Merkel cell carcinoma of the eyelid in association with chronic lymphocytic leukaemia

Merkel cell carcinoma (MCC) is a rare skin neoplasm. Tang and Toker first described MCC in 1978 and since then 19 cases in association with chronic lymphocytic leukaemia (CLL) have been reported. To the best of our knowledge, involvement of the eyelid by MCC has never been reported in the literature in association with CLL.

Case report
An 84 year old white man was referred with an 8 week history of a painless lump on his right upper eyelid (Fig 1A). He was complaining of visual obscuration secondary to a mechanical ptosis. Ophthalmic history was unremarkable and specifically there were no previous chalazions or trauma. On examination a firm lesion of the right eyelid measuring 2 x 1 cm with overlying telangiectatic vessels and sparing of the eyelashes was noted (Fig 1A). Further ophthalmic examination was unremarkable. General examination did not reveal any abnormalities.

General medical history revealed that the patient had been diagnosed with CLL 11 months previously and was being treated with pulsed chlorambucil. His condition was considered to be stable by his oncologist. At the time he had a white cell count of 15.7 x 10^9/l. These consisted of immature lymphocytes of a B cell lineage. A full thickness incisional biopsy was performed under local anaesthesia. Histopathological examination of the biopsy sample showed an intact epidermis with the underlying dermis being infiltrated by clumps of a small cell tumour (Fig 2A).

Immunostaining showed the tumour cells were negative for LCA (leucocyte common antigen), CD3 (T cell marker), CD20 (B cell marker), chromogranin, and S100 antigens. The tumour cells were positive forNSE (neuron specific enolase), EMA (epithelial membrane antigen) and CAM 5.2, which showed characteristic paranuclear accentuation (Fig 2B). Other staining techniques showed 50% of the tumour cells to be in cycle. All these features are consistent with the diagnosis of MCC.

Further investigation revealed no systemic metastasis. We opted for radiotherapy as the patient was reluctant to have surgical intervention. The patient was given a total of 40 Gy in 15 fractions. This caused the tumour to reduce in size relieving the mechanical ptosis (Fig 1B).

Comment
The recent surveillance, epidemiology, and end results (SEER) programme in the United States has estimated the incidence of MCC at 0.23/100 000. MCC is very rare below the age of 50 and is more common on sun exposed sites. It is an aggressive tumour with 12–45% being lymph node positive at presentation. This increases to 55–79% during the course of the disease. The 5 year survival has been reported at 30–64%. Involvement of the eyelid occurs in only 0.8% of MCC, and has not been reported in the literature in association with CLL.

Secondary tumours are common in B cell neoplasia with the relative risk of non-melanotic skin cancer being 4.7 in men and 2.4 in women. The frequency and aggressiveness of MCC and other skin neoplasms increases with immunosuppression, organ transplantation, as well as B cell neoplasia. The precise reason for such an association is not fully understood. Quaglino et al suggested that a depressed immunological system as well as exogenous oncogenic factors may, in various degrees, contribute to the development of neoplastic processes at different sites.

The treatment is wide local excision with or without adjuvant therapy consisting of block dissection of lymph nodes or radiotherapy. Adjuvant therapy reduces local recurrence and regional failure from 39% and 46% to 26% and 22% respectively. Most patients die from causes directly related to the disease. Potentially there is an increased risk of all skin tumours including MCC in patients suffering from CLL and this diagnosis should be considered when evaluating an eyelid lesion in such patients. In a patient with reduced immunity it would be best practice to send all surgical specimens for histology even if a simple chalazion is thought to be responsible for the lid lesion.

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References

Steatocystoma simplex of the caruncle

The caruncle has a non-keratinised epithelial lining similar to the conjunctival epithelium. However, unlike the conjunctiva, the caruncle harbours skin elements such as hair follicles, sebaceous glands, sweat glands, and accessory

Figure 1
(A) Merkel cell tumour involving most of the upper lid causing mechanical ptosis and visual obscuration. Biopsy site is seen laterally. (B) Merkel cell tumour after treatment with radiotherapy.

