Apopotosis mediates decrease in cellularity during the regression of Arthus reaction in cornea

Noriko Ozaki, Masamichi Ishizaki, Mohammad Ghazizadeh, Nobuaki Yamanaka

Abstract

**Background/aims**—The Arthus type allergic reaction is characterised by inflammatory cell infiltration and marked neovascularisation in the cornea. During the healing stages, inflammatory cells and newly formed microvessels gradually disappear. The aim was to establish whether apoptosis affected the regression of inflammatory cells and newly formed microvessels, in order to define more clearly the cellular mechanisms involved in the pathobiology of corneal diseases.

**Methods**—Albino male rabbits were injected subcutaneously with 5 mg/ml bovine serum albumin (BSA) incorporated in Freund's complete adjuvant twice weekly. Under the anaesthesia, 30 µl of a 0.5 mg/ml BSA solution was injected into the central corneal stroma to induce an Arthus type allergic reaction. The injured corneas were collected at various time points ranging from 3 to 20 days. Apoptotic cells were identified by both light microscopy using in situ Tdt-dUTP nick end labelling (TUNEL) method and electron microscopy.

**Results**—With increasing time after induction of the Arthus reaction, marked neovascularisation and infiltrated inflammatory cells such as polymorphonuclear cells (PMNs) and plasma cells were observed in the cornea. Thereafter, the inflammatory cells and newly formed microvessels gradually disappeared. Coincidently, the numbers of microvascular endothelial cells and infiltrated inflammatory cells undergoing apoptosis were increased. Apoptotic bodies were taken up by macrophages, PMNs, as well as myofibroblasts derived presumably from transformation of migrated keratocytes.

**Conclusions**—These data demonstrate that regression of the cellular infiltrates and microvessel endothelial cells associated with the Arthus reaction in the cornea occurs via apoptosis. This finding adds insights into the cellular mechanisms regulating the pathobiology of corneal diseases.

The avascular cornea has highly regular orientation of stromal collagen fibres that enhances optical transparency, and makes it distinct from other tissues in the body. Inflammation in the cornea differs in many respects from that in other body sites. Inflammatory cells may infiltrate the cornea from various sites, such as conjunctival and scleral vessels, via the limbus, and less commonly from the anterior chamber. When oedema is accompanied by inflammation, a portion of the disruptive light scattering is caused by infiltration of the stroma by inflammatory cells and by exudates derived from the limbal vessels. The degree of opacification and visual disturbance appears to be related to the number and density of inflammatory cells and their location, and to the presence or absence of associated enzymatically induced stromal damage, vascularisation, and calcium deposition. To maintain corneal transparency, it is necessary to eliminate the infiltrated inflammatory cells. On the other hand, corneal neovascularisation is a sequel of numerous inflammatory lesions of the ocular anterior segment. In addition, not only corneal neovascularisation but also ocular neovascularisation such as that occurring in diabetic proliferative retinopathy, neovascular glaucoma, and choroidal neovascularisation can cause severe disturbances of ocular function. Therefore, it is highly imperative to elucidate the mechanisms underlying the development, as well as, regression, of neovascularisation.

Recently, one form of cell death referred to as “programmed cell death” mediated by a process termed “apoptosis” has been recognised. Apoptosis plays an important part during tissue development, maintaining homeostasis, response to infection and wound healing by eliminating unwanted cells. Meeson et al reported that regression of capillary channels in pupillary membrane is caused by apoptosis. Wilson and Kim suggested that keratocyte apoptosis after corneal epithelial injury is an initiator of the wound healing response. A characteristic feature of apoptosis is recognised as DNA digestion at internucleosomal sites, producing small double stranded fragments of DNA that migrate in a ladder pattern (multiple of 185 bps) after electrophoresis in agarose gels. Moreover, apoptosis is identified morphologically by electron microscopy, or by light microscopy using the in situ Tdt-dUTP nick end labelling (TUNEL) method. Typical morphological changes in apoptosis include cell shrinkage and loss of normal contacts, dense chromatin condensation, cellular budding, and fragmentation. Finally, apoptotic cells are normally eliminated by macrophages or neighbouring cells without an inflammatory response.

The rabbit cornea provides an ideal site for studying the in vivo consequences of antigen-antibody reactions and diffusion of the newly formed microvessels as it is transparent and avascular. Germuth et al reported that the
of the unilateral eye under the anaesthesia with sodium thiopentone (Ravonal). Control animals were injected with 30 µl sterile normal saline in the central corneal stroma of the unilateral eye. The corneas were observed during the experimental period by direct visual examination. The rabbits were sacrificed with an overdose of sodium thiopentone intravenously at various time points ranging from 3 to 20 days. At each time point, three to four injured corneal buttons with adjacent sclera were collected and processed for light and electron microscopic examinations.

