LETTERS

Subconjunctival corticosteroid injection for the treatment of non-necrotising anterior scleritis

Scleritis commonly is a recurrent disease that requires long term immunosuppressive treatment that can be associated with significant adverse effects. Although topical and peri-orbital steroids are accepted therapeutic options for treatment of scleritis, subconjunctival administration of depot corticosteroids has been considered unsafe owing to the risk of scleral thinning and perforation. Recently, this has been challenged by reports describing the safe and effective use of subconjunctival depot steroid injections in patients with non-necrotising scleritis. Based on these reports we reviewed our experience using subconjunctival corticosteroid injections (SCI) in the management of non-infectious, non-necrotising anterior scleritis.

Case reports

A retrospective, non-comparative review of the clinical records of patients with scleritis evaluated at the National Eye Institute was performed and four patients treated with SCI were identified. Subconjunctival triamcinolone acetonide (Kenalog 40 mg/ml, Westwood Squibb Pharmaceuticals, Buffalo, NY, USA) (2–12 mg per injection) was given to an area of the eye at one time. The initial indication for SCI in all cases was unilaterally active non-necrotising anterior scleritis.

The initial indication for SCI in all cases was unilaterally active non-necrotising anterior scleritis. The patient was HLAB27 positive but did not have any associated systemic manifestations.

Approximately 37% of patients with scleritis have associated systemic diseases. A seminal review of scleritis cases showed that 26.1% of patients with scleritis require systemic immunosuppressive therapy and, of these, 37.5% experience a treatment complication.

Although associated with cataract and increased intraocular pressure, local steroid therapy offers the benefit of anti-inflammatory control without the side effects of systemic immunosuppressive drugs. However, because of the potential risk of scleral thinning and perforation subconjunctival steroid injections have been avoided for the treatment of scleritis.

Recently, two retrospective studies demonstrated rapid clinical improvement with SCI in all cases without evidence of scleral thinning or serious side effects in non-necrotising scleritis during follow up extending to 23 years. However, recurrences occurred in both studies requiring repeated injections. In a prospective study, patients with recalcitrant scleritis were able to discontinue all immunosuppressive therapy after treatment with SCI. Although SCIs were effective in rapidly controlling the active scleritis in our series, two patients developed steroid response and all developed recurrent scleritis requiring repeated injections. The major benefit of the SCI in this series was that they facilitated >50% reduction in each patient’s prednisone dose which could not be achieved or sustained before their use.

This study confirms the existing literature that SCI may be an effective adjunct for the treatment of active non-necrotising anterior scleritis.

Table 1 Clinical characteristics, treatment regimens, and adverse events with subconjunctival steroid injection

<table>
<thead>
<tr>
<th>Patient</th>
<th>Scleritis type</th>
<th>Associated disease</th>
<th>Systemic treatment before SCI</th>
<th>Systemic treatment after SCI (last follow up)</th>
<th>Side effects from systemic treatment</th>
<th>Prednisone dose followed by relapse before SCI</th>
<th>Duration of scleritis before SCI (months)</th>
<th>Prednisone dose followed by relapse before SCI</th>
<th>Duration of scleritis before SCI (months)</th>
<th>Systemic treatment after SCI</th>
<th>Follow up (months)</th>
<th>Side effects from SCI</th>
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<tr>
<td>1, 57, F</td>
<td>Nodular anterior</td>
<td>Mixed connective tissue disease (inactive)</td>
<td>Pred 15 mg, MTX 12.5 mg</td>
<td>None</td>
<td>Osteoporosis, Weight gain</td>
<td>7.5 mg, 63</td>
<td>None</td>
<td>22</td>
<td>None</td>
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<td>2, 41, M</td>
<td>Diffuse anterior</td>
<td>None</td>
<td>Pred 9 mg, MMF 2 g</td>
<td>Pred 4 mg, MMF 2 g</td>
<td>Pneumonia</td>
<td>9 mg, 14</td>
<td>Pred 4 mg, MMF 2 g</td>
<td>19</td>
<td>Transient IOP increase LE</td>
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<tr>
<td>3, 56, F</td>
<td>Diffuse anterior</td>
<td>Sjogren’s syndrome (inactive)</td>
<td>Pred 20 mg, MTX 20 mg</td>
<td>Pred 5 mg, MTX 25 mg</td>
<td>Weight gain, Myopathy, Recurrent URTI, GERD</td>
<td>15 mg, 36</td>
<td>None</td>
<td>17</td>
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<tr>
<td>4, 11, M</td>
<td>Diffuse anterior</td>
<td>None</td>
<td>Pred 50 mg, CSA 150 mg</td>
<td>Pred 10 mg</td>
<td>Weight gain, Persistent headache</td>
<td>50 mg, 11</td>
<td>Pred 10 mg</td>
<td>5</td>
<td>Subconjunctival haemorrhage, Transient IOP increase LE</td>
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</table>

SCI, subconjunctival injection of triamcinolone acetonide; CSA, cyclosporine; MMF, mycophenolate mofetil; Pred, prednisone; URTI, urinary tract infection; GERD, gastroesophageal reflux disease.

Doses in the table indicate daily oral dose for Prednisone and CSA and weekly dose for methotrexate.

*The patient was HLAB27 positive but did not have any associated systemic manifestations.
scleritis in appropriately selected patients and that this therapy is not unequivocally associated with the risk of scleral thinning or perforation. Local treatment with steroid injections may help reduce potential side effects from systemic therapy in patients with active scleritis in the absence of active systemic disease. As with all retrospective small case series our results should be interpreted with caution.

