A STUDY OF CLINICO-HISTOPATHOLOGICAL CORRELATION OF CUTANEOUS MANIFESTATIONS IN LEPROSY

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ABSTRACT

BACKGROUND

Leprosy is one of the oldest and chronic infectious diseases known to human beings caused by Mycobacterium leprae. The disease still carries a grave social stigma and ostracism, which compels the patients to hide the disease. Leprosy continues to be an important public health problem in most parts of Asia, especially India.1 Leprosy is a progressive, chronic granulomatous disease of the peripheral nerves and skin and other tissues such as mucous membranes, muscles and reticuloendothelial system. The disease presents in various clinico-pathological forms depending on the immune status of the host. The disease spectrum has been characterised in a number of classification systems, most widely being the Ridley-Jopling.

Aim- Fifty cases were taken to correlate clinical diagnosis with histopathological findings.

MATERIALS AND METHODS

All clinically suspected Leprosy patients attending Department of DVL were included in this study. History was taken in detail and complete examination of patients carried out, particularly with reference to skin, nerves and sensory disturbances. Slit skin smear was taken from the patients with specific findings. The biopsies were taken from the most active and untampered lesions including the margin of the lesion and sent to the Pathology Department in 10% neutral buffered formalin. The histopathological diagnosis was made based on the scheme put forth by Ridley and Jopling.

RESULTS

The present study comprised of 50 patients, 33 were male (66%) and 17 female (34%) with a male: female ratio of 1.9: 1. Table 1 shows the distribution of patients according to age group and gender. Majority of the patients (11 patients: 7 males and 4 females) were between 31-40 years of age, whereas least affected were below 10 years (2 female patients). The mean age of the patients studied was 41.34± 17.10. The range of youngest patient is 7 years old female and the oldest patient is 75 years old male. With regard to patient’s occupation, the largest group included are 20 farmers (40%) followed by 12 daily labourers (24%), students and housewife are equal percentage (12%), whereas only 2.0% of the patients are employees. The distribution of these cases based on Ridley and Jopling clinical and histopathological classification is shown in Table 4. It is clearly evident from Table 4 that clinically majority of the patients (40%) belonged to Lepromatous Leprosy (LL) group followed by borderline tuberculoid (BT) group (24%), tuberculoid leprosy (TT) group (14%) and mid-borderline (BB) group and indeterminate leprosy (IL) group with 6% to 4% each. Histopathologically, majority of the cases 32% belonged to Lepromatous Leprosy (LL) followed by BT (18%), BL (16%), TT and IL (12% each). Among cases with negative slit skin smear were 40 patients. 10 patient’s slit skin smears were positive in 20% of patients. Out of 20 patients, 2+ for 2 patients, 3+ for 7 patients and BI was maximum 4+ for 1 patient. The correlation between clinical and histopathological classification is shown in Table No. The overall concordance between the clinical and histopathological agreement was seen in 15 (34.5%) cases and maximum. Clinico-histopathological correlation was seen in BB (100%) followed by TT (57.1%), BT (50%), IL and BL (33.3%) and 0% in LL. The concordance rate was lower in the borderline groups with 33.3% in BL, 33% in IL and least concordance of 16.7% in LL. However, the concordance for TT was higher than the borderline groups with 57.1%. Histopathological analysis of the cases in the present study as shown in Table 3 was carried out with due attention to the epidermal atrophy, presence of clear subepidermal zone, dermal inflammatory infiltrate, presence and composition of granulomas, presence of giant cells and relative proportion of lymphocytes and foamy histiocytes in accordance with Ridley and Jopling histopathological criteria. Out of the 50 patients included in the study, 15 (30%) presented with a clinical suspicion of paucibacillary leprosy and 35 (70%) of multibacillary leprosy. From the chi-square output table, we see that no significance level has been achieved. McNemar Bowker test table showing no systematic association between the above two variables at 95% level of confidence. Hence, it concludes that there is no significant relationship between age wise patients and histopathological diagnosis.

