Solitary CD30+ anaplastic large cell lymphoma of the eyelid showing regression

CD30+ anaplastic large cell lymphoma (ALCL) belongs to the group of T cell non-Hodgkin’s lymphomas. The primary cutaneous variant of ALCL usually presents as a solitary, cutaneous, or subcutaneous reddish violet lesion, which can be superficially ulcerated. We present the case of a solitary CD30+ ALCL of the eyelid showing regression.

Case report
A 39 year old man presented with a 4 week history of a progressive painless ulcerating nodule on the right upper eyelid, unresponsive to oral flucloxacillin. He was systemically well and denied recent foreign travel or contact with animals.

A 17 mm diameter ulcer with rolled margins and serosanguinous exudate was evident over the right upper eyelid (fig 1). His cornea, conjunctiva, and anterior chamber were normal. Systemic examination was unremarkable.

Investigations including full blood count, urea and electrolytes, bone and liver profile, immunoglobulins and electrophoresis, autoantibody screening, and Treponema antibody were normal or negative. Tissue culture failed to demonstrate a bacterial, viral, or fungal pathogen. There was no clinical, radiological, or bone marrow evidence of extracutaneous disease.

Histology of the biopsy taken from the lid ulcer margin showed epidermal necrosis associated with ulceration (fig 2). The ulcer base showed haemorrhagic granulation tissue infiltrated by a mixture of lymphocytes, plasma cells, neutrophils, and eosinophils. There were also ill defined groups of large blast cells showing enlarged and pleomorphic nuclei and high mitotic activity. The immunohistochemical staining showed these cells to be of T cell lymphoid lineage. Many of the large blast cells were CD30 positive but negative for ALK-1 protein. The features were of a CD30 positive anaplastic large cell lymphoma (ALK negative).

Treatment options such as surgery and radiotherapy were discussed with the patient but as the lesion remained stable over a 10 day period, a conservative approach was agreed. A moderately potent topical corticosteroid (mometasone furoate 0.1% cream) was applied to the lesion once daily. When followed up 8 weeks later the ulcer had completely healed without scarring (fig 3).

Eighteen months has elapsed since presentation. There has been no recurrence of his disease, and he remains in good health.

Comment
ALCL represents a group of large cell lymphomas. They consist of a proliferation of predominantly large lymphoid cells with strong expression of the cytokine receptor CD30 (>75%). Using molecular and clinical criteria, three entities have been identified: primary systemic anaplastic lymphoma kinase (ALK) + ALCL, primary systemic ALK – ALCL, and primary cutaneous ALCL.1

Primary cutaneous ALCL arise de novo in the skin, commonly on the head and neck of older patients with a median age of 60 years and a male/female ratio of 3:2. Most patients present with solitary, asymptomatic nodules, which can be superficially ulcerated. Primary cutaneous ALCL has a more favourable prognosis than systemic ALCL, with an 5 year survival of approximately 90%. Partial or complete spontaneous regression can be observed in up to 25% of patients with primary cutaneous ALCL, accounting for the previous designation of “regressing atypical histiocytosis.”2 Treatment of localised lesions usually includes excision with or without radiation. However, patients with disseminated skin disease may benefit from systemic polychemotherapy.3 In our patient the lesion had resolved within 3 months of initial appearance. The application of a moderate potent topical steroid might have contributed to the regression of the ulcer.

We present a case of a primary cutaneous ALCL of the eyelid showing regression. Ophthalmologists should be aware of this sometimes self regressing entity and an expectant policy might be indicated in non-progressing tumours, thus avoiding potentially mutilating surgery or radiotherapy.

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References
Choroidal translocation with a pedicle following excision of a type 1 choroidal neovascular membrane

Excision of type 1 choroidal neovascular membranes (CNVM) in age related macular degeneration (AMD) have a poor visual outcome because of loss of retinal pigment epithelium (RPE). Simple replacement of the RPE may not work because the relation with Bruch’s membrane and choroidal complex is disturbed. Creating a free graft of these three layers detaches the choroid from its blood supply. Hence, restoration of these three layers to the subfoveal position while maintaining a connection to the adjacent choroidal blood supply is desirable.