H N Sen, R Ursea, R B Nussenblatt, R R Buggage
Laboratory of Immunology, National Eye Institute, National Institutes of Health, Bethesda, MD, USA

Correspondence to: H Nida Sen, MD, National Eye Institute, National Institutes of Health, Bldg 10, Room 10N112, Bethesda, MD 20892, USA; nidamd@yahoo.com
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References

Visual field defect in association with chiasmal migration of intraocular silicone oil

Silicone oil is used as a long term intraocular tamponade in vitreoretinal surgery for conditions such as rhegmatogenous retinal detachments. Complications such as silicone oil associated keratopathy, cataract formation, or emulsification with secondary glaucoma have been reported.1 We report the unusual case of intracranial silicone oil migration with subsequent visual field defect in the fellow eye and the results of neurosurgical intervention

Case report
A 66 year old male patient had undergone cataract surgery and lens implantation on both eyes 20 months earlier. Six months after cataract surgery a retinal detachment was seen in the left eye which was first treated with pars plana vitrectomy and gas tamponade (15% C<sub>2</sub>F<sub>6</sub>). After a period of 4 months, a ret detachment occurred and silicone oil (Acti.Sil-ol 5000, 5000 cps, Acti.Tec, Hennigsdorf, Germany) was injected. A redetachment was seen after 3 months which was managed with silicone oil injection (Acri.Sil-ol 5000, 5000 cps, Acri.Tec, Hennigsdorf, Germany) and gas tamponade (15% C<sub>2</sub>F<sub>6</sub>). One week after silicone oil injection a visual field defect of the right eye was noted. Visual acuity was 20/20. The optic disc of the left eye shows atrophy and glaucomatous damage, the cup-disc ratio is 1.0. Note the reflex on the retinal surface, caused by the silicone oil tamponade. (C) Axial T1 weighted MRI scans demonstrate intracranial silicone oil (S) which is hypointensive compared to the normal vitreous (V). Note the hypointensive signal in the area of the intracranial portion of the optic nerve and the chiasm (arrow). (D) T2 weighted scans demonstrate hypointensive signals in the silicone filled eye (S) as well as in the area of the optic nerve and chiasm.

Figure 1  (A) Temporal hemianopia of the right eye. Visual acuity was 20/20. (B) The optic disc of the left eye shows atrophy and glaucomatous damage, the cup-disc ratio is 1.0. Note the reflex on the retinal surface, caused by the silicone oil tamponade. (C) Axial T1 weighted MRI scans demonstrate intracranial silicone oil (S) which is hypointensive compared to the normal vitreous (V). Note the hypointensive signal in the area of the intracranial portion of the optic nerve and the chiasm (arrow). (D) T2 weighted scans demonstrate hypointensive signals in the silicone filled eye (S) as well as in the area of the optic nerve and chiasm.

Figure 2  (A) Left optic nerve (ON) before incision. Arrows point at translucent thinned perineural sheath with silicone oil beneath. OC, entrance to optic canal; ICA, internal carotid artery. (B) Silicone oil (arrow) evading from left optic nerve after incision of perineural sheath. (C) View into left optic nerve after complete removal of the silicone oil. The nerve has been hollowed out by the oil with just thin nerve walls remaining. (D) After intracranial surgery, a regression of the visual field defect was observed.
Intraocular pressure in the left eye was normal. Defect in the right eye was observed (fig 2D) and for treatment of elevated intraocular pressure in the left eye. At presentation, visual acuity was 20/20 in the right eye and light perception in the left eye. Intraocular pressure was in a normal range in the right eye and elevated to 35 mm Hg in the left eye despite local antiglaucomatous monotherapy using latanoprost (Xalatan). Emulsified silicone oil was detected in the anterior chamber angle during gonioscopy of the left eye. Funduscopy revealed a glaucomatous exudation and atrophy of the optic nerve in the left eye with a cup-disc ratio of 1.0 (fig 1B) and in the right eye with a cup-disc ratio of 0.7. The silicone oil filling of the globe being incomplete. Magnetic resonance imaging (MRI) of the brain was then performed for further evaluation. T1 weighted MRI scan revealed a hyperintense signal in the left vitreous cavity characteristic for silicone oil. An identical hyperintense signal was also observed in the left optic nerve and the left half of the optic chiasm (fig 1C); in contrast, in T2 weighted sections the silicone located in the optical system and in the vitreous cavity appeared hypointense (fig 1D). In order to prevent further progression of the visual field defect of the right eye, a decompression of the optic nerve seemed prudent. After informed consent, the patient underwent left subfrontal craniotomy and intracranial decompression of the optic nerve sheath and its close proximity to the compression caused by silicone oil within the left vitreous cavity characteristic for silicone oil. The case presented in this report is unique because of the additional affection of the fellow eye as a result of optic nerve compression caused by silicone oil within the optic nerve sheath and its close proximity to the chiasm and the intraventricular and visual field defect after neurosurgical intervention.

D Eckle, A Kampik, C Hintschich, C Haritoglou
Department of Ophthalmology, Ludwig-Maximilians-University, Mathildenstrasse 8, 80336 Munich, Germany

J-C Tonn, E Ulh
Department of Neurosurgery, Klinikum Großhadern, Ludwig-Maximilians-University, Marchioninistrasse 15, 81377 Munich, Germany

Comment
This case indicates that silicone oil can migrate intracranially under certain, yet unknown, conditions. Referring to the literature, there is only one case by Eller and co-authors describing the intracranial migration of silicone oil in a patient with AIDS who had undergone vitrectomy with silicone oil tamponade for treatment of retinal detachment secondary to cytomegalovirus infection of the retina. As in our patient, there was a coincidence of uncontrolled high intraocular pressure associated with atrophy and glaucomatous damage of the optic nerve. It seems likely that elevated intraocular pressure and optic nerve atrophy allowed intravitreal silicone oil to migrate intracranially, although a definite conclusion cannot be drawn from two case reports. However, a histopathological study previously demonstrated silicone cavities posterior to the lamina cribrosa in an silicone filled eye with glaucomatous nerve damage. The case presented in this report is unique because of the additional affection of the fellow eye as a result of optic nerve compression caused by silicone oil within the optic nerve sheath and its close proximity to the chiasm and the visual field defect after neurosurgical intervention.