CONCLUSION

Leprosy, though reported to be eliminated, still continue to be one of the common infectious diseases in India. Skin biopsy is the useful tool in clinical diagnosis of leprosy as well as therapeutic guide.

KEY WORDS

Mycobacterium Leprae, Leprosy, Histopathology, Ridley-Jopling Classification.

BACKGROUND
Leprosy is one of the oldest and chronic infectious diseases known to human beings caused by Mycobacterium leprae. The disease still carries a grave social stigma and ostracism, which compels the patients to hide the disease. Leprosy continues to be an important public health problem in most parts of Asia, especially India.1

Leprosy is a progressive, chronic granulomatous disease of the peripheral nerves and skin and other tissues such as mucous membranes, muscles and reticuloendothelial system. The disease presents in various clinico-pathological forms depending on the immune status of the host. The disease spectrum has been characterised in a number of classification systems, most widely being the Ridley-Jopling classification. In this classification, leprosy has been divided into five groups as Tuberculoid (TT), Borderline Tuberculoid (BT), Mid-Borderline (BB), Borderline Lepromatous (BL) and Lepromatous (LL).2,3

Diagnosis of leprosy is based on different clinical parameters, which involves detailed examination of skin lesions and peripheral nerves. Demonstration of acid-fast bacilli in slit skin smears by Ziehl-Neelsen’s staining also aids in diagnosis of leprosy. A reliable diagnosis hinges around a good histopathological diagnosis and demonstration of bacilli in histopathological sections.4,5 Clinical classification gives recognition only to gross appearances of the lesions, while the parameters used for the histopathological classification are well defined, precise and also take into account the immunological manifestations which enable it to successfully bridge the pitfalls in leprosy diagnosis. Histopathology provides confirmatory information for suspected cases, which can be missed in clinical practice or epidemiological studies and helps in exact typing.6

Aims and Objectives of the Study
1. To study the clinical pattern of the disease.
2. To correlate histopathological findings with AFB (Acid Fast Bacilli - M. Leprae) status.
3. To correlate clinical diagnosis with histopathological findings.

Source of Data
A hospital-based, clinical, observational study was conducted in the Department of Dermatology, Venereology and Leprosy. Fifty patients clinically diagnosed as having Leprosy were taken up for the study.

Sample Size
50.

RESULTS

The present study comprised of 50 patients, 33 were males (66%) and 17 females (34%) with a male: female ratio of 1.9:1. Table 1 shows the distribution of patients according to age group and gender. Majority of the patients (11 patients: 7 males and 4 females) were between 31 - 40 years of age, whereas least affected were below 10 years (2 female patients). The mean age of the patients studied was 41.34 ± 17.104. The range of youngest patient is 7 years old female and the oldest patient is 75 years old male.

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<td><strong>Total</strong></td>
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Mean Age: 41.34 ± 17.104

Table 1. Age and Gender Wise Classification of Patients Studied

NS- Not Significant, Chi-square test- P > 0.05

MATERIALS AND METHODS
Method of Collection of Data
All clinically suspected Leprosy patients attending Department of DVL were included in this study. The history was taken in detail and complete examination of patients carried out, particularly with reference to skin, nerves and sensory disturbances.

Slit skin smear was taken from the patients with specific findings. The biopsies were taken from the most active and untampered lesions including the margin of the lesion and sent to the Pathology Department in 10% neutral buffered formalin. The histopathological diagnosis was made based on the scheme put forth by Ridley and Jopling.

Inclusion Criteria
The study includes all clinically diagnosed cases of leprosy attending Department of DVL.

Exclusion Criteria
All cases of leprosy treated earlier. Patients with HIV infection.

Statistical Analysis
Statistical analysis of our study was performed using the software Statistical Package of Social Science version 20 (SPSS). The Clinical diagnosis of the leprosy cases (As provided by Department of Dermatology) using Ridley and Jopling scale correlated with the results of histopathological examination of their respective biopsies were analysed and matched using Kappa and McNemar test. A p < 0.05 was considered to be statistically significant. The data were reported as mean ± SD and frequency.
With regard to patient’s occupation, the largest group included are 20 farmers (40%) followed by 12 daily labourers (24%). Students and housewife are equal percentage (12%), whereas only 2.0% of the patients are employees.