Case report

A 74 year old woman presented with a 3 month history of a left central scotoma and visual acuity (VA) of counting fingers (CF). Clinical examination and fluorescein angiography (FFA) confirmed a type 1 subfoveal CNVM. The fellow eye was 20/30 with scattered soft drusen. Pars plana vitrectomy (PPV) and excision of the CNVM were performed as described previously. Atrophic choroidal vessels underlying the CNVM were not removed. A retinotomy was formed temporal to the fovea and vertical scissors inserted into the subretinal space. The RPE, Bruch’s membrane, and choroid were incised en bloc in the area temporal to the site of the CNVM to create a graft on a pedicle. The graft was manipulated to a subfoveal position. The pedicle and graft were equally sized to maximally exploit the rich choroidal vasculature and maintain continuity to the choroidal circulation. We were unable to unpredictable the position of choroidal vessels as indocyanine green angiography (ICG) was unavailable to us at the time of surgery. Surprisingly, little bleeding occurred and was easily controlled by increasing the infusion height. The patient required two subsequent operations for a rhegmatogenous retinal detachment with grade B proliferative vitreoretinopathy. The retina was flattened after inferior retinectomy and silicone oil insertion.

At review 4 years following initial surgery her vision was CF with a central scotoma on Goldmann field testing. The area of translocated RPE, Bruch’s membrane, and choroid was visible beneath the fovea with bare sclera demarcating its original site (fig 1A). At 4 years following surgery there was no recurrence of the CNVM on FFA (fig 1B) and ICG angiography demonstrated that the graft and pedicle were vascularised (fig 2).

Comment

Excision of type 1 CNVMs has a poor prognosis because of loss of RPE and atrophy of the choroid. Restoration of the normal anatomical relation between the retinal receptors and the underlying structures is essential for visual recovery. Retinal translocation with strabismic surgery for the movement of the retina to healthy RPE is prolonged and hazardous. Transplantation of homologous RPE cells alone to a subfoveal position has met with varied success. Aylward et al reported no visual improvement after transplantation of an autologous free graft, with fibrosis of the grafts at 10 months, perhaps because of loss of blood supply. Late revascularisation of some grafts has been reported at 1 year.

There was no visual improvement in our patient as she had a retinal detachment and additional procedures. We thought that the rich and redundant blood supply of the choroid allowed some freedom in the choice of graft harvest site. As proof of principle we have demonstrated that a choroidal/RPE graft with a pedicle is a feasible surgical technique, resulting in a sustained and vascularised graft. This technique is simpler than time consuming retinal translocation and does therefore merit further investigation.

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Intravitreal triamcinolone acetonide and central serous chorioretinopathy

Intravitreal injections of triamcinolone acetonide have increasingly been performed as treatment for intraocular diseases with intraretinal oedema and with subfoveal fluid accumulation, such as diffuse diabetic macular oedema, persistent pseudophakic cystoid macular oedema, central retinal vein occlusion, and exudative age related macular degeneration. In view of the widening spectrum of indications for intravitreal triamcinolone acetonide injections, it was the purpose of this study to evaluate whether intravitreal triamcinolone acetonide injections may be useful as treatment of longstanding central serous chorioretinopathy.

Case report

A 50 year old patient presented with a decrease in visual acuity to 1/20 in his right eye because of longstanding central serous chorioretinopathy. Six years earlier, visual acuity had started to deteriorate, and had remained at 1/20 for the past 2 years. Fluorescein angiogram showed a mottled appearance of the retinal pigment epithelium close to the fovea, and a leakage of dye in the late phase of the angiogram. There was...
no clear smoke stalk phenomenon (fig 1). In optical coherence tomography, the central retina was detached. Despite intensive topical treatment with prednisolone acetate eye drops and oral intake of carbonic anhydrase inhibitors, the morphological appearance of the fovea and visual acuity remained unchanged. Under topical anaesthesia, the patient received an intravitreal application of 20–25 mg of triamcinolone acetonide, which was transconjunctivally injected through the pars plana into the centre of the vitreous cavity. The technique has already been described in detail. The patient was fully informed about the experimental character of the treatment and had signed an informed consent. After the injection, all topical and systemic medication for his macular disorder was stopped.

Within the first 5 months after the injection, fluorescein angiograms and optical coherent tomograms did not show any marked changes in the macula (fig 2). Correspondingly, visual acuity remained at 1/20. Intraocular pressure increased up to levels of 30 mm Hg and was reduced to the normal values by topical application of a carbonic anhydride inhibitor. Thirteen months after the injection, the fovea was still slightly detached. Visual acuity remained at 1/20.

