Interlenticular opacification in piggyback AcrySof intraocular lenses: explantation technique and laboratory investigations

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Abstract

Background/aims—Interlenticular opacification (ILO) is a recognised complication of piggyback intraocular lenses (IOLs). The aetiology, histopathology, and treatment are not clearly defined, however.

Methods—Two pairs of AcrySof IOLs were explanted from a patient with bilateral ILO. The explantation technique and surgical challenges of IOL exchanges are described. The explanted IOL complexes and a sample of the anterior capsule were examined by phase, polarising, and immunofluorescence microscopy.

Results—A 50 year old man developed ILO bilaterally after piggyback AcrySof IOL implantation. A central contact zone was surrounded by a homogeneous para-central opacity possibly consisting of extracellular material previously laid down by proliferating lens epithelial cells (LECs). These opacities were in turn surrounded by interlenticular Elschnig pearl-type opacities contiguous with the same material filling the periphery of the capsular bag. The IOL complexes were very adherent to the capsular bag and they had to be separated with the help of high viscosity viscoelastic before a single piece PMMA IOL implantation via large limbal incisions. The sample of anterior capsule showed a ridge configuration from the piling of LECs in the site of apposition with the anterior capsule and cells showing different characteristics on either side of the ridge.

Conclusion—Cellular proliferation, deposition of ECM from proliferating LECs, and capsular changes induced by cell metaplasia may lead to ILO formation in piggyback AcrySof IOLs. Careful separation of the AcrySof IOL complex from the capsule, meticulous clean up of the proliferating material, and implantation of single or dual IOLs in the bag PMMA IOLs through a large incision with capsulorhexis enlargement may help in the prevention of recurrence of interface opacification.

Clinical details

A 50 year old man presented in October 1999 complaining of gradual deterioration of vision bilaterally over the preceding year and considerable annoying glare in bright light and at night. Initially his vision improved with hypermetropic spectacles, which needed to become progressively stronger. However this was not sufficient to alleviate the glare.

Two years earlier he was found to have bilateral cataracts and marked hypermetropic astigmatism. He underwent elsewhere a left phacoemulsification with implantation of piggyback acrylic IOLs (+15.5 and +16.50 dioptres (D)) AcrySof MA30BA, Alcon), followed by a right phacoemulsification with piggyback acrylic
(+15.0 D and +16.50 D AcrySof MA30BA) IOLs 1 week later. Approximately 1 month later he underwent bilateral paired arcuate keratotomies in the 7 mm corneal zone in order to reduce >3 D of corneal astigmatism. He was very pleased with his vision and was able to see very well for distance without spectacles (refraction not available).

When he presented to us in October 1999, his visual acuity was 6/18 unaided (6/5 with +2.25 +0.25 × 160) in the right eye and 6/24 unaided (6/9–3 with +2.75 +1.00 × 129) in the left. Both corneas showed superior and inferior arcuate keratotomies. The anterior chambers were deep and quiet and the pupils were normal. The anterior capsules did not show any evidence of fibrosis and the capsulorrhexes were overlapping the optic of the anterior IOLs. Both IOLs were in the capsular bag with their haptics almost aligned in the horizontal meridian in the left eye (Fig 1a and b) and about 30 degrees apart in the right eye (Fig 1c). Elschnig pearl-type opacities were present in the peripheral interlenticular spaces in both eyes extending up to the mid-periphery. There was a central area of contact measuring approximately 1 mm in diameter, which was free from opacities. A whitish ring of homogeneous membrane-like opacification surrounded the central contact zone. This ring was in turn separated from peripheral pearl-like opacities by clear areas at places. To improve vision, it was decided with the patient that the AcrySof IOLs were to be explanted and be replaced with a single IOL to prevent any possibility of ILO recurrence.

**Surgical technique**

In November 1999 the patient underwent the first explantation. A 6.5 mm superior two step limbal incision was made and the anterior chamber of the left eye was filled with a cohesive viscoelastic material (Healonid GV, Pharmacia & Upjohn). The anterior capsule was very taut against the optic of the anterior IOL. Using a bimanual technique, the anterior capsulorrhexis edge was lifted over the anterior optic (where it joined onto one of its haptics) with a Sinskey hook introduced through a side port, and a 27 gauge Rycroft cannula attached...
In April 2000 the patient underwent a similar procedure on the right eye with explantation of the AcrySof IOLs and implantation of a one piece +30 D PMMA IOL in the capsular bag. At the last postoperative visit (June 2000), 10 weeks after the right IOL exchange and 8 months after the left, the BCVA was 6/5 on the right eye with −1.50 +2.25 × 125 and 6/9 with −0.50 +1.50 × 45 on the left (Fig 1d).

