Lens Induced Secondary Glaucomas

Although recent advances in sutureless small incision cataract surgery and phacoemulsification and improved IOL designs have resulted in vastly superior outcomes with reduced complications related to wound repair and secondary glaucomas, several problems still remain. These require training of ophthalmologists in the emerging, new techniques and tackling complications peculiar to the new surgical methods, which may include glaucoma and inflammation secondary to retained lens fragments. These measures will also reduce complications such as pseudophakic glaucoma, for example, yet another significant cause of secondary glaucoma.

Phacomorphic glaucoma

The cataractous lens may become swollen (intumescent) which causes relative pupil block, the iris root is moved forward and this may result in blockage of outflow of aqueous fluid at the angle of the anterior chamber. This is a secondary form of angle-closure glaucoma.

Phacolectic glaucoma

Lens material may cause blockage of outflow of the aqueous at the drainage angle and this may occur after injury (including cataract surgery) or when lens material leaks through the lens capsule of a mature/hypermature lens. Macrophages, attempting to remove this abnormal mater-
secondary glaucoma. "Lens-induced" glaucoma, in the broadest sense of the word, can be prevented by excellent cataract surgery, by operating on unilateral dense cataracts, and on second eyes if it is considered likely that the patient will not return for follow-up.

Neovascular Secondary Glaucomas

Diabetic retinopathy and central retinal vein occlusion account for nearly two-thirds of patients with neovascular glaucoma. Secondary glaucomas associated with proliferative diabetic retinopathy and central retinal vein occlusion may occur in diabetics where abnormal new blood vessel formation has occurred causing disturbance to the outflow of aqueous at the angle of the anterior chamber. Ocular neovascularisation and glaucoma may develop in 33-64% of eyes with untreated, proliferative diabetic retinopathy. Diabetic retinopathy is a leading cause of blindness in persons aged 20-74 years. With improved treatment available for diabetes, life expectancy has been greatly increased, resulting in many more individuals with diabetic retinopathy. Effective metabolic control of diabetes in the population, efficient screening for early detection and treatment of diabetic retinopathy and retinal photoacoagulation of eyes with proliferative diabetic retinopathy are vital in prevention and management of neovascular glaucoma. Glaucomas which remain uncontrolled with lasers and conservative therapy may be treated with cyclophotocoagulation with the newer Nd: YAG or semiconductor diode lasers with the option of glaucoma filtering surgery and adjunctive antifibrosis agents. Neovascular glaucoma (rubeotic glaucoma) results from angle-closure secondary to a fibrovascular membrane in the anterior chamber due to ocular diseases characterised by retinal ischaemia and angiogenesis. Thrombosis of the central retinal vein will result in disturbance of the circulation within the eye and this may result in new vessel formation within the anterior segment. These abnormal blood vessels may affect the angle of the anterior chamber, where the blood vessels can be visualised, and secondary glaucoma can result.

Ischaemic central retinal vein occlusion (thrombotic glaucoma) is the second most common cause of ocular neovascularisation and glaucoma is seen in 58-86% of these eyes. Essential hypertension and primary open angle glaucoma remain the principle aetiological factors in the pathogenesis of central retinal vein occlusion.

Management of the neovascular glaucomas

Ideally, eyes with conditions where neovascular glaucoma is likely should be identified early, and prophylactic panretinal laser photoacoagulation given. Fundus fluorescein angiography (FFA) can be used to detect retinal capillary closure in eyes with central vein occlusion, and this should be done 6-8 weeks after the onset, once the retinal haemorrhages have cleared. All diabetics with retinal neovascularisation should have panretinal photoacoagulation. If facilities are not available for FFA, clinical signs of ischaemia should be used to detect eyes at high risk (i.e., extensive cotton wool spots).

Treatment by panretinal photoacoagulation of established ‘rubeotic’ glaucoma is often not successful, as permanent changes occur in the drainage angle.

Future research is directed at evolution of the inhibitors of angiogenesis to combat neovascular glaucoma. Antibodies to vascular endothelial growth factor (VEGF), the principle peptide involved in angiogenesis, have been successful in reversing new vessels in primate eyes. Suppressor gene therapy to prevent predisposition of individuals to neovascular glaucoma is also a distinct possibility in the future.

Eye Injuries and Secondary Glaucomas

Ocular injuries have been recognised as a common cause of monocular blindness in several studies and secondary glaucoma is one of the principle causes of visual impairment. Although there is no large population based series study on the prevalence of the traumatic glaucomas, the Aravind Comprehensive Eye Survey has found a prevalence of 0.2% of glaucoma in individuals with trauma (unpublished data). Most ocular trauma and ocular morbidity has been reported in males in the younger age group, accounting for severe economic burdens in terms of days lost in work and expenditure on treatment. Early recognition of trauma and elucidation of the mechanism of glaucoma is vital to prevent visual loss.

Haemorrhage into the anterior chamber (hyphaema) and angle recession

Degenerate red blood cells may block the trabecular meshwork at the angle of the anterior chamber and there may be a secondary rise of intraocular pressure. A total oralmostotalphachamomaybeassociated with a rise in pressure and, also, blood elements may penetrate the cornea resulting in corneal blood staining – which is very slow to clear. This type of hyphaema should be surgically released by paracentesis.