Figure 1 (A) Fundus photograph of the left eye showing an elliptical area of congenital retinal pigment epithelial (RPE) lesion temporal to the fovea. The lesion is depigmented in the nasal aspect with scalloped hyperpigmentation temporally. (B) Normal RPE cells, with prominent apical melanosome and basal nuclei. Normal Bruch’s membrane and choriocapillaris (haematoxylin and eosin ×330). (C) Hypertrophic, thickened retinal pigment epithelial cells from pigmented area of CHRPE. These are thickened, and packed with melanosomes, in both apical and basal areas (haematoxylin and eosin, ×500). (D) Atrophic, thinned RPE cells from depigmented areas of CHRPE lesion. Almost no melanosomes remain in these cells (haematoxylin and eosin, ×500). (E) Electron micrograph from segmentation of the cells from pigmented areas of CHRPE. Nuclei are both apical and basal. Melanosomes are dense and rounded. Bruch’s membrane is normal (electron microscopy ×2000). (F) Electron microscopy from the depigmented area of the CHRPE. RPE cells are atrophic and vacuolated, with very few small melanosomes. Bruch’s membrane consists of the RPE basement membrane (arrow), a lucent thickened inner collagenous layer (below) extending to the elastic layer (middle dark layer). Note the RPE cell (*) within the outer collagenous layer of Bruch’s membrane. The choriocapillaris (C) is separated from this by a thickened lucent outer collagenous layer (right), but choriocapillaris (and endothelial cell basement membrane) are missing from most areas (central and left) (electron microscopy ×2000).

References
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...depigmented in the nasal aspect in basal dimension and appeared flat. The...with Gardner's syndrome.

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Case report

A 62 year old woman with a large cilio-

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...in Bruch's membrane into the inner collagenous...pigmentation, starting with atrophy of RPE cells, followed by fragmentation of melanosomes progresses to gradual loss of melanosomes and RPE atrophy. Both the fragmentation of melanosomes and eventual atrophy appear to correlate with increasing changes in Bruch's membrane—namely, an increased thickness of the inner and outer collagenous layers, by electron lucent material may represent water, lipid or proteinaceous deposit. To our knowledge, this has not been reported previously, although review of an electron micrograph from the 23 year old patient of Buettner suggests that this feature may be present in a depigmented area of CHRPE. A close inter-reaction between the functions of RPE cells and Bruch’s membrane is well recognised. It remains possible that the depigmentation of RPE cells caused the changes in the Bruch’s membrane (‘‘ancient’’ CHRPE). It is also possible that, in our patient, changes affecting Bruch’s membrane are non-specific and are unrelated to CHRPE.

Acknowledgements

We are grateful to Mr Bart Wagner for assistance with electron microscopy and Mr Robin Farr with the graphics.

Table 1

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<thead>
<tr>
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<th>Age (years)</th>
<th>Type</th>
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FAP, familial adenomatous polyposis with or without extracolonic manifestations (Gardner’s syndrome).

Congenital hypertrophy of retinal pigment epithelium: a clinicopathological case report

Congenital hypertrophy of retinal pigment epithelium (CHRPE) is a peculiar congenital anomaly of the retinal pigment epithelium (RPE) diagnosed by its characteristic ophthal-
omicroscopic appearance.1 It is now realised that sporadic CHRPE is distinct from the similar appearing retinal lesions described in patients with Gardner’s syndrome.4 We recently elucidated an eye with a choroidal melanoma that also had a distinct area of solitary CHRPE with lacunae formation. This provided us with a unique opportunity to correlate clinical and histopathological features of a solitary CHRPE.

Case report

A 62 year old woman with a large cilio-

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A D Singh
Cole Eye Institute, Cleveland Clinic Foundation, Cleveland, OH, USA
Correspondence to: Arun D Singh, MD, Department of Ophthalmic Oncology, Cole Eye Institute, Cleveland Clinic Foundation, 9500 Euclid Avenue, Cleveland, OH 44195, USA; singha@ccf.org
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References

Finding malignant change in a necrotic choroidal melanocytoma: a clinical challenge
Melanocytic intraocular tumours can grow as to exceed their vascular supply, become necrotic, and induce inflammation. They present with atypical signs and become a diagnostic challenge. We present a case of a large melanocytic intraocular tumour that offered an atypical presentation, unexpected cytology, and finally diagnostic histopathology.

Case report
A 37 year old white woman presented with a painful right eye and vision loss. Examination showed no light perception, a relative afferent pupillary defect, a shallow anterior chamber, and an intraocular pressure of 58 mm Hg. Dense vitreous haemorrhage and tumour obscured her fundus.

Three dimensional ultrasonography revealed vitreous haemorrhage, a total retinal detachment, and a large choroidal mass (fig 1A, B). Computed tomography and magnetic resonance imaging (MRI) of the orbits showed a 2 cm intraocular mass with a collar-button extension (arrow) consistent with a choroidal melanoma (fig 1C).

The patient participated in a discussion of the risks and benefits of treatment, found primary enucleation unacceptable, and preferred fine needle aspiration biopsy (FNAB) before treatment.

Cytology
Transvitreal FNAB showed small, well preserved, spindle-shaped, dendritic, and epithelioid cells with evenly distributed melanin pigmentation (fig 2A) and no evidence of malignancy. Both cytology and subsequent immunohistochemistry were consistent with melanocytoma.

Though a combination of topical glaucoma medications and oral prednisone decreased her ocular pain, the patient consented to enucleation.

Macroscopic examination
A large evenly pigmented choroidal mass measuring 12 mm in height and 17x18 mm in greatest dimension was noted. The tumour extended from the ciliary body to the optic disc.

Light microscopy
Extensive infarction of the tumour was present. The cell outlines revealed a tumour composed of uniform large and heavily pigmented cells with round, small central nuclei. Many of these cells were magnocelectic and histopathologically consistent with melanocytoma. However, distinct cells with larger nuclei and prominent nucleoli were present singly and in small clusters. In these areas, rare atypical mitoses were noted. These features were diagnostic of malignant transformation of a large melanocytoma (fig 2B). At 6 months post-enucleation there is no evidence of metastatic disease.

![Figure 1](image1.jpg)

Figure 1 (A) B-scan ultrasound demonstrated a large intraocular tumour with scleral thickening and retrobulbar oedema. There was almost no intrinsic vascularity noted within the tumour. (B) A-scan ultrasonography of the right eye showed relative low internal reflectivity. (C) MRI images it displayed low signal intensity. Note the small eccentric collar-button (arrow). There was no evidence of extraocular tumour extension.