TT= Tuberculoid; BT= Borderline Tuberculoid; BB= Borderline Lepromatous; LL= Lepromatous, IL= Indeterminate Leprosy.

The distribution of these cases based on Ridley and Jopling clinical and histopathological classification is shown in Table 4. It is clearly evident from Table 4 that clinically majority of the patients (40%) belonged to Lepromatous Leprosy (LL) group followed by Borderline Tuberculoid (BT) group (24%), Tuberculoid Leprosy (TT) group (14%) and Mid-Borderline (BB) group and Indeterminate Leprosy (IL) group with 6% to 4% each. Histopathologically, majority of the cases 32% belonged to Lepromatous Leprosy (LL) followed by BT (18%), BL (16%), TT and IL are (12 % each).

Among cases with negative slit skin smear were 40 patients. 10 patients slit skin smears were positive in 20% of patients. Out of 20 patients 2+ for 2 patients, 3+ for 7 patients and BI was maximum 4+ for 1 patient.

Kappa= 0.566. The strength of agreement is considered to be above moderate p= 0.000 (Significant).

The correlation between clinical and histopathological classification is as shown in Table No. 6. The overall concordance between the clinical and histopathological agreement was seen in 15 (34.5%) cases and maximum clinico-histopathological correlation was seen in BB (100%) followed by TT (57.1%), BT (50%), IL and BL (33.3%) and 0% in LL.

The concordance rate was lower in the borderline groups with 33.3% in BL, 33% in IL and least concordance of 16.7% in LL. However, the concordance for TT was higher than the borderline groups with 57.1%. Histopathological analysis of the cases in the present study as shown in Table 3 was carried out with due attention to the epidermal atrophy, presence of clear sub-epidermal zone, dermal inflammatory infiltrate, presence and composition of granulomas, presence of giant cells and relative proportion of lymphocytes and foamy histiocytes in accordance with Ridley and Jopling histopathological criteria.

Out of 50 patients included in the study, 15 (30%) presented a clinical suspicion of paucibacillary leprosy and 35 (70%) of multibacillary leprosy. From the chi-square
output table, we see that no significance level has been achieved. McNemar Bowker test table showing no systematic association between the above two variables at 95% level of confidence. Hence, it concludes that there is no significant relationship between age wise patients and histopathological diagnosis.

Pic 1. Tuberculoid Leprosy with swelling of hand.

Pic 2. Histopathology of skin biopsy (H & E 400x) showing Langhans foreign body giant cells, black arrow

Pic 3. Borderline Tuberculoid Leprosy patient with lower lip swelling and hypopigmented patch on left cheek.

Pic 4. Histopathology of skin biopsy (H & E Stain 400x) showing focal collection of macrophages in the dermis

Pic 5. Lepromatous Leprosy with ENL reaction

Pic 6. Histopathology of skin biopsy (H & E stain 400x) showing dermal neutrophilic infiltrate amidst with macrophages
DISCUSSION

Due to its clinical diversity and resemblance to other diseases, leprosy is difficult to diagnose clinically. The sex ratio was heavily skewed towards males (66%). This is similar to other Indian studies undertaken by Gridhar M et al (77.6%) and Bhushan et al (72.34%). Mathur MC et al, however, observed 53.8% males in their study, while Moorthy et al observed 65.05% males.

The common age group affected in present study is between 31 - 40 yrs. (22%) and the second common age group affected was between 21 - 30 yrs. (20%) followed by 41 - 50 yrs. (18%), 51 - 60 yrs. (16%), 60+ yrs. (14%), 11 - 20 yrs. (6%) and less than 10 yrs. (4%). The mean age of the patients studied was 41.34 ± 17.1. In a study done by Moorthy BN et al, majority of patients were between 20 - 29 years (20.70%). Children below 9 years were least affected (6.45%). In present study also, youngest age affected was 7 years. In one series, age range was 6 - 72 years and mean age was 35.9 years. In another study done in Green Pastures Hospital, Pokhara, the mean age was 41 years.

