Serological response of leprosy patients to *Mycobacterium leprae* specific and mycobacteria specific antigens: possibility of using these assays in combinations

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Summary The serological response of 147 leprosy patients to 3 mycobacterial antigens, PGL-I, 35 kDa (*Mycobacterium leprae* specific) and LAM (which is a common mycobacterial antigen) were analysed. A stronger serological response was seen amongst the MB patients than the PB patients in all the assays. The 3 antibody levels correlated positively with each other in both MB and PB cases. An overlap of seropositivity was seen between anti-PGL-I and anti-LAM ($p > 0.05$). A progressive increase in seropositivity and a significant difference of absorbance or titre in antibody levels in all 3 assays over increasing grades of BI were seen in the MB patients ($p < 0.05$). A significant difference in seropositivity between untreated and treated groups of patients was observed for anti-PGL-I ($p < 0.05$) and anti-LAM ($p < 0.01$) antibodies. The sensitivity, specificity and efficiency of anti-PGL-I (50%; 99%; 70%), anti-LAM (43%; 95%; 64%) and anti-35 kDa (66%; 100%; 80%) assays taken individually were less than that of combinations of anti-PGL-I/anti-35 kDa (74%; 99%; 84%) or anti-PGL-I/anti-35 kDa/anti-LAM (80%; 94%; 86%). The difference in the efficiency of both sets of combination of assays were not statistically significant ($p > 0.05$).

Introduction

In recent years the serological response of leprosy patients to different mycobacterial antigens has been studied. Species-specific capsular phenolic glycolipid-I (PGL-I),\(^1\)\(^2\) protein antigens like 35 kDa,\(^3\) 36 kDa\(^4\) and 18 kDa\(^5\) of *Mycobacterium leprae* and common mycobacterial antigens like lipoarabinomannan (LAM),\(^6\) a potent B cell stimulator, have been used in enzyme-immunoassays to study serological response
amongst leprosy patients. Strong serological responses to these antigens have been shown in the lepromatous (LL) pole of the spectrum and a weaker response in the tuberculoid (TT) pole. These studies were carried out in a mixed group of treated and untreated\(^2,4,7-12\) and also selectively in untreated patients\(^13\) of multibacillary (MB) leprosy. Such studies have also been done in paucibacillary (PB) leprosy\(^3,7,14-17\). Here, a cross-sectional study involving both MB and PB leprosy patients has been carried out to analyse: (a) the serological response of both MB and PB patients to 2 _M. leprae_ specific antigens (PGL-I and 35-kDa) and a common mycobacterial antigen (LAM); (b) the correlation of the antibody levels with the bacterial index (BI) in MB patients; (c) the association of the antibody levels with treatment status; (d) the efficiency of 3 assays individually and in combination.

**Materials and methods**

A total of 147 leprosy patients attending CLT & RI were included in this study, comprising of 91 lepromatous (LL), 10 borderline lepromatous (BL), 26 tuberculoid–borderline (TT–BT) and 20 indeterminate (IND) leprosy cases, all who had been clinically diagnosed. They consisted of both untreated and treated patients. In addition, 100 healthy subjects working in different departments of CLT & RI were included in the study as controls. Sera from all patients and controls were collected and stored at \(-20^\circ\text{C}\) until used.

A routine slit-skin smear was done for all leprosy patients and the average BI was calculated according to the Ridley’s scale.

**SEROLOGICAL ASSAYS**

**AntiPGL-I ELISA**

IgM antiPGL-I antibodies were measured by using the procedure reported earlier.\(^18\) Briefly D-BSA antigen and BSA supplied by IMMLEP/WHO, diluted in carbonate-bicarbonate buffer (CBCB) were coated in duplicate in a 96-well flat bottom microtitre plate (Dynatec Micro-ELISA system, Germany). Sera at 1:300 dilution were added to both antigen and BSA coated wells. After incubation, antihuman IgM peroxidase conjugate (DAKO, Denmark) was added at 1:2000 dilution. Colour was developed with \(\alpha\)-phenylenediamine (SIGMA, USA) with \(\text{H}_2\text{O}_2\) in citrate-phosphate buffer, pH 5.0. The reaction was stopped with 5 \(\text{NH}_2\text{SO}_4\). The plates were read at 492 nm in an ELISA reader (MR 600 micro plate reader, Dynatec). Samples with a difference in the mean absorbance between the antigen and BSA coated wells equal to or more than 0.200 OD were considered positive.

