Comparison of the expression of interferon γ, IL2, IL4, and lymphotoxin mRNA in experimental autoimmune uveoretinitis

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Abstract
The aim of this study was to investigate the T lymphocyte subsets involved in experimental autoimmune uveoretinitis (EAU) by quantifying the numbers of cells expressing mRNA for each of the lymphokines interferon γ, interleukin 2, interleukin 4, and lymphotoxin throughout the disease process. Lewis rats were immunised with retinal S-antigen to provide a model of inflammatory eye disease. In situ hybridisation using cDNA probes specific for interferon γ, IL2, IL4, and lymphotoxin mRNA were utilised to localise lymphokine mRNA expression by infiltrating cells and the numbers of positive cells counted. Localisation of mRNA for all four probes was found on increasing cell numbers as the disease process progressed. Similar numbers of cells expressed mRNA for each lymphokine, generally a small percentage of the T lymphocyte total. Activated cells within the eye express mRNA for interferon γ, IL2, IL4, and lymphotoxin in EAU suggesting a mixed population of T lymphocyte subsets.

We have previously investigated the in vivo production of the lymphokines IL2, IFN-γ, IL4, and lymphotoxin in the Lewis rat model of EAU by in situ hybridisation analysis of the expression of lymphokine mRNA in the areas of T cell infiltration and tissue destruction. These studies have shown that all four lymphokines are produced in destructive foci in the retina and uveal tract, and that the cell numbers expressing the lymphokine mRNAs are found in areas of T lymphocyte infiltrate and appear to increase as the disease progresses. The aim of this study is to extend these investigations by quantifying the numbers of cells producing these lymphokines throughout the disease course to identify the potential T cell subsets involved and to provide information on the possible role of the infiltrating CD4+ T lymphocytes.

Materials and methods
The experimental methods used in these studies have been described in detail elsewhere. They are summarised as follows.

EAU MODEL
Female Lewis rats, weighing 100–150 g and 6–8 weeks old, were immunised in a hind footpad with 50 μg of purified bovine S-antigen in a 1:1 emulsion in complete Freund’s adjuvant supplemented with Mycobacterium tuberculosis organisms to a final concentration of 2.5 mg/ml. Animals were also given 5×10^6 heat inactivated Bordetella pertussis organisms in 150 μl phosphate buffered saline intraperitoneally. Eyes were rapidly removed, embedded in OCT (Shandon, Runcorn) and snap frozen in acetone and dry ice. Specimens were stored at −70°C.

IN SITU HYBRIDISATION
Four cDNA probes were used for the in situ hybridisation. The probe to rat IFN-γ was supplied by Dr T Kos, Rijswijk, The Netherlands. The rat IL2 probe and the rat IL4 probe were supplied by Dr A McKnight, MRC Cellular Immunology Unit, Oxford. The lymphotoxin probe was given by Dr N Ruddle, Yale University Medical School, New Haven, CT, USA. The cDNA probes were radiolabelled with 35S dCTP alpha (Amersham International, Amersham)
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using the random primer technique (Boehringer Mannheim, Lewes) and adjusted to 2×10^6 cpm/ml in hybridisation buffer.

Whole eye sections of 12 μm thickness were cut on a cryostat and mounted on specially prepared gelatin coated slides. Sections were fixed for 5 minutes in 4% glutaraldehyde, rinsed twice in hybridisation buffer and soaked in hybridisation buffer for 1 hour, rinsed in ethanol, and dried. The 35S labelled probe in hybridisation buffer was heated to 90°C for 10 minutes, cooled, and 100 μl applied to each slide under a parafilm coverslip. Sections were left to hybridise in a humidified chamber at room temperature for 72 hours. After hybridisation, slides were immersed in 2×SSC until the coverslips dislodged, rinsed in 2×SSC and washed at 40°C for 30 minutes in 1×SSC. Slides were then rinsed briefly in distilled water, in 70% ethanol for 5 minutes, in 95% ethanol for 5 minutes, and allowed to dry.

Control sections from each eye were fixed for 5 minutes in glutaraldehyde buffer, washed, and 100 μl of RNase A (Boehringer Mannheim) 1 mg/ml in 2×SSC were applied to each slide under a parafilm coverslip and incubated at 37°C for 1 hour. Slides were then washed in 2×SSC and simultaneous hybridisation with each probe carried out as above.

To determine the specificity of the lymphokine probe hybridisation, each section was also hybridised with a 35S labelled probe to the P53 oncogene which was considered to be irrelevant to the EAU disease process.

**Autoradiography**

The slides were dipped in K5 (Ilford, Mobberley) photographic emulsion diluted 1:1 in 0.5% glycerol, and left to expose at 4°C for 21–24 days over silica gel. Slides were then developed and counterstained with haematoxylin.