![Figure 2](image2.jpg)

Figure 2 (A) The cytological preparations from FNAB showed very small, pigmented dendritic and epithelioid cells with small dark oblong nuclei. No mitoses or necrosis was present. Some of the very deeply pigmented cells seen were melanophages. These cells were naevus-like, there were no cells that were large enough or that had appropriate nuclear morphology to be called melanoma (original magnification ×100). (B) Histopathology of the excised tumour showed a necrotic choroidal melanocytoma (left inset; original magnification ×100). Histopathology demonstrated spindle and epithelioid melanoma cells (right inset; original magnification ×100) and a melanoma with a mitotic figure in the centre (arrow). These findings suggested malignant change (original magnification ×100).
Comment
Malignant melanocytomas are darkly pigmented tumors that commonly arise at the optic nerve margin. They also occur in the iris, ciliary body, and the choroid. Though largely benign and stationary, malignant melanocytomas can (albeit rarely) undergo malignant transformation. When this occurs, it can be difficult to differentiate between melanocytomas and malignant melanomas by ultrasonography or fluorescein angiography. Adenoma and adenocarcinoma of the retinal pigment epithelium should also be considered.

In this case, FNAIB did not establish the diagnosis of a malignant melanocytoma and clearly demonstrates the potential dangers of relying on FNAIB (alone) for the diagnosis in atypical and necrotic intraocular tumors. Foci of malignant change were missed, leading to the diagnosis of melanocytoma. Fortunately, enucleation was performed.

This case also demonstrates that melanocytomas can undergo necrosis resulting in atypical features such as opaque media, pain, and inflammation. They can induce vaso-occlusive disease within the optic nerve, ischaemic necrosis of the tumour, ischaemic retinopathy, neovascular glaucoma, and melanoctytic glaucoma. Since both necrotic melanocytomas and necrotic malignant uveal melanomas present with the same acute symptoms (pain, inflammation, and an elevated intraocular pressure), these findings can detract the clinician’s ability to differentiate between these tumors.

Malignant change in a melanocytoma is a rare event. It is actually more common for a uveal melanoma to undergo spontaneous necrosis and present with intraocular inflammation, haemorrhage, secondary neovascularisation, and orbital inflammation. Such necrotising tumours have an increased incidence of penetration through the Bruch’s membrane, scleral invasion, and extrascleral extension. Clearly, large necrotic intraocular tumours can be a diagnostic challenge for both clinician and pathologist.

M Kurli, PT Finger, T Manor
New York Eye Cancer Center, New York, USA

M Kurli, S A McCormick
The New York Eye and Ear Infirmary, New York, USA

P T Finger
New York University School of Medicine, and New York Eye and Ear Infirmary, New York, USA

H E Grossniklaus
Emory University School of Medicine, Atlanta, GA, USA

Correspondence to: Paul T Finger, MD, The New York Eye Cancer Center, 115 East 61st Street, New York City, NY 10021, USA; pfinger@eyescancer.com
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References
3 Roith AM. Malignant change in melanocytomas of the uveal tract. Surv Ophthalmol 1979;23:404–12
Figure 2  Postoperative fundus photography, fluorescein angiography, and optical coherence tomography. (A) Fundus photography of the left eye exhibiting marked resolution of intraretinal haemorrhage and macular oedema, particularly in the foveal region. Arrowhead denotes site of previously complete occlusion. The course of the now ghost vessel is outlined by the white dots underneath. (B) Fluorescein angiography of the left eye at 10 seconds exhibiting noticeably decreased blocked fluorescence secondary to haemorrhage, continued hyperfluorescence of the vessel wall, and extensive but decreased capillary dropout estimated at approximately 5 disc diameters. Persistent, but less severe, vessel ‘‘pruning’’ was also evident. Arrowhead denotes site of previously complete occlusion. (C) Fluorescein angiography of the left eye at 6 minutes exhibiting decreased capillary dropout, persistent blocked fluorescence secondary to haemorrhage, and decreased leakage along capillary vessel walls. Decreased leakage is particularly noted in the foveal region. Arrowhead denotes site of occlusion. The course of the occluded vein is outlined in this angiogram by the underlying white dots. (D) Optical coherence tomography of the left eye exhibiting markedly decreased macular oedema and the absence of cystic spaces. Foveal thickness was calculated to be 133 μm.

At the 2 month postoperative visit, the patient’s visual acuity was 20/40. Fundus photography and fluorescein angiography revealed markedly decreased intraretinal haemorrhages and good macular perfusion, respectively. Macular thickness decreased from 512 μm to 133 μm on optical coherence tomography (fig 2). No cataract progression was noted and intraocular pressure remained stable throughout the postoperative course. The patient has remained stable for over 12 months.

Comment

Our group has previously published a case report of 25 gauge AAS with vitrectomy revealed markedly decreased intraretinal haemorrhages and good macular perfusion, respectively. Macular thickness decreased from 512 μm to 133 μm on optical coherence tomography (fig 2). No cataract progression was noted and intraocular pressure remained stable throughout the postoperative course. The patient has remained stable for over 12 months.

References


“Ecstasy” induced immunosuppression and herpes zoster ophthalmicus

It is uncommon for younger patients to develop herpes zoster ophthalmicus (HZO) without underlying immunocompromise. 3,4-Methylenedioxymethamphetamine (MDMA, “ecstasy”) is a widely abused psychomotor stimulant shown to cause a transient immunosuppression.1,4 In this report, we present the relation between MDMA induced immune dysfunction and the development of HZO in a previously healthy young male.

Case report

A 24 year old African-American male without significant past medical history presented with an eruptive, vesicular rash on his left forehead and eyelid consistent with the diagnosis of herpes zoster ophthalmicus (fig 1). Although he reported using safe sexual practice and denied injecting drug use, he admitted using an oral drug, ecstasy, three times a day for 4 days before the development of his symptoms. He tested negative for HIV and underlying malignancy, and slowly improved on intravenous aciclovir during his admission.