In present study among 50 cases, 34 (68%) cases showed correlation between clinical and histopathological diagnosis. Maximum correlation was observed in borderline Leprosy (100%) and borderline Lepromatous groups (100%) followed by Lepromatous Leprosy (70%), Indeterminate Leprosy (66.67%) and Tuberculoid Leprosy (57.14%). A poor correlation was seen in Borderline Tuberculoid leprosy.

Ridley and Jopling found agreement between clinical and histological types in 56 (68.3%) out of 82 patients. Agreement in present study is almost similar to Ridley and Jopling study (68%).

A similar study done by Jerath and Desai found agreement between clinical and histological types is 68.5%, which is slightly more than present study (68%).

A similar study done by Bhusan et al showed a concordance of 74.47% between the histological and clinical diagnosis out of 150 patients, which is more than present study (68%).

A similar study done by Vargas-Ocampo F et al found agreement between clinical and histological types in 2520 (42%) out of 6000 cases patients, which is less than present study (68%).

A similar study done by Kar et al showed a concordance of 70% between the histological and clinical diagnosis, which is less than present study (68%).

A similar study done by Bhutia AS et al found agreement between clinical and histological types in 878 (69%) out of 1276 cases patients, which is more than present study (68%).

A similar study done by Kar PK et al found agreement between clinical and histological types in 84 (70%) out of 120 patients, which is more than present study (68%).

A similar study done by Nadkarni NS et al found agreement between clinical and histological types in 2160 (81.8%) out of 2640 patients, which is more than present study (68%).

A similar study done by Sehgal VN et al found agreement between clinical and histological types in 34 (33%) out of 95 cases patients, which is less than present study (68%).

A similar study done by Manandhar U et al found agreement between clinical and histological types in 34 (45.33%) out of 75 patients, which is less than present study (68%).

A similar study done by Sharma et al showed a concordance of 53.44% between the histological and clinical diagnoses, which is less than present study (68%).

A similar study done by Mitra K et al found agreement between clinical and histological types in 1509 (57.16%) out of 2640 patients, which is less than the present study (68%).

A similar study done by Pandya AN et al found agreement between clinical and histological types in 29 (58%) out of 50 patients, which is less than present study (68%).

A similar study done by Moorthy et al showed a concordance of 62.63% out of 372 patients between the histological and clinical diagnosis which is less than present study (68%).

A similar study done by Kar et al showed a concordance of 70% between the histological and clinical diagnosis, which is less than present study (68%).

A similar study done by Vargas-Ocampo F et al found agreement between clinical and histological types in 2520 (42%) out of 6000 cases patients, which is less than present study (68%).

A similar study done by Kar et al showed a concordance of 70% between the histological and clinical diagnosis, which is less than present study (68%).
Maximum concordance was observed in LL type of leprosy in studies by Mathur MC et al, Gridhar M et al,9 Moorthy et al,10 Nandkarni NS et al15 and Bhatia AS et al,13 but in present study 70% concordance was observed in LL type of leprosy which is less than BB and BL.

Maximum concordance was observed in Indeterminate type of leprosy in studies by Kar PK et al14 and Kalla G et al,16 which is more than present study (57.14%).

Maximum concordance was observed in tuberculoid type of leprosy in studies by Jerath VP et al21 (88.8%), which is more than present study (66.67%).

As there is overlap in histopathologic features of different types of leprosy, morphology alone is not specific, thus adequate clinical data can help in good clinical pathologic correlation.

CONCLUSION
Leprosy, though reported to be eliminated, still continues to be one of the common infectious diseases in India. Skin biopsy is the investigation of choice for clinical diagnosis of leprosy as well as acts as a therapeutic guide.

REFERENCES