Aims—To examine the hypothesis that apoptosis of infiltrating cells contributes to spontaneous resolution of uveitis in clinically relevant rodent models.

Methods—Experimental melanin induced uveitis (EMIU) was induced in Fischer 344 rats by immunisation with 250 µg bovine ocular melanin. Endotoxin induced uveitis (EIU) was induced by injection of 200 µg Escherichia coli lipopolysaccharide. Formalin fixed, paraffin embedded ocular cross sections were stained by terminal deoxynucleotidyl transferase mediated deoxyuridine triphosphate nick end labelling (TUNEL) to identify apoptotic cells. Indirect immunoperoxidase staining of paraformaldehyde lysine periodate fixed tissue cross sections was used to demonstrate expression of inducible nitric oxide synthase (iNOS).

Results—TUNEL positive mononuclear cells were observed in the anterior uvea during both EMIU and EIU at all selected time points. However, whereas the majority of mononuclear cells appeared apoptotic from the outset of disease, neutrophils were notably TUNEL negative at all time points examined. Many infiltrating neutrophils expressed iNOS.

Conclusion—Apoptosis occurs early in the course of rat EMIU and EIU, and may contribute to resolution of these diseases. In general, infiltrating mononuclear cells die rapidly, while neutrophils survive, producing inducible nitric oxide synthase which may contribute to disease pathogenesis.

Materials and methods

Animals
Male and female Fischer 344 and DA rats were used aged 8 weeks or less. Animals were housed at 21°C and 50% humidity in a 12 hour light and 12 hour dark cycle, and were fed water and dried ration (New Joint Stock, Ridley Agriproducts, Murray Bridge, SA, Australia) ad libitum. Experimental protocols were developed in accordance with the National Health and Medical Research Council of Australia Statement for the Use of Animals in Research. All procedures and euthanasia were carried out under halothane (Zeneca Ltd, Macclesfield) inhalation anaesthesia.
INDUCTION OF EXPERIMENTAL MELANIN INDUCED UVEITIS

Ocular melanin was extracted from bovine choroids according to the protocol described by Broekhuyse and colleagues, and quantified as a dry weight. Rats received 125 µg of bovine ocular melanin in a 1:1 emulsion of sterile, non-pyrogenic 0.9% normal saline and Hunter’s TitreMax adjuvant (Sigma Chemical Company, St Louis, MO, USA) in a total volume of 60 µl by right hind footpad injection. Immediately afterwards, they were injected intraperitoneally with the same quantity of melanin mixed with 1 µg of pertussis toxin (Sigma Chemical Company) in a total volume of 40 µl normal saline.

INDUCTION OF ENDOTOXIN INDUCED UVEITIS

Rats were injected in one hind footpad with 200 µg of *Escherichia coli* 055:B5 lipopolysaccharide (Sigma Chemical Company) solubilised in 100 µl normal saline.

MONITORING OF UVEITIS AND COLLECTION OF UVEITIC EYES

Animals were examined daily at the slit lamp to identify the onset and course of uveitis. For TUNEL, groups of 3–7 animals were killed at predetermined time points. Rats with EMIU were killed at the time of injection, on days 1–2 (time of disease onset), days 3–7 (maximal inflammation), and during weeks 2–3 (resolution) of clinical uveitis. Rats with EIU were killed at the time of injection and at 24 hours (maximal inflammation) and 48 hours (resolution) after injection. For staining for iNOS, groups of 3–4 rats with EMIU were killed before development of uveitis, on days 1–2, 3–5, 7, and during weeks 2–3 of clinical disease.

INDUCTION OF ADJUVANT ARTHRITIS

The ankle joint of a rat with adjuvant induced arthritis was examined as a control tissue. Adjuvant arthritis was induced in a DA rat by subcutaneous injection of 0.1 ml of complete Freund’s adjuvant (Difco, Detroit, MI, USA) at the base of the tail. The rat was killed at day 12 when arthritis was florid.

