Studies of Human Leprosy Lesions

in Situ Using Suction-induced Blisters

1. Cellular Components of New, Uncomplicated Lesions

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Human immunity to Mycobacterium leprae is manifested by a wide spectrum of local inflammatory responses to M. leprae in the skin, producing the wide range of clinical and histologic changes observed in this disease \(^1\). Most studies of the immunology of leprosy, however, have evaluated in vitro the functional status of cells obtained from the peripheral blood of patients, rather than examining cells from tissue lesions \(^2\). The development of monoclonal antibodies against different lymphocyte membrane antigens has made it possible to examine the local infiltrate in situ in more detail \(^3\). Such studies have shown that the percentages of phenotypically distinct lymphocyte subpopulations are different in the peripheral blood and the cutaneous lesions, and between lesions of different types of leprosy \(^4\).

Although biopsies have the advantage of examining the site of immunologic activity and injury in situ, the trauma of the procedure does not readily permit the sequential, functional studies possible with peripheral blood cells in vitro. Moreover, if viable cells from biopsies are desired for functional studies, harsh physical or enzymatic methods are required to extract them from the connective tissue of the dermis.

We have attempted to combine the advantages of these two approaches by aspirating cells from blisters which have been induced by suction directly over skin lesions—a painless, nonscarring procedure \(^6\). We here report the results of immunocytologic studies of the cells obtained from such blisters over representative skin lesions in 27 untreated patients across the immunopathologic spectrum of leprosy.

MATERIALS AND METHODS

Patients. Volunteers were recruited from newly diagnosed patients at McKean Rehabilitation Institute, Chiang Mai, Thailand. Patients with active leprosy were classified according to the clinical features of the lesions by two experienced physicians (TCS and VS) and histologic criteria in representative biopsies (DMS) according to a five-part scale \(^11\). These included 11 lepromatous (LL), 9 borderline lepromatous (BL), 2 midborderline (BB), and 5 borderline-tuberculoid (BT) leprosy patients. Controls were recruited from cured patients with no evidence of reactivation, whose leprosy type was known from initial diagnostic examination by the same physicians in previous studies, confirmed in most cases by biopsy results \(^2\). These included 9 former LL, 2 former BL, 1 former BB, 12 former BT, and 5 former tuberculoid (TT) patients.

Blister formation and cell preparation. Four blisters were induced painlessly by gentle continuous suction for 1–3 hr through a plastic template taped to the skin, as previously described \(^6\). The template was placed directly over a representative skin lesion on the trunk or extremities in patients with active leprosy, or on the volar surface of the forearm in controls. Blisters were protected from injury by taping a small inverted cup over them.
Fluid was aspirated from blisters in a heparinized 1.0 cc “tuberculin” syringe in which the volume was measured. One blister was aspirated each day for 4 days; each blister was used only once.

The blister fluid was diluted to 1.0 cc with RPMI 1640 tissue culture medium, and the cells were collected on a 13 mm, 0.22 µm cellulose acetate filter (Millipore Corp., Bedford, Massachusetts, U.S.A.) (6). The filter preparations were fixed immediately in cold formol-acetone (14) for 30 sec, and washed in modified potassium phosphate-buffered saline (PBS) (17). While immersed in PBS, the filters were divided into several pieces, each of which was then stained with a different antibody or substrate.

**Cell staining and counting.** An indirect, two stage avidin-biotin method was used to stain individual pieces of each filter, using monoclonal antibodies Leu4, Leu3 or OKT4, and Leu2 or OKT8, to identify all T cells, T-helper and T-suppressor lymphocyte phenotypes, respectively. The secondary antibody was biotinylated sheep anti-mouse F(ab’), followed by an avidin-biotin-horseradish peroxidase complex (ABC). This substrate was developed using diaminobenzidine or aminoethyl carbazole (6). Monocytes/macrophages (MAC) were stained for nonspecific esterase (NSE) (19), and preparations were counterstained with hematoxylin or methyl green. (Attempts to stain monocytes in this system with a variety of commercially available monoclonal antibodies did not produce sufficiently reproducible results, for reasons which are not yet clear.)

Cytologic detail was well preserved, easily permitting discrimination between mononuclear cells and polymorphs. Similarly, the different patterns of NSE staining between lymphocytes and monocytes were easily discerned.

The percentage of positively stained cells (i.e., each subset) was determined by a differential count of at least 200 cells in each segment of the filter; the mean number of cells in 20 oil immersion fields was used to calculate the total number of cells on the filter, finally expressed as the number of cells per mm³ of blister fluid.

