Treascher Collins syndrome with novel ophthalmic findings and visceral anomalies

Treascher Collins syndrome (TCS) (mandibulofacial dysostosis (MFD) or zygomatico-mandibular dysplasia) is one of a group of congenital malformation syndromes that have in common maldevelopment of the first and second branchial arches. Clinical features typically include hypoplasia of the mandible and zygoma; a complex variety of ear abnormalities including malformed pinnae, atresia of the external auditory canals and anomalies of the middle ear ossicles; cleft palate; reeding chin; and sinus and choanal atresia. We present a pair of twins that possessed not only these typical visceral and previously unreported ophthalmic anomalies.

Case reports

A 32 year old, gravida 3 para 2-0-0-2 woman underwent elective abortion after fetal death at 20 weeks of gestation. Both twins had bilaterally hypoplastic zygomas, maxillae, and related muscles; severely hypoplastic and misplaced pinnae; bilaterally agenic external auditory canals; bilateral ocular clefts, and severe micrognathia (Fig 1). Twin A exhibited left sided choanal atresia, agenesis of the hard and soft palate, and multiple visceral anomalies, including dual superior venae cava, bilobed right and unilobed left lungs, bilateral renal and ureteral agenesis, rudimentary urinary bladder, and absent epididymes. Twin B was noted to have right sided choanal atresia, soft palate aplasia, hard palate hypoplasia, and a left sided cleft; twin B had no visceral anomalies. Ocular pathologi- cal dissection of twin A illustrated bilateral microphthalmia; corneal sceralisation; and maldevelopment of the uvea, lens, and retina. Ocular dissection of twin B revealed microphthalmia, aniridia, congenital cataracts, and bilateral vascularisation of the corneas.

Comment

These twins had multiple features characteristic of TCS, most notably hypoplastic zygo- mas, maxillae, and related muscles—perhaps the most characteristic features of TCS. In contrast, visceral anomalies, such as those of twin A, are rare. Only two cases have been previously reported: one with tracheoesopha- gial fistula, rectovaginal fistula, and anal atresia; another with achalasia. Renal agen- esis, found in twin A, has not been previously described in association with TCS. While ophthal- molological features in TCS are often exten- sive, they seldom involve the intraocular structures. Common findings include a defective inferior lateral angle of the orbit, caudal displacement of the superolateral orbit, true and pseudocolumbomas of the lids, lateral canthal dystopia, orbital lipodermoids, corneoscleral dermoids, and microphthalmos. Cata- racts, lenticul ar duct atresia, pupillary ectopia, distichiasis, and uveal colobomas have been reported less commonly; intravascular involvement in these twins is rare in TCS. Furthermore, aniridia, corneal sceralisation, and uvea, lens, and retinal maldevelopment are previously unreported.

TCS is an autosomal dominant disorder affecting one in 50 000 live births. The disorder appears to have arisen in these twins with no relevant family history, as occurs in 60% of cases. Expressivity is highly variable, ranging from clinically undetectable to perinatal death secondary to airway compromise; the disease severity that resulted in the fetal death of twin A is highly unusual. The responsible gene, TCOF1, has been mapped to 5q32–33.2 and the structure of its protein product, treacle, elucidated. To date, 51 disease causing mutations have been identified, nearly all resulting in a premature truncation of the protein and phenotype severity. The ophthalmic pathology observed in these twins may have resulted from a focal TCOF1 mutation and a yet to be defined role of treacle in (and renal) development. Considering the number and severity of abnormalities, the failure to identify a single "genetic hot spot," and the novel ophthalmic features, an alternate and perhaps more likely explanation is that a second gene, itself involved in ophthalmic embryology, was affected along with TCOF1. This unidentified gene may have been disrupted from a translocation involving 5q32–33.2 or from a deletion large enough to result in a contiguous gene syndrome. A final consideration is that these twins may not have had TCS but rather a new though closely related syndrome. Further genetic investiga- tion may shed light upon these speculations.

Acknowledgements

JLP and GB contributed equally to this work.

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Accepted for publication 7 September 2001

References


Acute postural drop in optic nerve perfusion after vitrectomy and gas in a patient with diabetic autonomic neuropathy

We report the case of an insulin dependent diabetic who suffered transient blindness as a result of a change of posture following vitrectomy surgery with injection of gas.

Postural hypotension in diabetics is secondary to autonomic neuropathy. The pathophysiology is not clear but it seems that the main factor is blunted catecholamine response to standing and failure of the lower limb vascular resistance to increase adequately. Consequently, the normal increase in POBF on standing is inadequate.

We advise particular caution in diabetics undergoing ocular surgery in which there may be a significant postoperative pressure rise. Similar problems can occur during surgery if hypotensive anaesthesia is either deliberately or inadvertently employed in diabetics. Beware of assuming poor postoperative vision to be purely the result of the presence of any previous reports describing optical coherence tomography (OCT) imaging of severe commotio retinae with an associated full thickness macular hole (FTMH).

Case report

A 15 year old boy presented 24 hours after blunt trauma to a football striking his right eye. On examination his best corrected visual acuity was counting fingers right eye and 6/6 left. Biomicroscopic examination revealed extensive commotio retinae over the posterior pole, no posterior vitreous detachment (PVD), and a FTMH. Colour photography and OCT imaging (OCT 2000 scanner, Zeiss-Humphrey) were performed (Fig 1). OCT confirms a FTMH and demonstrates extensive disruption of photoreceptor outer segments and retinal pigment epithelium (RPE).

Figure 1 (A) Right macula of 15 year old boy with extensive commotio retinae over posterior pole and an associated macular hole at 1 day after blunt injury. (B) Horizontal OCT scan through centre of macula confirms a full thickness macular hole and demonstrates extensive disruption of photoreceptor outer segment/retinal pigment epithelium layer. The optic disc is seen at the nasal edge of the scan.
He was treated conservatively with a short course of topical steroids. The colour fundus and OCT appearance at 1 month are shown in Figure 2. Despite spontaneous macular hole closure, visual acuity remained at counting fingers at 1 year follow up.

Comment
The major site of retinal trauma appeared on OCT to be at the level of the photoreceptor outer segment/RPE interface. The OCT images are consistent with fragmentation of photoreceptor outer segments and damaged cell bodies, as suggested by Sipperley et al in their study of the histological changes in commotio retinæ in primates.

The exact pathogenesis of macular holes remains uncertain. Ho et al outlined the three basic historical theories regarding aetiology—the traumatic theory, the cystic degeneration and vascular theory, and the vitreous theory. Of these, the latter has gathered the most support in the context of idiopathic macular holes.

In our case, the OCT imaging reveals that the edges of the macular hole are elliptical and irregular with no associated PVD, cortical vitreous condensation, or overlying prefoveal opacity. The characteristics suggest a different mechanism of hole formation from that proposed in idiopathic senile macular holes. We believe that mechanical distortion of the retina, relative to the vitreous and underlying sclera, created disruption of the photoreceptor outer segments and creation of a FTMH in this case. It is at the fovea and photoreceptor outer segment level that the retina has the least support from Müller cells and is therefore likely to undergo greatest deformation.

In the only previous report of OCT imaging in traumatic macular hole, a case with mild commotio retinæ was described in which extensive outer retinal disruption was not observed. There have been some encouraging reports suggesting that vitrectomy can successfully close traumatic macular holes as well as improve visual function in many cases. However, it seems unlikely that cases with severe commotio retinæ and associated photoreceptor/RPE damage, as demonstrated in our cases, would gain any benefit from surgical as opposed to spontaneous closure of a traumatic FTMH. The final visual prognosis is severely limited by the extent of initial photoreceptor damage, and the excessive pigmentary atrophy and clumping that follows.