The clinical course suggests that in this eye with longstanding central serous chorioretinopathy an intravitreal injection of a high dosage of triamcinolone acetonide was not accompanied by a fast resolution of the subfoveal fluid and an increase in visual acuity. For more than 5 months after the injection, the fovea remained clearly detached.

Vascular occlusion in serpiginous choridopathy

Serpiginous choridopathy (SC) is a rare disease inducing a permanent loss of vision, caused by a progressive destruction of the retinal pigment epithelium and chorioaipcillaris. Until now, no antigens or predisposing factors have been reported. SC usually affects both eyes and occurs in patients between the fourth and sixth decade; without any sex or race predilection. Clinically, deep cream-coloured lesions develop in the peripapillary region and then along the retinal vessels, centrifugally, inducing an atrophy of the retina. Other lesions may develop, isolated, in the posterior segment. The anterior segment is typically quiet; nevertheless, a mild anterior uveitis and/or vitritis have been observed. The course of the disease results in successive attacks and recurrences inducing permanent retinal atrophic changes and subsequently an irreversible loss of vision. Choroidal neovascularisation may occasionally develop. No specific diagnostic tests are available such that the diagnosis of SC is mostly clinical.

Case report

A 30 year old Indian man presented with a history of painless progressive visual loss affecting the right eye. No other ophthalmological or systemic complaints were present. His medical history was unremarkable.

Ophthalmological examination revealed a visual acuity of 20/50 in the right eye and 20/20 in the left eye without a correction in both eyes. Anterior segment examination revealed a mild inflammation with fine keratic precipitates on the inferior part of the right corneal endothelium. Intra-ocular pressure was 10 mm Hg in both eyes. Fundus examination of the right eye disclosed a moderate vitreous inflammation (cells: ++ ) and multiple deep cup choroidal lesions around the optic disc and along the superior and inferior retinal (temporal and nasal) vessels (fig 1). Our differential diagnosis was a white dot syndrome (APMPPE, SC), an infection (tuberculosis), or a sarcoidosis. We decided to hospitalise the patient.

A clinical examination revealed an erythrocyte sedimentation rate of 8 mm in the first hour (normal range 1–12), and a normal white blood count. Immunoglobulin electrophoresis, quantitative immunoglobulin levels, CD4+CD8 lymphocytes count, C3, C4 and CH50 examination were within the normal range. Tests for connective tissue disorders were negative and serum angiotensin converting enzyme was discretely elevated (74 U/L, normal range = 18–55) with a normal lysozyme level. Infectious serologies (toxoplasmosis, Borrelia burgdorferi, Treponema pallidum, HIV, herpes virus, Leptospira, Bartonella, rickettsiosis, brucellosis) were within the normal limits. An anterior chamber tap (polymerase chain reaction for herpes simplex virus (HSV) 1, HSV2, varicella zoster virus, cytomegatovirus, Epstein-Barr virus, toxoplasmosis, Mycobacterium tuberculosis) was negative. A lumbar puncture was normal (proteins 0.31 g/l, white cells 3 × 10^9/l, lymphocytes 74%), without oligoclonal bands on electrophoresis. PPD skin test was positive (15 mm) but chest x ray was normal. We have to consider that the patient has had a BCG vaccine in his childhood. The patient was HLA-B27 and A-29 negative but HLA B-7 positive. The initial clinical examination was completed by a neurological and a dermatological examination which were normal. A magnetic resonance image cerebral scan was normal.

The patient was given a course of methylprednisolone intravenously (4 × 250 mg/day for 3 days) followed by oral prednisone (1 mg/kg) at tapering doses, and aciclovir (3 × 10 mg/kg), intravenously for 10 days. We covered the patient with rifampicin, isoniazide, ethambutol, pyrazinamide, and B6 vitamin. Topical steroids and mydriatics were covered the patient with rifampicin, isoniazide, ethambutol, pyrazinamide, and B6 vitamin. Topical steroids and mydriatics were covered the patient with rifampicin, isoniazide, ethambutol, pyrazinamide, and B6 vitamin.
A regression of the inflammation in the right eye was noted as well as a “cicatrisation” of the choroidal lesions, which appeared as multiple geographical areas of atrophy of both the retina and pigmented epithelium between areas of normal retina. Our suspected diagnosis was a SC.