**Laboratory findings**

The IOLs and capsular fragment were observed by phase and polarising microscopy and photographed (Figs 2–5). The IOLs were separated, with difficulty, by inserting a fine blade between them. Some interlenticular material was left on each IOL, and cellular material was attached to the haptics. Indirect immunocytochemical staining was carried out for vimentin (an intermediate filament marker for LECs) and α smooth muscle actin (α-sm, a marker for epithelial-mesenchymal transition). Samples were also stained with phalloidin for F-actin and DAPI for cell nuclei.

When the fixed IOL assembly was examined by phase microscopy the regions of opacification appeared white due to light scatter (Figs 2 and 3a). The light scattering material was of two distinct types. A region with a regular cobblestone appearance surrounded the clear central area (Fig 3b). The regular hexagonal outlines (Fig 3b inset) gave the appearance of an epithelial sheet, but staining revealed no remaining cytoskeletal or nuclear elements. This material is likely to be the remains of extracellular matrix (ECM) secreted by a population of LECs that had migrated into the interlenticular space at an early stage of ILO development. The space between the outer edges of the IOL optics was filled with pearl-like and globular material (Fig 3c). This material stained for LECs in immunocytochemical studies with α-sm, a marker for epithelial-mesenchymal transition, and DAPI staining revealed the presence of large numbers of cell nuclei (Fig 3c) scattered throughout the globular material. Higher magnification (Fig 3f) showed that many of the nuclei were degenerate and some micronuclei were present.

The capsular fragment had a full covering of cells except for the rhexis edge, where cells may have been disrupted by the surgical explantation. The most striking feature (Fig 4a) was the presence of a prominent light scattering ridge running parallel to the curved capsulorrhexis edge. Polarising microscopy revealed that this ridge showed strong birefringence (Fig 4b) indicating alignment of cells and/or macromolecules within its structure. Similar birefringent structures with aligned cells have been observed at PMMA/capsule interfaces in donor capsular bags with single implanted IOLs.4–6

Viable cells were present within the ridge at the time of fixation, and cell nuclei were elongated and aligned along the length of the ridge (Fig 4d).

Cell morphology and α-sm expression were different on each side of the ridge. Cells on the IOL flap between the ridge and the rhexis edge

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**Figure 2** Photomontage of low magnification light micrographs to show overall structure of the fixed IOL pair before separation. Cellular and extracellular material between the optics appears white owing to light scatter. A central clear zone is surrounded by an area of homogeneous light scatter and this in turn is surrounded by a region of intense light scatter from Elschnig pearl-like material.

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The IOLs and capsular fragment were placed in fixative (4% formaldehyde in phosphate buffered saline). After 1 hour, the fixative strength was reduced to 1% for transport of specimens to the laboratory.
were variable in shape and size (not shown), had irregular nuclei, and had formed multilayers (Fig 4d). All the cells in this region were positive for $\alpha$-sma expression (Fig 4c). In contrast, on the capsule between the ridge and the equatorial region the epithelial-like cells formed a regular monolayer with round or ovoid nuclei (Fig 4e) and hexagonal mid-cell borders of F-actin (Fig 4f, boxed area). These features are characteristic of normal human LECs. Only a small number of scattered cells in this monolayer showed $\alpha$-sma expression (Fig 4c and f). Cells clinging to the haptics, which had been pulled out from the inner regions of the bags were positive for vimentin (not shown), indicating their true lens epithelial nature. A schematic diagram summarises the anatomy and cell biology (Fig 5).

Discussion

Interlenticular (or interpseudophakos) opacification can develop in some eyes with “in the bag” piggyback IOLs. The true incidence is not known, as it tends to occur more than 1 year after surgery. Gayton and Apple have reported an incidence of 43% in 30 eyes with acrylic IOLs and 22% in 31 eyes with PMMA lenses followed for at least 2 years. Roy observed ILO in six out of 27 eyes with AcrySof IOLs and Shugar and Schwartz in one of 14 eyes with plate haptic silicone IOLs. ILO is associated with hyperopic shift, the
mechanism of which is not clearly elucidated. The amount of the refractive shift is higher in acrylic than in PMMA IOLs and it has not been reported yet in silicone IOLs. Several reasons have been suggested to explain this phenomenon.