Figure 2
(A) Haematoxylin and eosin stain Merkel cell tumour. (B) Immunostaining with CAM5.2 showing characteristic para-nuclear accentuation.
Figure 1 Clinical appearance of the pale yellow lesion of the right caruncle (arrowed).

lacral tissue. Consequently, the caruncle may develop a tumour or cyst similar to one found in the skin, conjunctiva, or lacrimal gland.

Case report
A 26 year old woman presented with a 2 mm, asymptomatic, pale yellow lesion of the right caruncle, present for 8 months (Fig 1). It was excised intact under local anaesthetic and histological examination revealed a cyst lined by stratified squamous epithelium and containing sebaceous glands in its wall (Fig 2). These communicated directly with the cyst lumen. No associated hair follicles were seen. An eosinophilic, crenulated cuticle was present on the inner aspect of the cyst wall in some areas. The patient had no other skin lesions of note. The nails, teeth and hair were normal. There was no family history of similar lesions. At the 6 month follow up there has been no signs of recurrence or development of other cysts.

The diagnosis was of steatocystoma simplex (SS) of the caruncle.

Comment
Steatocystoma simplex, the non-hereditary solitary counterpart of steatocystoma multiplex (SM), was first described as a distinct entity by Brownstein in 1982.1 It is a benign adenocutaneous tumour, thought to originate from a naevus malformation of the pilosebaceous duct junction.2 Lesions are described on the forehead, nose, scalp, neck, axillae, chest, upper limbs, back, legs, and even intraorally.3 To our knowledge though, steatocystoma has not previously been reported in the caruncle.4,5

Thirty two cases of SS are reported in the literature, divided evenly between men and women and ranging in age from 15 to 70 years.4,6 Clinical and histological features in SS are usually identical to those seen in the individual lesions of SM. Lesions are described as asymptomatic, flesh coloured or yellowish, intransigent, well circumscribed, soft, mobile, and non-tender. On incision they are found to contain an oily substance composed of sebum.7

However, it is important to confirm the solitary nature of a steatocystoma. SM can be familial and autosomal dominantly inherited (steatocystoma multiplex congenita).7 Several familial cases have been linked to pachyonychia congenita and ectodermal dysplasia through a mutation in keratin 17.8 It has been associated with hypothyroidism, hypohidrosis, hypotrichosis and hypertrophic lichen planus, ichthyosis, and kolonichysta.9 SM may be progressive with inflamed cysts, rupturing, and healing with scarring. Steatocystoma is histologically characterised by a cystic structure with sebaceous glands within the cyst wall and epithelium that displays an eosinophilic cuticle. It is possible to make a diagnosis of steatocystoma if the characteristic hyaline luminal cuticle is present, even in the absence of sebaceous elements.

The differential diagnosis included sebaceous gland hyperplasia, sebaceous gland adenoma, and lipogranuloma. Clinically they are also characterised by a yellow, nodular appearance.10 Rarely, sebaceous gland carcinoma of the caruncle may occur.11 Along with hidrocystoma and eruptive vellus hair cyst they can usually be excluded histologically. Oncocytomas are asymptomatic, slowly progressive, solid or cystic masses but usually described as reddish blue/taupe.

Most treatment regimens for steatocystoma reflect the multiplicity and widespread extent of lesions of SM. Oral isotretinoin has been used to reduce associated inflammation, Cryosurgery, carbon dioxide laser, and incision with removal of the cyst wall have been employed.12,13 We felt the best way to manage this solitary lesion arising in the caruncle was by simple excision with removal of the cyst wall intact, thereby reducing the risk of recurrence. We were able to confirm the unique nature of this lesion and rule out malignancy.

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References

Preliminary results with posterior lamellar keratoplasty for endothelial failure

We describe the technique and the results of three cases where we performed a posterior lamellar keratoplasty.