**Serum Antibody Competition Test—ELISA (SACT-E)**

Antibodies to the _M. leprae_ specific 35-kDa protein were detected by the protocol of J. Ivanyi, using peroxidase conjugated ML-04 monoclonal antibody (kindly supplied by J. Ivanyi, MRC, the Royal Postgraduate Medical School, London). Microtitre plates (Immulon, M129 B Dynatec) were coated with 10 \(\mu\text{g/ml}\) (50 \(\mu\text{l/well}\)) of _M. leprae_ soluble
Serological response of leprosy patients to specific antigens

extract (kindly supplied by R. J. W. Rees, IMMLEP M. leprae bank) in phosphate-buffered saline (PBS), pH 7.2, and incubated overnight at 4°C. After washing once with Tris-buffered saline, pH 7.4, containing Tween 20 (TBST), blocking was done with 150 μl/well of 1% skimmed milk powder in TBST (TBSTM). Sera in 10-fold dilution of 1:10, 1:100 and 1:1000 in TBSTM (25 μl/well) were added in duplicate and incubated for 1 hr. After removing the sera, 25 μl/well of appropriately diluted peroxidase conjugated ML-04 was added and incubated for 2 hr. After washing with TBST, colour was developed with o-phenylenediamine substrate solution (50 μl/well for 20 min). The reaction was stopped with 5NH4SO4 and reading was done at 492 nm in an ELISA reader. Positive control wells incubated with ML-04 conjugate without inhibiting sera were included in all the plates.

The dilution of the sera causing 50% inhibition of binding of ML-04 to the antigen (ID50) was calculated. Samples with ID50 titres equal to or more than 10 were considered SACT-E positive. All the 100 control sera tested were SACT-E negative.

AntiLAM ELISA

IgG antiLAM antibodies were measured by coating with LAM from M. tuberculosis H37Ra (kindly supplied by D. Chatterjee, Colorado State University, Fort Collins, Colorado, USA) at a concentration of 1 μg/ml, diluted in CBCB, pH 9.6 in duplicate. Control wells were coated with buffer (CBCB) alone. Sera at a dilution of 1:1000 were added (50 μl/well) to both antigen and control wells. After incubation for 2 hr, antihuman IgG peroxidase conjugate (DAKO, Denmark) at 1:6000 dilution was added. o-phenylenediamine with H2O2 in phosphate citrate buffer, pH 5.0, was used as a substrate solution. The reaction was stopped with 5 NH4SO4 and plates were read at 492 nm in an ELISA reader. Samples with a difference in the mean absorbance between the antigen and control wells equal to or more than 0.500 OD (mean ± 3 standard deviations of 100 controls) were considered positive.

STATISTICAL ANALYSIS

The Mantel–Haenszel Chi-squared method (cases, control as strata) was used to study the significance of differences between the antibody assays. The coefficients of correlation among 3 assays and between each assay and BI in individual patients were determined by Pearson’s correlation coefficient. The Mann–Whitney U test was done for the differences in the distribution of absorbances or titre between the different BI groups. The Kruskal–Wallis test was applied when more than 2 groups were compared. The significance of differences of seropositivity between MB and PB groups and between smear positive and smear negative MB cases were derived from 2 × 2 contingency tables by applying Yate’s corrected Chi-squared test. Probability (p) values <0.05 were considered as significant.

All the statistical tests were carried out using SSPS/PC+ and EPI-INFO software packages.

Results

SEROLOGICAL RESPONSE

The serological response among the MB group of patients was found to be uniformly
Table 1. Serological response of both MB and PB leprosy patients to 3 mycobacterial antigens.

<table>
<thead>
<tr>
<th>Patients (N)</th>
<th>MB (101)</th>
<th>LL (50) BI (+)</th>
<th>LL (41) BI (−)</th>
<th>BL (10) BI (+)</th>
<th>PB (46)</th>
<th>TT-BT (26)</th>
<th>IND (20)</th>
<th>Control (100)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PGL</td>
<td>58 (59)</td>
<td>74 (37)</td>
<td>34 (14)</td>
<td>80 (08)</td>
<td>33 (15)</td>
<td>27 (07)</td>
<td>40 (08)</td>
<td>1 (01)</td>
</tr>
<tr>
<td>LAM</td>
<td>44 (44)</td>
<td>52 (26)</td>
<td>24 (10)</td>
<td>80 (08)</td>
<td>41 (19)</td>
<td>35 (09)</td>
<td>50 (10)</td>
<td>5 (05)</td>
</tr>
<tr>
<td>35 kDa</td>
<td>90 (91)</td>
<td>100 (50)</td>
<td>76 (31)</td>
<td>100 (10)</td>
<td>13 (06)</td>
<td>12 (03)</td>
<td>15 (03)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Any of 3</td>
<td>93 (94)</td>
<td>100 (50)</td>
<td>83 (34)</td>
<td>100 (10)</td>
<td>52 (24)</td>
<td>46 (12)</td>
<td>60 (12)</td>
<td>6 (06)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>50</td>
<td>99</td>
</tr>
<tr>
<td></td>
<td>43</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>66</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td>94</td>
</tr>
</tbody>
</table>