After 3 weeks, the patient developed the same lesions in the left eye with an occlusion of the superior temporal vein (fig 2). At that time the patient was on prednisone 40 mg/day and anti-TB treatment. A complete clinical examination was done again, but still all results were within the normal limits. The same treatment was introduced (methylprednisolone, intravenous aciclovir).

As the relapse occurred under steroid therapy (prednisone 40 mg/day), the administration of a monospecific anti-inflammatory drug was discussed. The patient was given mycopheneolate mofetyl (Cellcept, 2 g/day) and oral prednisone for 1 year, at tapering doses. Anti-TB treatment was continued too. No secondary effects were noted.

We followed the patient during 12 months; his visual acuity returned to 20/20, without a correction, in both eyes. The anterior segment was normal. The posterior segment examination disclosed permanent geographic chorioternal atrophic lesions along the vessels in both eyes, confirming the diagnosis of SC.

Comment
SC induces a progressive loss of the retinal pigment epithelium and choriocapillaris. The cause of this disorder is still under investigation but some studies suggest that an inflammatory or a vascular factor are involved in the pathogenesis. Histopathological studies have shown the presence of extensive lymphocyte choroidal infiltrates but in eyes chronically affected, the distribution of the lesions and their angiographic features (fluorescein and green indocyanine) may suggest a choroidal occlusion. Genetic studies demonstrated an increased frequency of HLA-B7 in this affection, our patient was positive. SC affects not only white people but also oriental and blacks people. There is no sex predilection and the patients are middle aged when the diagnosis is made. The patient described in this report was 30 year old which is uncharacteristic. The aetiology remains unknown although recently tubercular serpinious-like choroiditis has been reported.

Clinically, inflammatory signs may be noted both in segments (anterior uveitis, vitritis) along with the classic whitish choroidal lesions in the acute stage. The most frequent ocular complication of SC is sub-retinal neovascularisation which affects 13% to 20% of the eyes. Retinal vasculitis is also observed.

Gupta et al reported a case of SC with a branch vein occlusion in the acute phase. Our patient developed a vein occlusion while under treatment (steroids, aciclovir, and anti-TB therapy) which is rare and was never reported to our knowledge. Haemorrhages are sometimes observed in inflammatory diseases (Belchet’s syndrome, sarcoidosis) or in infectious posterior uveitis (syphils, viral infections, toxoplasmosis).

This case demonstrates that SC can affect young patients and that HLA-B7 can be found in Indian patients with SC. The clinical features (vascular occlusion in this case) and the development of new lesions while under treatment let us suspect that the cause of this disease is still not clear.

References

Figure 2 Fundus of the left eye. Presence of multiple deep creamy choroidal lesions along the retinal vessels with a vascular occlusion along the superior temporal vein. The patient was treated with steroids, aciclovir, and anti-tuberculosis treatment.

Persistent acanthamoeba keratitis in a non-contact lens wearer following exposure to bird seed dust
Acanthamoeba keratitis is a serious and vision threatening disease. It is commonly associated with contact lens wear (up to 93%). Early diagnosis and treatment are essential to improve the visual outcome. Devastating ocular damage can be attributed to various factors such as misdiagnosis, incorrect treatment, excessive topical steroid before diagnosis, and resistance.

Acanthamoeba keratitis in non-contact lens wearers is rare and poses a diagnostic challenge. We present a case of acanthamoeba keratitis in a non-contact lens wearer following accidental exposure to bird seed dust. The strain of acanthamoeba obtained from this patient appeared to show in vitro resistance to polyhexamethylene biguanide, PHMB and chlorhexidine after a good clinical response initially.