From each filter, one control segment was incubated with secondary antibody followed by the ABC reagent, and another with ABC reagent only, prior to development with substrate and indicator. In addition, control preparations of peripheral blood leukocytes, collected on filters and washed and fixed in identical fashion, were stained with every blister preparation. Background staining (for both blister and peripheral blood preparations) was usually slight, and presented no problem in interpretation. If controls (or blister samples) showed evidence of high background, low or irregular positive staining, or were otherwise technically unsatisfactory for any antibody, the data for that antibody were not included in the analysis.

**Statistical analysis.** Quantities and ratios were ranked and tested using the nonparametric Wilcoxon two-sample test.

**RESULTS**

The volume of fluid obtained from blisters at all times after induction ranged from 30–150 µl (median = 60; mean = 66); the range and distribution of volume did not differ significantly between the active patients and the controls. In blisters induced in the skin of cured, control volunteers the median total cellularity varied from 80–400 cells/mm³ (ranging from 50–550 cells/mm³) over all times studied (Fig. 1). Cellularity was higher during the first 24 hr, as part of the nonspecific inflammatory response to blister induction (6). No statistically significant differences were seen among patients who had had different types of leprosy earlier.

To determine the reproducibility of cell sampling using the blister technique, two blisters were aspirated at the same time from a number of control patients. The paired results gave a correlation coefficient of 0.613 (p = 0.003) (Fig. 2).

In active leprosy lesions, total cell counts varied much more widely than in the controls, ranging from 1–2800 cells/mm³ (Fig. 3). Median values for the different immunopathologic types of leprosy were similar at all times studied.

Mononuclear cells predominated in active lesions, with median percentages of 70%–91% over all times studied (Fig. 4). Polymorphs accounted for more than 30% of the cells in blisters over some BL lesions (3 patients) and LL lesions (2 patients), but not in BB or BT lesions. Neither eosinophils...
Fig. 1. Cellularity of blisters in cured, inactive control patients. Four blisters were induced (as described in text) on the volar surface of the forearm in patients with no evidence of active disease. Blisters were aspirated at different intervals after induction. Each point = cell count from one blister; bar = median value.

Fig. 2. Correlation of cell counts in paired blisters. Four blisters induced on the forearm of cured, inactive patients were sampled in pairs at indicated intervals after blister induction. Total cell counts were determined independently for each blister. Correlation coefficient for these paired samples is 0.613; slope of the curve was determined by linear regression.

Fig. 3. Total cell counts in blisters over active, uncomplicated leprosy lesions. Four blisters were induced directly on active lesions which were classified by clinical and histologic criteria. Blisters were aspirated only once, at different intervals after induction. For each type of lesion: point = median value; vertical bar = range; N = number of observations.

nor basophils were present in substantial numbers.

Among mononuclear cells, the median ratio of T cells: monocytes (Leu4+: NSE+ cells) remained between 0.5-1.5 for all types of leprosy at all times studied, except for a predominance of T cells at 48 hr in BT lesions (Fig. 5). This difference was statistically significant (p < 0.005), and it is notable that this was observed in blisters with similar total cell counts (Fig. 3), indicating an absolute as well as a relative increase in T cells in BT lesions.

At 48 hr, the median helper: suppressor (H:S)(CD4:CD8) ratio was 3.3 in BT lesions and 1.3 in BL and LL lesions, a statistically significant difference (p < 0.005). The median H:S ratio ranged from 1.0-3.0 at other times, but the range of values was also great and at none of these times were the differences statistically significant (Fig. 6).

**DISCUSSION**

This study addresses the feasibility of using suction-induced blisters to evaluate the
relative numbers and types of cells in the cutaneous infiltrates of leprosy, with minimal trauma to the patient or the cells. Rebuck’s “skin windows” (10) were the conceptual forerunner of this approach. Other investigators attempted to apply Rebuck’s technique to leprosy (4, 5), but the cutaneous abrasion required is difficult to standardize, and meaningful data concerning mononuclear cells was obscured by the brisk polymorphonuclear responses induced by cover slips. As noted by the originators of the blister technique (4), this method introduces no foreign material of any kind and therefore approximates more closely the conditions of the intact skin. The suction used to induce the blisters does elicit a mild, nonspecific inflammatory response which peaks during the first 24 hr in healthy volunteers (4). In this study, samples were obtained from 24 hr onward to minimize the effects of this nonspecific response.

The total cellularity of the blisters over active lesions varies greatly, not unlike the variations in cellularity observed in the lesions in histologic sections (11). The cellularity of blisters itself is therefore not a valuable indicator of the immunologic status of the underlying lesion, although it is also notable that many blisters over active lesions had many more cells than the highest numbers seen in blisters in cured, inactive patients. The marked kinetic changes which may be seen with this technique during an accelerating immune response (4) were neither expected nor observed in these lesions.

