Launching Of Immunization with the Vaccine
Mycobacterium Indicus Pranii for Eradication of Leprosy in India

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Received: 16 June, 2017; Accepted: 26 June, 2017; Published: 29 September, 2017

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Abstract

Vaccines are developed to mobilize the immune system of the recipient to react against an invading micro-organism and thus prevent infection to occur in the body. Besides immuno-prophylaxis, several vaccines have also revealed their precious immuno-therapeutic properties. Yet another highly important function of vaccines is their utilization for “Eradication” of a disease. Eradication of small pox could not have been achieved without a vaccine. The same is being achieved for elimination of Poliomyelitis in several countries with the massive immunization of the children with the available vaccines, oral and/or injectable.

Keywords: Tuberculosis; Ano-genital warts ; Myeloma ; Adjuvant

Introduction

This article reports a historically important step that the Ministry of Health, Govt of India, has taken on May 7, 2017, to launch an immunization project to eradicate leprosy from India.

Leprosy is not native to India. The causative micro-organism Mycobacterium leprae was in fact discovered by Armauer Hansen in Norway two centuries back. Although armadillos can host M. leprae, humans appear to be the main reservoir of infection. M. leprae grows in multibacillary leprosy patients, who shed the bacteria to infect others around them. Luckily 99% of humans are resistant to invading M. leprae (because of their competent immune system). The few who fall victim, sustain M. leprae in the community.

India is one of the few countries of the world, where leprosy is still prevalent. In fact in 2015, a total of 127,326 new cases were identified which account for 60% of the global total of new cases (www.who.int/lep/en). The Govt of India has been employing a multi-drugs regime to treat free of charge leprosy patients. This has brought down the prevalence rate significantly over the years.

However since 2006, the prevalence rate is static despite intensive programme of treatment with drugs. Moreover the incidence of new cases is on increase (Figure. 1).

An expert committee of the Indian Council of Medical Research along with the Directorate of Health Services has decided to employ the vaccine based on autoclaved Mycobacterium indicus pranii (MIP) in addition to Drugs for treatment of leprosy patients. Their family members and immediate contacts will also be immunized with 2 doses of the vaccine at 6 months interval. The Project will initially be launched in 5 districts of high endemicity and extended to other districts on basis of the observed efficacy of immunization with this vaccine.

Mycobacterium Indicus Pranii (MIP)

MIP vaccine was developed by me along with my co-workers to overcome the inability of leprosy patients to react to some key antigens of M. leprae. Extensive studies were carried out to understand the nature of immune deficit in those who contract the disease on exposure to M. leprae. The Golden Jubilee issue of Leprosy in India is entirely devoted to these studies [1]. The patients respond normally to Cholera or Typhoid vaccines [2]. Their immune deficit is primarily their inability to react against M. leprae. Their lymphocytes do not undergo blast transformation in presence of M. leprae and do not generate cytokines influencing macrophages to permit or not permit the multiplication of ingested M. leprae.

Leprosy patients manifest a spectrum. The multi-bacillary lepromatous leprosy (LL) patients have extreme immune deficit and are loaded with M. leprae, whereas the tuberculoid leprosy (TT) patients have limited lesions with hardly any bacilli. (Table 1) describes an experiment which we performed employing lymphocytes isolated from lepromatous (LL) patients or from tuberculoid (TT) leprosy patients. M. leprae require host cells, which are the macrophages to multiply. Macrophages were developed from glass adhering monocytes in peripheral blood. It hardly mattered whether these originated from LL, TT patients or normal human volunteers. In order to assess the permissive multiplication of M. leprae, we took advantage of the fact that macrophages do not multiply in vitro, and do not synthesize...
DNA, whereas any microorganism, such as \textit{M. leprae} must synthesize DNA for replication. Thus a pulse of 3H-Thymidine could be employed to decipher the multiplication of \textit{M. leprae} in a given situation. (Table 1) shows the important role of cytokines generated by ‘competent’ lymphocytes in response to \textit{M. leprae} to limit or permit DNA synthesis as an index of permissive multiplication of \textit{M. leprae}. While lymphocytes of LL patients do not restrict \textit{M. leprae} to multiply, lymphocytes of TT patients restrict their multiplication.

For development of vaccine against leprosy, we adopted a heterologous approach, as the very nature of immune deficit is the inability of susceptible individuals to react to \textit{M. leprae}. Thus using \textit{M. leprae} as the basis of vaccine was illogical. We collected 16 strains of atypical mycobacteria, many were non-pathogenic and all of them were cultivable in vitro. Each one of them was investigated for its ability to cause blast transformation of lymphocytes from lepromatous leprosy patients. Five strains of mycobacteria, namely \textit{M.vaccae}, \textit{M. phlei}, \textit{M.gordonae}, ICRC bacillus and unidentified mycobacteria from atypical collection, which was coded by us as Mw during.