Case report
A 57 year male patient presented with pain, blurring of vision, and photophobia of his left eye. Two weeks before the presentation he had an accidental exposure to bird seed dust (brand name Trill, manufactured by Master Foods, Hungary) for his budgies while cleaning the seed pot. It was a seemingly trivial injury not likely to have caused a breach of epithelium. Examination revealed a visual acuity of 6/6 of the right eye and 6/18 for the left eye. The left eye showed multiple punctate epithelial erosions with epithelial and stromal infiltrates. There was no retained debris at the time of presentation. Initially he was treated as a case of viral keratitis with topical aciclovir and steroid. Although there was an early improvement, the keratitis relapsed after 2 weeks. At that stage a typical ring infiltrate suggestive of acanthamoeba keratitis developed and epithelial culture grew Acanthamoeba polyphaga. He was started on intensive treatment with PHMB, Brolene, and neomycin. His symptoms improved and his visual acuity recovered to 6/9 over a period of 3 weeks. Topical steroids were then added. The antimicrobial treatment was given for 2–3 months and withdrawn gradually over next 4–6 weeks after complete resolution. But following complete cessation of all drops he developed a recurrence with positive cultures. We restarted the intensive treatment with PHMB and chlorhexidine, but the clinical response was poor. Pain was severe with intense limbal inflammation and signs of sceritis. A corneal biopsy was performed which showed persistence of infection. Resistance to PHMB (minimum inhibitory concentration (MIC), 3.125 µg/ml) and chlorhexidine (MIC, 6.25 µg/ml) was demonstrated in the culture obtained from the biopsy. The strain showed in vitro sensitivity to propamidine (MIC, 3.9 µg/ml). A change of treatment to topical propamidine isethionate 0.1% (Brolene) and neomycin led to a rapid response with a decrease in symptoms. Six months after initial diagnosis he is comfortable on maintenance treatment with propamidine isethionate 0.1% and neomycin, but unfortunately he has developed a dense central corneal scar (fig 1) and vision of hand movements.

Comment
Acanthamoeba keratitis not related to contact lens wear has been reported before and risk factors include trauma, dirty water splash, and exposure to leaf juice. Exposure to feather dust has to our knowledge not been reported previously as a known risk factor. Unfortunately, an attempt to culture acanthamoeba from the actual bird seeds and tray was unsuccessful. The second
PostScript

uncommon feature in our case is the demonstration of in vitro resistance of this strain of acanthamoeba to two of the modern first line acanthamoeba drugs (PHMB and chlorhexidine) while showing a good sensitivity to propamidine. This is contrary to what has been reported by other authors.\(^a\)\(^b\) We are unable to say whether resistance developed during treatment or was pre-existent, as sensitivity profiles of the earlier isolates were not obtained. This patient's initial good clinical response was achieved with a combination of PHMB and propamidine with the latter tapered early during the course of treatment, indicating at least partial in vivo sensitivity to PHMB in the earlier stages. A poor association between in vivo and in vitro resistance has been described for bignonanides,\(^b\) but this case shows that in vitro MIC can be useful information in the management of persistent acanthamoeba keratitis.

Other authors have stressed the need for long term treatment\(^c\) and this case also underscores the importance of prolonged effective antimicrobial treatment in order to prevent recurrences.

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References


Brinzolamide induced reversible corneal decompensation

Topical carbonic anhydrase inhibitors (CAIs) such as brinzolamide 1% (Alphagan; Alcon Laboratories, Fort Worth, TX, USA) attenuate bicarbonate efflux, and this may lead to corneal oedema. To our knowledge, this is the first report of complete resolution of corneal oedema after cessation of topical brinzolamide 1%.

Case reports

A 57 year old African-American man with primary open angle glaucoma (POAG) presented with painless blurry vision left eye 1 hour after instilling brinzolamide 1% in both eyes. He had been on brinzolamide 1% both eyes twice daily, brimonidine tartrate 0.2% (Alphagan) in both eyes twice daily, and latanoprost 0.005% (Xalatan) in both eyes once at night for 2 years. On presentation, best corrected visual acuity (BCVA) was 20/25 right eye and 20/50 left eye. The left eye had mild corneal oedema. Descemet’s folds, and whitish fleck-like debris on the corneal endothelium (fig 1A). Intraocular pressures (IOPs) were 15 mm Hg and 16 mm Hg. The brinzolamide 1% in both eyes was discontinued. Timolol maleate 0.5% (Timoptic) in both eyes twice daily and topical predniso- lon acetate 1% (Pred Forte) left eye four times daily were started. By 1 week follow up, the cornea was clear (fig 1B). Specular microscopy revealed endothelial cell counts (ECC) of 1355 cells/mm\(^2\) right eye and 648 cells/mm\(^2\) left eye with enlarged pleomorphic endothelial cells left eye (fig 2). Central corneal thickness (CCT) was measured as 512 \(\mu\)m right eye and 505 \(\mu\)m left eye.

