Activation of peripheral blood mononuclear cells in bronchoalveolar lavage fluid from patients with sarcoidosis: visualisation of single cell activation products

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Abstract

Background – Interstitial lung diseases are characterised by the recruitment of mononuclear cells to disease sites where maturation occurs and activation products, including lysozyme (LZM), are released. Analysis of in vitro cell culture supernatants for activation products masks the functional heterogeneity of cell populations. It is therefore necessary to examine the secretion of activation products by single cells to assess whether the activation of newly recruited mononuclear phagocytes at the sites of disease in the lung is uniform and controlled by the local microenvironment.

Methods – The reverse haemolytic plaque assay was used to evaluate, at a single cell level, the ability of bronchoalveolar lavage (BAL) fluid from seven patients with sarcoidosis to activate Ficoll-Hypaque-separated peripheral blood mononuclear cells by comparison with BAL fluid from six normal volunteers and nine patients with systemic sclerosis. Monolayers of peripheral blood mononuclear cells and sheep red blood cells were cultured either alone or in the presence of 20% (v/v) BAL fluid with a polyclonal anti-LZM antibody. LZM/anti-LZM complexes bound to red blood cells surrounding the secreting cells were disclosed following complement lysis of red blood cells and quantification of plaque dimensions using microscopy and image analysis.

Results – Bronchoalveolar lavage fluid from all the patients with sarcoidosis increased LZM secretion by peripheral blood mononuclear cells compared with unstimulated mononuclear cells. By contrast, BAL fluid from the other individuals had no effect on LZM secretion.

Conclusions – Single cells activated by BAL fluid can be evaluated by the reverse haemolytic plaque assay. BAL fluid from patients with sarcoidosis, but not from patients with systemic sclerosis or normal individuals, contains components capable of activating mononuclear phagocytes to secrete lysozyme.

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Sarcoidosis is characterised by the accumulation at disease sites of mononuclear cells resulting in the formation of epithelioid cell granulomas. Circulating monocytes which are attracted to disease sites are subjected to the influence of local proinflammatory signals resulting in phenotypic and functional changes as they undergo maturation and activation. One of the activation products of stimulated mononuclear phagocytes is lysozyme (LZM), and measurements of LZM concentration have been used as an index of active disease. The primary function of LZM has always been considered bactericidal, even though previous work has shown that it may play a part in the modulation of monocyte-lymphocyte interactions and depression of neutrophil chemotaxis.

If monocytes attracted to disease sites in sarcoidosis are stimulated to produce increased amounts of LZM, we hypothesised that factors within the local microenvironment are responsible for this change in function. To test the hypothesis we have used a method of identifying secretory products at the single cell level using the reverse haemolytic plaque assay to examine the effects of bronchoalveolar lavage (BAL) fluid on LZM secretion. The ability of the reverse haemolytic plaque assay to visualise cytokine products by single cells allows the modulation in function of cells to be assessed in the absence of cell–cell interaction and minimises paracrine effects of proinflammatory mediators from other cells in culture, thus enabling the direct effect of added BAL fluid on cell activation to be analysed. We therefore examined BAL fluid from patients with sarcoidosis, patients with systemic sclerosis, and normal individuals and compared the capacity of each to activate “naive” monocytes to synthesise and secrete LZM.

Methods

Patients

Three groups were studied: (1) patients with sarcoidosis (n = 7; M:F 4:3; age 30-26 (1.6) years), (2) patients with fibrosing alveolitis associated with systemic sclerosis (n = 9; M:F 4:5; age 44-6 (1.7) years), and (3) normal volunteers (n = 6; M:F 3:3; age 26-3 (0.46) years). Of the patient groups, two patients with sarcoidosis were current smokers and three with systemic sclerosis were receiving treatment (prednisolone).

A diagnosis of sarcoidosis was confirmed by the presence of non-caseating granulomas in...
transbronchial lung biopsies or a positive Kveim reaction in a clinical setting consistent with sarcoidosis. Patients with systemic sclerosis met the American Rheumatism Association preliminary criteria for this diagnosis. A diagnosis of pulmonary fibrosis in systemic sclerosis was made by the finding of typical changes on high resolution computed tomographic scanning.

All patients underwent bronchoalveolar lavage (BAL) as part of their clinical evaluation. The normal volunteers chosen were non-smokers with no evidence of pulmonary disease. Approval for lavage of the normal volunteers was given by the Royal Brompton Hospital ethics committee.

