Massive proliferation of lens epithelial remnants after Nd-YAG laser capsulotomy

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Abstract
Eight eyes exhibited massive proliferation of lens epithelial remnants following Nd-YAG posterior capsulotomy. All eyes had pre-existing retinal pathology. Six had undergone vitrectomy (four for proliferative diabetic retinopathy) before extracapsular cataract extraction with posterior chamber intraocular lens implantation. The other two eyes had familial exudative vitreoretinopathy or retinopathy of prematurity, respectively. Five eyes required removal of the lens proliferations via a pars plana approach. High levels of growth factors in the posterior segment associated with proliferative disorders of the retina may play a role in lens cell proliferation.

(Br J Ophthalmol 1995; 79: 261–263)

Retention and sequestration of lens epithelial elements (Soemmerring’s ring) and their subsequent metaplasia and proliferation causes opacification of the posterior lens capsule in up to 50% of patients after extracapsular cataract extraction (ECCE) with or without intraocular lens (IOL) implantation. In the majority of such patients, central clearance of the opacification using the neodymium-yttrium/aluminium/garnet (Nd-YAG) laser is straightforward and effective. It is occasionally complicated by damage to the IOL, raised intraocular pressure, cystoid macular oedema, retinal detachment, or spreading of endcapsular low grade endophthalmitis. The need for further laser or surgical capsulotomy is rare, though Holz et al described pearl formation following Nd-YAG capsulotomy after removal of a traumatic cataract, requiring further Nd-YAG clearance. We report a series of eight patients, with some features in common, all of whom developed massive proliferation of lens epithelial elements following Nd-YAG capsulotomy, and discuss the possible cause of this phenomenon.

Patients and methods
Eight patients who had earlier undergone ECCE+IOL followed by Nd-YAG capsulotomy, went on to develop massive proliferation of lens epithelial remnants. The associated clinical features, previous surgery, and the further surgical interventions reviewed were reviewed.

Results
All eight patients had associated posterior segment problems. Four had proliferative diabetic retinopathy (PDR) while the remainder had each presented with acute retinal necrosis, idiopathic vitreous haemorrhage with a fibrocellular epimacular membrane, familial exudative vitreoretinopathy, or the late sequelae of retinopathy of prematurity (ROP) (Table 1). Vitrectomy had been undertaken in six eyes before cataract surgery; in three of these, silicone oil had been injected and had subsequently been removed before ECCE+IOL (Table 1). The four vitrectomised diabetics with massive lens proliferation represent a small minority of the patients under our care who have undergone phakic diabetic vitrectomy followed by ECCE+IOL and later Nd-YAG capsulotomy. One of the two patients in whom vitrectomy had not been undertaken had nevertheless undergone (in both eyes) encirclement and cryotherapy for progressive retinal detachment, and the other had Nd-YAG capsulotomy followed in all eyes by massive capsule proliferation, maximal at the edges of the capsulotomy.

In all eight eyes ECCE+IOL had been uneventful. In four, a heparin surface modified IOL had been implanted. Nd-YAG capsulotomy was followed in all eight eyes by massive capsule proliferation, maximal at the edges of the capsulotomy. There was no associated rubecrosis in any of the eyes at this stage. In two eyes the capsulotomy was wide and the visual axis was unaffected, but six required further capsule management. In three, further Nd-YAG capsulotomy was attempted but in only one case (case 3) was this successful. The bulbous, gelatinous nature of the globular proliferations was consistent between patients as was the ability to absorb Nd-YAG energy without disruption. Five patients ultimately required surgical corticocapsulotomy via the pars plana (Table 1). We include three illustrative case reports:

Table 1 Summary of the features of eight patients with massive proliferation of lens remnants

<table>
<thead>
<tr>
<th>Case</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
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<td>M/56</td>
<td>M/18</td>
<td>F/59</td>
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<td>ARN</td>
<td>ERM</td>
<td>PEVR</td>
<td>PDR</td>
<td>ROP</td>
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<td>No</td>
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<tr>
<td>Heparin IOL</td>
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<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
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</tr>
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<td>Interval between Nd-YAG capsulotomy and observation of lens proliferation (months)</td>
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<td>3</td>
<td>6</td>
<td>12</td>
<td>18</td>
<td>9</td>
<td>8</td>
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<tr>
<td>Surgical corticocapsulotomy</td>
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<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
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</tbody>
</table>

ARN=acute retinal necrosis; ERM=epiretinal membrane; PEVR=familial exudative vitreoretinopathy; PDR=proliferative diabetic retinopathy; ROP=retinopathy of prematurity.
CASE 1
A 38-year-old insulin dependent diabetic of 23 years standing presented with bilateral severe PDR. In his left eye, surgical treatment was unsuccessful. In the right eye, an extramacular tractional retinal detachment from severe fibrovascular proliferation slowly progressed though the visual acuity was 6/6. The eye underwent vitrectomy, membrane delamination, endolaser, and sulphur hexafluoride gas (SF₆) injection. One month later a further procedure was necessary, including a further delamination, endolaser, encirclement and buckling, and silicone oil injection. Three months later the silicone oil was removed. The development of a cataract led to ECCE with implantation of a heparin surface modified posterior chamber IOL. A visual acuity of 6/9 was achieved. Deterioration of vision due to posterior capsular opacification was treated by Nd-YAG laser capsulotomy 5 months later. A visual acuity of 6/6 was only temporary owing to the development of massive lens/capsule 'pearls', encroaching on the visual axis. Further Nd-YAG laser treatment was again temporarily successful but a third treatment was ineffective owing to the apparently gelatinous nature of the material, with poor laser uptake. Pars plana corticocapsulectomy was therefore performed. The patient maintained a visual acuity of 6/6 2 years later.