Varicella zoster reactivation is a potential complication of immunosuppression. Presdisposing factors to reactivation include increasing age, neoplastic diseases, HIV, immunosuppressive therapy, or debilitating systemic disease. Since zoster in patients under 50 years old is uncommon, a search for predisposing factors is indicated. It has been shown, in short term studies, that ecstasy use results in immune suppression.4 Therefore, we believe that our patient’s prolonged use of ecstasy facilitated the development of HZO.

3,4-Methylenedioxymethamphetamine is a widely abused psychomotor stimulant with behavioural effects similar to those elicited by amphetamine and hallucinogens. MDMA has become a popular drug of abuse over the past two decades. A study in 1988 found that 39% of students on a college campus admitted to having taken the drug at least once in the previous year. Surveys have also shown a steady increase in MDMA use in 8th, 10th, and 12th graders.5,6 MDMA produces a profound emotional warmth, and self acceptance. Cardiac arrhythmias, hepatotoxicity, and many psychiatric disorders have been associated with MDMA use.7,8 Therefore, we believe that our patient’s prolonged use of ecstasy facilitated the development of HZO.

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provided evidence that a single administration of MDMA induces a rapid and sustained suppression of induced lymphocyte proliferation and a decrease in circulating lymphocytes. The alterations in immune function were accompanied by a significant increase in plasma corticosterone concentrations. They hypothesised that users of MDMA may have reduced immunocompetence and therefore display increased susceptibility to both infectious and neoplastic disease.

In a study by Pacifici et al. it was shown that MDMA administered to humans at doses compatible with its recreational use caused rapid changes in certain immunological parameters. They demonstrated that 1 hour after administration, there was a significant reduction in CD4 T cell count and an increase in natural killer cell count.

Consequently, MDMA administration produces a rapid and sustained suppression of immune function in the rat (ImmunoPharmacology: 1998;38:253–60).

References


Reversible posterior leukoencephalopathy syndrome: a cause of temporary cortical blindness

Reversible posterior leukoencephalopathy (RPLS) describes a syndrome of headaches, confusion, seizures, and visual disturbances associated with transient characteristic lesions on neuroimaging, predominantly affecting the posterior region of the brain. RPLS affects patients with hypertension, eclampsia, renal failure, and those on immunosuppressants and chemotherapeutic agents. We discuss a case of RPLS that presented primarily with visual symptoms.

Case report

A 10 year old boy with B cell acute lymphoblastic leukaemia affecting his maxilla was commenced on chemotherapy with intravenous vincristine and cyclophosphamide, oral prednisolone and folic acid, together with intrathecal methotrexate, cytarabine, and hydrocortisone for central nervous system extension.

Ten days after the onset of chemotherapy he complained of sudden bilateral visual loss. On examination his visual acuity was perception of light in both eyes. His pupils were equal and reactive to light and fundoscopy was normal; there was no other neurological deficit. His blood pressure was elevated at 150/102 mm Hg. Over the next 24 hours his speech and conscious state became disturbed, his Glasgow coma scale (GCS) score dropped to 7/15 and he developed increased tone and clonus on his left side. A computed tomography (CT) scan showed symmetrical areas of low density involving both occipital and parietal regions associated with mild swelling. A magnetic resonance imaging (MRI) scan, 24 hours after the onset of symptoms, revealed extensive high signal intensity changes on T2 weighted sequence involving the white matter of the occipital and parietal regions (fig 1). Cerebral arteriography revealed a normal intracranial circulation.

Nifedipine was prescribed to control the patient’s mean arterial blood pressure to approximately 90 mm Hg. The chemotherapy regimen was not changed because he was between courses. His GCS score and speech improved. Two weeks later, his visual acuity had improved to 6/12 in both eyes. A repeat MRI scan, 23 days after the first study, showed a dramatic improvement with slight residual signal change in the parieto-occipital region on T2 weighted images (fig 2). Nine months after the onset of symptoms, and having completed his chemotherapy, he maintains a visual acuity of 6/6 in the right eye and 6/9 in the left eye, with full visual fields and normal colour vision.

Comment

RPLS is characterised by altered mental status, headache, seizures, and visual disturbance, often occurring in patients receiving chemotherapy. Seventy six per cent of children with RPLS have at least three of these four signs. The most common visual abnormality is cortical blindness but homonymous hemianopia, visual neglect, and blurred vision can also occur. Patients are commonly hypertensive at presentation, but this may be mild.

As illustrated, areas of temporarily increased signal intensity (representing white matter oedema) in the posterior regions of the cerebral hemispheres on T2 weighted MRI sequences are characteristic. Sparing of the calcarine fissure and paramedian occipital lobe structures aids in the differentiation of RPLS from bilateral infarction of the posterior cerebral artery territory. Although the parieto-occipital region is principally affected, the syndrome may affect the frontal lobes, basal ganglia, brainstem, or cerebellum.

While not completely understood, the cerebral oedema is thought to result from increased vascular permeability and cerebral hyperperfusion as a result of a combination of autoregulatory failure in the cerebral vessels and hypertension. This patient was treated with methotrexate and cytarabine.
both of which have been reported to cause RPLS. These drugs may have a direct cytotoxic effect on vascular endothelial cells, may induce and exacerbate hypertension and may lower seizure threshold. Intrathecal chemotherapy may cause cerebral vasospasm, contributing to cerebral vascular autoregulation impairment. The parieto-occipital region may be preferentially involved because of less extensive sympathetic innervation of the posterior cerebral circulation compared to other intracranial arteries, which reduces the ability of an already impaired cerebral autoregulation to compensate. This case illustrates that the neurological impairment from RPLS is substantially reversible with prompt treatment of the associated hypertension and the dose reduction or discontinuation of inciting drugs. It highlights the importance of the recognition of RPLS and the prompt start of appropriate treatment to prevent irreversible brain injury.

**References**


**Morquio syndrome: electron microscopic findings**

Mucopolysaccharidoses (MPS) are a group of hereditary storage diseases secondary to deficiencies of lysosomal enzymes. MPS type IV is known as Morquio syndrome. In “classic” or type A Morquio syndrome the deficient enzyme is N-acetylgalactosamine-6-sulphate-sulphatase. Morquio syndrome has been associated with cataract, optic atrophy, tapetoretinal pigmentary degeneration, and corneal clouding. We report a case of type A Morquio syndrome with electron microscopic findings.