Percentage of seropositive (absolute number in parentheses)

Significant difference of seropositivity of IgM antiPGL-I and anti35 kDa antibodies between MB and PB patients ($p < 0.01$).

Significant difference of seropositivity of IgM antiPGL-I and IgG antiLAM and anti35 kDa antibodies between smear positive and smear negative MB cases ($p < 0.001$).

Significant difference of seropositivity between antiPGL-I and 35 kDa in total ($p < 0.05$).

No significant difference of seropositivity between antiPGL-I and antiLAM in total ($p > 0.05$).

Higher than on the PB group of patients (Table 1). In MB cases, 58% had IgM antiPGL antibodies, 44% antiLAM antibodies and 90% anti35 kDa antibodies; 93% of patients in the MB group had any of the 3 antibodies. In the PB cases, 33% had antiPGL antibodies, 41% antiLAM antibodies and 13% anti35 kDa antibodies; 52% of patients in the PB group had any of the 3 antibodies. In the MB group, smear positive patients had a higher percentage of seropositivity (antiPGL-I, 75%; antiLAM, 57%; anti35 kDa, 100%) compared with smear negative (antiPGL-I, 34%; antiLAM, 24%; anti35 kDa, 76%) patients. In general, the 35 kDa antigen was found to elicit a serological response in a higher proportion of different types of the MB patients compared to the other 2 antigens. Within the PB group, apparently a higher percentage (antiPGL-I, 40%; antiLAM, 50%; anti35 kDa, 15%) of IND type patients showed a serological response to all the 3 antigens compared to the TT-BT patients (antiPGL-I, 27%; antiLAM, 35%; anti35 kDa, 12%).

Out of 100 controls employed in the study 1% had antiPGL-I antibodies, 5% antiLAM antibodies and none had anti35 kDa antibodies.

ASSOCIATION WITH BI

Of the 101 MB cases studied, 60 were smear positive with a BI ranging from 0.16 to 3.67.
Table 2. Relationship between serological responses to 3 mycobacterial antigens and BI in MB leprosy patients (% seropositive (mean absorbance/titre)).

<table>
<thead>
<tr>
<th>Bacterial index</th>
<th>No. patients</th>
<th>PGL</th>
<th>LAM</th>
<th>35 kDa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>41</td>
<td>34% (0.462)</td>
<td>24% (0.855)</td>
<td>76% (174.5)</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>20</td>
<td>65%* (0.622)</td>
<td>45%* (0.913)</td>
<td>100%* (223.9)</td>
</tr>
<tr>
<td>&gt; = 1 to &lt; 2</td>
<td>21</td>
<td>76%* (0.514)</td>
<td>52%* (0.946)</td>
<td>100%* (284.4)</td>
</tr>
<tr>
<td>&gt; = 2</td>
<td>19</td>
<td>84%* (0.579)</td>
<td>74%* (1.032)</td>
<td>100%* (673.6)</td>
</tr>
</tbody>
</table>

* Significant differences in the distribution of absorbance (or) titres of antibodies with BI positive group compared to BI negative group (Mann–Whitney, p < 0.05).

The increase in the absorbances (or) titres over the range of BI was significant for all the 3 antibody assays (p < 0.001) by Kruskal–Wallis analysis.

All the 3 assays showed progressive increase of seropositivity with an increase in BI. Such a trend was also observed in mean antibody titres (Table 2). All the 3 levels of the antibodies positively correlated with BI (anti35 kDa vs BI, r = 0.4638, p < 0.001; antiPGL-I vs BI, r = 0.3119, p < 0.01; and antiLAM vs BI, r = 0.3689, p < 0.001).