**BRONCHOALVEOLAR LAVAGE FLUID**

Bronchoalveolar lavage fluid was collected from patients with sarcoidosis, systemic sclerosis, and normal individuals during routine bronchoscopy and lavage. Briefly, 100 ml of warmed normal saline was instilled in 20 ml aliquots in three subsegmental bronchi (the right middle lobe, right lower lobe, and lingula) and aspirated immediately. The cells were separated from the BAL fluid by centrifugation at 200 g, 4°C, seven minutes. The BAL fluid was then subjected to a further centrifugation at 600 g for 10 minutes and stored in 5 ml aliquots at −80°C until further analysis. Before use in the reverse haemolytic plaque assay the BAL fluid was passed through a filter with a pore diameter 0.2 μm.

**ISOLATION OF HUMAN PERIPHERAL BLOOD MONONUCLEAR CELLS**

Peripheral blood mononuclear cells were isolated from heparinised venous blood from the same normal individual using a Ficoll-Hypaque gradient (Pharmacia, Milton Keynes, UK) according to the method of Boyum. After three washes in Hank’s buffered salt solution (Gibco, Uxbridge, UK) the peripheral blood mononuclear cells were resuspended to a final concentration of 4 × 10⁶ cells/ml in RPMI 1640 media (Gibco, Uxbridge, UK) containing 1% bovine serum albumin (BSA) and penicillin (100 U/ml)/streptomycin (100 μg/ml) (complete media) for use in the reverse haemolytic plaque assay.

**REVERSE HAEMOLYTIC PLAQUE ASSAY**

The assay was performed using a modification of the method of Lewis et al. In brief, Cunningham chambers were prepared by coating microscope slides with poly-L-lysine and constructing a 30 μl chamber using double-edged tape and a coverslip. Human peripheral blood mononuclear cells and protein-A coated sheep erythrocytes were mixed in equal volumes and aliquoted into Cunningham chambers. The cells were allowed to settle on the poly-L-lysine-coated slides for 40 minutes in an atmosphere of 5% CO₂ at 37°C. Excess unattached cells were removed by rinsing each chamber with complete media leaving a confluent monolayer of erythrocytes and peripheral blood mononuclear cells on the poly-L-lysine-coated surface. The chambers were then filled with complete media containing 1/50 dilution of rabbit anti-human LZM (Dakopatts Ltd, High Wycombe, UK) from which sodium azide had been previously omitted, by using an Amicon Concentrator (Amicon Ltd, Gloucestshire, UK), and 20% (v/v):final dilution in culture chamber) of either normal saline (control) or BAL fluid from patients with sarcoidosis, systemic sclerosis, and normal subjects (the optimum concentration of BAL fluid determined by preliminary experiments). The chambers were incubated for three hours at 37°C in 5% CO₂ and then washed with complete media and filled with complete media containing 1/30 dilution of guinea pig complement (Gibco, Uxbridge, UK). After 25 minutes incubation at 37°C in 5% CO₂, during which period plaques of haemolysis were formed around the LZM-secreting cells, the chambers were washed with complete media without BSA and either fixed with 4% glutaraldehyde in phosphate buffer (pH 7.4) and stored at 4°C under Tris-buffered saline (TBS) prior to estimation of plaque dimensions, or infused with 0.5% (v/v) Trypan Blue solution and incubated for five minutes at 37°C to test the viability of peripheral blood mononuclear cells. Cell viability was always >95% at the end of three hours incubation in the reverse haemolytic plaque assay.

The specificity of plaque formation for LZM secretion was confirmed since no haemolytic plaques were observed when (a) mononuclear cells were omitted from the assay; (b) when the anti-LZM antibody was omitted and replaced with 1:50 normal rabbit serum; (c) when complement was omitted; (d) when uncoated erythrocytes were used; (e) when the antibody was preabsorbed at 4°C overnight with >3 μg/ml human LZM (Calbiochem, Nottingham, UK).

To minimise the day to day variations that exist during culturing of the peripheral blood mononuclear cells in the reverse haemolytic plaque assay, even though culturing conditions were stringently observed, each assay contained at least 2 × 10⁶ cells/ml in the fluid chamber. Each study group examined and a control in which the volume of BAL fluid was substituted with normal saline.

**IMMUNOSTAINING OF MONOCYTES**

At the end of the assay the plaque-forming cells were identified as monocytes by immunolabelling with KP1 mouse monoclonal antibody (gift from Dr DY Mason, Oxford, UK) which recognises proteins of 110, 70, and 40 kDa expressed in the lysosomal fraction of mononuclear phagocytes. Visualisation of positively stained perinuclear areas was achieved using the alkaline phosphatase/anti-alkaline phosphatase (APAAP) immunocytochemical method employing a 1/50 dilution of rabbit anti-mouse IgG, 1/50 dilution of APAAP complex, alkaline phosphatase substrate, and fast red with haematoxylin counterstaining. After colour development the chambers were dried, the coverslip removed, and a new coverslip mounted.
LYSOZYME CONTENT IN THE BAL FLUID
Concentrations of free LZM in the BAL fluid were measured by radial immunodiffusion (The Binding Site, Birmingham, UK).