**Immunohistochemistry**

Sections of each eye were stained by a standard avidin-biotin-complex method (Vector, Peterborough) using primary monoclonal antibodies to pan rat T cells (OX 19, Serotec, Oxford), IL2 receptor (OX 39, Serotec), and MHC class II expression (OX 6, Serotec).

**Results**

**EAU**

Animals uniformly developed histological evidence of disease on day 12 post-induction. The histological changes were characterised of rat EAU. There was a marked mixed leucocyte infiltrate of the anterior and posterior segments of the eyes with patches of retinal oedema progressing to a total retinal destruction in which lymphocytes were the predominant infiltrating cell.

**Localisation of Lymphokine mRNA**

Eyes from 24 immunised and four control (non-

Figure 1 (A) Multiple foci of positive localisation of IL4 mRNA in choroid (C) (arrows to examples) and retina (R) (arrowheads to examples) at day 14 post-immunisation. Haematoxylin counterstain, ×120. (B) Adjacent section to (A) pre-treated with RNase. No localising signal with IL4 probe. Haematoxylin counterstain, ×120.
imunised) rats were studied by in situ hybridisation. The time points examined were as follows: days 10 and 11, two animals from each day; days 12, 13, 14, 17, and 21, four animals from each day. A total of 398 individual hybridisations were carried out to determine the pattern and reproducibility of the results. The results of all hybridisations were integrated to produce an overall analysis of the findings.

Levels of background signal (the amount of scattered dark grains in the photographic emulsion) varied on the numerous hybridisation procedures carried out but in all experiments, for all four probes, discrete collections of dark grains were identifiable over the background signal. The collections of autoradiographic signal were seen in section from eyes at day 12 post-immunisation onwards and occurred over mononuclear cells in the areas of the T lymphocyte (OX 19+) infiltrate. These foci were interpreted as positive localisation of expression of lymphokine mRNA by lymphocytes. Figure 1 provides an example of the localisation autoradiographic signal which was obtained.

To provide an estimate of the relative numbers of cells expressing mRNA for an individual lymphokine at a given time point in the disease, the number of positive foci in the retina and choroid were counted on each eye section and the mean value taken at each time point. Standard errors of these counts were calculated. These figures were grouped together as early disease (days 12 and 13 post-immunisation), established disease (days 14 and 17), and advanced disease (day 21). Tables 1–4 document the results obtained for each probe.

Figure 2 illustrates the overall comparative results for cell numbers for each probe found in the uveal tract and retina. Interferon γ mRNA was found localised to small numbers of cells in the retina and choroid in early disease. As the destructive process progressed fewer positive cells were found in the retina and increasing numbers were seen in the choroid. Moderate numbers of cells expressing IL2 mRNA were seen in the retina throughout the course of the disease and increasing numbers of positive cells were found in choroid in late disease. Lymphotxin mRNA was expressed by increasing numbers of cells in both the retina and choroid as the disease progressed. Cells expressing IL4 mRNA increased initially in the retina but declined as the retinal destruction advanced whereas positive cells increased in numbers in the choroid as the disease progressed.

The numbers of positive cells for each probe were generally a small percentage (less than 10%)
Comparison of the expression of interferon γ, IL2, IL4, and lymphotoxin mRNA in experimental autoimmune uveoretinitis of the total T lymphocyte number as defined by the OX19 antibody. On the frozen sections examined immunohistochemically it was not possible to make accurate cell counts of the T cell numbers because of the difficulty in defining individual cells with positive immunohistochemical stain in areas of marked accumulation of infiltrating mononuclear cells. Hence it was not feasible to calculate the percentage of cells giving positive hybridisation signal within the T cell population. It was, however, possible to compare the mean overall number of positive cells for each lymphokine probe within the eyes (in all areas) throughout the disease process; this is illustrated in Figure 2.

The data illustrated by Figure 2 show that increasing numbers of cells express mRNA for all four lymphokines as the disease progresses. The increase in numbers expressing IL2 mRNA is seen earliest. In general, fewer cells expressed lymphotoxin mRNA at each stage of disease process.

Discussion

We have previously demonstrated the presence of mRNA for IFN-γ, IL2, lymphotoxin, and IL4 in areas of T lymphocyte (OX 19+) infiltration in the active stages of EAU in the Lewis rat model.11 The results presented in this paper provide an analysis of the relative numbers of cells expressing mRNA for each lymphokine throughout the disease. Increasing numbers of cells were found to express mRNA for each lymphokine as the disease progressed. The marked increase in positive cells in the late stages of the disease generally reflects an increase in cell numbers in the uveal tract since the retina is often markedly necrotic in end stage disease in EAU. The numbers of cells in the retina positive for the IFN-γ and IL4 probes declined in late disease. Because the experimental procedure was not rigorously standardized between the probes a statistical comparison of the differences in the results for the four probes would not be valid.

