Acid-Fast Bacilli Found in Sphagnum Vegetation of Coastal Norway Containing Mycobacterium leprae-specific Phenolic Glycolipid-I

Jindrich Kazda, Lorentz M. Irgens, and Arend H. J. Kolk

The existence of noncultivable acid-fast bacilli (AFB) in sphagnum moss vegetation of former leprosy regions in coastal Norway has previously been reported (1). After the inoculation of these bacilli into nine-banded armadillos, antibodies reacting with Mycobacterium leprae-specific antigen 7 (as tested by Prof. Harboe, Oslo) were identified, strongly suggesting that these mycobacteria were M. leprae.

In an attempt to further identify these mycobacteria, additional samples of sphagnum vegetation were collected in previously leprosy-endemic areas of western Norway. The AFB in these samples were examined with monoclonal antibodies against the phenolic glycolipid-I (PGL-I) of M. leprae (2) and in the nude mouse model. In addition, to assess the possibilities of M. leprae propagating in such biotopes, M. leprae was inoculated into samples of this sphagnum vegetation and its survival was monitored over a period of time.

MATERIALS AND METHODS

In Naustdal Parish of Sunnfjord, Norway (61°30’N, 5°30’E), five sphagnum biotopes were selected according to criteria established by Irgens, et al. (4) for the selection of sphagnum biotopes in which a high concentration of AFB might be expected. Each sample consisted of all the grey layer from an area of about 1 square meter of well-growing Sphagnum rubellum, S. magellanicum, and S. imbricatum. The samples were handled with sterile plastic gloves, and transported within 14 days to the laboratory where the material was stored at +4°C until processed.

From the five biotopes, a total of nine samples was collected in the beginning of July 1986. Sterile plastic syringes were filled with approximately 100 cc of the grey layer, compressed, and the fluid collected. The compressed vegetation was weighed for the calculation of AFB count/g. For further processing, the fluid was centrifuged at 8000 × g × 30 min, and the sediment diluted in 2 ml of M/30 phosphate buffer, pH 7.0.

The first examination of the sediment took place within 4 weeks after the collection; the second, after the vegetation had been incubated for a period of 2 months at 34°C and 22°C in 12-hr pulses to simulate conditions optimal for the growth of noncultivable AFB (6). In both examinations, the number of AFB were counted. In the second examination, the AFB showing a positive reaction with M. leprae-specific monoclonal antibodies (MAbs) were counted separately. In this reaction, an indirect immunofluorescence technique (IFT) was used as previously described (7).

In the IFT procedure 10 spots were used on each slide, in duplicate, for each experiment. The control was coated with a suspension of armadillo-derived M. leprae (spots 1 and 2). The negative control contained M. avium serotype 2 (spots 3 and 4). Spots 5, 6, 7, and 8 were used for the experiments. They were coated with the suspensions to be examined. The last two spots served as further negative controls coated with M. leprae (spot 9) or M. avium serotype 2 (spot 10) but without any MAbs.

In the experiments, the M. leprae-specific MAb F47-21 against the PGL-I was used for spots 1, 3, 5, 6, and 7. In addition, MAb F85-2, specific for M. avium serotype 2, was used in spots 2, 4, and 8. For the evaluation
TABLE 1. Total acid-fast bacilli (AFB) counts per gram of fresh compressed sphagnum (grey layer).a

<table>
<thead>
<tr>
<th>Sample</th>
<th>AFB count before incubation</th>
<th>AFB count after incubation</th>
<th>Of which PGL-I-positive AFB</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Total</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>$3.3 \times 10^5$</td>
<td>$9.6 \times 10^5$</td>
<td>$1.7 \times 10^4$</td>
</tr>
<tr>
<td>2</td>
<td>$3.1 \times 10^5$</td>
<td>$8.4 \times 10^5$</td>
<td>$1.3 \times 10^4$</td>
</tr>
<tr>
<td>3</td>
<td>$3.3 \times 10^5$</td>
<td>$2.8 \times 10^5$</td>
<td>$1.4 \times 10^4$</td>
</tr>
<tr>
<td>4</td>
<td>$1.3 \times 10^5$</td>
<td>$1.9 \times 10^5$</td>
<td>$2.2 \times 10^4$</td>
</tr>
<tr>
<td>5</td>
<td>$3.1 \times 10^5$</td>
<td>$3.0 \times 10^5$</td>
<td>$2.4 \times 10^4$</td>
</tr>
<tr>
<td>6</td>
<td>$1.1 \times 10^5$</td>
<td>$1.7 \times 10^5$</td>
<td>$1.1 \times 10^4$</td>
</tr>
<tr>
<td>7</td>
<td>$2.0 \times 10^5$</td>
<td>$8.1 \times 10^5$</td>
<td>$1.7 \times 10^4$</td>
</tr>
<tr>
<td>8</td>
<td>$1.2 \times 10^5$</td>
<td>$1.6 \times 10^5$</td>
<td>$1.5 \times 10^4$</td>
</tr>
<tr>
<td>9</td>
<td>$2.8 \times 10^5$</td>
<td>$3.0 \times 10^5$</td>
<td>$1.4 \times 10^4$</td>
</tr>
</tbody>
</table>

a Total AFB counts and counts of AFB containing PGL-I as detected by MAbs in IFT after incubation for 2 months.