ASSOCIATION WITH THE TREATMENT STATUS

The multibacillary group consisted of 19 untreated and 82 treated cases. Amongst untreated cases, 84% had antiPGL-I antibodies, 88% antiLAM antibodies and 100% anti35 kDa antibodies. Amongst treated cases 52% had antiPGL-I antibodies, 38% antiLAM antibodies and 88% anti35 kDa antibodies (Table 3). The paucibacillary group consisted of 30 untreated and 16 treated cases. Amongst untreated cases 43% had antiPGL-I antibodies, 57% antiLAM antibodies and 13% anti35 kDa antibodies. Amongst treated cases 13% had antiPGL-I antibodies, and the same percentage of cases were also positive for the other 2 assays. Though the untreated cases had a higher percentage of seropositivity in all the 3 assays than did the treated, the differences were significant for antiPGL-I (p < 0.05) and antiLAM (p < 0.01) assays but not for the anti35kDa assay (p > 0.05).

CORRELATION BETWEEN THE ANTIBODY ASSAYS

In the MB group, all the 3 assays positively correlated with each other. There was a strong correlation between antiPGL-I and antiLAM antibodies (r = 0.5048, p < 0.001) and also between antiPGL-I and anti35kDa antibodies (r = 0.3917, p < 0.001). A moderate correlation was seen between antiLAM and anti-35kDa antibodies (r = 0.3172, p < 0.01).

In the PB group, a moderate positive correlation between the assays was observed (antiPGL-I vs anti-35 kDa, r = 0.4385, p < 0.01; antiPGL-I vs antiLAM, r = 0.3495, p < 0.05 and antiLAM vs anti35 kDa, r = 0.4183, p < 0.01).

Multibacillary patients with antiPGL-I antibodies were also more likely to have antiLAM antibodies (p < 0.05), but such an overlap was not seen between anti35 kDa and
Table 3. Serological responses to 3 mycobacterial antigens in untreated and treated groups of patients.

<table>
<thead>
<tr>
<th>Patients</th>
<th>Treatment status</th>
<th>PGL</th>
<th>LAM</th>
<th>35 kDa</th>
</tr>
</thead>
<tbody>
<tr>
<td>MB (101)</td>
<td>UNT (19)</td>
<td>84% (16)</td>
<td>88% (14)</td>
<td>100% (19)</td>
</tr>
<tr>
<td></td>
<td>TRT (82)</td>
<td>52%* (43)</td>
<td>38%† (31)</td>
<td>88%‡ (72)</td>
</tr>
<tr>
<td>PB (46)</td>
<td>UNT (30)</td>
<td>43% (13)</td>
<td>57% (17)</td>
<td>13% (04)</td>
</tr>
<tr>
<td></td>
<td>TRT (16)</td>
<td>13% (02)</td>
<td>13% (02)</td>
<td>13% (02)</td>
</tr>
</tbody>
</table>

* Differences in seropositivity of antiPGL between treated and untreated groups (p < 0.05).
† Differences in seropositivity of antiLAM between treated and untreated groups (p < 0.01).
‡ Differences in seropositivity of anti35 kDa between treated and untreated groups (p > 0.05).
UNT, untreated; TRT, Treated.

the other 2 antibodies. In the PB group as well, an overlap of seropositivity was only seen between anti-PGL-I and anti-LAM antibodies (p > 0.05). The seropositivity for any of the antibodies, when taken in combinations like antiPGL-I/antiLAM, antiPGL-I/anti35 kDa, or antiPGL-I/anti35 kDa/antiLAM, showed sensitivity and specificity higher than the corresponding assays taken individually (Figure 1). Thus, when the seropositivity for any of the 2 assays using the M. leprae specific antigens, namely, PGL-I and 35 kDa, were taken together, the combined sensitivity (74%) and specificity (99%), and thereby the

Figure 1. Sensitivity, specificity and efficiency of the assays, alone and in combinations. ■, sensitivity; □, specificity; △, efficiency,* Difference of seropositivity between combinations of antiPGL-I/anti-35 kDa and antiPGL-I/antiLAM (p > 0.05); difference of seropositivity between combinations of antiPGL-I/anti-LAM and antiPGL-I/anti-LAM/anti35 kDa (p < 0.01); difference of seropositivity between combinations of antiPGL-I/anti-35 kDa and antiPGL-I/antiLAM/anti35 kDa (p > 0.05).