STAINING BY TUNEL

TUNEL was performed according to a modification of a published method. Eyes were fixed in 10% buffered formalin for a minimum of 24 hours and paraffin embedded. The ankle joint was decalcified as previously described before similar fixation. Sections of 5 µm were deparaaffinised, hydrated through graded alcohols to distilled water, and permeabilised with 0.16 units/ml proteinase K (Merck, Darmstadt, Germany) for 10 minutes. Nick end labelling was accomplished by incubating sections with 0.11 units/ml terminal deoxynucleotidyl transferase in pH 6.6 buffer containing 5 mM cobalt chloride, 200 mM potassium cacodylate, 25 mM TRIS-HCl, 250 µg/ml bovine serum albumin, 0.375 nM biotinylated deoxy-uridine triphosphate (dUTP) and 75 µM deoxy-adenosine triphosphate (all from Boehringer-Mannheim, Mannheim, Germany) for 60 minutes at 37°C. The reaction was terminated by washing in 0.6 M sodium chloride 0.06 M sodium citrate buffer for 15 minutes, in Dulbecco’s A phosphate buffered saline (PBS) for 1 minute, and in PBS containing 2% volume/volume (v/v) fetal calf serum (Trace Biosciences, Sydney, NSW, Australia) and 0.2% v/v Triton X100 (Sigma Chemical Company) for 30 minutes. Non-specific binding sites were blocked with 10% v/v heat inactivated (56°C, 30 minutes) normal swine serum (Commonwealth Serum Laboratories, Melbourne, VIC, Australia) for 10 minutes, and the sections were washed for 10 minutes with PBS containing 0.2% weight/volume (w/v) gelatin (PBS gelatin) before endogenous peroxidase was blocked with 0.07% v/v hydrogen peroxide in methanol for 20 minutes. Sections were washed again in PBS gelatin for 10 minutes and then incubated with a 1 in 300 dilution of horseradish peroxidase conjugated streptavidin (Dako Corporation, Carpinteria, CA, USA) in PBS for 1 hour. Sections were washed in PBS gelatin, and developed for 5 minutes in 9 mM TRIS-HCl buffer at pH 7.6 with 40 mM sodium azide, 20 mM 3,3-diaminobenzidine tetrahydrochloride, 9 mM imidazole, and 0.07% v/v hydrogen peroxide (all from Sigma Chemical Company). Finally, sections were counterstained with haematoxylin. Negative control sections were stained in an identical manner, but with omission of biotinylated dUTP from the nick end labelling mixture. Positive control sections were treated for 10 minutes with 10 U/ml DNase I (Boehringer-Mannheim) before TUNEL staining.

VIDEO IMAGE ANALYSIS OF TUNEL SECTIONS

Ocular cross sections from selected rats were scanned at a magnification of 200X using a computer assisted colour video image analysis system (Video-Pro 32, Leading Edge, Adelaide, SA, Australia). Images were captured using a Panasonic CCD video camera and digitised with a PV 100 16 bit colour video digitiser card in an Intel 80486 DX processor based personal computer. The digitised image was displayed on a SVGA monitor in a 640 × 480 pixel variable window with 21 bit resolution. Transmitted light intensity and stability of light output were standardised. Video image analysis measurements were made of the total area stained blue (by haematoxylin) and brown (by the TUNEL chromogen product). The total number of haematoxylin stained cell nuclei (representing nuclei of both uveal cells and infiltrating cells) was determined using feature counting.