Removal of interlenticular opacities is a challenging process. Elschnig pearl-type material can be easily removed from the interface. However, membrane-type opacities, which can be stripped surgically from the PMMA IOLs without difficulty, are very adherent to the AcrySof IOLs necessitating lens exchange. Creation of a large corneal or limbal wound and removal of the IOLs en bloc avoids the more time consuming and traumatic intraocular separation or division of the lenses. Explantation of AcrySof IOLs can be difficult owing to the adherence to the anterior capsule. This tight apposition ties in with the ridge configuration

![Image of the anterior capsule specimen.](http://bjo.bmj.com/)

**Figure 4** Images of the anterior capsule specimen. (a) Low magnification phase image of the whole fragment before staining. Three regions with quite different characteristics were identified. Region 1 extends from the capsule rhexis (arrows) to a distinct thickened ridge (2) which forms a bright region along the length of the specimen. This ridge marked the edge of contact with the IOL. Region 3, from the ridge towards the equatorial (non-IOL) region of the capsular bag was evenly covered with cells. (A tissue fragment (starred), which appeared to have been attached to another tissue within the capsular bag, was present at one end of the specimen.) Frame size 1 × 0.75 mm. (b) Polarising microscopy pattern obtained from the capsular fragment when viewed through a polariser and analyser set at 90° and a first order red plate set at 45° to both axes. The blue birefringence colour results from the ordered alignment of cells (and probably molecules such as collagen) in the region of the prominent ridge (region 2 in (a)). In the red areas there is no significant degree of order in the system. Arrows mark the rhexis edge. Frame size 1 × 0.75 mm. (c) Whole fragment stained for α-smooth muscle actin (α-sm). The distinct ridge (between arrowheads; see also Fig 4c) marks the interface between capsule overlying the IOL optic (region 1 in (a)) and capsule facing the interior space between capsular leaflets (region 3 in (a)). Cells in region 1 are all α-sm positive, whereas only are small number in region 3 are positive. Frame 3.7 × 2.5 mm. (d) DAPI staining of nuclei of the ridge region (between arrowheads in (c)). The ridge passes diagonally through the centre of the micrograph (between arrows) and contains multilayers of elongated, aligned nuclei. A row of round epithelial nuclei and other out of focus nuclei are on the inner side (top left and region 3 in (a)), and multilayers of elongated/oversize/misshapen nuclei on the outer side (bottom right and region 1 in (a)). Frame size 145 × 100 μm. (e) DAPI staining of the monolayer of nuclei of epithelial cells in region 3. The oval regular nuclei appear identical to those found on a freshly excised normal donor lens epithelium. Frame size as (d). (f) Immunofluorescence, pseudocolour confocal image of cells from the epithelial layer in region 3. F-actin is encoded red and α-sm green. Note the regular–hexagonal F-actin outlines (boxed area) of the normal epithelial cells, and these are exactly as they would have appeared in a natrese, unoperated lens capsule. A few scattered cells (see also Fig 3f) show α-sm expression. Bar = 25 μm.
seen in our capsular specimen where cells had clearly accumulated parallel to the optic edge, formed multilayers, and deposited ECM. Cellular proliferation and deposition of ECM around the four haptics also create firm adhesions with the capsular equator, which requires careful handling in order to maintain the integrity of the zonular apparatus. Adhesion of the posterior IOL to the capsular bag may necessitate the use of viscoelastic injection behind the former. Separation of the two IOLs may be difficult as the AcrySof IOLs are fused together possibly by ECM produced by LECs proliferated in the interlenticular space. Insertion of a lens glide facilitates the safe removal of the lens complex protecting the capsular bag from accidental tearing by the trailing haptics.

Prophylaxis and prevention of recurrence of ILO is a problem already addressed by several surgeons and various suggestions have been put forward. For IOL power requirement of +30D or less, we favour a single implant in the bag. For higher power requirements, we favour dual in the bag PMMA implants with equal power split to reduce optical aberrations and dual in the bag PMMA IOLs in the capsular bag (unpublished data). In the bag implantation avoids the problems associated with IOL-ciliary muscle contact.