* Efficiency (index of usefulness) = \[ \frac{\text{No. of true positives} + \text{No. of true negatives}}{\text{Total no. of subjects studied}} \]
efficiency (84%), was higher. Similarly, when the seropositivity for any of the 3 assays, using both the species specific (PGL-I and 35 kDa) and the genus specific (LAM) antigens was taken together, the combined sensitivity (80%) and efficiency (86%) was even higher, though there was a decline in specificity (94%). Yet the utility of both the sets of combination assays was comparable (p > 0.05).

Discussion

Many mycobacteria-specific and M. lepraespecific antibodies have been demonstrated in the sera of leprosy patients. Though they are not known to be protective, their presence in the sera of leprosy patients may be used as markers of bacillary load, monitors of prognosis of the disease and predictors of occurrence of future relapse or reactions. Several studies have been done against this background. In the present study, a cross-sectional analysis of 147 leprosy patients showed a stronger serological response in the MB group of patients than the PB group to all the 3 antibodies, but significantly to both M. lepraespecific antigens (PGL-I and 35-kDa). The response to 35 kDa antigen was higher in both smear-positive and smear-negative MB patients, with sensitivity being greater than in the other 2 assays. These findings are consistent with the observations of other researchers. In the PB group the sensitivity of 35 kDa was less, though others have shown higher sensitivity. However, all the 3 assays, when individually analysed, had a higher specificity with considerably lower sensitivity.

All the 3 antibodies studied correlated positively with each other, both in the MB and the PB cases. Further, an overlapping of positivity was seen only between antiPGL-I and antiLAM antibodies, but not between anti35 kDa and other antibodies. Thus, this specific 35 kDa epitope of M. leprae was observed to pick up, at least in the MB group, those leprosy patients who were negative for antibodies against the other 2 antigens.

All the 3 antibody assays showed a progressive increase of seropositivity and mean antibody titres over the increasing grades of BI, all the 3 antibodies positively correlated with BI as well. This correlation was highly significant for anti-35 kDa antibody. Thus, this positive association of antibody titres with bacterial load can be used to monitor the prognosis of patients in conjunction with BI. Hence, these assays would be complementary to BI in detecting additional cases of multibacillary patients.

Both in the MB and the PB groups, the seropositivity among untreated patients was higher than in the treated patients. This difference was significant for antiPGL-I and antiLAM antibodies only. But a relatively steep fall in the anti35 kDa antibody titre with least individual variability on treatment has been observed by others. Thus the antibody assays would seem to supplement the clinical evaluation in assessing the prognosis of patients on chemotherapy. Further prospective studies may throw some more light on this observation.

Amongst the PB group, ‘IND’ type patients showed apparently high seropositivity for all the 3 antibodies in comparison to ‘TT–BT’ type. This might suggest that most ‘IND’ leprosy patients are likely to progress to the MB type of leprosy, as has been observed by other reporters.

An assay using lipoarabinomannan of M. tuberculosis was used in this study because of its similarity with the LAM of M. leprae. Despite not being species-specific this antigen appears to be useful in the serology of leprosy. The sensitivity of this assay is
comparable to that of antiPGL-I assay. Further, this assay could differentiate between smear positive and smear negative MB cases and also between treated and untreated cases like antiPGL-I assay. AntiLAM antibodies correlated positively with M. leprae-specific antibodies in MB and PB cases. AntiLAM assay, when combined with other assays, increased the efficiency in both antiPGL-I/antiLAM and antiPGL-I/antiLAM/anti35 kDa combinations. Hence the value of antiLAM assay in leprosy serology.

Thus this analysis on the serological responses of leprosy patients to M. leprae-specific (PGL-I and 35 kDa) and a genus specific antigen (LAM) revealed that these antibody assays are complementary to clinical evaluation of leprosy patients. This was evidenced by a strong immunological response amongst leprosy patients to the 3 antigens, significant correlation with BI and with the treatment status of the patients. The assays, though individually highly specific, were less sensitive. This could be compensated by analysing the seropositivity for any of 2 M. leprae-specific antibodies (antiPGL-I/anti35 kDa) or any of the 3 antibodies (antiPGL-I/anti35 kDa/antiLAM) in combinations. In this way we obtain an improved application of the serological assays in the management of leprosy patients, until an antigen which is highly sensitive in detecting both MB and PB cases equally is developed.

