Presence of α smooth muscle actin in lens epithelial cells of aphakic rabbit eyes

Daijiro Kurosaka, Katsuhiro Kato, Toshiyuki Nagamoto

Abstract

Aims—To determine whether α smooth muscle actin (α-SMA), a marker for myofibroblastic cells, is present in lens epithelial cells (LECs) in rabbit aphakic eyes.

Methods—Phacoemulsification was performed in rabbit eyes, which were enucleated after surgery. Immunohistochemical methods were used to detect α-SMA in LECs.

Results—Five days after surgery, the presence of α-SMA positive LECs was observed mainly around the adhesive portion of the anterior capsule margin and the posterior capsule. The posterior capsule was wrinkled at the adhesive portion. The α-SMA positive LECs were flattened with spindle-shaped cross sections. Seven days after surgery, the α-SMA positive LECs covered most of the central posterior capsule. They disappeared 10 days after surgery. On the other hand, the cuboidal LECs in the capsular bag were negative for α-SMA.

Conclusion—The flattened LECs with spindle-shaped cross sections observed 5 days after cataract surgery contained α-SMA. Such LECs were distinguished biochemically from the cuboidal LECs, which lacked α-SMA.

(Br J Ophthalmol 1996;80:906–910)

Following cataract surgery, the residual lens epithelial cells (LECs) migrate and proliferate.1–4 Owing to their contractile ability, some LECs produce fibrotic capsular opacification and contraction of the lens capsule.6,7 This directly causes visual disturbances6 and also brings about postoperative complications such as a narrowing of the anterior capsular opening,6–10 retinal detachment,11,12 decentration of an intraocular lens,6 and postoperative hypotony.10

Alpha smooth muscle actin (α-SMA) is an isoform of actin and a marker for myofibrodiferentiation13–14 that is involved in the contractility of various cells.15–17 The contractility of the fibroblasts depends on α-SMA expression and is inhibited by the microinjection of an α-SMA antibody into these cells.15 The ability of retinal pigment epithelial cells to exert tractional forces on their matrix increases with continued passage in vitro, and is correlated with the de novo expression of α-SMA.16 We recently reported that the contractility of LECs induced by growth factors also depends on the expression of α-SMA in vitro.17

The LECs from anterior subcapsular cataract (ASC), but not those from the normal lens, express α-SMA in vivo.18 The LECs involved in ASC are flattened rather than cuboidal, and are surrounded by collagen fibrils.19,20 They cause the lens capsule to contract.19,20 Following cataract surgery, the LECs that produce capsular contraction become flattened and are surrounded by collagen fibrils.4,17 Ultrastructurally, these post-cataract surgery LECs resemble those of ASC.7 These observations suggested that the flattened LECs found after cataract surgery also express α-SMA in vivo. The presence of α-SMA has been reported only in the intercellular stroma of the LECs observed in the fibrotic membrane removed from the surface of the optic of the intraocular lens of the patient.21 We conducted the immunohistochemical study in aphakic rabbit eyes to better understand the relation between α-SMA and the LECs after cataract surgery.

Material and methods

A total of 48 young Japan albino rabbits, weight 1.0–1.5 kg, were studied. Each of the following agents was applied topically three times on the day of surgery to one eye of each rabbit: diclofenac sodium, tropicamide, phenylephrine hydrochloride, and nofloxacin. Animals were anaesthetised with intravenous pentobarbitone sodium (40 mg/kg) and atropine sulphate (0.02 mg/kg). A superior corneal incision was made with a 3 mm keratome, and a viscoelastic material (hyaluronic acid, Pharmacia, Uppsala, Sweden) was injected into the anterior chamber. Continuous circular capsulorhexis of the anterior capsule was carried out using a capsulorhexis forceps. The lens nucleus was emulsified and the residual cortex was removed with a phacoemulsifier (Cavitron 9001, Alcon Surgical, Fort Worth, TX, USA). The corneal incision was closed with a continuous 10-0 nylon suture. At the end of the operation, gentamicin (10 mg) was injected subconjunctivally and an antibiotic ointment (oxytetracycline hydrochloride) was applied to
the eye. Procedures adhered to the guidelines of the Association for Research in Vision and Ophthalmology Resolution on the Use of Animals in Research.