Fig. 4. Percentage of mononuclear cells in blisters over different types of leprosy lesions. Percentage of mononuclear cells (lymphocytes and monocytes) was determined for blisters aspirated at different times after induction on different types of leprosy lesions. Each point = one blister; horizontal bars = median values.

Fig. 5. T-cell monocyte ratios in leprosy lesions. Among cells obtained from blisters at different times after induction over leprosy lesions, total T-cell numbers were determined using anti-Leu4 antibody; monocytes were identified by staining for nonspecific esterase. For each type of lesion: point = median value; vertical bar = range; N = number of observations. Difference between BT and BL-LL 48 hr after induction is statistically significant (p < 0.005).
Fig. 6. T-helper:suppressor ratios in uncomplicated leprosy lesions. Among cells obtained from blisters at different times after induction over leprosy lesions, T-helper and -suppressor phenotypes were determined by immunostaining with anti-Leu3 and OKT8 antibodies, respectively. For each type of lesion: point = median value; vertical bar = range; N = number of observations. Difference between BT and BL-LL 48 hr after induction is statistically significant (p < 0.005).

since no antigenic stimulation was used to induce an immune response in this study. The result is, instead, a profile of the "baseline" status of different types of leprosy lesions.

The method shows an acceptable degree of reproducibility from blister to blister, as evidenced by the data on paired blister samples. Some of the scatter which is seen is probably due to imperceptible differences in trauma during blister induction, or to variations in the quantity of red blood cells, which is usually negligible but was not quantitated in this study.

The high percentage of mononuclear cells within the blisters is in agreement with the histologic features of these lesions (11), and indicates that the method is suitable for the study of the cells involved in immunologic responses in leprosy. The percentage of polymorphs infrequently exceeded 25%–30% of the total and was, in most instances, far less. Other investigators have demonstrated that the introduction of very few bacteria is sufficient to induce an acute inflammatory response in a blister (6), and we view the occasional occurrence of numerous polymorphs in one blister (out of a set of 3-4 in a volunteer) as likely to be the result of such contamination due to a crack or puncture of the blister. Interestingly, however, among patients in whom 3–4/4 blisters contained over 30% neutrophils, the majority developed a reaction (either type 1 or 2) soon thereafter. The extent and significance of this observation are under investigation.

Monocytes/macrophages were a major cellular component in blisters over active lesions, consistent with their prominent place in all types of leprosy lesions. However, monocytes/macrophages were also numerous in blisters in cured, control patients, as they were in earlier studies of healthy volunteers (6 and Scollard, unpublished results). The activation status and functional activity of the monocytes/macrophages may differ in these different situations, but these parameters were not assessed in this study.

In BT lesions, T-helper/inducer phenotypes outnumbered T-suppressor/cytotoxic phenotypes at least 3 to 1 at 48 hr after blister induction, in agreement with the patterns observed in biopsy studies (6). Recruitment of T cells into the blister appeared to continue after this time, probably nonspecifically, and the trend was no longer evident at 96 hr. The statistically significant results at 48 hr, however, indicate that the method does provide a sample of the cutaneous infiltrate, reflecting the status of the lesion.

Phenotype alone does not indicate the functional status of the cells. Preliminary studies suggest that direct assessment of some functional capabilities of blister exudate cells is possible, however, and further work is in progress.

The blister technique has proved to be very acceptable to patients, enabling multiple sequential studies of the same patient. Another distinct advantage of this method over biopsies is the ability to make more precise determinations of the total number of cells present in the blister—an accurate
clear cells predominated at all times studied in BT than in BL and LL lesions at 48 hr. Suction blisters offer a quantitative reproducible method, minimally traumatic and amenable to multiple sampling, with which cells can be obtained which are representative of the cutaneous infiltrates in leprosy lesions.

SUMMARY

The cellular contents of blisters induced by suction over new, uncomplicated leprosy lesions, and in the skin of cured, control patients, have been examined with enzymo- and immuno-histochemical staining over a period of 4 days. The total cellularity of the blisters varied over a wide range, not correlated with the type of leprosy. Mononuclear cells predominated at all times studied, with nearly equal percentages of monocytes and T lymphocytes. The T-helper: suppressor ratio was significantly greater in BT than in BL and LL lesions at 48 hr. Suction blisters offer a painless, quantitative, reproducible, multiple-sampling method for obtaining cells from the cutaneous infiltrates of leprosy for phenotyping or functional analysis.

REFERENCES

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