Acknowledgment

We thank IMM/WHO for the supply of D-BSA antigen; Dr R. J. W. Rees for the provision of M. leprae soluble extract through IMM/WHO/UNDP/World Bank Tropical Disease Research Programme; Professor J. Ivanyi for providing ML-04 conjugate; and Dr D. Chatterjee for supplying LAM antigen. We give sincere thanks to Mr L. Rathinavel and Mr K. Kuberan, Laboratory Division, CLT & RI, for their dedicated technical assistance, and we thank Dr P. Krishnamoorthy, Epidemiology and Statistics Division, CLT & RI for the assistance in statistical analysis and Dr N. Elangeswaran, Laboratory Division, Dr A. V. Paul, Clinical Division and Dr P. S. Rao, Director-In-Charge of CLT & RI for their encouragement.

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Serological response of leprosy patients to specific antigens


Réponse sérologique des patients lépreux aux antigènes spécifiques de *Mycobacterium leprae* et spécifiques des mycobactéries: possibilité d’utiliser ces essais en association

B. Sekar, R. N. Sharma, G. Leelabai, D. Anandan, B. Vasanthi, G. Yusuff, M. Subramanian et M. Jayasheela

**Résumé** Nous avons analysé la réponse sérologique de 147 patients lépreux à 3 antigènes mycobactériens, PGL-I, 35 kDa (spécifique de *Mycobacterium leprae*) et LAM (qui est un antigène mycobactérien commun). Une réponse plus forte a été observée parmi les patients MB que parmi les patients PB dans tous les essais. Les taux des 3 anticorps étaient en corrélation positive entre eux à la fois dans les cas MB et PB. Une séropositivité croisée a été observée entre anti-PGL-I et anti-LAM \( p > 0.05 \). Une augmentation progressive de la séropositivité et une différence significative entre l’absorption ou le titre des taux d’anticorps dans les 3 essais sur des degrés croissants de BI ont été observés chez les patients MB \( p < 0.05 \). Une différence significative de la séropositivité entre les groupes de patients non-traités et traités a été observée avec les anticorps anti-PGL-I \( p < 0.05 \) et anti-LAM \( p < 0.01 \). La sensibilité, spécificité et efficacité des essais avec anti-PGL-I (50%; 99%; 70%), anti-LAM (43%; 95%; 64%) et anti-35 kDa (66%; 100%; 80%) pris séparément étaient inférieures à celles des associations anti-PGL-I/anti-35 kDa (74%; 99%; 84%) ou anti-PGL-I/anti-35 kDa/anti-LAM (80%; 94%; 86%). La différence entre l’efficacité des deux séries d’associations d’essais n’était pas statistiquement significative \( p > 0.05 \).

La respuesta serológica de los pacientes leprosos a antígenos específicos para *Mycobacterium leprae* y Micobacterias: la posibilidad de usar estos ensayos en combinaciones

B. Sekar, R. N. Sharma, G. Leelabai, D. Anandan, B. Vasanthi, G. Yusuff, M. Subramanian y M. Jayasheela

**Resumen** Se analizaron las respuestas serológicas de 147 pacientes leprosos a 3 antígenos micobacterianos, PGL-I, 35 kDa (específico a *Mycobacterium leprae*) y LAM, un antígeno micobacteriano común. Se observó una respuesta serológica más fuerte entre los pacientes MB que entre los PB, en todos los ensayos. Los tres niveles de anticuerpo correlacionaban positivamente entre sí, tanto en los casos MB como los PB. Se observó un traslape en los resultados de seropositividad entre anti-PGL-I y anti-LAM \( p > 0.05 \). Se observó un aumento progresivo de seropositividad y una diferencia significativa de absorbcencia o titulación de los niveles de anticuerpo en todos los 3 ensayos con crecientes tipos de BI en los pacientes MB \( p < 0.05 \). Se observó una diferencia significativa de seropositividad entre los grupos de pacientes sin tratar y tratados con anticuerpos anti-PGL-I \( p < 0.05 \) y anti-LAM \( p < 0.01 \). La sensibilidad, especificidad y eficiencia de los ensayos anti-PGL-I (50%; 99%; 70%), anti-LAM (43%; 95%; 64%) y anti-35 kDa (66%; 100%; 80%) tomados individualmente eran menores que las combinaciones de anti-PGL-I/anti-35 kDa (74%; 99%; 84%) o anti-PGL-I/anti-35 kDa/anti-LAM (80%; 94%; 86%). La diferencia de eficiencia entre ambos juegos de combinación de ensayos no era significativa \( p > 0.05 \).