We believe OCT imaging provides additional information both on the pathogenesis of commotio retinæ and in the assessment of outer retina disruption following ocular trauma. This information may help in the selection of patients likely to benefit from surgical intervention.

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Accepted for publication 26 September 2001

References

Acute ocular ischaemia and orbital inflammation associated with systemic lupus erythematosus
We report a patient with systemic lupus erythematosus (SLE) who developed bilateral ocular ischaemic syndrome in association with orbital inflammation leading to devastating visual loss.

Case report
A 73 year old white woman presented with unilateral acute anterior uveitis and polarynethropathy of the hands, knees, and neck. Over the next 4 years she suffered recurrent anterior uveitis, marginal keratitis, and episcleritis, which were treated with topical steroids. She had bilateral age related cataract and underwent left cataract surgery. Postoperatively, her visual acuities were 6/60 right eye and 6/9 left eye.

She was admitted 7 months later with extensive ischaemic macular oedema, hyphaema, and subconjunctival haemorrhage. Her visual acuities were 6/30 right eye and 6/60 left eye.

She was discharged on December 19, 2001 but on January 17, 2002 she developed recurrent left uveitis, hyphaema and 3 mm retinal detachment.

Figure 2
(A) Right macula 1 month after injury with extensive retinal pigment epithelium disruption, marked epiretinal membrane, and spontaneous closure of the macular hole. (B) Horizontal OCT scan through centre of right macular confirms resolution of commotio retinæ, spontaneous closure of macular hole, and disruption of the photoreceptor/retinal pigment epithelium reflex.
The family did not consent to histopathological examination of the eyes.

Comment
Acute orbital inflammation and orbital myositis leading to proptosis are rare manifestations of SLE.1,2 To our knowledge acute ocular ischaemic syndrome with orbital inflammation is a novel complication of SLE. The ophthalmic manifestations may have resulted from a vasculitis involving the vessels supplying the globe and extraocular muscles. The limitation of ocular movements was most probably the result of mechanical restriction. Although the proptosis, lid and conjunctival oedema improved with immunosuppression, the visual acuity did not recover. This may have been because of irreversible visual loss from optic nerve dysfunction due to ischaemia, compression from the acute orbital inflammation, or secondary to raised intraocular pressure. Close cooperation between ophthalmologist and rheumatologist is essential in the management of these patients, in order to try and prevent blindness.

References

Chickenpox neuroretinitis in a 9 year old child

Chickenpox in children is usually thought of as a benign infectious disease with few ocular complications. Posterior segment involvement from primary varicella zoster infection has rarely been reported in children. We describe the clinical features and visual outcome of an unusual case of neuroretinitis presenting in a 9 year old child.

Case report
An immunocompetent 9 year old boy acquired primary varicella zoster virus (VZV) infection from his sibling and developed the characteristic exanthematous vesicular rash. Four days after the onset of the rash he woke with discomfort in his right eye and described his vision as being “all grey” on that side. He presented to the emergency department the same day and was found to have a visual acuity of 3/6 on the right and 3/3 on the left (LogMAR). A relative afferent pupillary defect (RAPD) was present on the right. His anterior segment was quiet with no vitritis; however, he had slight macular thickening and a subtle cherry red spot on funduscopescope, along with some mild peripapillary swelling and disc haemorrhage. On review in the ophthalmology clinic 2 days later his vision had reduced to 1/60 (Sheridan Gardiner singles) on the right. He had no new skin lesions and all those present had crusted. No lid lesions were present. He had a marked RAPD, red desaturation, and mild conjunctival injection. His anterior segment and vitreous remained clear. The right disc was hyperaemic with peripapillary swelling and haemorrhage. The macular area was pale and oedematous (Fig 1). Examination of the left eye was completely normal.

Considering the onset of ocular symptoms and signs following the appearance of the typical VZV skin lesions, a presumptive diagnosis of chickenpox neuroretinitis was made. He was admitted and commenced on intravenous aciclovir (250 mg x 3 per day). Confirmation IgM titre for VZV were unfortunately not performed. No change in his acuity was observed over the next few days; however, his right disc was noted to become slightly pale 2 days after treatment of the skin. At this point intravenous methyl prednisolone was instituted at a dose of 5 mg/kg per day. During a gradual resolution of the macular and peripapillary oedema over the next 5 days, his disc remained pale (Fig 2) and his acuity measured as 3/30 (LogMAR) after 7 days of intravenous aciclovir and 5 days of methyl prednisolone. Systemically he remained completely well and afebrile on treatment. He was discharged with a further 3 day course of oral aciclovir and a 6 day reducing course of oral prednisolone.

Over 5 months of follow up his acuity has not improved beyond 3/30 (LogMAR). The right optic disc is pale and a yellow lipid deposit is present at the macula with some reticular macular pigmentation. The left eye had been normal throughout.

Comment
Posterior segment involvement as part of primary VZV infection in children has only been reported twice in our two case reports. Copenhaver1 reported a 3 year old with bilateral papillitis and a unilateral macular lesion associated with encephalitis following Varicella Zoster virus (VZV) infection. His patient presented with an acute exotropia 24–48 hours before the onset of cutaneous VZV. Funduscopy revealed papillitis, phlebitis, and a macular lesion which had resulted in a macular haemorrhage. His patient recovered vision and resolution of the macular lesion within 3 weeks of presentation. Capone and Meredith1 describe a case of unilateral central visual loss in a 2 year old child with chickenpox who presented with optic neuritis resulting in a poor visual outcome. Their patient presented with an acute exotropia and bilateral papillitis following VZV. Funduscopy revealed optic disc swelling, a macular haemorrhage, and retinal opacification just outside the arcades and scattered intraretinal haemorrhages and were also described. In these two cases sequential acuity measurement or photography were not possible because of the young age of the subjects.

Our case is particularly interesting, not only because these are the first published fundal photographs of VZV neuroretinitis in a child, but also because of the relatively mild ocular findings which have resulted in no visual loss. The young age of the patient is atypical of ocular VZV infection.1 Adults who contract primary VZV infection tend to run a more severe course than children.2 Ocular complications in children are extremely rare.3

The typical posterior segment involvement of VZV is acute retinal necrosis (ARN).1 The youngest case of ARN in association with chickenpox has been reported in a 4 year old.4 In adults, ARN is described as being less severe when presenting at the time of primary zoster infection than as a result of secondary reactivation of latent, previously acquired VZV.5 The changes typical of ARN were absent in this case. Unilateral papillitis and retinitis confined to the macular area were the main features. Optic neuritis has been reported by several authors in association with primary VZV.6 Many of these cases are bilateral and coincident with encephalitis1 or occurring in those who are immunocompromised.7 Unilateral optic neuritis has been described in an 18 year old several weeks following a varicella rash which remitted without sequelae following the administration of corticosteroid.8

The mainstay of treatment of VZV retinitis is with intravenous aciclovir. Whether any advantage is gained in administering systemic steroid with the aciclovir is controversial.9 We do not know if a more positive visual outcome may have been achieved if intravenous therapy had been commenced on presentation. It is therefore suggested that prompt treatment of VZV retinitis with intravenous aciclovir be started in patients, particularly in a child, presenting with any posterior segment signs.

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A 35 year old white man, who fulfilled the international Study Group criteria for Behçet's disease, attended the Behçet's clinic at the Birmingham and Midland Eye Centre. He complained of four red areas on his left arm (Fig 1). Two days earlier he had undergone acupuncture for what he described as “tennis elbow.” These red areas corresponded to where the acupuncture needles had been inserted. Examination revealed these areas to be pustules (inset) that were characteristic of a positive pathergy test.