A 77 year old white man, who had had cataract extraction 46 years earlier and subsequent aphakia right eye, had been followed for open angle glaucoma in both eyes for 25 years. He was on timolol maleate 0.5% in both eyes twice daily, latanoprost 0.005% in both eyes once at night, and pilocarpine hydrochloride 4% gel (Pilopine Gel HS) in both eyes once at night. His visual acuities were hand movement right eye and counting fingers at 1 foot left eye. An IOP of 19 mm Hg right eye and 10 mm Hg left eye necessitated the addition of timolol maleate 0.5% annual rate seen in normal subjects.\(^a\)

Corneal oedema is also reached 1 hour after dosing, such as brinzolamide 1% (Azopt; Alcon Laboratories, Fort Worth, TX, USA) attenuate bicarbonate efflux, and this may lead to corneal oedema. To our knowledge, this is the first report of complete resolution of corneal oedema after cessation of topical brinzolamide 1%.

Comment

The Merck Worldwide Adverse Experience System database for dorzolamide 2% includes 25 reports of corneal oedema. Nearly all of these cases had a history of multiple ocular surgeries and compromised corneas.\(^c\) Dorzolamide is a reversible inhibitor of carbonic anhydrase II and does not accumulate in the cornea with repeat dosing,\(^a\) so any corneal effect from a similar medication should indeed be potentially reversible.\(^b\) Dorzolamide’s peak concentration in rabbit corneas is also reached 1 hour after dosing, and the half life in the cornea is approximately 2 hours.\(^b\) This is consistent with case 1 having blurry vision 1 hour after instilling brinzolamide 1%.

Patients with primary open angle glaucoma or ocular hypertension and a baseline ECC of greater than 1500 cells/mm\(^2\) have an average 3.6% decrease in ECC after a year of dorzolamide 2% three times daily.\(^b\) This endothelial loss is much higher than the 0.6% annual rate seen in normal subjects.\(^a\)

Topical CAIs can cause corneal oedema in compromised corneas—for example, those...
with cornea guttata, but the oedema is reversible if identified early. Before initiating brinzolamide 1%, an ECC may be needed for high risk patients (that is, monocular, previous ocular surgery, corneal disease, etc). Because of the potentially irreversible nature of the corneal decompensation, topical CAIs may be relatively contraindicated in patients with significant corneal disease.

Although dorzolamide and brinzolamide are both topical CAIs, their chemical formulas are different, and a side effect that is associated with dorzolamide may not necessarily be assumed to be associated with brinzolamide. It is important to be aware that brinzolamide can also potentially cause cornea oedema.

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An unusual cause of acquired horizontal diplopia in a young adult

Cysticercosis is caused by infection from the larval form of Taenia solium, is endemic to regions with poor sanitation. Human cysticercosis more commonly affects the central nervous system, with less common involvement of ocular tissues. Myocysticercosis is a subset of orbital cysticercosis and is considered a rare entity.

Case report

A 19 year old Nepalese housewife presented with left sided headache that had been present for 1 year. She had been treated for migraine headache in another hospital. Ocular examination revealed orthoptia in primary gaze with Snellen acuity of 6/6 bilaterally. Both the Humphrey visual fields and the colour vision testing were normal in both eyes. There was no relative afferent pupillary defects detected. Ocular motility testing revealed left abduction deficit with the resulting horizontal diplopia. Her tel’s exophthalmometer reading revealed no proptosis. Her fundi examination were both normal and there were no optic disc swelling.

Magnetic resonance image (MRI) (brain/orbit) with gadolinium contrast (fig 1) showed a cystic enhancing mass measuring 1.2 cm×0.6 cm adjacent to and including left medial rectus muscle, sparring the muscle tendon. Further assessment with B-scan ultrasonography (fig 2) revealed an intraocular cyst within the left medial rectus muscle located within mid-orbit.

Full blood count found no eosinophilia; systemic cysticercosis involvement was excluded by negative radiological findings (chest x-ray and computed tomography (CT) of brain and abdomen were all normal).