Rabbits were killed with an overdose of pentobarbitone sodium at 0 hours, at 1, 3, 5, 7, and 10 days, and at 2 weeks, 1, 3, and 6 months after the operation. At least four rabbits were studied at each specified time. Eyes were enucleated and immersed in 10% neutral buffered formalin. After fixation, the globe was sectioned at the equator. Specimens were dehydrated through a graded series of alcohols and embedded in paraffin. Sections were cut from specimens.

The labelled streptavidin-biotin method using a Histostein-SP kit (Zymed, South San Francisco, CA, USA) was used to detect α-SMA immunohistochemically. Sections were deparaffinised with xylene, rehydrated in a graded series of alcohols, and immunostained for α-SMA according to the manufacturer’s instructions, with the following modifications. Endogenous peroxidase activity was quenched by treating the sections with 3% hydrogen peroxide in methanol for 20 minutes at room temperature. Non-specific background staining was eliminated by incubating the sections with non-immune goat serum (reagent 1A; H-SP kit) at room temperature. A primary mouse monoclonal antibody directed against α-SMA (IgG2a, clone 1A4, code no M851, Dakopatts, Denmark) was diluted 1:100 with primary antibody diluting buffer (Biomedia Corp, Foster City, CA, USA). Sections were incubated with this diluted antibody in a moist chamber at room temperature for 30 minutes. Sections were then rinsed three times with phosphate buffered saline (PBS), incubated with a biotinylated secondary antibody (goat anti-mouse IgG; reagent 1B; H-SP kit) at room temperature for 10 minutes, and washed three times with PBS. Streptavidin-peroxidase was added (reagent 2; H-SP kit), and the sections were incubated at room temperature for 15 minutes, then rinsed three times with PBS. Finally, peroxidase was visualised by the addition of a solution containing 3-3’-diaminobenzidine hydrochloride (0.3 mg/ml), 0.005% hydrogen peroxide, and 50 mM TRIS-HCl buffer at room temperature for 4 minutes. Sections were then counterstained with haematoxylin.

Mouse monoclonal IgG2a antibody (clone Dak-G05, code no X943, Dakopatts) was used as a negative control. No immunoreaction was detected in the negative control. The sphincter and dilator muscles of the iris were used as internal positive controls.22

**Results**

Some residual LECs were observed on the peripheral anterior capsule immediately after cataract surgery. Some cortical lens fibres also persisted. These LECs and lens fibres were negative for α-SMA (data not shown).

On postoperative day 1, the anterior capsule at the capsulotomy margin rolled up and adhered to the posterior capsule. This adhesion divided the postoperative lens into three portions: the central posterior capsule, the adhesive portion, and the capsular bag. The capsular bag consisted of the peripheral posterior capsule and anterior capsule. In the capsular bag, LECs on the anterior capsule began to flatten and migrate to the posterior capsule via the lens equator and the adhesive portion. However, LECs did not migrate to the central posterior capsule from the adhesive portion. LECs were negative for α-SMA (Fig 1).

On postoperative day 3, a monolayer of LECs covered the inner surface of the anterior and posterior capsules in the capsular bag. Some of these LECs were vacuolated. The capsules were attached to one another like a zipper. The LECs at the adhesive portion were multilayered but not vacuolated, and were sur-
rounded by a fibrous material. Few LECs migrated to the central posterior capsule. These LECs were negative for α-SMA (Fig 2).

On postoperative day 5, the multilayered LECs at the adhesive portion were flattened with spindle-shaped cross sections, whereas the LECs in the capsular bag were monolayered and vacuolated. The fibrous material that had surrounded the LECs at the adhesive portion had disappeared and the LECs had begun to migrate to the central posterior capsule. Only the flattened LECs at the adhesive portion were positive for α-SMA (Fig 3).

On postoperative day 7, the flattened LECs were α-SMA positive, and covered almost the entire central posterior capsule (Fig 4 (top and middle)). The capsular bag was occupied by vacuolated LECs that were negative for α-SMA. After the seventh postoperative day, clumps of α-SMA positive, flattened LECs were occasionally visible in the capsular bag (Fig 4 (top and bottom)).

On postoperative day 10, most of the flattened LECs that earlier had covered the central posterior capsule had now disappeared, whereas the flattened LECs at the adhesive portion remained. The remaining flattened LECs were positive for α-SMA (Fig 5).