Case reports
The following surgical technique was performed in all cases. The donor posterior button was obtained from an entire fresh globe. We made sure that intraocular pressure was 0.1 in the right eye and 0.06 in the left. The donor posterior button was obtained from an entire fresh donor eye and closed for posterior keratoplasty.

With our microkeratome an 8.5 mm in diameter, nasal hinge and 250 µm flap was obtained. The trephination was made with a 7 mm Barron trephine and completed with conical scissors, under viscoelastic protection.

After the intraocular injection of acetylocline the posterior donor button was placed on the recipient eye under viscoelastic protection. Six 10-0 Nylon interrupted sutures were used to secure and close the wound. Immediately after, the flap was put back and fixed with six interrupted 10-0 Nylon sutures and the knots were buried. The viscoelastic anterior chamber was exchanged with BSS using an automatic pressure controlled irrigation-aspiration system.

Case 1
This was a 36 year old woman with Fuchs' endothelial dystrophy. Preoperative BSCVA was 0.4 in the right eye and 0.6 in the left. Silt lamp examination showed diffuse corneal oedema clearly affecting the anterior layers of the cornea. Endothelial cell count (ECC) was below 900 cells/mm² in both eyes. Surgery was performed on the right eye (Fig 1A). The follow up was done for 12 months (Fig 1B).

Case 2
Case 2 was a 57 year old man with Fuchs' endothelial dystrophy. Preoperative BSCVA was 0.1 in the right eye and 0.06 in the left. Silt lamp examination showed diffuse stromal corneal oedema in the left surgical eye. ECCs were difficult to perform because of the light.
This was a 57 year old man with history of myopia in both eyes (right eye −4.00, left eye −9.00). There was a history of subretinal macular neovascularisation and cataract extraction in his left, surgical, eye, with an ECC of 950 cell/mm. Preoperative BSCVA was 0.3 in the right eye and 0.2 in the left. The follow up was done during 14 months (Fig 1D).

Examinations in all cases were at day 1, 1 week, 3, 6, 9, and 12 months. All Nylon sutures were removed before the 6 month control. All surgeries were technically uneventful. The immediate and late postoperative controls showed transparency of the cornea and no signs of rejection. In case 1 at the time of removing the superior suture (3 months post-operatively), a separation between the anterior cap and the edge of the corneal recipient eye was observed (because of the stromal flap oedema) and two interrupted 10–0 Nylon suture were placed for 3 more months.

Uncorrected and BSCVA did not improve in all cases in spite of corneal transparency (Table 1). We observed a significant increase in astigmatism in all cases during the follow up and also after the suture removal, but it was not the main cause of reduced vision (Table 2). We checked vision changes with refraction over rigid gas permeable lenses but the results were lower than expected.

**Comment**

Many attempts have been made to independently replace the endothelial layer. First Moly5 and McCulley6 used eyes of the animal models and obtained successful results. Later, Melles et al described a surgical technique in which through a scleral tunnel incision a mid-stromal pocket was dissected to separate and transplant the posterior stroma7–9 using a microkerate to access the posterior cornea, also had similar results. We used the open technique as described by Busin, but suturing both corneal layers, and our cases showed a significant astigmatism and very low visual results.

Reviewing our experience during the past 6 years in performing penetrating keratoplasty (PK) for Fuch’s dystrophy, and obviously understanding that this is not a comparative study, we realised that our mean improvement in best corrected visual acuity was 3.1 lines (range 0–8), with a mean postoperative time for visual rehabilitation of 8 months (range 3–18 months).

The recovery time was slower when compared with PK, perhaps because of the optical distortion of the interface. We must also not forget that we sutured both the donor button and the superficial lenticule, perhaps inducing interface distortion. Also it is important to mention the risk of wound leakage and interface aqueous humour dissection.