The patient was prescribed treatment with albendazole 15 mg/kg per day for 8 days. Unfortunately, she had intolerable side effects (nausea, vomiting, and distressing nocturnal left eye pain) to the medication which she used for only 3 days. She was reluctant to continue with albendazole. Her symptoms settled after a short course of oral analgesics.

She has remained asymptomatic. Repeated Hess and diplopia charts B-scan ultrasonograph re-evaluation at 6 months did not reveal any cysts in the muscle and her ocular motility had returned to normal.

Comment

The finding of “scolex” within the intramuscular cyst and her status of Nepalese native lend strongly to a diagnosis of myocysticercosis. Enzyme linked immunosorbent assay (ELISA) to detect the antibody to cysticercosis has been suggested to reduce the secondary inflammation.

Surgical excision of an extraocular muscle cyst had been described. However the potential risk of damage to adjacent tissue and adhesion from surgical exploration should not be taken lightly, particularly when effective medical therapy is available.

Stool tests should be done for all the members of the family to detect asymptomatic carrier because the treatment with systemic antihelmintic treatment is highly effective. It also serve to break the life cycle of the parasite.

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Use of scanning laser ophthalmoscopy in visual conversion reaction

Visual conversion reaction is a psychosomatic anomaly that manifests as reduced visual acuity (VA) and visual field defects. Scanning laser ophthalmoscopy (SLO) can detect a scotoma and VA under direct fundus observation. However, there have been no reports of patients diagnosed with visual conversion reaction using SLO. We report a patient with visual conversion reaction using SLO.

Case report

A 20-year-old woman presented with a sudden bilateral loss of vision. She reported being under severe stress at work. The best corrected visual acuity (BCVA) was counting fingers in both eyes. The external eye examination and pupillary responses were normal in both eyes. Conventional ophthalmoscopy, funduscopy, and fluorescein angiography were unremarkable. A general medical examination showed no abnormalities. There were no scotomas (based on Goldmann size III stimulus on the retina), and the stability of fixation was central and stable using SLO microperimetry (fig 1B). The VA using SLO was 20/20 in both eyes. We diagnosed visual conversion reaction in this case.

Comment

This is the first report of a patient with visual conversion reaction using SLO. In this case, the BCVA was counting fingers in both eyes at the first visit. However, the VA using SLO was 20/200 both eyes and better than the conventional examination. The visual fields were constricted in both eyes to within 5° of fixation using Goldmann perimetry (fig 1A). The results of magnetic resonance imaging, computer tomography of the brain and orbits, visual evoked potentials, and electroretinography were unremarkable. A general medical examination showed no abnormalities. There were no scotomas (based on Goldmann size III stimulus on the retina), and the stability of fixation was central and stable using SLO microperimetry in both eyes (fig 1B). The VA using SLO was 20/200 in both eyes. We diagnosed visual conversion reaction in this case.

Ten months later, the BCVA was 20/20 in both eyes. The visual fields in both eyes using Goldmann perimetry were normal (fig 2A). There were no scotomas, and the stability of fixation was central and stable in both eyes using SLO microperimetry (fig 2B). The VA using SLO was 20/20 in both eyes. We diagnosed visual conversion reaction in this case.

References

We welcome the article by Loon et al.1 Earlier, we published our finding of the SARS coronavirus in specimens collected by the novel technique of conjunctiva upper respiratory tract irrigation (CURTI), but not in paired nose and throat swabs, very early in the course of the disease.2 In designing CURTI, we considered safety to medical personnel and, also, finding a method that samples all three portals of entry for upper respiratory viral pathogens—the eyes, nose, and mouth. Loon et al’s findings complement our study by showing that SARS coronavirus can indeed be found in tears.

While we agree with their conclusion that the ability to isolate the virus early in the course is important, we do not think that the eyes are important organs that propagate the virus, other than to ophthalmologists and to unwary close contacts. For instance, the eyes cannot generate infectious aerosol. Rather, we feel that the eyes are important portals of entry and have not been given sufficient attention—witness medical personnel in full personal protection gear and N95 masks but without watertight goggles, and sometimes without splashguards.