**Case report**

A 58 year old man was referred for consideration of left corneal grafting for progressive corneal hazing of several years’ duration. He was diagnosed with Morquio syndrome in childhood on the basis of severe growth retardation, skeletal dysplasia, and facial dysmorphism. Subsequently, a diagnosis of type A Morquio syndrome was confirmed on enzyme assay of skin fibroblasts. On examination visual acuities were 6/12 right eye and 6/18 left eye. There was bilateral corneal oedema and stromal haze, greater on the left, precluding fundal examination. His intracocular pressures were normal and his angles, although poorly visualised, appeared open.

Following an uneventful left penetrating keratoplasty the host cornea was examined by light, transmission and scanning electron microscopy.

On light microscopy, the basal cells of the epithelium of the epithelium were swollen as a result of oedema but both the epithelium and Bowman’s membrane were of uniform thickness. In paraffin sections the spaces around the keratocytes were enlarged, but this was because of fixation artefact and this abnormality was not seen in the toluidine blue sections. The stroma around the enlarged spaces contained a thin dense border which stained with Alcian blue but not with colloidal iron/periodic acid Schiff or mucicarmine. Descemet’s membrane and the endothelial monolayer were of uniform thickness. In the toluidine blue sections the stroma appeared hypercellular and had a disorderly lamellar pattern. Both the keratocytes and the endothelium contained numerous intraepithelial bodies.

At the ultrastructural level, the epithelium contained abnormal inclusion bodies. In areas of the epithelium, the apical portion of the basal cells was packed with small clear membrane-bound vacuoles which decreased in numbers in the wing cell and superficial cell layers. In other sectors the epithelium appeared normal. There were sectors where the basement membrane was markedly thickened and Bowman’s membrane was discontinuous in small areas where fibrocytic and/or inflammatory cells were invading beneath the basal layer. Numerous small unmyelinated nerves were distributed throughout the basal cell layer.

![Figure 1](http://www.bjophthalmol.com)

**Figure 1** (A) Within the cytoplasm an irregular electron dense mass (arrowhead) is surrounded by concentric membranes (arrow). A granular deposit is present in the stroma around the cell membrane (*A* × 120 000). (B) Normal keratocytes were absent from the site of the cytoplasm of each cell contained inclusions although the content was variable. Some cells were disrupted by the volume of the inclusions (*A*); many of the cells were surrounded by a layer of electron dense granular material (arrowheads). The corneal lamellae were distorted by the swollen cells (* × 3800).*
Every keratocyte, to a varying degree, appeared to be involved in the overproduction of mucopolysaccharide and glycolipids. Intracytoplasmic inclusions in the form of multilamellar bodies, fingerprint whorl patterns, fibrillar granular inclusions, small lipid vacuoles, and clear vacuoles were all observed (figs 1 and 2). The collagen bundles in the stromal lamellae appeared normally aligned but scattered throughout the stroma were distended spaces or lacunae occupied by disrupted keratocytes and these distorted the lamellae. In the stromal lamellae around the keratocyte lacunae, there was deposition of an encircling layer of granular material and bundles of wide banded collagen fibres were present in the surrounding corneal stroma (fig 2).

The endothelial cell cytoplasm contained similar abnormalities to those described in the keratocytes with small clear membrane bound vacuoles which had fused to form large empty cytoplasmic spaces causing the cell membranes to collapse. By scanning electron microscopy the endothelial cells possessed a cobblestone appearance because of numerous small nodular bulges on the apical surface but the majority were of normal size and shape and the hexagonal integrity was maintained. Small linear groups of degenerate cells with cytoplasmic disruption were scattered throughout the monolayer. The graft remained clear with no evidence of recurrence 2 years postoperatively with a best corrected visual acuity of 6/6.

**Comment**

This patient had corneal opacification but none of the other common ocular associations of Morquio syndrome. As with the two previous studies which investigated the corneal opacification type A Morquio syndrome with electron microscopy, we found the most obvious abnormality to be mucopolysaccharide inclusions in the form of intracytoplasmic, multilamellar concentric bodies particularly within keratocytes but also affecting the epithelium and endothelium. The degree of keratocyte disruption was striking and possibly related to the late stage of the disease in this case. We confirmed the previous findings of bundles of abnormal collagen fibres, which were present in the corneal stroma around the lacunae and of areas of epithelial membrane bound vacuoles.

We identified areas of basement membrane thickening and areas where Bowman’s layer was discontinuous with fibrocytic and/or inflammatory cells invading beneath the basal layer. Numerous small unmyelinated nerves were distributed throughout the basal cell layer. These epithelial changes have not been commented on previously and are possibly secondary to the epithelial oedema which had been identified in this case by light microscopy.

Although early recurrence of opacification has been reported the corneal graft in this case remains clear at the 2 year follow up.

**References**


**Importance of early morning intraocular pressure recording for measurement of diurnal variation of intraocular pressure**

Intraocular pressure (IOP) is subject to cyclic fluctuations through the day. Diurnal variation in glaucoma was first reported in 1898. Duke-Elder and others reported high IOP on awakening. There is therefore a chance of missing a pressure elevation with single readings.

Phasing is mainly carried out from 0900 to 1800 hours, thereby missing any early morning spikes of IOP. Any delay in the first measurement will miss variations in IOP that are present immediately on awakening.

We carried out a retrospective study, where patients under the care of the senior author (CL) are routinely admitted for phasing in order to obtain an early morning recording of IOP immediately on awakening.

**Report**

A total of 93 patients (mean age 73.2 years) were admitted for phasing under the care of the senior author between December 1997 and August 2002. The first measurement of IOP took place between 0800–1000 hours. Further measurements were taken every 2 hours until 2000 hours. The first measurement was taken early morning (between 0600–0800 hours) immediately on awakening. Goldmann applanation tonometry was carried out by experienced staff.

