

**Mycobacterium lepromatosis**

*Mycobacterium lepromatosis* is a recently emerged leprosy-causing acid fast bacterium (Figure 2). Preliminary phylogenetic analysis of 16S rRNA and few other gene segments revealed significant divergence from *Mycobacterium leprae* as a well-known causative agent of leprosy that justifies the status of *M. lepromatosis* as a new emerged species [13]. *M. lepromatosis* caused not only all diffuse lepromatous leprosy (DLL) cases specifically but also more cases of lepromatous leprosy and other clinical forms of leprosy [14].

It is emerging day by day as one of the major infectious diseases all over the world including developing countries.

**Genetic and Environmental Factors**

Genetic and environmental factors are playing vital role in leprosy infection. There are different approaches to investigate genetic mechanisms of resistance which includes pedigree analysis, genetic marker analysis, twin studies, investigation of particular genes, etc. Twin studies are playing vital role and they are showing higher incidence of leprosy in monozygotic when compared to dizygotic twins. This study is indicating the important role of genetics in causing different types of infectious diseases [3].

Linkage studies and association studies are very useful in the investigation of different types of genetic diseases. Linkage studies are mainly dealing with gene mapping experiments, whereas association studies are dealing with allele frequencies comparison of a particular gene. This analysis generally occurs between patients and healthy individuals. Most of these studies are using single-base polymorphisms as markers for investing genetic disorders. SNP is a single nucleotide polymorphism which can lead to alterations in the functions and structures of a protein [3].

**Immunology of Leprosy Disease**

Leprosy is a chronic disease caused by an acid fast bacterium named *Mycobacterium leprae* (*M. leprae*) which shows a broad spectrum of clinical features. Tuberculoid leprosy is at one end and lepromatous leprosy is at the other end of the spectrum. T-lep patients will generally show high levels of cell-mediated immune responses against *M. leprae*, which will results in resistance to leprosy infection with very less clinical manifestations. On the other hand, L-lep patients will show very low cell-mediated immune responses against the leprosy bacteria as well as the developed form of the disease [17].

**Symptoms**

The main symptoms of leprosy disease include disfiguring of skin sores, bumps or lumps that do not go away after several months. These symptoms will become permanent if they left untreated. It will mainly affect skin region. There are three main types of the disease: Tuberculoid (a mild form of leprosy); Lepromatous (a more severe form of the disease, where kidneys and male reproductive organs may also affect); and Borderline (patients with symptoms of both tuberculoid and lepromatous forms) [18].

**Transmission**

We cannot treat leprosy as a highly infectious disease. It is probably transmitting through droplets from the mouth and nose during close and frequent contacts with untreated patients. Nasal secretions from the lepromatous patients can yield ten million viable organisms per day. All the people who become infected will not develop leprosy disease (genetic factors are influential). People who are living in endemic areas are at highest risk with poor conditions such as insufficient diet, contaminated water and inadequate bedding. These
people will also subject to other diseases that decrease or compromise immune function [19].

Loss of Sensibility

These bacteria mainly infect skin macrophages and Schwann cells in peripheral nerves [16,20]. The involvement of autonomic fibers causes alteration in glandular functions. It will lead to dry mucous membrane and dry skin and which is responsible for the loss of tactile, thermal and pain sensibility. The damaged motor fibers are responsible for the abolition of muscular response or muscle atrophy. The detection of protective sensory losses becomes very important to identify peripheral neuropathy and also to avoid plantar ulcer development and eventual amputation of lower limbs. The injury of sensory fibers is not a unique feature of leprosy. Other diseases such as vascular disorders, diabetes and different neuropathies of trauma can also generate plantar sensory loss [19].

Symptoms during Incubation period

Incubation period of leprosy is usually two to four years with major manifestations of skin lesions, muscle atrophy, numbness, photophobia, nasal stuffiness and blurred vision [21].

Diagnosis

Semmes-Weinstein test

The Semmes-Weinstein technique is a widely used technique to evaluate plantar sensibility. This test consists of nylon wires of same size and of different diameters with a variation of strength 0.05 g to 300 g and the order of associated colors are: green, blue, violet, dark red, orange, magenta and red. The protective sensation loss in the feet and hands is primarily indicated by the lack of response to the stimulus of the violet, blue and green filaments [19].

*M. leprae* which is infectious to human can be considered very critical microorganism as it cannot be cultured in artificial media. Resistance to anti-leprosy drugs such as rifampicin, dapsone and ofloxacin evolves by an amino acid substitution at the binding sites of these drugs [19]. There is an urgent requirement to discover new anti-leprosy agents. Although the available treatment for leprosy is effective, it is also quite expensive. Recent advances in biological sciences and computational approach are allowing our researchers to understand and also to design drug targets which are safe and inexpensive. Pharmacophore development, structure based optimization and virtual screening techniques are playing vital role in designing drugs [2].

Treatment

One of the major problems encountered for reducing the incidence of leprosy disease is late diagnosis which will lead to active infectious diseases [3]. MDT and early diagnosis are the key elements in eliminating the leprosy disease as a concern of public health. The approach of MDT consists of three drugs: rifampicin, dapsone and clofazimine. Dapsone (diaminodiphenylsulfone) was the first developed leprosy drug and it remained effective for a long time until the bacteria develops resistance. MDT required to be taken over a twelve month period for the active treatment. In addition, to suppress the cellular immunological response, Immunosuppressants such as cyclosporine A and azathioprine can be used in association with corticosteroids [19]. MDT protocol depends on combinatorial antibacterial effect of three chemotherapeutic agents, clofazimine, dapsone and rifampicin. This combination treatment is administered for six to twenty four months under partial medical supervision (Table 1) [21].

**Table 1: MDT protocol for MDT leprosy** [21].

<table>
<thead>
<tr>
<th>Clinical diagnosis</th>
<th>Drugs</th>
<th>Dose</th>
<th>Mode of administration</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pauicbacillary</td>
<td>Rifambicin</td>
<td>600 mg</td>
<td>Once a month/ supervised</td>
<td>1-2 years and could be extended</td>
</tr>
<tr>
<td></td>
<td>Dapsone</td>
<td>50 mg</td>
<td>Once a month/ supervised</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>50 mg</td>
<td>Daily/self administered</td>
<td></td>
</tr>
<tr>
<td>Multibacillary</td>
<td>Rifambicin</td>
<td>600 mg</td>
<td>Once a month/ supervised</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Clofazimine</td>
<td>300 mg</td>
<td>Once a month/ supervised</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>50 mg</td>
<td>Daily/self administered</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dapsone</td>
<td>100 mg</td>
<td>Once a month/ supervised</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>100 mg</td>
<td>Daily/self administered</td>
<td></td>
</tr>
</tbody>
</table>

**Table 1: MDT protocol for MDT leprosy** [21].

Conclusion

One strand of leprosy research is focused on detection of disease to enable earlier treatment and the other strand of research is focused towards the assessment of patients for post-treatment. The ultimate aim for the leprosy research is the development of a prophylactic vaccine which will protect against both drug-resistant and drug-susceptible strains. However, immunoprophylaxis for the leprosy disease continues to be largely speculative. This is due to problems with culturing the infectious agent. The research in the area of leprosy remains an active area of scientific research [18].

References