We think that the time of graft deswelling was not as expected because at the time of suture removal a separation was noted between the anterior cap and the recipient eye in cases 1 and 2. We placed sutures in this site but the time of suture removal was extended to 12 months. Another contributing factor would be host-graft disparity, trephination, and suture technique.

In our experience this technique shows that it is possible to change only the posterior layers of the cornea with successful anatomical result. Nevertheless, from a functional perspective penetrating keratoplasty has been a much better and faster approach and, in fact, in both techniques we are replacing the endothelium using an open sky technique.

### Table 1

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### Table 2

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Persistent accommodative spasm after severe head trauma

Spasm of accommodation is the sudden occurrence of difficulty in focusing the eyes. The term "accommodative spasm" is used to describe the phenomenon of sudden onset and recovery of difficulty in accommodation. The spasm may be unilateral or bilateral, and it may be transient or persistent.

Case report

Case 1

A 34-year-old woman presented to the ophthalmology service with a history of severe head trauma 9 years prior. She had been wearing corrective lenses for myopia and astigmatism. She had been treated with cycloplegic drops for 6 months after head trauma. She had not worn glasses in the past years.

Cycloplegic refraction was prescribed, and the patient was instructed to wear the glasses at home and when outdoors. She was also prescribed atropine drops every 12 hours, which she took for 6 months. She reported improved vision with glasses.

The patient was referred to the neurology service for further evaluation. MRI of the brain showed multiple lesions in the peri-cerebellar region. The lesions were marked by T2-weighted images (TR 3570, TE 120 ms) spin echo sequences, 3 mm thickness, with normal findings in the cerebellar vermis (Fig 2).

The cycloplegic refraction was prescribed, and the patient was instructed to wear the glasses at home and when outdoors. She was also prescribed atropine drops every 12 hours, which she took for 6 months. She reported improved vision with glasses.

Figure 1 Case 1. T2 weighted (TR 3570, TE 120 ms) spin echo sequences, 3 mm thickness, with normal findings in the mid-brain.

References


Comment

In our patients, a relation of accommodative spasm and the head trauma seems well established because it appeared soon after they have recovered from severe head injury. Persisted for several years despite the prolonged use of cycloplegic drops, and the patients were not trying to obtain any benefit by complaining of blurred vision.

The supranuclear control of accommodation is poorly understood. In cats, neurons that discharge in temporal correlation with accommodation were found in the lateral suprasylvian area. Electrical stimulation on ipsilateral interpositus nuclei and on contralateral lateral interpositus and fastigial nuclei in the cerebellum are known to induce accommodation. These nuclei are connected to parasympathetic oculomotor neurons in the mid-brain.

Very little accommodative dysfunction resulting from central lesions has been reported in humans. Ohtsuka et al. studied a patient with left middle cerebral artery occlusion who had reduced accommodative responses and markedly lowered accommodation velocity. Their patient had low density lesions on computed tomograph (CT) scan involving the left temporal lobe, near the Sylvian fissure. Kawasak et al. reported a patient with normal accommodation amplitude but increased accommodation and relaxation times. Their patient recovered normal accommodation 10 days after removal of a large subtentorial arachnoid cyst and the authors suggested that the cerebellum might have a role in the organisation of the human central control system of accommodation.

Bohmann and France described a patient with persistent spasm of accommodation after head trauma. CT scan revealed a skull base fracture without intracranial abnormalities and the authors suggested that a possible
Extradural plasmacytoma of the eyelid

A 74 year old man presented with a foreign body sensation in the right eye. He had had recurrent infections, blepharitis, and right eyelid swelling. On examination, there was a firm, painless mass at the lateral canthus with internal shielding of the eye. The mass was excised surgically and histopathology revealed a large firm lesion in the right upper eyelid with no palpable lymph nodes. The clinical diagnosis was of a chalazion.