In all, 17 patients requiring treatment or change in management for their glaucoma were identified. Of these, 13 were identified as having POAG. Four patients out of 13 (30.8%) showed peak IOP (23–28 mm Hg, including one patient with a best corrected visual acuity of 6/6).

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**Correspondence to:** T Leslie, Department of Ophthalmology, Grampian University Teaching Hospitals, Aberdeen AB25 2ZD, UK; tleslie@grampian.scot.nhs.uk

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Based on our study we believe that more cases of glaucoma could be identified by early morning measurement of IOP during phasing.

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P P Syam, I Movrikakis, C Liu
Sussex Eye Hospital, Eastern Road, Brighton BN2 8BF, UK
Correspondence to: Me C S C Liu, Sussex Eye Hospital, Eastern Road, Brighton BN2 8BF, UK; CS Liu@aol.com
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References


Autoimmune retinopathy associated with intravesical BCG therapy

Intravesical BCG is used as adjunctive immunotherapy for superficial carcinomas of the urinary bladder. Endophthalmitis and uveitis are the reported ocular complications.1–3 We report an unusual case of autoimmune retinopathy in a 58 year old man treated with intravesical BCG. The clinical features resembled CAR (cancer associated retinopathy) but there was no serum reaction to the CAR autoantigen.1 We believe that this is the first reported case of autoimmune retinopathy caused by BCG treatment.

Case report

A 58 year old white man was referred with reduced vision and photophobia for 4 months following a 6 week (one instillation per week) course of intravesical BCG immunotherapy for recurrent transitional cell bladder carcinoma. On examination, after 4 months of complaints, the visual acuity was 6/24 in each eye and N18 for near. Anterior segment examination revealed bilateral early nuclear cataracts. The anterior chamber and vitreous were quiet. The fundus examination showed mildly attenuated arterioles (RE>LE) with a mild pigmentary disturbance in the mid-periphery. Intraocular pressure was normal. Colour vision showed a protan-deutan axis in both eyes. There was no family history of colour blindness. Full field electroretinogram (ERG) amplitudes were subnormal for photopic and scotopic conditions in both eyes. Latency was normal. An electrooculogram (EOG) was borderline normal in both eyes. Automated perimeter revealed bilaterally enlarged blind spots. Fundus fluorescent angiography showed narrowed arterioles. The choroid was normal. Serum analysis revealed no reaction to 23 kD CAR autoantigen.1 However, an unusual reaction which may represent a form of autoimmune degeneration was reported by the laboratory.

The photophobia increased over the next 2 months. A repeat full field ERG revealed that photopic responses were indistinguishable from noise in the right eye and just detectable in the left eye. On examination visual acuity in each eye was 6/24. Oral prednisolone 20 mg daily was commenced. After a year it was decided to stop it as there was no improvement in vision. On his last follow up visit, the visual acuity was counting fingers in each eye and N36 at 10 cm. This had remained stable for 3 years. The photophobia was stable owing to dense cataracts and was well controlled with dark glasses.

The fundus remained unchanged. The primary bladder carcinoma was unchanged and no secondaries were found.

Comment

BCG has been used for the treatment of bladder cancers for many years. Ocular toxicity caused by BCG resulting in granulomatous uveitis with vitiligo has been reported.1 The first case of endogenous endophthalmitis after use of BCG for metastatic bladder carcinoma was reported in 1988. The patient developed bilateral infiltrative retinits and vitritis. Vitrectomy revealed Mycobacterium leprae1–2.

CAR is associated with melanomas, cancers of lung, cervix, colon, bladder, prostate, and breast. Progressive visual loss, night blindness with ring scotoma, and markedly abnormal ERG findings were described. Fundus revealed mild retinal pigmentation.1 Histopathology of CAR affected eyes revealed loss of retinal receptors and atrophic changes in the retinal pigment epithelium (RPE), most marked in the macula. An abnormal state of RPE hypersensitivity contributing to retinal degeneration has been reported. Grunwald et al, Kornguth et al and Kehnert et al found antibodies to normal retina in serum of cancer patients with associated retinopathies.1 One key autoantigen was identified as a 23 kD photoreceptor component “recoverin,” later found to be expressed by small cell carcinomas. Other CAR antigens have also been identified.1,2 Corticosteroids, gamma globulin, and plasmapheresis do help in some cases.3

Induction of autoimmune reactions within the eye by extracellular diseases brings into question the possibility of a sensitisation involving the M lviv component of the BCG vaccine. The live attenuated mycobacterium of intravesical BCG infects normal and cancerous cells.
bladder epithelial cells. The infection then initiates a cascade of immune events including a rapid influx of CD4+ T cells, cytokine release (IL-2, IFN-gamma and alpha) with an increase of gamma/delta T cells. These cells are not MHC type II restricted. We propose this is how activation of cellular immunity may lead to autoimmune phenomena in the eye.

Our case demonstrated the clinical features of an autoimmune retinopathy but the reaction to CAR antigen was absent. However, the unusual reaction reported suggests the presence of an immune mechanism which has the potential to cause a progressive type of retinopathy, the prognosis of which is unknown and which does not respond to steroids.

We may not have presented enough evidence to link the use of intravesical BCG with the ocular changes, although the temporal relation to the BCG therapy suggests that BCG is the most likely trigger of the autoimmunity. We suggest the bladder neoplasm is also a possibility. Ophthalmologists need to be aware of this rare possibility in order to improve their diagnostic yield in investigations of patients with similar clinical profiles. Patients also need to be warned of this rare but serious complication. This ocular complication in a previously normal patient necessitates regular ophthalmic screening in potential patients.

S Sharan
Save Sight Institute, University of Sydney, Department of Ophthalmology, Sydney Eye Hospital, 8 Macquarie Street, Sydney 2000, Australia

C E Thirkill
Department of Ophthalmology, UC Davis Health System, University of California, Sacramento, CA 95817, USA

J R Grigg
Save Sight Institute, University of Sydney, Department of Ophthalmology, Sydney Eye Hospital, 8 Macquarie Street, Sydney 2000, Australia

Correspondence to: Sapna Sharan, Save Sight Institute, University of Sydney, Department of Ophthalmology, Sydney Eye Hospital, 8 Macquarie Street, Sydney 2000, Australia; sapnasd@yahoo.co.uk
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References

Preoperative videotape sessions and patient satisfaction with cataract surgery
We read with interest the article by Pager.1 The study showed that a preoperative videotape session describing the experience of day stay cataract surgery resulted in a significant increase in overall satisfaction during the immediate postoperative period.