Figure 1 Arrows indicating pustules corresponding to a positive pathergy test. Inset: magnified image of one pustule.
A positive pathergy test is an important diagnostic sign of Behçet’s disease. 7 Its prevalence varies by geographic region, being less common in patients from Northern Europe. Nevertheless, patients with Behçet’s disease should be made aware of this potential complication if they intend to undergo acupuncture.

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Accepted for publication 22 October 2001

Reference


Corneoscleral fibrous histiocytoma

Fibrous histiocytomas of the corneoscleral limbus are rare tumours. We present a case report and a review of the clinical and histopathological findings from cases presented in the literature.

Case report

A 19 year old white male presented with a 6 month history of a painless growth on the inferior corneoscleral limbus of the left eye (Fig 1A and B). There were no other ocular symptoms. He had no medical history of note.

His vision was 6/5 unaided in the right eye, and 6/6 unaided in the left eye. The growth was approximately 5 mm in diameter, vascularised, and yellowish in appearance. There were no other ocular abnormalities. The lesion was excised by conjunctival excision and sent for histopathological opinion. Histopathological appearance of a benign fibrous histiocytoma includes a mixture of fibroblastic and histiocytic cells that are often arranged in a cartwheel or storiform pattern, and accompanied by varying numbers of inflammatory cells, including foam cells and siderophages. No atypical nuclei or mitotic figures are present. Although some authors regard these tumours as reactive proliferations of fibroblasts, others do not accept this view because the lesions tend not to regress spontaneously. Recurrence is rare, with less than 5% of cutaneous benign fibrous histiocytomas recurring after local excision.

In contrast, malignant fibrous histiocytomas of the corneoscleral limbus characteristically appear in later life, between the ages of 50–70 years, with an equal distribution of males to females. They are highly aggressive tumours, and have been reported to have a local recurrence rate of 100% if a limited excision is performed. Recurrence can occur within a few months of excision. There are seven reported cases of corneoscleral malignant fibrous histiocytoma. Two of the cases had an enucleation and two cases underwent orbital exenteration.
Follow up of these patients ranged from 18 months to 3 years and all were free from recurrence. The other three patients had a local excision. One patient developed local scleral recurrence 2 months later. (It is not evident from the literature if the borders of excision were clear of tumour.) He was found to have a metastasis in the “parotid gland” 4 months later, and even tumour.) He was found to have a metastasis in literature if the borders of excision were clear of recurrence. Malignant fibrous histiocytomas have a broad range of histological appearances; storiform-pleomorphic, myxoid, giant cell, and inflammatory. The storiform-pleomorphic type is the most common. The cells are predominantly plump pleomorphic spindle-shaped with occasional large, ovoid histiocyte-like cells. Modest amounts of inflammatory cells, such as lymphocytes and plasma cells may be present. The differential diagnosis of a malignant fibrous histiocytoma includes pleomorphic carcinoma, malignant melanoma, and other sarcomas. For those limbal fibrous histiocytomas with a benign histopathological appearance, the management should be local surgical excision. Malignant fibrous histiocytomas need to be managed cautiously, preferably by wide local excision and cryotherapy at the earliest opportunity. If necessary, enucleation should be considered to fully excise a limbal malignant fibrous histiocytoma.

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Accepted for publication 22 October 2001

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diseases such as herpetic keratitis and atopic keratoconjunctivitis. To ensure appropriate treatment the organism must first be identified by cultures in enriched media.

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Accepted for publication 31 October 2001

References


Blinking and operating: cognition versus vision

The difference in the refractive indices between the air and the tear film results in the tear film having the greatest optical power of any part of the eye. Eyelid blinks are important for maintenance of the tear film. Between blinks there is progressive thinning of the tear film, which becomes non-uniform on the ocular surface and may break up. This produces an irregular air/tear interface, with a reduction in image quality. The longer the period between blinks the greater the effect on the tear film and reduction in vision. Restoration of the tear film occurs immediately following a blink so that the ability to maintain a regular tear film is dependent on the blink rate. A reduction in the blink rate such as, for example, a pause between blinks of 15 seconds has been associated with a change in the shape of the profile of the corneal tear film and up to a 6% reduction in visual acuity. More importantly, however, a reduction in blink rate leads to a reduction in contrast visual acuity. The ability to distinguish between layers during surgery, such as the posterior capsule of the lens and anterior vitreous face, or peeling a layer of the retinal surface in vitreoretinal surgery, is dependent on the surgeon having and maintaining good contrast acuity. Blink rates and blink amplitude vary according to vision task, stated behaviour and a reduction in the blink rate occurs with tasks of increasing visual difficulty. For example, visual tasks requiring concentration, such as video display terminal use, result in a decrease in average blink rate from 18.4/min to 3.6 blinks/min.

In order to determine whether the blink rate of ophthalmic surgeons alters during intraocular surgery, the blink rate patterns of nine ophthalmic surgeons were recorded. Two observers recorded the blink rate during casual conversation and when the surgeons were using the operating microscope. None of the surgeons were aware that their blink rates were being recorded, which was done by two medical students during their ophthalmic attachment—that is, their presence in the theatre was accepted as part of their ophthalmic training. The blink rate for each surgeon was recorded in each condition for four and 10 times. The mean blink rate for each surgeon during casual conversation and while operating are presented in Table 1. There was a significant reduction in the average blink rate between both conditions (16.69/min and 4.75/min, p<0.0001, paired t test), on average a third and a half fold decrease occurred while operating. It was also noted that the onset of conversation such as the request for an instrument or demonstration of an intraocular surgical layer was associated with the onset of a blink response.

The reduction in blink rates observed in this study—that is, 16.69 to 4.75, are similar to that found by Patel et al for visual tasks such as video display terminal use. A reduction in blink rate to 4.75 translates to a reduction in contrast acuity of approximately 10% with the onset of an intraocular surgical layer. Such a reduction in contrast acuity is likely to have an effect on the ability to differentiate between different intraocular surgical layers. Blinking occurs between visual fixations and may be timed so as not to interfere with significant visual input. Blink rate is low when information memory is operating, and cognitive processes utilising display areas accessible to visual input are disrupted during the blackout period of a blink. Blinking is thus suspended during certain cognitive activities to avoid disrupting these processes.

Frequent instillation of tears and contact lenses may reduce the blink rate. A reduction in the blink rate to the scleral flap during the procedure. Postoperatively, subconjunctival injections of steroids such as corticosteroids or intracameral injections of mitomycin C or 5-flurouracil may be used to treat the surgical site. The potential problem so that they can train themselves to blink during parts of the operation or investigation where good contrast acuity is not essential and where disruption of the cognitive processes is likely to have a minimal effect.

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Accepted for publication 21 November 2001

References


Should we vaccinate for glaucoma surgery?

Trabeculectomy is the most common non-laser surgical procedure performed for treatment of all forms of glaucoma. It involves the fashioning of a fistula from the anterior chamber of the eye to the subconjunctival space. This allows for extra drainage of aqueous humour to the subconjunctival space. This produces a localised elevation of the conjunctiva in the area of the trabeculectomy called a ‘filtering bleb’. Antimetabolites may be used intraoperatively and perioperatively to increase the success of glaucoma filtering surgery by their action on wound healing. 5-Fluorouracil or mitomycin C is administered to the scleral flap during the procedure. Post-operatively, subconjunctival injections of
S. pneumoniae may both be found on healthy normal eyes.