STAINING FOR INDUCIBLE NITRIC OXIDE SYNTHASE

Enucleated eyes from Fischer 344 rats were fixed in paraformaldehyde lysine periodate for 4 hours and dehydrated in 7% and 15% v/v sucrose in PBS. They were then embedded in Tissue-Tek OCT (Miles Incorporated, Elkhart, IN, USA) and snap frozen in liquid nitrogen. Ocular cross sections were cut by cryostat at 8 µm thickness. Sections were incubated for 10
minutes at room temperature with 10% v/v normal swine serum in PBS, then with a monoclonal murine anti-rat iNOS antibody (IgG2a isotype) (Transduction Laboratories, Lexington, KY, USA) or SAL5, an isotype matched irrelevant control antibody (the gift of Dr L Ashman, Hanson Centre for Cancer Research, Adelaide, SA, Australia) for 18 hours. At this point, and subsequently, the sections were washed with PBS gelatin. They were incubated for 30 minutes with biotinylated affinity isolated goat anti-mouse immunoglobulin (Dako Corporation) diluted 1:500 in PBS containing 1% v/v normal rat serum, washed, and then incubated for a further 30 minutes with horse-radish peroxidase conjugated streptavidin (DAKO Corporation) diluted 1 in 1000 in PBS. Sections were developed and counterstained as described above.

Results

APOPTOSIS DURING EU AND EMIU
In the initial stages of EMIU, many TUNEL negative neutrophils were observed infiltrating the uvea and anterior chamber, but most mononuclear cells were TUNEL positive (Fig 1). During the period of florid inflammation, there were many positive staining mononuclear cells infiltrating the anterior uvea, the aqueous, and vitreous, and sometimes the limbus and choroid. In one representative, severely inflamed eye, 79% (SD 4%) of iris cells, 75% (3%) of ciliary body cells, and 88% (7%) of aqueous cells were TUNEL positive by video image analysis. As the disease began to resolve, apoptotic mononuclear cells were still detectable, although numbers of infiltrating cells were reduced.

Figure 1  Sections of a Fischer 344 rat eye during early EMIU (at time of disease onset), stained by TUNEL.  
(A) Ciliary body, negative control: no biotinylated dUTP.  
(B) Ciliary body, test, showing many TUNEL positive (apoptotic) mononuclear cells.  
(C) Anterior chamber, test, showing TUNEL positive mononuclear cells and TUNEL negative neutrophils.  
Haematoxylin counterstain.  
Magnification ×1400.
The Fischer 344 rat showed moderate anterior segment inflammation and vitritis in response to systemic endotoxin injection. At 24 hours (the time of maximal inflammation), infiltrating cells were seen in the aqueous and vitreous. The majority were TUNEL negative neutrophils, although a moderate number of TUNEL positive mononuclear cells were also present (Fig 2). In one representative eye, 40% (8%) of aqueous cells were TUNEL positive, but readings taken from the uveal tissues did not vary significantly from baseline. At 48 hours, a moderate infiltrate was still present in some eyes, but these cells were generally in the vitreous. TUNEL negative neutrophils predominated, but some TUNEL positive mononuclear cells were also present.

No TUNEL positive nuclei were observed in negative control sections (biotinylated dUTP omitted from the nick end labelling mixture). All nuclei were TUNEL positive in the positive control sections (treatment with DNase I before TUNEL staining). In test sections from rats killed before the onset of clinical and histological uveitis, an unexpectedly high background of TUNEL positive cells was noted. When quantified by video image analysis, this background amounted to 58% (20%) and 56% (18%) total cells in the iris and ciliary body, respectively. However, cells infiltrating the aqueous or vitreous during uveitis were plainly distinguishable from the background.

In contrast with the results obtained with the ocular sections, very few apoptotic nuclei were visible in the infiltrate present in the ankle joint of a DA rat with florid adjuvant arthritis, included as a control for severely inflamed tissue infiltrated with leucocytes (Fig 3).

**Discussion**

Staining by TUNEL showed that apoptosis was a prominent feature in both melanin induced and endotoxin induced models of uveitis and that cell death began very early, at the time of onset of clinically apparent and histologically evident inflammation. However, whereas infiltrating mononuclear cells were found to die rapidly in situ, infiltrating neutrophils were found to survive. If apoptosis is FasL mediated, which seems most likely given the constitutive expression of this molecule within the anterior uvea,23 and given that FasL could readily be detected in the rat eye by immunoperoxidase staining (data not shown), the neutrophil must become refractory to this death signal. Watson and colleagues have recently demonstrated that the cross linking of the adhesion molecule CD11b on the neutrophil surface, the endothelial transmigration of neutrophils into inflammatory sites and, of particular relevance to EIU, the presence of lipopolysaccharide in the local microenviroment...
Apoptosis is a prominent feature of acute anterior uveitis in the Fischer 344 rat.