Patient satisfaction has drawn increasing attentions in all field of medicine for medical, financial, and litigation reasons. Age, types of aphasic correction, information received by patients, ocular co-morbidity, and postoperative visual acuity have been identified as important factors affecting the overall satisfaction in patients undergoing cataract surgery.2 While this study addressed the psychosomatic response in the perioperative period, it would be interesting to know how the videotape session can affect the perception of surgical outcome. It would be even more worthwhile to assess how such a videotape session can modify the patient response to unfavourable outcomes when surgical complications occur. Further information on patient satisfaction in either group during the follow up period would be relevant.

It was shown that the majority of patients could not recall relevant information after verbal consent.3 Remembering the information deteriorated significantly after the operation, even more so in those of an advanced age and with lower educational level.4 Videotape has been used in the informed consent process in other medical fields. Patients having gastrointestinal endoscopy were found to be more satisfied with videotape followed by physician discussion than either method alone.5 It has also been demonstrated to lead to higher knowledge scores, especially in patients with lower education levels.6 Since cataract patients are usually old and come with very high expectations, we think that further research is required to explore the use of videotape in order to achieve a better informed consent.

K S C Yuen, A C K Cheng, W-M Chan
The Chinese University of Hong Kong, Department of Ophthalmology and Visual Sciences, Prince of Wales Hospital, Shatin, Hong Kong, China

Correspondence to: Dr Kenneth S C Yuen, The Chinese University of Hong Kong, Department of Ophthalmology and Visual Sciences, Prince of Wales Hospital, Shatin, Hong Kong, China; ksuyen@gmail.com
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Our study is less than the percentage of normal eyes with stage 1 or 2 PVD in the Uchino study.3 This suggests that the vitreomacular configuration defined as VMHA by Sahni et al is probably not a finding specific to the AMD patients included in the study.

J Witkin, J S Duker
New England Eye Center, Tufts-New England Medical Center, Tufts University, Boston, MA, USA

Correspondence to: J S Duker, New England Eye Center, Tufts-New England Medical Center, Tufts University, Boston, MA, USA; jsd@tufts.nemc.org
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Optical coherence tomography of the vitreomacular interface in photodynamic therapy

We would like to comment on the excellent article by Sahni et al.1 In their paper, a number of descriptive terms for optical coherence tomography (OCT) analysis are defined in patients with age related macular degeneration (AMD) with subfoveal choroidal neovascularisation undergoing treatment with photodynamic therapy. One term, “vitreomacular hyaloid attachment (VMHA),” was used to refer to incomplete separation of the posterior hyaloid from the macula. Twenty of 56 patients (35.7%) included in the study had VMHA on OCT. We would like to point out that this vitreomacular configuration is identical to that described in a previous report of normal eyes.2

The study by Uchino et al reported OCT findings at the vitreoretinal interface in 209 normal eyes. In their study, they defined five individual stages of posterior vitreous detachment (PVD). Two stages represented partial PVD with persistent macular attachment, identical to VMHA as defined by Sahni et al. Stage 1 was defined as focal perifoveal PVD in one to three quadrants with persistent vitreofusal attachment, and stage 2 was defined as perifoveal PVD in all four quadrants with persistent vitreofusal attachment. Of the 209 normal eyes, 47.8% had stage 1 PVD and 12.6% had stage 2 PVD. Mean age of the patients in the study was 52.3 years (range 31–74 years).2

We find it interesting that the percentage of AMD patients with VMHA in the Sahni study is less than the percentage of normal eyes with stage 1 or 2 PVD in the Uchino study.3 This suggests that the vitreomacular configuration defined as VMHA by Sahni et al is probably not a finding specific to the AMD patients included in the study.

Author’s reply

We thank Mennel et al and Duker and Witkin for their interesting comments regarding our article.

Mennel et al make some interesting points on the immediate structural changes that occur after photodynamic therapy (PDT), a topic that we thought was outside the scope of our study. I agree that the short term and long term changes after treatment are important and need to be taken into account in future studies on patients undergoing PDT.

We are grateful to Duker and Witkin for pointing out the article by Uchino and colleagues.1 Our patients were older (mean 76 years (range 59–94)) than those reported by these authors (mean 52.3 years). Uchino et al had 20 patients in the same age group and only two patients were above 70 years of age. Both these had a complete posterior vitreous detachment (PVD). All our patients had associated pathology and the majority had undergone PDT, all of which may have influenced the outcome. While the finding of vitreomacular attachment may be more common in normal eyes (>50%), our study7 suggests that the incidence may be lower in patients with exudative age related macular degeneration (35.7%).

NOTICES

EVE 2005 meeting

This will take place on 5-8 October 2005 in Vilamoura, Portugal. For further details please contact: Christy Lacroix, EVER Secretary, Kapucijnenvoer 33, B-3000 Leuven, Belgium (tel: +32 (0)16 233 849; fax: +32 (0)16 234 097; email: ever@skynet.be).

World Ophthalmology Congress 2006 – Brazil

The World Ophthalmology Congress (which is replacing the International Congress of Ophthalmology) is meeting in February 2006 in Brazil.

For further information on the congress and committees, scientific program and coordinators of different areas are available at the congress website www.ophthalmology2006.com.br

Red eye

The latest issue of Community Eye Health (No 53) discusses the role of primary care in the treatment of red eye. For further information please contact: Journal of Community Eye Health, International Resource Centre, International Centre for Eye Health, Department of Infectious and Tropical Diseases, London School of Hygiene and Tropical Medicine, Keppel Street, London WC1E 7HT, UK (tel: +44 (0)20 7612 7964; email: Anita.Shah@lshtm.ac.uk; online edition: www.jchef.co.uk). Annual subscription (4 issues) UK £28/US$45. Free to developing country applicants.