**LIGHT MICROSCOPY**

Corneal tissue specimens were fixed in 4% buffered paraformaldehyde for 16–24 hours at 4°C. Each tissue was divided into two halves. One semicircular half was embedded in paraffin and the other was immersed in OCT compound and frozen in dry ice-acetone. A section from each paraffin block was stained with haematoxylin and eosin for histological examinations.

**MORPHOLOGICAL IDENTIFICATION OF APOPTOSIS**

Apoptotic cells were assessed morphologically by both light and electron microscopy. For light microscopy, a modification of the TUNEL method was used to detect the apoptotic cells, and a simultaneous immunostaining for thrombomodulin (TM) was performed to identify the microvessels. Briefly, 4 µm thick frozen sections were incubated with 20 µg/ml proteinase K for 15 minutes at room temperature and washed in distilled water. The sections were rinsed in TdT buffer (30 mM TRIS-HCl, pH 7.2, 140 mM sodium cacodylate, 1 mM cobalt chloride) and incubated with 0.5 U/µl TdT (Gibco BRL; Rockville, MD, USA) and 10 µM fluorescein-12-dUTP (Boehringer Mannheim, Germany) in TdT buffer for 60 minutes at 37°C. The sections were washed with PBS four times each for 5 minutes and incubated with goat anti-rabbit TM (provided by Dr David Stern, Columbia University) at 1:800 dilution. After washing three times with PBS, the sections were incubated with swine anti-goat IgG labelled with Texas red (EY Laboratories, San Mateo, CA, USA) at 1:50 dilution, washed three times with PBS, the sections were incubated with goat anti-rabbit IgG (30 mM TRIS-HCl, pH 7.2, 140 mM sodium cacodylate, 1 mM cobalt chloride) and incubated with 0.5 U/µl TdT (Gibco BRL; Rockville, MD, USA) and 10 µM fluorescein-12-dUTP (Boehringer Mannheim, Germany) in TdT buffer for 60 minutes at 37°C. The sections were washed with PBS four times each for 5 minutes and incubated with goat anti-rabbit TM (provided by Dr David Stern, Columbia University) at 1:800 dilution. After washing three times with PBS, the sections were incubated with swine anti-goat IgG labelled with Texas red (EY Laboratories, San Mateo, CA, USA) at 1:50 dilution, washed three times with PBS, and cover slipped using Vectashield H-1000 mounting medium (Vector Laboratories, Inc, Burlingame, CA, USA).

**INTERPRETATION OF STAINING**

Examination of the stained sections and photography were performed with a fluorescence microscope (Model BX 60, Olympus, Tokyo, Japan). The number of apoptotic cells was assessed semiquantitatively. TUNEL positive endothelial cells of microvessels and inflammatory cells in each area were counted in five microscopic fields at 400x magnification and averaged.

**ELECTRON MICROSCOPY**

For the electron microscopic study, the excised corneas were cut into small pieces and fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffered solution and frozen in dry ice-acetone. A section from each paraffin block was stained with haematoxylin and eosin for histological examinations.

**Methods**

**INDUCTION OF ARTHUS REACTION IN CORNEAS**

Albino male rabbits weighing 2.5–3 kg were purchased from Saitama Experimental Animal Supply (Saitama, Japan). Animal experiments were performed in compliance with the experimental animal ethics review committee of Nippon Medical School, Japan. As described previously, animals were injected subcutaneously weekly twice with 5 mg/ml bovine serum albumin (BSA) incorporated in Freund’s complete adjuvant. Thereafter, the rabbits were injected with 2 mg/ml BSA as a booster weekly two or three times. Rabbits whose antibody titre for BSA had risen more than 16-fold by the Ouchterlony method were used for this study. To induce the Arthus type allergic phenomenon, 30 µl of an 0.5 mg/ml BSA solution was injected in the central corneal stroma of the sensitised rabbit eye under the anaesthesia with sodium thiopentone (Ravonal). Control animals were injected with 30 µl sterile normal saline in the central corneal stroma of the sensitised rabbit eye. The sensitised rabbit induced an immune reaction as a result of injection (arrow) at day 0 (0d), immuno-ring (arrow) at 2 days (2d) after injection and induction of neovascular response. Newly formed microvessels (arrow) reached to the centre of the cornea at 12 days (12d). These microvessels (arrow) gradually diminished at the healing stage on day 20 (20d).