CASE 2
A 55-year-old woman presented with unilateral ocular discomfort and deterioration of vision to hand movements owing to acute retinal necrosis with combined tractional and rhegmatogenous retinal detachment.

Vitrectomy was performed, including retinotomy, internal drainage, silicone oil injection, endolaser, and encirclement. Three months later the silicone oil and a macular epiretinal membrane were removed and further endolaser applied. The retina remained attached and a visual acuity of 6/18 was attained. Sixteen months later, following visual loss to counting fingers, ECCE and IOL implantation (heparin surface-modified) was performed uneventfully, and a visual acuity of 6/18 was regained. Ten months later Nd-YAG posterior capsulotomy was necessary. Proliferation of lens remnants occurred but the capsulotomy was sufficiently wide that the visual axis was uninvolved (Fig 1). A visual acuity of 6/18, owing to cystoid macular oedema (which has been present before the capsulotomy was performed), was maintained 1 year later.

CASE 3
A 36-year-old insulin dependent diabetic of 32 years' standing presented with bilateral severe PDR. The right eye, treated by vitrectomy, membrane segmentation, and endolaser photocoagulation, retained a visual acuity of 6/18. The left eye had ruberosis and a combined tractional and rhegmatogenous retinal detachment, and underwent vitrectomy, membrane delamination, internal drainage of subretinal fluid, encirclement, endolaser photocoagulation, and intraocular gas (SF₆) injection. An initially satisfactory result was followed by the development of cataract. Two years after vitrectomy, ECCE was performed with insertion of a heparin surface modified posterior chamber IOL. Visual acuity, initially recovering to 6/9, then deteriorated owing to posterior capsule opacification. Nd-YAG laser capsulotomy was followed by massive proliferation of lens epithelial remnants, encroaching upon the visual axis. This material was partially cleared by further Nd-YAG laser treatment, and although remnants persisted, there was a functionally clear visual axis and a visual acuity of 6/9 a year later.

Discussion
Massive proliferation of lens epithelial remnants after Nd-YAG capsulotomy has, to our knowledge, been described in only one patient,¹⁰ and has not been reported in patients with PDR, including those who had previously undergone vitrectomy.¹¹¹² In this series of patients, massive proliferation of lens remnants occurred within months of the capsulotomy, reducing or closing the capsular aperture and sometimes reducing vision. All patients had had posterior segment pathology including epiretinal proliferations whether fibrovascular or fibrocellular.

We postulate a role for growth factors in the initiation and perpetuation of the lens cell proliferation. It is well documented that vitreous aspirates from patients with PDR or proliferative vitreoretinopathy have significant mitogenic activity compared with control.
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1 Sormmerring D. Beobachtungen nderigen mechanischen Veränderungen des Auges nach Laserschnitt. Frankfurt a/M, Wesche, 1828.

vitreous13-15, elevated levels of potent mitogenic factors such as basic fibroblast growth factor,16 insulin-like growth factor I,17 and platelet derived growth factor18 have been identified. In vitro studies have shown that lens epithelial cell proliferation is stimulated by these factors,19 and indirect evidence for an in vivo effect is provided by the finding of increased lens size in insulin dependent diabetes.20 It is plausible that it is these or other growth factors which induced the massive proliferation of lens cells in our patients. That such growth factors are present, and are prevented from reaching the anterior segment by an intact posterior capsule, is said to account for the increased incidence of iris ruberosis in diabetics after intracapsular cataract surgery or posterior capsulotomy.11 21 Furthermore, the lens capsule, rather than simply being a barrier, may act as a 'sponge' for growth factors of retinal origin22 and this could lead to an increased concentration of these molecules in the region of the posterior capsule.

Disruption of the posterior capsule by the Nd-YAG laser could act in one or more of a number of different ways to promote lens proliferation. Firstly, disruption of the capsular bag may result in mitogen containing fluid from the posterior segment bathing the lens epithelial remnants. Secondly, the laser may directly provoke cellular proliferation, a phenomenon observed following Nd-YAG laser to trabecular meshwork cells.23 Thirdly, the Nd-YAG laser may 'activate' lens epithelial cells to produce proteolytic enzymes capable of releasing otherwise inactive growth factors bound to matrix molecules in the remaining capsule. However, we do not believe that Nd-YAG capsulotomy is an essential prerequisite for such lens proliferation. Another two insulin dependent diabetics under our care developed gross lens proliferation causing wide separation of the IOL from an intact posterior lens capsule following vitrectomy and ECCE+IOL.

Our patients had complex eye problems, often requiring more than one operation. The need for further surgical intervention in the form of corticocapsulotomy was therefore unfortunate, though there were no untoward sequelae. Maximisation of cell clearance from the posterior capsule during cataract surgery is advocated, though the technical difficulty of dealing with the capsule after previous vitrectomy is an important consideration. If required, a particularly large Nd-YAG capsulotomy should be performed in such patients so that, even should massive proliferation occur, there is less chance of involvement of the visual axis.