Streptococcus endophthalmitis may only be present tran-
conjunctival swabs were performed for cul-
incidence of blebitis. In most reported cases
blebs seems to be increasing.
eculectomy and trabeculectomy without
thalmitis is higher when antimetabolites are
use.

dose if the strains of S pneumoniae prevalent in the community are reflected in the poly-
saccharides contained in the vaccine. Its cost to the NHS is £420 per 0.5 ml. There have been no reports of epidemics of infective blebitis. If it was contagious, there would have been epidemics or clustering in our glaucoma clinics. We can find no evidence of case to case transmission. In fact, all reported cases and series appear sporadic. The second most common is H influen-
zae. The second most common is Haemo-
philia type b at over 23%.

S pneumoniae. The second most common is Haemoph-
libits, conjunctivitis, upper respira-
tive tract infection, hibernal occurrence, trauma, and vitreous wicks. With the increased use of antimetabolites in glaucoma surgery, the incidence of thin walled cystic blebs seems to be increasing. These blebs are more prone to leakage. Some studies con-
clude that bleb related endophthal-
mitis is higher when antimetabolites are used.
This is more common with inferior lim-
bal trabeculectomy. However, some studies show equal incidence in augmented trab-
eculectomy and trabeculectomy without
antimetabolite augmentation.

There are few data available for the incidence of blebitis. In most reported cases conjunctival swabs were performed for cul-
therefore, the need for topical antibody protection is negated by the presence of systemic antibodies against the specific bacte-
ria. We believe that by minimising the possibility of systemic infections with these agents we diminish the likelihood of blebitis.

It is possible that these vaccines could be given to patients with a history of blebitis and trabeculectomy. The cost for both vaccines would be less than £200.00.

Two hundred and fifty vaccinations could be paid for by the price of a single episode of bleb associated endophthalmitis. Assuming a long term infection rate of 2%, these vaccines could possibly prevent two cases of bleb asso-
ciated endophthalmitis, representing a saving of £5000 to the NHS.

Apart from the cost, vaccination has the potential to prevent significant ocular morbidity.
At the very least, these vaccines should be considered in high risk patients undergoing augmented trabeculectomy. We plan to con-
duct a prospective study of the effect of these vaccinations upon the incidence of blebitis and bleb related endophthalmitis.

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Accepted for publication 21 November 2001

References

The diagnosis of retinitis pigmentosa (RP) is made on the basis of characteristic retinal pigment migration changes, visual field defects, and reduction in amplitude or loss of rod scot-
topic standard electroretinography (ERG) responses, with a possible history of night blindness and a positive family history of RP. Multifocal electrophysiology (mERG), is a new technique that constructs a topographi-
cal map reflecting retinal function. Recent studies have suggested that the spatial resolution of mERG is sufficient to detect focal changes in retinal function as RP progresses.

We describe a case of early RP in which the amplitude and implicit times of the patient's standard ERG rod and maximal responses were normal in the right eye and equivocal in the left eye. However, the peripheral retinal mERG amplitude and implicit times were reduced and delayed. These abnormalities were obtained using a custom built wide field mERG, which facilitates assessment of a 90° degree retinal field.

Case report
A 29 year old woman was referred to the eye clinic by her optometrist. Abnormal retinal pigmentation was found in both fundi on routine examination. She had no visual prob-
lems and was otherwise systemically well. Her 51 year old mother is known to have RP.

On examination, she had 0.5 diopters of hypermetropia in both eyes. Her best cor-
rected logMAR visual acuities were 0.075 in the right eye and 0.025 in the left eye. Her colour vision, anterior segment examination, and intraocular pressures were normal in both eyes. Fundal examination revealed semicircu-
lar arcs of intraretinal “bone spicule” pigmen-
tation in the inferior mid-periphery of each retina. Her optic discs appeared normal and there was no evidence of attenuation of the retinal vasculature.
A Humphrey 120 point threshold related perimetry test was performed and the patient maintained fixation throughout the test. There was an arc of absolute visual field defect (Fig 1). The rod b-wave response of patient’s right and left eye compared with normal control (normal range of rod b-wave amplitude 72–367 nV) [A]; the maximal b-wave response of patient’s right and left eye compared with controls (normal range of maximal b-wave amplitude 241–709 nV) [B]; the cone b-wave response of patient’s right and left eye compared with normal control (normal range of cone b-wave amplitude 68–222 nV) [C].

The implicit times and amplitude of the scotopic rod, photopic cone and flicker responses of the patient were normal. The scotopic maximal b-wave amplitude was reduced by 14% in the left eye and normal in the right eye (Fig 1).

Wide field mfERG was performed, using a technique previously described. The amplitudes of the central and peripheral mfERG responses were grouped and averaged (Fig 2A) and compared with similar responses from normative data (Fig 2B). The average amplitude of the central mfERG response was 75 nV in the right and 101 nV in the left (normal range 74–122 nV) (Fig 2A). The average peripheral retinal mfERG responses were 29 nV in the right eye and 45 nV in the left eye (normal range 61–108 nV). The normal range is derived from a group of 40 controls, aged 20–40 years. In addition, the mfERG responses were reduced in areas that had normal visual field sensitivities.

Comment
Retinitis pigmentosa in its early stages of evolution is characterised by rod dominated photoreceptor dysfunction. Although mfERG is a photopic response, thought to predominantly reflect cone function, the nature of mfERG stimulation (that is, stimulus frequencies from 5 Hz to 75 Hz), indicates that this composite response may contain contributions from rods, in addition to cones and peripheral receptor cells.

The global nature of the Ganzfeld ERG requires approximately 30% of the retina to be dysfunctional before abnormalities can be detected. In this case report, the standard ERG did not help to confirm the diagnosis of RP. However, the spatial resolution of the peripheral wide field mfERG indicated peripheral retinal dysfunction, suggestive of RP.

We conclude that wide field mfERG may have advantages over Ganzfeld ERG in the electrophysiological diagnosis of some forms of retinitis pigmentosa. A Humphrey 120 point threshold related perimetry test was performed and the patient maintained fixation throughout the test. There was an arc of absolute visual field defect.

**References**

**Day 1 review following cataract surgery: are we seeing the precise details?**

The Royal College of Ophthalmologists published cataract surgery guidelines in February 2001. This document includes protocols relating to postoperative visits suggesting that there are no additional risks to patients who are not reviewed on the first postoperative day. This is a change in recommendation from previous college guidelines in 1995 suggesting a review within 48 hours.

There may follow a growing impetus for ophthalmologists to dispense with the first day review, given the reduced demand on clinical time and the corresponding accrual of staffing and financial resource benefits. While we applaud the dissemination of practice guidelines, they constitute “merely tools, not rules” to aid clinical decision making. They may have inherent limitations in particular circumstances and may require evaluation for effective application in clinical settings.

Four studies were quoted by the guideline authors, three of which advocated the omission of day after review and one of which was equivocal, suggesting that it was unsafe to abandon this practice unless raised intraocular pressures (IOP) were controlled. The numbers of patients included ranged from 100 to 387. The results of these studies are shown in Table 1.