Further, if the haemorrhage has been the result of a severe blunt injury, for example, with damage to the trabecular meshwork and the angle of the anterior chamber, later healing with fibrosis may cause a severe

---

Table 2: Common Causes of Secondary Glaucomas seen in a Tertiary Eye Care Hospital in South India.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Number of Individuals with Secondary Glaucoma</th>
<th>Percentage of Total Glaucoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lens induced glaucomas</td>
<td>158</td>
<td>2.50</td>
</tr>
<tr>
<td>Neovascular glaucomas</td>
<td>58</td>
<td>0.95</td>
</tr>
<tr>
<td>Pseudophakic glaucomas</td>
<td>38</td>
<td>0.62</td>
</tr>
<tr>
<td>Retinal glaucomas</td>
<td>25</td>
<td>0.48</td>
</tr>
<tr>
<td>Traumatic glaucomas</td>
<td>16</td>
<td>0.26</td>
</tr>
<tr>
<td>Steroid-induced glaucomas</td>
<td>12</td>
<td>0.20</td>
</tr>
<tr>
<td>Secondary glaucomas of unspecified cause</td>
<td>60</td>
<td>1.0</td>
</tr>
</tbody>
</table>

*Retrieved from the statistics of the Glaucoma Services of the Aravind Eye Hospital, Madurai, India in the year 2000.
Secondary Glaucomas

Type of secondary raised intraocular pressure (post-traumatic angle recession).

Drug-induced Secondary Glaucoma

Corticosteroid-induced glaucoma

Longer term use of topical and systemic corticosteroids can result in a rise of intraocular pressure, which is usually reversible once the medication is discontinued. Glaucoma due to the indiscriminate use of topical corticosteroids for allergy and spring catarrh has left children blinded from glucocorticoid optic atrophy. Ophthalmologists have a pivotal role to play in preventing such needless blindness by educating health workers and the general public, together with control of the availability of corticosteroids.

Uveitis and Secondary Glaucomas

In uveitis, cells and proteins in the anterior chamber disturb the normal outflow of aqueous fluid through the trabecular meshwork, causing raised intraocular pressure. Using a focal light and magnification a ‘flare’ may be seen in the anterior chamber—like a shaft of sunlight streaming into a room full of dust. As a result of the inflammatory reaction within the eye there may be adhesions between the pupil margin and the anterior lens surface (posterior synchiae) and/or in the angle of the anterior chamber (peripheral anterior synchiae). The iris bows forward as adhesions form and this further embarrasses the drainage angle of the anterior chamber—described as ‘iris bombé’.

Glaucoma secondary to uveitis is an important clinical entity, often with severe visual impairment. The management is complex since complicated cataract, macular oedema and media haze largely contribute to ocular morbidity apart from glaucomatous optic nerve damage. It has been reported that between 5.2% and 15% of eyes with uveitis develop secondary glaucoma. Though most uveitic entities are idiopathic, known causes include infections like leptospirosis, toxoplasmosis, AIDS, onchocerciasis and drug-resistant tuberculosis. Adequately combating these microbial infections can significantly reduce ocular morbidity due to uveitis—apart from addressing the cause of elevated ocular pressures and the institution of appropriate therapy.

Pigment Dispersion Syndrome/ Pigmentary Glaucoma

In certain eyes, pigment particles may circulate abnormally in the aqueous fluid, and these in turn may cause blockage at the drainage angle. There is some debate as to whether this form of glaucoma should be described as primary or secondary glaucoma.

Exfoliation Syndrome/ Pseudoexfoliative Glaucoma

Abnormal accumulation of particles (not unlike dandruff in appearance) may accumulate in the anterior eye. This abnormal material can cause blockage of the drainage angle. Pseudoexfoliative glaucoma is particularly found in Sudan, Somalia, Ethiopia and Tanzania. It is less common in West Africa. Some consider this to be a form of primary glaucoma.

Epidemic Dropy

This acute toxic disease is caused by the ingestion of Argemone mexicana oil, an adulterant of cooking oils. It has been reported in India, Mauritius, Fiji, Bangladesh and southern Africa. Rash, oedema of the lower limbs, gastrointestinal and cardiovascular disturbances may be accompanied by a secondary form of glaucoma and retinal vascular abnormalities.

Prevention of Secondary Glaucomas

- Accessible, affordable cataract services of high quality to prevent lens induced glaucoma
- Good management of hypertension to reduce retinal vein occlusions
- Good control of diabetes to prevent non-neovascular glaucoma
- Early detection and good management of conditions associated with the potential for retinal ischaemia and neovascularisation
- Increased awareness among eye care professionals, the public and pharmacists of the dangers of topical (and systemic) steroids
- Health education about avoiding eye injuries

Treatment

The management of eyes with secondary glaucoma depends on whether there is the potential for useful vision. If there is, then treatment should be aimed at lowering the IOP, reducing any associated inflammation, and treatment of the underlying cause(s) (i.e., removal of a hyphaema; removing a hypermetropic lens), plus other interventions to restore sight. Long term treatment to control the IOP may be required, or glaucoma surgery once the eye has become quiet and stable.

If there is not the potential for useful vision (i.e., secondary glaucoma due to CRVO or end stage diabetic retinopathy), the aim of management is to give symptomatic pain relief (e.g., mydriatics and steroids, injection of retrobulbar alcohol).

Comment

Though secondary glaucomas numerically represent a smaller percentage than the primary forms of the disease, they nevertheless cause significant ocular morbidity and visual impairment. Early identification of the primary ocular and systemic diseases that predispose to the secondary glaucomas would play a significant role in limiting the burden of needless blindness.

References


TEACHING EYE HEALTH
Detlef Prozesky
MBChB MCommH PhD
The seventh article in the Teaching Eye Health series on Evaluation of Courses
will be in the next issue of the Journal
Volume 14, Issue No. 40
Editor

Community Eye Health Vol 14 No. 39 2001