The lesion was removed surgically and histopathology (Fig 1) revealed an incompletely excised extramedullary plasmacytoma with a high proliferative index and amyloid change. Immunocytochemistry was positive for IgG kappa light chains. Further investigations including full blood count, liver function tests including lactate dehydrogenase (LDH), protein electrophoresis, skeletal survey, and bone marrow aspiration were normal with no evidence of multiple myeloma.

The whole of the upper eyelid was treated with radiotherapy using a customised lead cutout with internal shielding of the eye (Fig 2). 120 kV x rays were used giving a dose of 30 Gy in 10 fractions over 2 weeks.

On follow up (at 2 years) there has been no evidence of local recurrence or the development of myeloma, and lacrimation in the eye appears normal.

Comment

Solitary plasmacytomas are rare tumours. They are classified as either solitary plasmacytomas of bone (SPB) or extramedullary plasmacytomas (EMP) of soft tissue. The majority of EMPs (about 80%) involve the upper air passages of the head and are thought to arise in the submucosa, where plasma cells are numerous. Other sites include lymph nodes, spleen, skin and subcutaneous tissues, gastrointestinal tract, thyroid, and testes.

There is a relation between solitary plasmacytomias and subsequent development of multiple myeloma. About 44-69% of patients with a solitary bone plasmacytoma will develop multiple myeloma within a median of 3 years. Although EMPs recur in almost 50% of cases, this is usually in bone but unlike multiple myeloma it remains circumscribed within the bone with no predilection for the axial skeleton. However, progression to myeloma does occur though at a lower rate than for SPB. Alexiou et al reported a rate of progression to myeloma for both upper aerodigestive tract and non-aerodigestive tract extramedullary soft tissue plasmacytomas of 16.1% and 14.6% respectively. As no predictors of progression have been identified patients probably need indefinite follow up.

Eye abnormalities such as cysts of the ciliary body and vascular lesions have been described in multiple myeloma but primary plasmacytoma involving the eye is rare. Nine cases affecting the orbit have been described in the literature but this is only the fourth case of a primary plasmacytoma arising from the eyelid that has been reported. Most of the earlier reports of the plasmacytomias arising in the eye are not true plasmacytomas and are in fact granulomas due to chronic inflammation. Usual symptoms are progressive painless swelling of the eyelid, proptosis, and diplopia. They can occur at any age but the mean age of onset is in the sixth to seventh decade. The youngest reported age case was that of an 11 year old who had plasmacytoma of the orbit.

Of the three previously reported cases, all were treated with surgical excision. Their immunocytochemistry was IgG lambda chain, kappa light chain, and IgG lambda chain respectively. Our case is similar but was treated successfully with radiotherapy after incomplete excision.

Solitary extramedullary plasmacytomas can be controlled with radiotherapy alone. Response rates with radiotherapy are as high as 94% and 93% for SPB and EMP respectively. The optimal dose of radiotherapy has not been defined, although it appears that a dose of at least 30 Gy is required. Many centres use doses of between 40-50 Gy. The extent of radiotherapy portals is also a subject of debate with many recommending inclusion of regional lymph nodes if possible. The median survival of patients with EMP treated with radiotherapy was 8.5 years in one study with most patients dying of causes unrelated to their EMP.

Surgery is also an option, with Alexiou et al reporting a lower rate of progression to myeloma for those treated surgically (7%) compared with those treated with radiation (17.5%). The conversion rate for patients treated with both modalities was 13.5%. These results may reflect differences in the size of lesions, with small extramedullary plasmacytomas in easily accessible sites being amenable to surgical excision.

Chemotherapy is used for those patients who progress to multiple myeloma.

 references

A 71 year old white woman was referred by her optician after attending for routine glasses update. On questioning she did complain of a “slight blurring of vision” gradually for several months. She had a history of left amblyopia. Her medical history included asthma, osteoarthritis, lymphoedema, fibromyalgia, and hiatus hernia. Her only medications were inhalers and paracetamol. She had no headaches. Her medical history included oesophagitis and hiatus hernia. Her only medication was antidepressant. She had previously taken calcium supplements. Her unaided visual acuity was 6/12 right eye improving to 6/9 with pinhole, and 6/18 left eye improving to 6/12 with pinhole. She had bilateral nuclear sclerotic cataracts and both fundi revealed numerous pale elevated lesions clustered around the superotemporal and inferotemporal arcades (Fig 2A).