**Comment**
In our view, deriving meaningful conclusions that may underpin clinical practice are difficult, owing to the varying methodological...
Table 1  Day 1 postoperative complications noted in clinical studies

<table>
<thead>
<tr>
<th>Study/in patients</th>
<th>Cornea oedema</th>
<th>Raised IOP</th>
<th>Uveits</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tufail et al 1897</td>
<td>4 (1%)</td>
<td>8 (2%)</td>
<td>–</td>
<td>12 (3.1%)</td>
</tr>
<tr>
<td>Whitefield et al 1990</td>
<td>10 (10%)</td>
<td>3 (3%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Cohen et al 1991</td>
<td>26 (13%)</td>
<td>12 (6%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Tan et al 1992</td>
<td>5 (4.4%)</td>
<td>3 (1.3%)</td>
<td>1 (0.4%)</td>
<td>3 (1.3%)</td>
</tr>
</tbody>
</table>

approaches used in these studies. In Tufail’s study, extracapsular cataract extraction was the predominant surgical technique used. Cohen et al excluded more than 50% of patients with complicated ocular histories or complicated surgery and Whitefield et al had similar extensive exclusion criteria, although the number excluded was not mentioned.

We would draw attention to a recently published study by McKellar and Elder, which to our knowledge is one of the largest cohort studies, aside from national cataract surveys, reporting on first and seventh day complications of cataract surgery. Of 1000 patients, the study found that on the first postoperative day complications were observed in 10% of eyes, of which 88% was raised IOP. Unlike most of the previous studies, all patients with available records were eligible, including those with preoperative risk factors and those with surgical complications. These figures align more closely with our “gold standard” of the United Kingdom (10%) may have an undetected early postoperative complication. In this case, we observed choroidal abnormalities in a patient with neurofibromatosis (Fig 2) at the corresponding region in the posterior pole in both eyes. There were no scotomas in those regions using SLO microperimetry.

Comment
In this case, we observed choroidal abnormalities in a patient with neurofibromatosis type 1. The conventional fundus examination, including biomicroscopic examination and fundus colour photography, did not show remarkable changes. However, the SLO examination showed regions of bright patches with infrared imaging using the direct confocal mode and dark patches with the indirect mode at both posterior poles. Infrared light penetrates the retina into the choroid more than visible light. Therefore, the bright patch regions seem with confostral infrared imaging and the absence of such regions under helium-neon light examination indicates that the patchy regions are of choroidal origin, as reported by Yasunari and colleagues. In addition, we observed dark patchy regions in the corresponding area using the indirect mode of infrared imaging (which also can obtain images of the deeper retinal layers non-invasively) instead of using indocyanine green fundus angiography. Yasunari and colleagues reported that choroidal abnormalities (100%) occurred more frequently than plexiform neurofibroma (29%) and Sakurai-Lisch nodules in the iris (76%).

Choroidal abnormalities in neurofibromatosis type 1 with non-invasive infrared imaging
Retinal abnormalities have been reported worldwide in patients with neurofibromatosis type 1. However, there have been few reports of choroidal abnormalities. We report a patient with choroidal abnormalities, associated with neurofibromatosis type 1, using a scanning laser ophthalmoscope (SLO).

Figure 1  SLO image of the right eye using the infrared direct confocal mode.

Figure 2  SLO image of the right eye using the infrared indirect mode.
The authors have no proprietary interest in any aspect of this report.

References


Corneal melt and perforation secondary to floppy eyelid syndrome in the presence of rheumatoid arthritis

Floppy eyelid syndrome (FES) is an uncommon condition that is often underdiagnosed or misdiagnosed owing to the somewhat trivial and non-specific symptoms with which it often presents. Association with the dry eye of rheumatoid arthritis it can, however, have devastating effects.

Case report

A 60 year old moderately obese man with well controlled rheumatoid arthritis (RA) presented to the eye clinic with recurrent red and gritty eyes. A diagnosis of dry eye syndrome with blepharitis was made. He was also found to have a mucocele of the left lacrimal sac. Lid hygiene and ocular lubricants yielded an initial encouraging response and he was discharged. He re-presented 5 years later with similar symptoms and reduced visual acuity (VA) of 6/36 in the left eye. A diagnosis of dry eye syndrome with secondary corneal epithelial changes was made. Topical lubricants failed to relieve the condition satisfactorily and he was therefore scheduled for punctal occlusion. However, upon admission for this 8 weeks later, he was found to have an asymptomatic perforation of the left cornea, with a VA of 6/60. The perforation was treated with glue and a bandage contact lens, topical antibiotic, steroids, and lubricants. Systemic immunosuppression was considered in view of the history of RA, but was withheld as a heavy growth of staphylococcus had been cultured from the cornea. The right eye demonstrated signs of dryness but was otherwise healthy with VA of 6/9.

Bilateral punctal occlusion was undertaken as planned, and in addition a left dacrocystorhinostomy (DCR) to eliminate the mucocele as a potential reservoir of infection. The eye, however, continued to slowly deteriorate, with persisting mucopurulent discharge, despite the DCR. Eventually uncontrolled endothalmitis developed requiring evisceration.

He re-presented 8 months later with reduced VA of 6/60 in the right eye secondary to a corneal melt (Fig 1A). A chronic mucopurulent discharge had also developed in the right eye, but no lacrimal sac mucocele was identifiable. On this occasion, however, it was noted on examination that while evertting the eyelids, all four lids exhibited excessive laxity (Fig 1B). This, together with a florid papillary tarsal conjunctival reaction and the chronic mucus discharge, led to a diagnosis of RA associated dry eye syndrome exacerbated by FES.

All four eyelids were immediately subjected to considerable shortening by pentagonal tarsorrhaphy; the corneal melt was treated with a bandage contact lens, with topical antibiotic, steroids and lubricants. The surgery was dramatic with complete resolution of discharge and gradual spontaneous repair of the corneal melt (Fig 2). The VA eventually recovered to 6/9.

Comment

FES occurs most frequently in middle aged obese males, although it has been described in young, slim males, females, and one child. Typically, the upper tarsus is rubbery and the upper eyelid everts easily with gentle upward pressure. A florid papillary conjunctivitis and chronic mucus discharge are common. Severe corneal involvement is rare, with only four reports in the literature of ulceration in association with FES and only two cases of perforation.

Although the exact pathophysiology of FES is uncertain, a sequence of events may lead to its development and to the secondary corneal changes has been proposed. Unidentified predisposing factors, possibly congenital, create a floppy upper tarsus. Whereas examination of post-excisional specimens have revealed normal tarsal collagen, elastin fibres are nearly absent. It is unclear whether this finding is causative or secondary. During sleep, a local pressure induced ischaemia may develop in the tarsus that, when relieved, results in a reperfusion injury which could injure tarsal elastin. In addition, there is a high incidence of obstructive sleep apnoea in FES patients and nocturnal diurnal pressure, could further contribute to the local ischaemia and subsequent elastin damage.

Corneal involvement may occur through one or more mechanisms. Spontaneous nocturnal lid eversion resulting from pressure of the pillow on the upper lid may lead to repeated trauma of the corneal epithelium. Lash ptiotis may contribute to this direct trauma. The cornea, however, may be damaged from a more subtle but important mechanism. Affected lid specimens demonstrate a marked polymorphonuclear infiltrate, which may be the sequelae of the reperfusion injury described above; this tarsal infiltrate and the associated papillary response may have direct toxic effects on corneal epithelium and stroma. It is perhaps intuitive that the corneal complications found in FES may be more severe when, as in our case, co-existing pathologies are present. Blepharitis and RA associated dry eye may both independently cause significant corneal pathology.

This case serves as a reminder that multiple pathologies may contribute to the clinical picture. If FES is not to be missed, ocular examination must include lid eversion and inspection of the tarsus.

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References


Ocular trauma with small framed spectacles
Penetrating injuries are widely reported with spectacle related eye trauma, particularly in car accidents. The use of high grade plastics and secure frames have been shown to reduce the incidence of spectacle related eye trauma. Spectacle safety may be compromised in the trend for small frames and frameless spectacles and may place patients at risk of serious ocular injury.