We also think that employing the services of ophthalmologists for the purpose of collecting tear specimens for the diagnosis of SARS will fail to achieve its most medical environments. On the other hand, our method of CURTI is entirely self help, deployable in quarantine locations, and avoids unnecessary contact between an infectious source and susceptible individuals. The finding of SARS CoV in tears raises several additional questions:

(1) How does the virus end up in the tear? Was it the result of direct inoculation at the time of infection into permissive conjunctival epithelial cells, either by hand or aerosol, or was it the result of secretion from a lacrimal gland infected haematogenously? The lacrimal glands are not very different anatomically from the salivary glands. Yet saliva has been shown to be a poor specimen for the diagnostic detection of severe acute respiratory syndrome coronavirus in tears.3,4,5,6

(2) Is there any means or advantage in using lacrimitis, or evidence of infection of the lacrimal gland infected haematogenously? The lacrimal gland is not generally considered choosy, nor does the eye have a dense network of lymphatic vessels, important in virus spread. While it is true that the gland is rich in trigeminal nerve fibers, this is not unique to the eye, nor does it provide a dense lymphatic plexus.

(3) Is there any means or advantage in using any of the apparently effective methods described.


References

Comments on using fibrin glue in pterygium surgery

I read with great interest the article by Koranyi and coworkers, who evaluated a new technique for pterygium surgery using a fibrin tissue adhesive (Tissel Duo Quick).1 In their randomised trial the authors concluded that using the glue instead of sutures caused less postoperative pain and shortened the surgical time. Nevertheless, the timing of the randomisation is not clearly stated in their report.

Whether or not the surgeon knew the patient’s group (sutures or fibrin glue) at the time of pterygium removal and conjunctival graft harvesting may have influenced the extent of the removal and the size of the graft. Therefore, the differences in postoperative pain and/or recurrence could be related to those initial steps and not only to the final step, as the authors suggest. Ideally, the surgeon should be informed if the conjunctival graft should be sutured or glued after harvesting it.

Additionally, in their discussion the authors did not mention the risk of infection when using fibrin glue. Some viruses, such as parvovirus B19 (HPV B19) are particularly difficult to remove or inactivate, and human infection has been reported after the use of fibrin glue.2 In thoracic surgery, epidemiological evidence suggests that more than 20% of uninfected people were subsequently infected with HPV B19 by use of fibrin during the procedure.3 Prions are also of concern. The direct application of any of the apparently effective methods of prion decontamination to plasma products is inappropriate because they are harsh and denaturing.4 Although the risks for both diseases are minimal, the authors should have addressed this issue in their discussion.

Authors’ response

We thank Dr Alvarenga for his comment on our article about using fibrin glue for pterygium surgery.

We write in the “Material and methods” section that only the thickened and keratized portion of the conjunctiva was excised and that the graft was prepared to have the same size as the nasal conjunctival defect, regardless of which group the patient was randomised to. Thus, the knowledge of the surgery method did not influence the results or conclusions of the study. It is also impossible to randomise the patients without the surgeon’s knowledge, because the grafts were handled somewhat differently in the different methods described.

The fibrin adhesives in the references mentioned by Alvarenga are clearly not Tissel Duo Quick (Baxter). In the article by Hino et al, the specific trade name of the fibrin sealant is not given, but the text states that the sealant in question is treated by dry heat to inactivate viruses—Tisseel is vapour heated, so it cannot be Tissel. In the article by Kawamura et al the product is Beriplast (Aventis Behring). Nevertheless, we will pay attention to this matter in the future.

We have not been able to find any report of HPV B19 infection when fibrin glue was used in minor surgery. To date we have used the fibrin glue in more than 700 eye procedures since 1999. Any clinical infection with HPV B19 has not yet been detected in our patients.

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Korean kindergarten vision screening programme

Lim et al report a large scale effort to mix home plus health centre acuity screening in preschool children.5 We are very encouraged by the work of Lim et al, particularly concerning the frequency of ocular symptoms in the Korean preschooler, the number of children who were not dismissed from specified follow up (presumed amblyopia risk), and the inclusion of a simple, home administered test which over 97% of children were able to pass. It is of high merit that parents were carefully instructed to place tissue and tape over the non-tested eye,
though this does not preclude peeking if the parent is not paying particular attention. Positive answers to the parental questionnaire were not very specific for eye disease and therefore could greatly increase societal cost if used as a screening method. We have a few points of clarification for these authors: How was the home acuity test initially reported child is the eighth documented child. The issue of the possible association of assisted reproductive techniques (ART) with bilateral disease and three with unilateral disease."