In this study lymphokine mRNA was not localised to individual cells and therefore the exact nature of the cells involved has not been identified. Several lines of evidence would, however, suggest that T lymphocytes are expressing the mRNAs we have investigated. As documented by our previous studies11,12 the expression of lymphokine mRNA was found in areas of T lymphocyte (OX 19+) infiltrate as defined immunohistochemically on adjacent sections. Others have shown that T lymphocytes are the most common cell type in the retinal destructive lesions, can adoptively transfer disease, and are therefore likely to be producing lymphokines as part of active role in EAU. Moreover, IL2 and IFN-γ are almost exclusively T cell cytokines and IL4 and lymphotoxin are common T cell products.11,12

Given the above evidence for lymphokine production by infiltrating T cells in EAU it is worth considering the potential cellular subsets involved. Subsets of CD4+ T lymphocytes have been defined based on their pattern of lymphokine secretion and these subsets appear to correlate with the functional properties of the activated T cells.11,12 Th1 cells produce IL2, IFN-γ, and LT and are thought to function as cytotoxic or inflammatory effector cells and to mediate delayed type hypersensitivity (DTH) responses. Th2 cells produce IL4, IL5, and IL6 (DTH responses). Th2 cells also are considered to act as helper cells—for example, upregulating B cell and antibody responses. In addition, murine T cell clones have been described which have an unrestricted lymphokine secretion profile;13,14 these have been termed Th0 and it has been proposed that they represent a precursor stage in the development of the restricted Th1 and Th2 patterns. CD4+ T cells are the predominant infiltrating cell type in the early stages of EAU.15 In relation to the Th1/Th2 dichotomy the expression of mRNA for the lymphokines IFN-γ, IL2, lymphotoxin, and IL4 by, in general, similar numbers of cells implies the presence of a mixed infiltrate of both subsets. Alternatively, the CD4+ T cells could be of the Th0 subtype, not yet committed to a Th1 or Th2 lymphokine secretion pattern.

The active tissue destruction seen in the more advanced stages of EAU is characterised by a mixed infiltrate of lymphocytes and other inflammatory cells such as macrophages. It is likely that macrophages are also actively involved in local cytokine production in EAU. CD8+ T cells, which are present in greater numbers in the late stages of EAU are also probably involved in the process of cytokine production.

It has been demonstrated that the majority of infiltrating T lymphocytes in EAU do not appear to be S-antigen specific16 and it would be of interest to determine the antigen specificity of the T lymphocytes involved in lymphokine production in EAU.

The demonstration by in situ hybridisation of cytokine mRNA on a tissue section indicates that individual cells have been stimulated to express the cytokine gene but does not necessarily mean that they are actively producing cytokine protein. This has been demonstrated for IFN-γ production by rat CD4+ T cells16 where a greater frequency of CD4+ T cells positive for IFN-γ mRNA did not correlate with the level of secreted IFN-γ protein. It has been shown that stimulation of peripheral blood T cells resulted in expression of mRNA for IL2, IL2 receptor, and IFN-γ which correlated well with the levels of the respective proteins observed.17 Likewise, a good concordance has been demonstrated between IL2 and IL4 mRNA and secreted lymphotoxin as detected by bioassay.18 Recent work, however, has shown that cellular IL2 mRNA expression is not necessarily reflected by production of bioactive IL2.19 In view of the conflicting results of these studies, experimental work demonstrating lymphokine mRNA must be interpreted with caution with regard to production of bioactive
lymphokine. The work described in this paper does, however, illustrate that the T lymphocytes within the eye in EAU are activated to express lymphokine mRNA.

It is at present uncertain whether the lymphokine secretion patterns of rat T cells parallel those of human T cells although there appear to be basic similarities between the T cell systems. With regard to EAU the cellular infiltrate at the destructive sites in the experimental model is analogous to that found in human disease and an understanding of the effector cells involved has potential therapeutic applications in human disease. Demonstration of in vivo production of IFN-γ, IL2, lymphokinin, and IL4 implies that these lymphokines are potentially involved in the local pathobiology of EAU. Furthermore, the presence of these four lymphokines would allow CD4+ and CD8+ T lymphocytes and macrophages to act as effector cells in the destructive process in uveitic eyes. Further experimental work is necessary to define the biological roles of locally produced lymphokines in intraocular inflammatory disease.

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17 Cher DJ, Mosmann TR. Two types of murine helper T cell clone. II. Delayed type hypersensitivity is mediated by TH1 clones. J Immunol 1987; 138: 3688-94.