of PGL-I-positive bacilli, a pronounced immunofluorescence in spots 5, 6, and 7 together with a characteristic rod-shaped appearance was considered. The remaining spots were negative except for spot 1 (positive control for the known M. leprae suspension) and spot 4 (positive control for M. avium). Furthermore, 0.03 ml of the diluted sediment was injected into the right hind foot pads of nude mice (CD-1, nu/nu SPF) using five animals for each sample. At 1-month intervals, the swelling of the foot pads was measured and compared with the noninoculated left hind foot pads. Six, 8, and 9 months after inoculation, 1-2 animals were autopsied, and the inoculated foot pad was homogenized (*) and examined for the presence of AFB in IFT with MAb 47-21 of living cells (*) and of cultivable mycobacteria (Löwenstein-Jensen and Middlebrook 7H10 solid media). In three instances, IFT-positive homogenates were inoculated into nude mice.

To assess the potential of M. leprae to survive and multiply in the grey layer of the sphagnum, a sample from biotope 1 was chosen. Altogether, five series of examinations were performed in which 20-ml sterile plastic syringes were filled with sphagnum. In the first series, the syringes were inoculated with armadillo-derived M. leprae (*) from the liver of nine-banded armadillo no. 115 (originating from the Leprosy Tissue Bank, Research Institute, Borstel) suspended in 2 ml of sterile-filtered sphagnum fluid in a concentration of $5.9 \times 10^7$ M. leprae/cc of sphagnum. In the second series, the syringes were inoculated with M. leprae at a concentration of $5.9 \times 10^8$ M. leprae/cc sphagnum. Viability of the M. leprae cells was tested using the fluorescein diacetate (FDA) and ethidium bromide (EB) reactions (*). The third series was not inoculated and served as a control. In the fourth and fifth series, the sphagnum was heated for 30 min at 121°C and then inoculated as in the first and second series. From each series, two syringes were examined at 1, 2, 5, 6, 10, 12, and 16 weeks after inoculation.

RESULTS

In all nine sphagnum samples, AFB were found (Table 1). During incubation, the average AFB count increased from $2.35 \times 10^3$ to $3.15 \times 10^4$ of compressed vegetation. After incubation, PGL-I-positive AFB were
found in all samples with morphology and fluorescence similar to that of *M. leprae* (The Figure).

When inoculated into nude mice and examined over 9 months, no foot-pad swelling was observed. However, the concentration of PGL-I-positive AFB increased in 3 samples, was stable in 2 samples, and decreased in 2 samples. In 2 out of the 9 samples, no PGL-I-positive AFB were found after 6 months. FDA-positive AFB, implicating viable cells, were found at one or more examinations in all samples, except for the two samples without PGL-I-positive AFB after 6 months. In three of the samples, cultivable mycobacteria were present 6 months after inoculation; however, after 9 months no cultivable mycobacteria were found.

After inoculation into the grey layer of sphagnum vegetation, the count of *M. leprae* diminished during the first week (Table 2). During the next 15 weeks, a more than tenfold increase was observed which was more or less independent of the number of *M. leprae* inoculated. After 16 weeks of incubation, when inoculated into nude mice these AFB produced swelling similar to the pattern of *M. leprae*. The count of PGL-I-positive AFB in noninoculated sphagnum vegetation remained about the same throughout the culture period. These PGL-I-positive AFB did not produce foot-pad swelling in nude mice. When inoculated into heated sphagnum, *M. leprae* disappeared within 6 weeks.

**DISCUSSION**

The present results confirm, by a more specific method, our previous findings (5, 6) in sphagnum vegetation of viable microorganisms which cannot be differentiated from *M. leprae*. Certainly, the specificity of the method based on PGL-I may be questioned. However, until now, mycobacteria other than *M. leprae* have not been found to contain PGL-I on the cell surface, and the specificity of the monoclonal antibodies against PGL-I used in this study have been confirmed by a World Health Organization workshop (7). To assess the possibilities of *M. leprae* surviving in sphagnum vegetation, the growth of *M. leprae* was monitored in such vegetation over 16 weeks. It appeared that *M. leprae* incubated in nonheated vegetation multiplied more than tenfold and retained its pathogenicity for nude mice. Lack of foot-pad swelling in nude mice together with a limited multiplication suggest that these AFB may be of lower pathogenicity than the *M. leprae* obtained from clinical cases. Still, these AFB survive and to some extent multiply in nude mice. Whether a range in pathogenicity is a characteristic of *M. leprae*, perhaps also relevant to the differentiation in various clinical forms, needs further clarification.