We present the case of an aphakic patient who sustained a blunt injury following ocular compression by her spectacles. Her injuries could have been avoided if larger framed spectacles had been worn.

Case report
A 79 year old aphakic woman sustained a non-penetrating injury to her left globe by walking into a door. She noted a sharp pain and sudden loss of vision. The globe was compressed by her spectacles, which were smaller than her orbital rim. Her glasses were not damaged and there was minimal periocular soft tissue injury.

She was aphakic, following bilateral cataract extraction for congenital cataracts. The spectacle refraction was +9.00 with a short back vertex distance of 5 mm (Fig 1).

The pinhole acuity was 6/36, a quiet, deep anterior chamber was noted with no aqueous leak, the intraocular pressure was 0 mm Hg. Funduscopy revealed a light vitreous haemorrhage, peripapillary choroidal ruptures, and a 360 degree suprachoroidal haemorrhage (Fig 2).

Hypopyon following traumatic ciliary artery spasm was diagnosed and the patient was treated conservatively, with topical atropine twice daily and dexamethasone four times daily. After 4 days the hypopyon resolved and the intraocular pressure returned to 14 mm Hg. The suprachoroidal haemorrhages resolved over 2 weeks and the visual acuity improved to 6/24.

Comment
The potential ocular damage from framed and frameless spectacles has been highlighted in a number of reports. These often result from minor road traffic accidents with inflation of air bags that damage the spectacles. The trauma is usually sufficient to break the lenses in the spectacles and the resultant globe laceration is the main cause of morbidity. The recent trends towards smaller framed spectacles has not been reported as a potential risk to the patient; however, in this case, with small framed spectacles (with a short back vertex distance), minor trauma was sufficient to cause serious eye injury. The patient’s previous larger framed spectacles would have prevented such an injury as the lenses would have been supported by the orbital margin and not the globe.

This case demonstrates the previously unconsidered risk of small framed spectacles in aphakic patients. The back vertex distance may be short, increasing the risk of blunt injury.

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References

Late opacification of SC60B-OUV acrylic intraocular lenses
Optical clarity of the intraocular lens (IOL) is paramount in maintaining visual improvement after cataract extraction. While the advent of newer foldable IOLs have revolutionised cataract surgery by the smaller incisions required, their long term safety has been established only with time. There have been recent reports of two separate groups of hydrophilic IOLs that have shown various degrees of opacification, 6 months to 2 years after implantation. These include the Hydroview lens (Bausch & Lomb Surgical, Claremont, CA, USA) and the model SC60B-OUV (Medical Developmental Research, Clearwater, FL, USA). We report our experiences with two patients who had permanent reduction in visual acuity 1–2 years after implantation of the SC60B-OUV IOL.

Case reports
Case 1
An 82 year old man with visual acuities of 6/60 in both eyes because of a right macular hole and left posterior subcapsular cataract underwent routine left phacoemulsification and intraocular lens implantation in March 1998. A foldable hydrophilic acrylic IOL (Model SC60B-OUV, MDR, Inc) was implanted in the capsular bag. Postoperative recovery was uneventful, with left visual acuity improving to 6/4 with correction, 6 weeks after the surgery.

He was referred back to the clinic in March 2000 with symptoms of intermittent binocular diplopia, which was relieved with Fresnel prisms. His left visual acuity was 6/6, but the intraocular lens was found to be uniformly cloudy (Fig 1A). As the patient was not experiencing any symptoms from his cloudy IOL, no intervention was advised. Over the next 6 months the brownish discoloration of the IOL increased in intensity and the patient complained of “foggy vision.” His visual acuity dropped to 6/12 and he was disturbed by the distorted, but clear images from his right eye and hazy images from his left eye. Fundus examination of the left eye has also become progressively difficult. IOL exchange is being considered, bearing in mind the presence of the macular hole in the fellow eye and the risks of removing a posterior chamber IOL 3 years after implantation.

Case 2
A 78 year old woman with bilateral Fuchs’ corneal endothelial dystrophy and long standing central retinal vein occlusion in the left eye, underwent uneventful right phacoemulsification with intraocular lens implantation (Model SC60B-OUV, MDR, Inc) in August 1998. Her right visual acuity improved to 6/9, 6 months after the surgery.

Figure 1 (A) Diffuse opacification of the IOL 2 years after implantation in case 1. (B) The explanted opacified IOL optic from case 2 is compared to a normal acrylic IOL.
but approximately 1 year after the surgery there was gradual deterioration of vision to 6/24 in August 2000. At this stage the IOL was noted to be cloudy; there was also progression of her endothelial dystrophy. She underwent uneventful right penetrating keratoplasty in July 2000 and has a current right visual acuity of 6/12. The explanted opacified IOL is compared to a normal clear acrylic IOL in Figure 1B.

Comment

The safety and efficacy of AcrySof polyacrylic IOLs has been reported to be equal to or better than poly(methylmethacrylate) IOLs. The unexpected late opacification of the acrylic IOL (SC60B-OUV), implanted in the only “good” eye of both our patients, resulted in significant visual disability and clinical dilemma. This model of IOL was first produced in June 1997 by Medical Development Research (MDR, Inc). More than 60 000 of these lenses have been implanted worldwide, but only outside the United States. Reports on opacification of the IOL started coming through to the manufacturer in May 1999. (Summary of SC60B-OUV lens opacification investigation, personal communication from MDR, Inc, 20 July 2001.)

Several theories have been put forward to explain the late clouding of the IOL optic.1-3 Analysis of 23 explanted IOLs of the same model has shown that degeneration of the ultraviolet filtration material and calcium deposits within the optic biomaterial are responsible for the opacification of the IOL.4 Werner et al analysed nine explanted IOLs of the same model and demonstrated the presence of calcium phosphate salts in the deposits within the optics of the IOL.5 Investigations by the manufacturer identified four lots of polymeric biomaterial formulated and prepared by Vista Optics (London) and used by MDR, Inc in the IOL manufacture, that correlated with opacification complaints. (Summary of SC60B-OUV lens opacification investigation, personal communication from MDR, Inc, 20 July 2001.)

Ninety two of the estimated 60 000 SC60B-OUV IOLs implanted were explanted and returned to the company. MDR, Inc ceased exporting SC60B-OUV lenses in June 2000 and claims that the opacification represents only 0.15% of total SC60B-OUV IOLs implanted. (Summary of SC60B-OUV lens opacification investigation, personal communication from MDR, Inc, 20 July 2001.) However, this does not account for those patients who have not yet had their opaque IOLs identified or explanted. The lateness of the onset of opacification and resulting visual disability may mean that we are seeing only the tip of the iceberg.

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Accepted for publication 3 December 2001

References


Treatment of superior limbic keratoconjunctivitis with a unilateral bandage contact lens

The typical patient with superior limbic keratoconjunctivitis (SLK) is a woman aged between 20 and 60 years of age with chronic red and irrigated eyes.1 Although both eyes are usually affected, the condition may be asymmetrical.2 After episodes of exacerbation and remission it usually resolves. The patient may also have abnormal thyroid function.3 SLK has been treated with silver nitrate or thermal cautery of the superior bulbar conjunctiva, pressure patching, and large diameter bandage contact lenses (BCL), topical trans-retinoid acid 0.1%, and resection of the superior bulbar conjunctiva.3 Over 50% of patients with SLK are said to have keratoconjunctivitis sicca4 and recently upper punctal plugs have been used to treat SLK.

We report two cases in which a unilateral BCL wear ameliorated the symptoms of bilateral SLK and a possible explanation is discussed.