After postoperative day 14, the LECs that covered the inner surface of the anterior capsule exhibited a normal cuboidal appearance. The LECs gradually became differentiated. The capsular bag was now filled with fibres and had formed Soemmerring's ring. However, the LECs at the adhesive portion were flattened and positive for α-SMA. The posterior capsule was wrinkled at the adhesive portion, to which the α-SMA positive LECs were attached (Fig 6). At this time, the LECs did not migrate to the central posterior capsule.

Discussion
After the fifth postoperative day, flattened LECs were observed mainly around the adhesive portion of the anterior capsule margin and the central posterior capsule, and were present only transiently on the central posterior capsule. These flattened LECs could be distinguished morphologically from the monolayered LECs in the capsular bag, which showed early vacuolation and later became cuboidal. The flattened LECs related to the wrinkling of the lens capsule. This behaviour of the flattened LECs resembled that described previously by other investigators.6–8 However, the present study also demonstrated that these flattened LECs were α-SMA positive.

Flattened LECs, especially around the adhesive portion of the anterior capsule margin and the central posterior capsule, are involved in capsular contraction and are thus considered to have contractile ability.6–7 Ultrastructurally, they reportedly exhibit the specific microfilaments that are the morphological features of the myofibroblasts.6–5

Myofibroblasts possess morphological and biochemical features that are between those of the fibroblast and the smooth muscle cell, and are involved in retractive phenomena and the accumulation of the extracellular matrix.13–14 α-SMA is the isoform that is present in the smooth muscle cells, and is a hallmark of the myofibroblastic phenotype.13–14 Our results suggested that 5 days after surgery, flattened LECs possessed the biochemical features of the myofibroblast.

Five days after cataract surgery, the α-SMA positive LECs were observed mainly around the adhesive portion of the anterior capsule margin and the posterior capsule, and were present only transiently on the central posterior capsule. Active transforming growth factor
Presence of α-smooth muscle actin in lens epithelial cells of aphakic rabbit eyes

β2 (TGF-β2) is present in the aqueous humour of humans and rabbits. It was recently found that TGF-βs increase the expression of α-SMA in cultured LECs. These observations suggest that the TGF-β2 in the aqueous humour may play a role in the presence of α-SMA in LECs.

In this study, the α-SMA-positive LECs disappeared from the central posterior capsule 10 days after the operation although they had almost covered the central posterior capsule 3 days earlier. They were not observed thereafter.

In aphakic dog eyes, LECs are present in the central posterior capsule 7 days after surgery, but are not observed 10 weeks after surgery. Liu et al. reported that the majority of rat LECs disappear from explants that were exposed to TGF-β for 5 days. These observations suggest that TGF-β may also be involved in the disappearance of LECs from the central posterior capsule. However, in aphakic human eyes, LECs have been found in the central posterior capsule as long as 2 years postoperatively. The reason for these species differences is not known.

The present study showed the presence of α-SMA in the LECs but not in the intercellular stroma. However, Frezzotti et al. described the presence of α-SMA filaments in the intercellular stroma, but not in the LECs, of the fibrotic membrane that was removed surgically from the surface of the optic of the intraocular lens of the patient. In a study of monkey eyes, most LECs around the anterior capsule margin showed signs of degeneration 12 months after cataract surgery; ultrastructurally, cellular debris was observed around these cells. We suspect that some of the cellular debris in the stroma may have shown positivity for α-SMA, although Frezzotti et al. did not specify the period following cataract surgery at which the membrane had been removed.

In conclusion, the present study showed that the flattened LECs seen 5 days after cataract surgery and that are related to a wrinkling of the posterior capsule were positive for α-SMA. Cuboidal LECs were negative for α-SMA. Thus, one can distinguish the flattened LECs from the normal LECs biochemically by the presence of α-SMA.

We thank Professor Y. Oguchi for his valuable suggestions on the manuscript.

This work was supported in part by a grant in aid (No 08771527) for scientific research from the Ministry of Education, Science, and Culture of Japan.

Presence of alpha smooth muscle actin in lens epithelial cells of aphakic rabbit eyes.

D Kurosaka, K Kato and T Nagamoto

doi: 10.1136/bjo.80.10.906