The present findings add to the etiological role of environmental *M. leprae*. It has been found previously that in the health districts with the highest incidence rates of leprosy in Norway, a statistically significant association existed on a farm between the incidence of leprosy and the conditions relevant to the occurrence of mycobacteria in its environment (8). The gradual decline of leprosy in Norway certainly relates to a series of factors (9), but the lack of pathogenicity of naturally occurring organisms for nude
mice found in the present study is consistent with no new cases of leprosy having been recorded in Norway since the 1950s. If this mechanism was responsible for the decline, antibodies against PGL-1 might still be expected to exist in the Norwegian population. The demonstration of such antibodies would be evidence to support the hypothesis, and examinations aimed at the clarification of this issue should be pursued.

SUMMARY

In the grey layer of sphagnum vegetation originating from former leprosy-endemic regions of coastal Norway, acid-fast bacilli (AFB) containing *Mycobacterium leprae*-specific phenolic glycolipid I (PGL-I) on the surface have been found. These AFB survived in foot pads of nude mice with multiplication but without swelling. This contrasts to experimental leprosy with clinically derived *M. leprae* where swelling and unlimited multiplication takes place. The naturally occurring AFB may be of a lower pathogenicity than *M. leprae* obtained from clinical cases. The possibility of *M. leprae* surviving in sphagnum vegetation was assessed by inoculation of clinically derived *M. leprae* into the grey layer of the sphagnum. It multiplied more than tenfold and retained its pathogenicity in nude mice for 16 weeks, the duration of the experiment. The lack of pathogenicity of sphagnum-derived, *M. leprae*-like mycobacteria may be relevant to the decline of leprosy in Norway.

RESUMEN

Se han encontrado bacilos ácido-resistentes (BAAR) conteniendo en su superficie el glicolípido fenólico-1 específico del *Mycobacterium leprae* en la capa gris de la vegetación musgosa encontrada en las regiones anteriormente endémicas en lepra de las costas de Noruega. Estos BAAR sobrevivieron en las almohadillas plantares del ratón desnudo, con multiplicación pero sin hinchamiento. Esto contrasta con la lepra experimental causada por el *M. leprae* derivado de casos clínicos, donde ocurre hinchamiento y multiplicación ilimitada. Los BAAR encontrados en el musgo pueden tener una menor patogenicidad que los *M. leprae* obtenidos de casos clínicos. La posibilidad de que *M. leprae* sobreviviera en el musgo se analizó inoculando *M. leprae* de casos clínicos en la capa gris del musgo. Aquí, el *M. leprae* se multiplicó más de 10 veces y retuvo su patogenicidad en el ratón desnudo durante 16 semanas, la duración del experimento. La falta de patogenicidad de la micobacteria derivada del musgo, podría ser relevante en la disminución de la lepra en Noruega.

RÉSUMÉ

On a décédé des bacilles acido-résistants (AFB) contenant l’antigène phénoglycolipidique-1 (PGL-I) spécifique pour *Mycobacterium leprae*, à la surface de la couche grise d’une végétation de spheignes provenant de régions auparavant endémiques pour la lépre dans la région côtière de la Norvège. Ces bacilles acido-résistants ont survécu dans les coussinets plantaires de souris nues, en se multipliant, mais sans témoigner de gonflement. Ces observations ne correspondent pas à ce qu’ont observé dans la lépre expérimentale avec *M. leprae* obtenus chez des malades où le gonflement et une multiplication illimitée prennent place. Les bacilles acido-résistants naturels pourraient témoigner d’une pathogénicité plus faible que *M. leprae* recueilli à partir de lésions cliniques. La possibilité d’une survie de *M. leprae* dans la végétation de spheignes a été évaluée par l’inoculation au niveau des couches grises des spheignes de bacilles de la lépre obtenus chez des malades cliniques. Ces bacilles se sont multipliés plus de 10 fois et ont gardé leur pathogénicité chez la souris nue pendant 16 semaines, ce qui correspond à la durée de l’expérience. L’absence de pathogénicité des mycobactéries ressemblant à *M. leprae* et obtenues dans les spheignes pourrait avoir joué un rôle dans le déclin de la lépre en Norvège.

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REFERENCES