Case reports

Case 1

A 38 year old woman presented with a 3 month history of Irritated photophobic eyes that were unresponsive to preserved lubricants. Her right eye was amblyopic. On systemic review she reported weight loss, heat intolerance, and insomnia. Slit lamp examination revealed bilateral superior conjunctival hyperaemia, superior punctate epithelial erosions, and four to five filaments and micropapillae on the superior cornea of each eye. Both superior tarsal conjunctivae had moderate papillary reactions. Schirmer’s test without anaesthesia was 14 mm on the right and 15 mm on the left at 5 minutes. Non-preserved lubricants every 1–2 hours and Lacrilube ointment at night were prescribed. Thyroid function tests revealed hyperthyroidism and she was referred to an endocrinologist who commenced carbimazole.

Three months later she returned still complaining of persistent severe discomfort, photophobia, and a burning sensation in both eyes. The ocular examination was unchanged. A silicone hydrogel BCL (Pure Vision, Bausch and Lomb, 36% water content) was inserted into the right eye and within an hour she had symptomatic relief in both eyes. Non-preserved lubricants were continued for the left eye.

On review 2 months after insertion of the BCL she remained asymptomatic. Mild superior limbal hyperaemia on the right and punctate staining of the superior conjunctiva and adjacent cornea on the left were found on examination.

Case 2

A 54 year old woman was referred with a 3 year history of sore, gritty eyes, worse on the left. The tear break up time was <10 seconds and Schirmer test without anaesthesia was right 0 mm and left 1 mm after 3 minutes. There was some relief from lubricants, although on occasion the pain was so severe that she required oral analgesia.

On examination punctate epithelial erosions were found on the superior bulbar conjunctiva of both eyes, but were more marked on the left (Fig 1). A silicone hydrogel BCL (Pure Vision, Bausch and Lomb) was inserted into the left eye, which rapidly ameliorated the symptoms in both eyes. Lubricants were continued for the right eye.

Three months later she remained asymptomatic and no fluorescein staining was seen. The BCL was removed. Within a month her bilateral ocular discomfort returned and the left BCL was refitted with immediate symptomatic relief in both eyes. Two months later the BCL was lost and her bilateral ocular discomfort recurred within 2 days. Since then her BCL has been replaced every 3 months. Occasionally she has used lubricants for the right eye.

One year following her presentation she became tachycardic and hyperthyroidism was diagnosed. Her endocrinologist commenced carbimazole and B blockers. Her treatment was later changed to propylthiouracil after she suffered from carbimazole induced arthralgia. Thyroid function tests were normal 6 months later and the propylthiouracil was stopped.

Comment

The pathogenesis of SLK is unclear. It may be the result of mechanical irritation from increased pressure of the upper eyelid against the globe and/or increased motility of the upper bulbar conjunctiva from hypothyroidism or ageing.1 Increased upper eyelid tightness may be the result of thyroid eye disease or chronic inflammation and, in addition, may...
impair the normal turnover of bulbar conjunctival epithelial cells. This may be aggravated, in some patients, by blepharospasm, which increases the force on the globe. 1, 2 Therapeutic lenses can produce rapid symptomatic relief in SLK. 1, 3 They may be helpful in the treatment of SLK as they relieve pain, facilitate healing of punctate epithelial erosions by protecting the ocular surface from the eyelids, reduce upper lid pressure on the globe and alter tear dynamics. 3 During blinking as the upper lid moves downwards to meet the lower lid, significant forces are exerted on the globe. The lens can reduce the force on the superior limbus from blinking as it has a lower mechanical stiffness and elastic modulus. 3

In the first week of contact lens wear tear production increases dramatically and tear film instability increases. 4 Tear production then normalises and tear tonicity rises as evaporation increases. 5 A lens may then aid aqueous tear deficiency, which can accompany SLK, by ensuring a continuous precorneal tear film. 6

There is little certainty of the mechanism of bilateral symptomatic relief from unilateral BCL wear in SLK. One possible explanation is that BCL wear reduces the tactile corneal reflex. 7, 8 During the first week after lens insertion, and this reduction, which increases with the duration of lens wear, would then decrease bilateral reflex blinking. 7, 9 This would protect the superior limbus in both eyes from the friction associated with blinking and may also break the cycle that leads to blepharospasm in SLK. 8 It should not be forgotten that continuous BCL wear carries risks including microbial keratitis and corneal vascularisation. 10 New extended wear silicone BCL, as used in our patients, increases oxygen transfer and has been shown to reduce such risks. 11

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Accepted for publication 17 December 2001

References


MAILBOX

Keratctasia after PTK

Takahashi and colleagues have elegantly described an interesting and rare complication of phototherapeutic keratectomy (PTK) in their recent report of an unusual case of keratctasia after PTK. 1 The hypothesis that risk of ectasia is proportional to residual stromal base, or depth of ablation, fits with the assumed biomechanical aetiology of this recently reported complication of laser refractive surgery. The generally accepted empirical minimum thickness of 250–300 µm of corneal stroma, excluding flap thickness, remains speculative, as we do not understand the underlying pathophysiology. Indeed, although Holland et al. 2 highlighted the association of thin residual stromal thicknesses, post-PRK and LASIK, with keratctasia, they also described this complication after surface ablated hypermetropic PRK ablation, where the centre was minimally ablated and residual stromal thickness was greater than 360 µm. 2 The authors suggest, in the reported case, that band-shaped keratopathy (BSK) may have compromised the tensile strength of the cornea. This seems unlikely as this condition generally affects the superficial anterior cornea, and usually does not penetrate deeper than Bowman’s layer and its suitability for treatment by PTK. However, further clinical detail which the authors have not provided might reveal underlying corneal pathology with secondary “rough” BSK rather than “smooth” BSK.

However, there are a number of reasons, other than simple biomechanical compromise, for keratctasia following PTK in this case: (1) forme fruste keratoconus—as no preoperative topography or surface asymmetry values are presented to enable the reader to rule this out; (2) clinical keratoconus, which seems less likely in respect of patient’s age and a preoperative cylindrical error of only −1.50 D; (3) idiopathic keratctasia, possibly secondary to widespread deregulated keratocyte apoptosis. The latter has been demonstrated after LASIK, with a considerable and longstanding decrease in keratocytes in the peri-ablation area. 3 Also, Helena et al. 4 demonstrated apoptosis to a depth of at least 50 µm after all of the following procedures: epithelial scrape, corneal scrape PRK, transepithelial PRK, and LASIK. Epithelial scrape and LASIK demonstrated keratocyte apoptosis to depths of up to 75 µm and 100 µm, respectively. The authors have recently identified a keratocyte free zone 160 µm of corneal stroma following LASIK, and theoretically more widespread apoptosis as a response to excimer laser photorefractive surgery, may contribute to keratctasia.

While it is difficult to ascertain why keratctasia occurs, in this case with a residual stromal thickness of over 500 µm, from the data provided the most likely aetiologies would seem to be either undiagnosed forme fruste keratoconus or idiopathic keratctasia. Currently, recent reviews illustrate the dearth of substantial information available regarding idiopathic keratctasia (iatrogenic keratoconus), with a little over 60 cases published. At this point, although some are likely to be due to over-ablation, for many cases such as this the exact aetiology remains unknown and is likely to be multifactorial, and one of these factors is residual corneal thickness. The fact that keratctasia can occur, after what would be considered minimal ablation, highlights the unpredictability of occurrence, but with over a million cases of LASIK or PRK occurring each year, the stimulus to identify contributing factors is significant.

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BOOK REVIEWS

Age Related Macular Degeneration—Current Treatment Concepts.