The aetiology of ILO is not clearly understood to date. A small capsuleorrhesis edge overlapping the anterior lens surface of the anterior in the bag IOL creates a closed system where LECs are allowed to proliferate in the interlenticular space between the lenses. The truncated edge of the AcrySof IOLs along with the adhesion of AcrySof in the posterior capsule do not allow migration of the lens epithelial cells on the posterior capsule, which may divert the migrating cells into the interlenticular space. This may explain the higher incidence of ILO in AcrySof piggyback IOLs. Inadequate cortical clean up may be another contributing factor. Examination of exfoliated membranes, based on haematoxylin and eosin studies, has shown two histological types and we were able largely to confirm these findings. Two distinct histological types of ILO were observed in these specimens—that is, the Elschnig pearl-type consisting of proliferating/retained LECs in the interlenticular space and the amorphous membranous-type consisting of ECM deposits with the same cobblestone appearance given by LECs. To the best of our knowledge no detailed immunological studies of cell types/viability have been reported before.

The pearl-type material in the present specimens was found to consist of swollen and globular live cells with an F-actin cytoskeleton and nucleus. They showed evidence of some form of transformation/differentiation in that some expressed α-sma, and some had degenerative nuclei. While these cells would be unlikely to undergo cell division the process of globular swelling could be ongoing, increasing the amount of material. The membranous region of opacification was not amorphous but showed a distinct hexagonal pattern with similar dimensions to LECs. Since no live cells were present this most likely represents ghost cells—that is, devitalised cells, within a sheet of extracellular matrix produced by the previously live lens epithelial cells.

The cellular material in the anterior capsule fragment reveals viable LECs despite the time interval from the operation, similar to those seen in eyes with single IOLs. The close apposition of the anterior capsule and anterior IOLs periphery leads to distortion of epithelial cell alignment, which in turn results in the build up of multiple layers of cells and ECM at the interface between the IOL edge and the capsule, forming the thickened ridge. Interestingly, the cells residing in the area between the ridge and the capsule are irregular and only a few cells express α-sma. These cells have the potential to proliferate and colonise any surfaces and spaces within the capsular bag leading to further accumulation of opacified material. Piling of the cells at the ridge could lead to a diversion of the cells into interlenticular space and further proliferation to form Elschnig pearl and globules. These structures are very similar to those seen in capsular opacification with single IOL implantation.

It is very likely that an initial small central contact zone is created at the time of implantation by the compression of the IOLs owing to limited space in the capsular bag. This compression would inhibit encroachment of LECs into this central region. However, LECs are able to proliferate and lay extracellular matrix in the interlenticular space up to the edge of the contact zone. Opacification of the anterior capsule overlapping the optic and fusion of the peripheral anterior and posterior capsule by proliferating cells result in added anterior-posterior contraction forces, further compression of the IOLs, and hence expansion of the contact zone. In addition to the
compression forces exerted on the IOLs by the capsular bag, ECM laid by the proliferating cells may act as a biological “glue” attracting the optic surfaces even closer causing more entrapment and death of the epithelial cells within the expanded contact area. Expansion of the central contact zone with the progressive flattening of the opposing surface of the two IOLs would lead to gradual reduction of the refractive power in the central area of the IOL complex and hyperopic shift. As the paracentral and peripheral vision are obliterated by the opacification in these areas (Fig 2) the patient is forced to use the central contact area for vision. Owing to flattening this area has less refractive power resulting in hyperopic shift. The opacified areas also cause glare.

Conclusions
The tight apposition of the anterior lens capsule to the anterior optical surface of the anterior piggyback IOL results in the formation of a closed system, bounded by the edge of the IOL/capsular ridge and encompassing within it the remainder of the capsule and the interlenticular spaces and surfaces. This closed environment between the anterior and posterior capsules is much larger than that formed by a single IOL and so there is a greater volume in which material can accumulate. Significantly, there are also more surfaces upon which material can be deposited. Cellular proliferation, deposition of ECM from proliferating LECs and capsular changes induced by cell metaplasia may lead to ILO formation and hyperopic shift in AcrySof IOLs. Careful separation of the AcrySof IOL complex from the capsule, meticulous clean up of the proliferating material, and implantation of a single or dual in the bag PMMA IOLs through a large incision with capsulorrhesis enlargement may help in the prevention of recurrence of interface opacification.

A free paper entitled “Interlenticular opacification in piggyback AcrySof intraocular lenses: a histopathological study” and a video entitled “Interlenticular opacification in piggyback AcrySof intraocular lenses: surgical technique for exchange of IOLs” containing parts of the submitted manuscript were presented at the UKISCRS 2000 annual meeting in Chester, UK.

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