Investigations were shortlisted for possessing the requisite capability \cite{4}. Lepromin like preparations were made from these mycobacteria and evaluated for eliciting Delayed Hypersensitivity skin responses in both tuberculoid (TT) and lepromatous leprosy (LL) patients \cite{5}. On basis of its ability to evoke not only positive response in TT but also in LL patients, Mw was chosen from these 5 strains for further investigations. Can it convert a lepromin negative LL patient who continue to be lepromin negative to lepromin positivity status? Dr. Choudhary at the School of Tropical Medicine, Kolkata, conducted investigations.
in 32 individuals who were initially patients suffering from lepromatous leprosy and who were rendered bacterial negative after prolonged treatment with drugs. All of them continued to be lepromin negative, as is usually the case with LL patients. Drugs kill bacteria, but do not improve the immune status of the patient. After immunization of these patients with autoclaved Mw vaccine, surprisingly 20 of them became lepromin positive and what was more, this trait of positivity was retained on retest after several months. [5]

**Properties of MIP**

MIP as adjunct to the standard Multidrugs regime accelerates bacterial clearance and shortens the recovery period. [6] What is further impressive is that it clears granulomas. Patients recover into almost normal beings without the usual blemishes of the disease. (Figure. 2) shows a few patients who were treated with MIP along with the drugs.

MIP clears also the bacilli resident in peripheral nerves restoring normal sensitivity.

**Figure 2:** Some representative cases of LL/BL multibacillary patients treated with MDT plus MIP (*Mycobacterium indicus pranii*) (6).
Additional Properties of MIP

i. Tuberculosis

MIP shares antigens with not only *M. leprae* but also *M. tuberculosis*. It has been employed successfully in treatment of Category II “Difficult to treat” tuberculosis patients. [7]

ii. Potent invigoration of both cellular and humoral immune response

MIP being used as adjuvant in our Birth Control Vaccine against hCG, where its inclusion enhances the antibody titres substantially. [8]

iii. Clearance of ugly ano-genital warts and lesions elsewhere on the body

Given intralesionally, MIP clears dramatically ano-genital warts [9] and also lesions elsewhere such as feet. (Figure 3) [10]

iv. Preventive and therapeutic action on SP2/O Myelomas and other cancers

Prof. Dipankar Nandi at Indian Institute of Sciences, Bangalore, has reported both preventive and therapeutic action of MIP on SP2/O myelomas in mice. [11]

![Figure 3: Cure by MIP of warts on feet. (A) Before treatment and (B) After 5 months of treatment with MIP. (10).](image)

Being an approved product by the Drugs Controller General of India (DCGI), and marketed by a Company to whom technology has been transferred, it is being used by patients suffering from various types of cancers.

Genome sequence of MW (now named MIP)

A network of 3 laboratories in India headed by Prof. SE Hasnain, Prof. Anil Tyagi and Prof. Akhilesh Tyagi have determined the genome sequence of Mycobacterium w [12,13]. Being a hitherto un described microorganism in International Collection, it has been named as *Mycobacterium indicus pranii*. Pran is my familiar name and nii is the National Institute of Immunology, New Delhi, of which I was the Founder Director, where much of the work on MIP was done under my guidance.

Summary

Described briefly in this communication is the background leading to the development of an immunotherapeutic cum immuno-prophylactic vaccine against leprosy. The vaccine is an autoclaved suspension of non-pathogenic mycobacteria, whose genome sequence is now known and has been named as *Mycobacterium indicus pranii* (MIP). Along with drugs, it accelerates bacterial clearance and shortens recovery period of leprosy patients. It converts nearly 98% of normal lepromin negative healthy contacts to lepromin positivity status. On May 7, 2017, the Government of India has launched a field programme to eradicate leprosy in leprosy endemic districts where the patients will be immunized with the vaccine besides the usual drugs. Family members and contacts of the patients will receive 2 doses of the vaccine at 6 months interval.

Besides leprosy, MIP has been employed beneficially for treatment of Category II “Difficult to treat” tuberculosis patients. MIP given intralesionally clears ugly warts on various parts of the body. MIP is a potent invigorator of tuberculosis response and is being used as adjuvant in a vaccine against hCG for preventing unwanted pregnancies. MIP has also demonstrated preventive and therapeutic action against some cancers. MIP is approved by the DCGI and is being made available to public by a company to which it is licensed.
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References