This book is one in a large series of medical radiology textbooks concentrating on diagnostic imaging and radiation oncology. There are 71 authors in all, the majority of whom work either in the eastern United States or Germany, and the book contains 24 chapters. The text is aimed at specialists in radiotherapy rather than ophthalmology. The majority of the chapters are concerned with radiotherapy of age related macular degeneration (AMD) with contributions from experts in this field. Some of the chapters contain results of controlled studies and are of good value, whereas others have less scientific merit in that they are long term follow up studies without controls. For ophthalmologists wishing to look into the subject of

www.bjophthalmol.com
radiotherapy for ARMD this would be a good source of material and is well referenced.

There are a few chapters on the clinical manifestations, diagnosis, and surgery of ARMD but there is no real mention of laser treatment. Many of these chapters are of limited scope and do not provide a comprehensive overview of the ophthalmic assessment and management of ARMD.

Although not stated in the book it reads as if it is the proceedings of a clinical meeting. The chapters do not read in a coordinated way and essentially present the results of individual units describing their methods and results of radiotherapy. As such it is a useful source of information for those with an interest in this topic but it is of limited value for ophthalmologists wishing to obtain a balanced view of current treatment of ARMD.

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Clinical Ophthalmic Pathology.

This is an excellent, easy to read, well illustrated book. It is one of the first of its kind to bring pathology alive by describing diseases via pathogenesis as opposed to anatomy. For the trainee in ophthalmology, optometry, and visual science it, therefore, provides a more logical approach to the understanding of ocular diseases. As the book attempts to cover many subjects it sometimes does not do justice to any one. It would have been better to have had the reading lists at the end of each chapter. However, as it stands the book is an excellent introduction to pathology complementing clinical textbooks. If read together with clinical texts it certainly will broaden the knowledge base of all trainee ophthalmologists. As a result of its logical and simple approach I was left frustrated at times with the lack of background knowledge. However, going through pathologically based chapters including injury and repair, immunity, genetics, growth, degeneration, vascular disorders, and disorders of the nerve and muscle, I was left entertained, as a clinician, with a greater understanding of pathological processes.

The final chapter for the clinician in the laboratory I felt could have been expanded, delivering more detail, particularly, on the current molecular methods used in pathological practice today.

A Dick

NOTICES

Childhood blindness

The latest issue of Community Eye Health (No 40) discusses new issues in childhood blindness, with an editorial by Clare Gilbert, senior lecturer at the International Centre for Eye Health. For further information please contact: Journal of Community Eye Health, International Centre for Eye Health, Institute of Ophthalmology, 11–43 Bath Street, London ECIV 9EL, UK (tel: +44 (0)20 7608 6910; fax: +44 (0)20 7250 3207; email: eyesresource@ucl.ac.uk; website: www.jchc.co.uk). Annual subscription (4 issues) UK£25/US$40. Free to workers in developing countries.

International Centre for Eye Health

The International Centre for Eye Health has published a new edition of the Standard List of Medicines, Equipment, Instruments and Optical Supplies (2001) for eye care services in developing countries. It is compiled by the Task Force of the International Agency for the Prevention of Blindness. Further details: Sue Stevens, International Centre for Eye Health, 11–43 Bath Street, London ECIV 9EL, UK (tel: +44 (0)20 7608 6910; email: eyesresource@ucl.ac.uk).

Second Sight

Second Sight, a UK based charity whose aims are to eliminate the backlog of cataract blind in India by the year 2020 and to establish strong links between Indian and British ophthalmologists, is regularly sending volunteer surgeons to India. Details can be found at the charity website (www.secondsight.org.uk) or by contacting Dr Lucy Mathen (lucy.mathen@yahoo.com).

Specific Eye Conditions (SPECS)

Secific Eye Conditions (SPECS) is a not for profit organisation which acts as an umbrella organisation for support groups of any conditions or syndrome with an integral eye disorder. SPECS represents over fifty different organisations related to eye disorders ranging from conditions that are relatively common to very rare syndromes. We also include groups who offer support of a more general nature to visually impaired and blind people. Support groups meet regularly in the Boardroom at Moorfields Eye Hospital to offer support to each other, share experiences and explore new ways of working together. The web site www.eyescondition.org.uk acts as a portal giving direct access to support groups own sites. The SPECS web page is a valuable resource for professionals and may also be of interest to people with a visual impairment or who are blind. For further details about SPECS contact: Kay Parkinson, SPECS Development Officer (tel: +44 (0)1803 524238; email: k@eyesconditions.org.uk; www.eyesconditions.org.uk).

XXIXth International Congress of Ophthalmology

The XXIXth International Congress of Ophthalmology will be held on 21–25 April 2002 in Sydney, Australia. Further details: Congress Secretariat, C/- ICMS Australia Pty Ltd, GPO Box 2609, Sydney, NSW 2001, Australia (tel: +61 2 9241 1478; fax: +61 2 9231 3522; email: ophthalm@icmsaustralia.com.au; website: www.ophthalmology.aust.com).

12th Meeting of the European Association for the Study of Diabetic Eye Complications (EASDEC)

The 12th meeting of the EASDEC will be held on 24–26 May 2002 in Udine, Italy. The deadline for abstracts is 15 February 2002. Three travel grants for young members (less than 35 years of age at the time of the meeting) are available. For further information on the travel grants, please contact Pr CD Agardh, President of EASDEC, Malmö University Hospital, SE-205 02 Malmö, Sweden (tel +46 40 33 10 16; fax: +46 40 33 33 66; email: carl-david.agardh@endo.max.lu.se). Further details: NORD EST CONGRESISS, Via Aquilea, 21–33100 Udine, Italy (tel: +39 0432 21399; fax: +39 0432 50687; email: nordest.congresi@ul.net.uno.it).

3rd Interdisciplinary Symposium on the Treatment of Autoimmune Disorders

The 3rd Interdisciplinary Symposium on the Treatment of Autoimmune Disorders will be held in Leipzig, Germany on the 6–8 June 2002. Topics to be covered include: basic aspects of autoimmune diseases, experimental therapeutic concepts, and clinical studies providing novel concepts or novel focus on established therapies. There will also be the presentation of the Niels-Ilja-Richter Award (application deadline is April 2002, further details on the web site). Further details: Prof. Dr. med. Michael Sticherling, Department of Dermatology, University of Leipzig (email: stiemi@medizin.uni-leipzig.de; website: www/autoimmum.org); Fördergesellschaft zur Therapie von Autoimmunerkrankungen e.V. (email: autoimmun.org@gmx.de).

International Society for Behçet’s Disease

The 10th International Congress on Behçet’s Disease will be held in Berlin 27–29 June 2002. Further details: Professor Ch Zouboulis (email: zoubbec@zedat.fu-berlin.de).

Singapore National Eye Centre 5th International Meeting

The Singapore National Eye Centre 5th International Meeting will be held on 3–5 August 2002 in Singapore. Further details: Ms Amy Lim, Organising Secretariat, Singapore National Eye Centre, 11 Third Hospital Avenue, Singapore 168751 (tel: (65) 322 8374; fax: (65) 227 7290; email: Amy_lim@snec.com.sg).

BEAVRS Meeting

The next BEAVRS meeting will be held in the Dalmahoy Hotel near Edinburgh on 31 October to 1 November 2002. Further details: Susan Campbell, Medical Secretary, Gartnavel General Hospital (email: susan.j.campbell.wg@northglasgow.scot.nhs.uk).

Correction

The authors of the letter “Recurrent corneal ulceration as late complication of toxic keratitis”, appearing in the February issue of BJO (2002;86;245–6), would like to add an author, SH Santander.