ALBUMIN CONCENTRATION IN BAL FLUID
Albumin concentrations in the BAL fluid samples were measured by bromocresol green colorimetric assay (Sigma Chemical Co Ltd, Poole, Dorset, UK) and performed in duplicate. Briefly, a standard solution of human serum albumin (Sigma Chemical Co Ltd), 500 g/l in normal saline, was diluted to a range covering albumin concentrations of 0·05–0·5 g/l. Duplicate 1 ml aliquots of each standard and BAL fluid sample were pipetted into 3·5 ml polystyrene test tubes (Sarstedt Ltd, Beaumont Leys, Leicestershire, UK) and, after the addition of an equal volume of bromocresol green solution, the reaction was allowed to proceed at room temperature for 10 minutes. The absorbance of each sample was measured at 630 nm on a Gilford 2600 spectrophotometer (Corring Ltd, Essex, UK) and the concentrations of albumin in the BAL fluid samples were estimated from the linear relationship demonstrated between albumin concentration and absorbance as a standard curve (r = 0·999, p < 0·0001).

STATISTICAL ANALYSIS
Results of single replicate experiments were analysed with the Student's t test. Group data are distributed non-parametrically and comparisons between groups were made using the Mann-Whitney U test. A p value of <0·05 was considered to be significant for all analyses. The data in the text and in the figures are shown as mean (SE).

Results
EVALUATION OF REVERSE HAEMOLYTIC PLAQUE ASSAY
To evaluate the reverse haemolytic plaque assay as a means of identifying LZM secretion by single cells, preliminary experiments were conducted with peripheral blood mononuclear cells cultured in the absence of stimulation to determine spontaneous secretion of LZM and the specificity of plaque formation for LZM.

Easily identifiable plaques were seen after incubation of peripheral blood mononuclear cells for three hours, at which time point >95% of all cells were viable (fig 1). Plaque formation was completely suppressed if antibody to LZM was incubated with human placental LZM at concentrations >3 µg/ml at 4°C for 12 hours prior to being added to the culture medium (data not shown).

KP1 immunostaining for mononuclear phagocytes confirmed that all plaque-forming cells were monocytes.

INFLUENCE OF BAL FLUID FROM PATIENTS WITH SARCOIDOSIS ON THE SECRETION OF LZM
In order to identify the concentration of BAL

PLAQUE QUANTIFICATION
Plaque area was measured using Leitz microscopy, Apple Macintosh computer, and image analysis package (Improvision UK). Fifty plaques per chamber were measured in five replicate chambers (250 plaques in total).

The following criteria were applied when measuring plaque area: (a) lysis of sheep erythrocytes around the plaque-forming cell had to be visible and well defined; (b) cells were chosen for plaque measurement by examining randomly selected non-overlapping microscope fields; (c) only plaques formed by single cells were counted; (d) plaque counting was performed “blind” without knowledge of the identity of the chambers. The coefficient of variation for intraobserver variability was 4·3% for 600 plaques.
**HETEROGENEITY OF LZM SECRETION WITHIN THE MONOCYTE POPULATION**

Because it is recognised that the mononuclear phagocyte population is heterogeneous, the population of plaque-forming cells was examined for heterogeneity of LZM secretion. There was considerable variation amongst individual peripheral blood mononuclear cells in the amount of LZM secreted after three hours of incubation. This variation of plaque size ranged from 250 μm² to 3000 μm²; fig 3 illustrates the results from one such assay.

Incubation of normal peripheral blood mononuclear cells with lavage fluid obtained from patients with sarcoidosis produced an increase in the number of cells that formed larger size plaques, shown as a shift of the distribution curve to the right indicating a much wider distribution in plaque sizes. Plaque size distributions after incubation with lavage fluid from normal individuals or patients with systemic sclerosis were no different from controls using normal saline (fig 3). While 33% of plaque-forming cells stimulated by fluid from patients with sarcoidosis produced plaques of area greater than 1250 μm², on average only 4% of plaque-forming cells stimulated by BAL fluid from other groups produced plaques of this size.

**AUGMENTATION OF LZM SECRETION BY BAL FLUID FROM PATIENTS WITH SARCOIDOSIS**

To assess the influence of BAL fluid from normal subjects, patients with systemic sclerosis and patients with sarcoidosis on the secretion of LZM by mononuclear phagocytes from a single normal individual, replicate assays were performed using BAL fluid from at least a single individual from each group in each group of assays. The results from one such assay are shown in fig 4. It can be seen that BAL fluid from patients with sarcoidosis increased the secretion of LZM by approximately 50% (p<0.01).