**Figure 1** Sensitised rabbit corneas showing opacity as a result of injection (arrow) at day 0 (0d), immuno-ring (arrow) at 2 days (2d) after injection and induction of neovascular response. Newly formed microvessels (arrow) reached to the centre of the cornea at 12 days (12d). These microvessels (arrow) gradually diminished at the healing stage on day 20 (20d).
buffer, pH 7.4, overnight at 4°C. After washing in 0.1 M phosphate buffer, the tissue blocks were postfixed with 1% OsO₄ in 0.1 M phosphate buffer, pH 7.4, for 2 hours, dehydrated, and embedded in Epon 821.

Ultrathin sections were stained with uranyl acetate and lead citrate, then examined with a Hitachi H-7100 electron microscope at an accelerating voltage of 75 kV.

Results
GROSS EXAMINATION
Immediately after injection of BSA solution, the cloudiness of the cornea as a result of the injection was observed (Fig 1, 0d). Several hours after injection, the cloudiness was resolved but not cleared up completely. At 2 days after injection, an immuno-ring was induced by precipitation of immune complexes (Fig 1, 2d). At day 3, the neovascular response originating from the limbal vessels was evident. At day 9, extensive newly formed microvessels extended towards the immuno-ring, and at the end of second week reached the centre of the cornea with an increase in the central portion and a decrease around the limbus (Fig 1, 12d). In the areas through which the network of newly formed microvessels had transited, the cloudiness disappeared. By the end of 3 weeks, these microvessels were also gradually diminished (Fig 1, 20d).

LIGHT MICROSCOPY
At 2 days, the corneas that formed immuno-ring were oedematous and demonstrated the migration of inflammatory cells. By the fluorescent antibody method, rabbit IgG and C3 deposits were present coincidently with the immuno-ring (data not shown). By TUNEL assay, no cell stained positively. At day 3, infiltration of inflammatory cells was noted in association with the immuno-ring (Fig 2a), and at the limbus extensive newly formed microvessels were observed (Fig 2b). TUNEL positive cells were present in the stroma (Fig 3, arrowheads), but there were no such cells in the microvessels (Fig 3, arrows). The area of hypercellularity including capillaries and inflammatory cells gradually translocated from the periphery to the centre of the cornea. After several days, TUNEL positive cells in the microvessel wall began to appear (Fig 4, arrowheads). They were observed frequently in the area circumjacent to the limbus side of the cornea.
capillary accumulation, and less at the periphery of this area. TUNEL positive cells could still be observed in the stroma at the end stage of inflammation. The accumulation of plasma cells surrounding the microvessels was remarkable, but then decreased in the area through which the network of capillaries had transited.

**ELECTRON MICROSCOPY**

By electron microscopy, various types of apoptotic cells and bodies were detected. From 3 days after the challenge, apoptotic neutrophils and bodies containing fragments of chromatin condensation were observed. Excessive apoptotic bodies beyond the capacity of take up by phagocytes were occasionally observed (Fig 5). From day 9, plasma cells showed apoptotic changes (Fig 6). From day 12, newly formed microvessels containing endothelial cells with initial apoptotic changes such as crescent condensation of the chromatin were observed (Fig 7). Apoptotic bodies were found in phagocytic cells (Fig 8). Apoptotic PMNs were phagocytised by macrophages (Fig 8A) and apoptotic neutrophils by PMNs (Fig 8B). Apoptotic bodies were also taken up by fibroblastic cells (Fig 8C).

**Discussion**

The cornea is an immune privileged site.\(^{16}\) Immune privilege is a term applied to several organs including the brain, testis, and ovary and these sites prohibit the spread of inflammation, since even minor episodes can threaten organ integrity and function.

Recently Griffith et al showed that the induction of apoptosis is a potent mechanism of immune privilege.\(^{17}\) In the present study, we observed the developmental process and regression of infiltrated inflammatory cells and newly formed microvessels in the rabbit cornea sensitised with BSA and focused on changes in the corneal cellularity and the relevance of apoptosis in this respect.

At 2 days after the challenge, immuno complex was formed by antigen-antibody reaction, which was observed as an immuno-ring with the naked eyes. Subsequently, the complement system could be activated by the antigen-antibody reaction or by the resulting complexes. Such an activation could release complement C3a and C5b fragments and they could induce infiltration of PMNs from the limbal vessels. Previous studies have shown that the membrane attack complex (MAC) which is composed of complement C5b-9 fragments promoted the secretion of IL-8 which induces infiltration of PMNs from human umbilical vein endothelial cells.\(^{18}\) This process is thought as a mechanism of PMNs infiltration in the cornea.