Haematological investigations and chest x-ray were unremarkable. Ultrasound scanning revealed that these were areas of calcification with high reflectivity (Fig 2B).

Case 3
A 71 year old man attended routinely for review 2 weeks after cataract extraction. He had no past medical history and vision was 6/9 right eye and 6/6 left eye. He was noted to have a small optic disc haemorrhage coincidentally and therefore dilated fundal examination was performed. He had bilateral pale yellow elevated lesions along the superotemporal arcades and also nasally in both fundi. Again electrolytes were investigated and were within the normal range.

Comment
Sclerochoroidal calcification has been described in the literature as an uncommon condition found usually in older white patients. Patients have been unnecessarily investigated in the past and even treated for tumours unnecessarily. Calcification can be dystrophic or metastatic but in these idiopathic cases it is the former. The calcification is believed to be deposited at the sites of insertions of the oblique extraocular muscles in a similar way that Cogan scleral plaques are calcification at the insertions of the horizontal recti muscles. Reports have been made of sclerochoroidal calcification associated with Barter syndrome, Gitelman syndrome, hyperparathyroidism, and hypomagnesaemia. It is important to exclude any electrolyte abnormality when a patient presents with this condition, but prolonged investigations are unnecessary. Autofluorescence and ultrasound appearances are very useful to diagnose this condition. Patients rarely develop visual disturbance with sclerochoroidal calcification, but infrequent follow up is advised as cases of associated choroidal neovascular membranes and serous detachments with the lesions have been documented.

Acknowledgements
ACON kindly sponsored the publication costs of the figures.

References
Stereotactic irradiation of biopsy proved optic nerve sheath meningioma

The role of conventional external beam radiotherapy in the management of optic nerve sheath meningiomas (ONSM) has been controversial because of limited radiation sensitivity of these tumours and radiation damage to surrounding tissues. Recently, in a study of 64 patients with ONSM managed with observation, surgery, radiotherapy, or surgery and radiotherapy, Turbin and colleagues found that patients treated by (conventional) radiotherapy alone demonstrated the best long term visual outcome, and suggested fractionated external beam radiation (5000–5500 CGy) as the initial treatment in selected cases, when preservation of visual function is a reasonable goal. The collateral damage secondary to conventional radiotherapy may be minimised by better focusing and shaping of the radiation beams, as in stereotactic radiotherapy (SRT). We report on a woman whom we treated with fractionated SRT for a biopsy proved, large ONSM.

In April 2000 a 41 year old woman was referred with a 1 month history of proptosis of her left eye (Fig 1, top left). She had been treated for a presumed orbital “pseudotumour” with oral prednisone (initial dose 90 mg/day) without effect. At referral, she had no history of diplopia or retrobulbar pain. On examination, the visual acuity (VA) was 1.25 (unaided) of the right eye and 0.8+ (cc S+2) of the left eye. The intraocular pressure was 18 and 21 mm Hg in the right eye and left eye, respectively. There was a left relative afferent pupillary defect (RAPD). The motility of the left eye was limited in all quadrants. Visual field testing (Humphrey) showed mild periocular swelling and conjunctival translucency of the sclera.