In addition to the Fas-FasL interaction, there are other mechanisms by which apoptosis may be induced. The tumour necrosis factor receptor (TNFR), TNFR1 or p55, which is structurally related to FasL, may trigger apoptosis after binding tumour necrosis factor alpha (TNF-α)\(^8\). It follows that TNF-α, a pro-inflammatory cytokine believed to be expressed in the anterior uvea of rats with uveitis,\(^9\)\(^-\)\(^10\) is possibly signalling the apoptosis of infiltrating leucocytes which we observed. Yet, why mononuclear cells should be susceptible to this signal whilst neutrophils are resistant to it, cannot be explained on the basis of current knowledge.

Our findings led us to speculate that the neutrophil may play an important effector part in uveitis, even in T lymphocyte mediated disease as exemplified by EMU. Neutrophils release an enormous range of pro-inflammatory mediators including eicosanoids, platelet activating factor, cationic proteins, and...
cytokines. They also produce iNOS, an enzyme which catalyses the generation of the free radical, nitric oxide (NO). There is substantial evidence for a pathogenic role for nitric oxide in EIU, particularly in relation to breakdown of the blood-aqueous barrier. Expression of iNOS mRNA and its translated product by infiltrating and local cells is detected during EIU, and drugs which inhibit iNOS activity significantly reduce the severity of the inflammation. That iNOS is not obligatory for disease expression is highlighted by our finding that EIU occurs even in iNOS gene knockout mice. However, our observation that iNOS is expressed during the early stages of EMIU lends weight to the suggestion that nitric oxide is at least one of the molecular mediators of uveitis, produced by the neutrophil which has managed to avoid FasL-mediated apoptosis.

What then is the role of the mononuclear cell in uveitis? In a recent study, reverse transcriptase polymerase chain reaction was used to determine mRNA expression of a range of cytokines in the iris and ciliary body during EMIU. Only mRNA for TNF-α, a cytokine that can be produced by neutrophils, rose to high levels. In contrast, mRNA levels of interferon gamma, interleukin (IL) 2, IL-4, IL-6, IL-10, IL-12, cytokines which are produced predominantly by lymphocytes and/or monocytes, showed little or no increase. In EIU, mRNAs for a variety of cytokines are detected before infiltrating leukocytes are seen, but resident tissue macrophages responding directly to lipopolysaccharide could contribute to the production of many of these cytokines. In situ hybridisation has, in fact, localised TNF-α and IL-1 mRNA to “histiocyte-like” cells in the anterior uvea. These findings fit with our observation that most infiltrating mononuclear cells seen in the eye during EMIU and EIU are either dead or dying, and are not releasing a cascade of cytokines. Although the phenotype of these mononuclear cells was not determined in this study, previous work indicates they are lymphocytes and macrophages/monocytes. In marked contrast with the findings in the eye, relatively few apoptotic cells of any phenotype were observed in the ankle synovium during florid adjuvant arthritis, a finding that was to be expected given that FasL is not constitutively expressed in this tissue.

In summary, the available evidence leads us to hypothesise that apoptosis of mononuclear cells may certainly contribute to the resolution of uveitis, but that neutrophils refractory to the Fas-FasL mediated death signal may be responsible for tissue damage. Treatment directed specifically against the neutrophil or neutrophil products may have significant therapeutic value both in T lymphocyte controlled disease and in predominantly neutrophil mediated inflammation. Clinical similarities between the two animal models of acute anterior uveitis and acute anterior uveitis in humans lead us to consider the possibility that there are parallels for disease pathogenesis at both cellular and molecular levels. Treatments which specifically target the neutrophil may be useful in the management of human acute anterior uveitis.

We thank Professor S Wesselingh and Ms M Lewis for expert advice, and Mr R Yates for animal husbandry. This work was supported by The National Health and Medical Research Council of Australia, The Ophthalmic Research Institute of Australia, and The Flinders Medical Centre Foundation.

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