When evaluating the group data as a whole the average plaque size measurements were expressed as percentage increase of the average plaque size of the control. The group data are summarised in fig 5. There was a uniform significant augmentation (p<0.01) of LZM secretion by peripheral blood mononuclear cells from the same normal individual cultured with 20% (v/v) BAL fluid from patients with sarcoidosis but not with BAL fluid from patients with systemic sclerosis or normal individuals. Importantly, all seven BAL fluid samples from patients with sarcoidosis increased plaque sizes by an amount greater than BAL fluid samples from any of the other groups.

**INFLUENCE OF LZM CONCENTRATIONS IN BAL FLUID ON PLAQUE FORMATION**

Because varying amounts of LZM are present within BAL fluid, reflecting in vivo secretion, LZM concentrations were evaluated to determine whether they influenced plaque formation.
Bronchoalveolar lavage fluid concentrations of LZM were similar in all three groups (table) and average plaque size did not correlate with LZM concentrations in BAL fluid. Three lavage fluid samples containing LZM levels >15 μg/ml (final concentration in culture chambers 3 μg/ml after 20% (v/v) dilution) were the only samples which abolished plaque formation—that is, native LZM bound antibody in the culture medium leaving no free antibody binding sites for LZM released by cultured monocytes. This is consistent with our previous experiments which showed that LZM concentrations >3 μg (>15 μg/ml after 20% (v/v) dilution in the final culture mix) blocked the formation of plaques. There was no correlation between LZM levels in the BAL fluid samples and the size of the average plaque area (p>0.1).

**Discussion**

This study has shown that normal peripheral blood mononuclear cells exposed to lavage fluid from patients with sarcoidosis, but not lavage fluid from patients with systemic sclerosis or normal individuals, synthesised and secreted approximately 40% more LZM than cells incubated in the presence of normal saline. Furthermore, the use of a reverse haemolytic plaque assay to visualise mediator production by single cells has shown that there is a considerable heterogeneity in LZM secretion by mononuclear cells, and that this heterogeneity is enhanced by BAL fluid from patients with sarcoidosis but not those with systemic sclerosis or normal individuals.

Heterogeneity among the mononuclear phagocyte population at disease sites has long been recognised and, in diseases like sarcoidosis or cryptogenic fibrosing alveolitis, this heterogeneity can be further visualised using panels of monoclonal antibodies. Furthermore, in sarcoidosis diversity of cell surface phenotype is also recognised within the mononuclear phagocyte population which make up the granulomas. Surface phenotype heterogeneity has not yet been related to functional status because of the constraints applied by traditional cell culture methodologies. Studies of peripheral blood monocyte subpopulations separated by density gradient do, however, show a diverse secretory repertoire probably reflecting the in vivo tight local control exerted over inflammatory cells such as those of the mononuclear phagocyte series. Activated cells within granulomas in sarcoidosis are known to synthesise and secrete LZM and serum LZM levels have been used in some studies as a marker of an active disease process. In this regard, our study has shown that BAL samples of the epithelial lining fluid from the lower respiratory tract of patients with sarcoidosis contain macromolecules capable of enhancing LZM secretion by normal monocytes. This is consistent with other studies which have identified stimulating factors in lavage fluid specific for sarcoidosis. Although the nature of these factors is not known, interferon gamma, a lymphokine which is spontaneously released from cells obtained from the lungs of patients with sarcoidosis, is known to stimulate LZM secretion in cultures of macrophage cell lines. It remains possible, therefore, but not proven by this study, that interferon gamma is the stimulating factor in the lavage fluid.

As well as a marker of biological activity in sarcoidosis, LZM may have an immunomodulatory role. Although traditionally considered to be a macromolecule which has bactericidal activity, LZM is now known to reduce neutrophil chemotaxis and also to exert an upregulatory effect on monocyte-depleted mixed lymphocyte cultures. Furthermore, LZM can reduce the in vitro response of lymphocytes to mitogens. This implies a specific control of monocyte function within the local microenvironment in sarcoidosis which might have a direct effect on restoring tissue homeostasis.
In systemic sclerosis, a condition in which mononuclear phagocytes also traffic to disease sites and also release potent mediators—including tumour necrosis factor alpha and interleukin-8\textsuperscript{23,24}—which can modulate the disease process, other control mechanisms are likely to be involved in stimulating monocytes to produce a quite distinct repertoire of cytokine mediators which are, therefore, disease dependent.

The reverse haemolytic plaque assay has a number of advantages over traditional culture methods. Firstly, although methods.

Furthermore, we have shown that there is heterogeneity within the monocyte population in terms of amount of LZM produced per cell, and that the stimulatory activity is found only in lavage fluid obtained from patients with sarcoidosis. The reverse haemolytic plaque assay will allow the further exploration of subpopulations of mononuclear phagocytes to be undertaken with particular regard to their secretory repertoire, and this may provide greater insights into the regulation of mononuclear phagocyte function in the lung at disease sites.

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