The half life of circulating PMNs is estimated to be 5 hours, but that of infiltrated PMNs in tissue is unclear. In either case the ageing PMNs eventually undergo apoptosis.\(^{19}\) Furthermore, it is well known that in acute inflammation G-CSF, GM-CSF, interferon \(\gamma\), and TNF-\(\alpha\) suppressors delay apoptosis of PMNs.\(^{20} 21\) In this study, the evidence for PMNs undergoing apoptosis was documented at 3 days, but not earlier, such as at 1 day. The same mechanism may be responsible for delayed apoptosis of neutrophils in the present study.
We observed plasma cells infiltrating around the newly formed microvessels. In general, plasma cells represent the final differentiation stage of B lymphocytes after encountering antigenic agents and interacting with various cell types. The number of plasma cells has to be tightly controlled to avoid overproduction of antibodies. Accordingly, it seems reasonable to speculate that unnecessary plasma cells undergo apoptosis to maintain the balance. In this study, we confirmed in vivo that plasma cells in the cornea underwent apoptosis, particularly in the areas through which the newly formed capillary network had transited. This finding further suggests that apoptosis may participate in the control of antibody producing cells.

From 3 days after the challenge, corneal neovascularisation was observed resulting in more increase in corneal cellularity and consequently the loss of corneal transparency. If corneal neovascularisation persists for a long period of time, it may result in lipid leakage and haemorrhage. After the inflammation subsided, the newly formed microvessels and corneal hypercellularity gradually regressed. We previously confirmed that the endothelial cells of newly formed microvessels in the injured cornea expressed TIMP-2, which is an inhibitor of MMPs (unpublished data). It was also reported that TIMP-2 inhibits neovascularisation.\(^{22}\) It is possible that the expression of TIMP-2 may contribute to the regression of newly formed microvessels. Pollman et al\(^{11}\) demonstrated that endothelial cell apoptosis is a prominent feature of in vitro capillary network regression and formation.

In this study, electron microscopy revealed that the endothelial cells of newly formed microvessels underwent apoptosis. This finding suggests that apoptosis reduces the hypercellularity due to the infiltration of PMNs, appearance of plasma cells, and neovascularisation in the cornea, leading to the recovery of the corneal transparency.

At present, it is not clear whether total regression of inflammatory cells in our study is through apoptosis. However, studies on apoptosis of normal cells in the developing rat kidney have shown that uptake of apoptotic cells by phagocytes and degradation of ingested apoptotic cells beyond histological recognition occurs very rapidly (within an hour or so) as a means of cell clearance.\(^{23}\) Therefore, low frequencies of histologically demonstrable apoptotic cells reflects clearance of large numbers of cells, particularly since a macrophage can ingest many apoptotic cells. This factor may account for the low frequency of apoptotic cells observed in the histological sections in this study.
It is usually stated that the apoptotic bodies are often taken up by neighbouring cells, and macrophages do not induce an inflammatory response. Tsuji et al. reported that apoptotic human umbilical vein endothelial cells can activate the alternative pathway of the homologous complement and that the complement is related to the clearance of apoptotic cells by phagocytes. On the other hand, myofibroblasts that are characterized by the expression of α-smooth muscle actin contribute to the healing of wounds. Recently, it has been reported that during corneal wound healing, the keratocytes that migrate to injured stroma transform into myofibroblasts. In our study, it was confirmed by electron microscopy that the apoptotic bodies were taken up by macrophages, PMNs, and myofibroblasts. In this process, it is possible that an alternative pathway may be involved, and apoptosis may contribute to the recovery of the corneal transparency. On the other hand, we were unable to detect DNA degradation by electrophoresis, probably due to the overall gradual occurrence of apoptosis. Meanwhile, in the occasionally observed areas with several apoptotic bodies, excessive apoptotic bodies beyond the capacity of take up by phagocytes may undergo secondary necrosis and participate in the prolonged inflammatory sequel there.

Several factors have been proposed as regulators of apoptosis. Fas ligand and tumour necrosis factor (TNF), bind to their receptors and induce apoptosis, while overexpression of bcl-2 blocks it. Griffith et al. showed that in the mouse cornea, Fas ligand is expressed on the epithelium and endothelium, suggesting its importance in controlling inflammatory cells entering from the conjunctiva or anterior chamber. Stuart et al. reported that Fas ligand expressed on human cornea is a critical factor determining the success of orthoptic corneal allografts by inducing cell death in inflammatory cells. It has been reported that all plasma cell lines expressed Fas at a low to moderate level. As also in the present study, it is possible that by Fas-Fas ligand interaction, inflammatory cells and endothelial cells of newly formed microvessels may undergo apoptosis.

In conclusion, our data demonstrate that regression of the cellular infiltrates and microvessel endothelial cells associated with the Arthus reaction in the cornea occurs via apoptosis. This finding adds insight into the cellular mechanisms regulating the pathobiology of corneal diseases.

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