After surgery we observed the patient for 9 months. During this period her left (corrected) VA deteriorated to 0.2 and her left visual field showed progression of her scotomas. This prompted us to treat her with fractionated SRT in March 2001. The radiation, delivered with a 6 MV linear accelerator (Varian), was given 5 days a week at 1.8 Gy per fraction, with a cumulative dose of 54 Gy. Treatment planning was based on orbital MRI matched with CT scans. A non-invasive stereotactic frame was fixed with an external coordinate system (one isocentre). Target and surrounding tissues at risk were defined as volume of interest on contrast media enhanced T1 weighted MRIs and transferred to CT by the stereotactic localisation technique using a three dimensional planning system (X Plan, Radionics). Portals were optimised using a beam’s eye view technique. Five irregularly shaped non-coplanar beams (arcs) per treatment were used. Beam shaping was done with a mini-multileaf collimator (Radionics). No early complications of the radiation treatment were noted. At 6 months after SRT, the (corrected) VA of her left eye had recovered to 0.8, while no RAPD was observed.

Figure 1 Top left. Appearance of a 41 year old woman with a biopsy proved optic nerve sheath meningioma before SRT. Note the left exophthalmos and periorbital swelling. Top right. Post-treatment appearance. Note the decrease of the fullness of the left eyelids. Also note the left upper lid retraction secondary to left upper lid ptosis. Bottom left. Orbital MRI scan (T1 weighted with fat suppression and gadolinium contrast enhancement) at presentation. Bottom right. Six months after radiotherapy. A decrease of both tumour size and proptosis is clearly visible.

Figure 2 Top left. Ultrasound examination at presentation. A large, heavily vascularised retrobulbar mass is visible. Top right. Six months after radiotherapy, the tumour has diminished in size and vascularisation. Note that a different depth setting of the ultrasound system has been used. Bottom. Histopathology of the optic nerve tumour, showing whirls of meningothelial cells, with small nuclei and inconspicuous nucleoli, consistent with a meningioma (Fig 2, bottom).

Circle M. The collateral damage secondary to conventional radiotherapy may be minimised by better focusing and shaping of the radiation beams, as in stereotactic radiotherapy (SRT). We report on a woman whom we treated with fractionated SRT for a biopsy proved, large ONSM. In April 2000 a 41 year old woman was referred with a 1 month history of proptosis of her left eye (Fig 1, top left). She had been treated for a presumed orbital “pseudotumour” with oral prednisone (initial dose 90 mg/day) without effect. At referral, she had no history of diplopia or retrobulbar pain. On examination, the visual acuity (VA) was 1.25 (unaided) of the right eye and 0.8+ (cc S+2) of the left eye. The intraocular pressure was 18 and 21 mm Hg in the right eye and left eye, respectively. There was a left relative afferent pupillary defect (RAPD). The motility of the left eye was limited in all quadrants. Visual field testing (Humphrey) showed mild periocular swelling and conjunctival translucency of the sclera.

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Her periorcular swelling had markedly diminished (Fig 1, top right). Compared to previous measurements, the protrusion of the left eye had diminished by 4 mm. Funduscopy, however, showed mild pallor of the left optic nerve head. Visual field testing showed unchanged loss of the left visual field compared to pretreatment values, with a higher false negative threshold. VEP measurements showed improved amplitudes, but prolonged latency compared to previous examination. Orthoptic examination showed ductions similar to those before treatment. Post-treatment MRI revealed a markedly decreased tumour size and a decrease of exophthalmos (Fig 1, bottom right). Colour Doppler ultrasonography showed a decrease in tumour size with markedly diminished vascularisation (Fig 2, top right). At the last follow up visit, 16 months after treatment, her left VA and visual fields were stable.

Comment
As in the recent report on a presumed ONSM by Moyer et al., fractionated SRT in our biopsy-proved case gave a remarkable visual recovery without detectable side effects. Both the size and the blood flow of the tumour regressed within the first 6 months, leading to reduced exophthalmos and periorcular swelling. The effect of restored cosmesis was important to this young woman whose main complaint was her unilateral exophthalmos. Since our follow up is limited to 16 months, no conclusions with regard to long term outcome can be made. More cases of SRT for ONSM need to be studied over a longer period of time to assess the efficacy of this treatment.

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References

Severe interferon associated retinopathy
Interferon alfa is used in various human malignancies for its antitumour activity. One of its ocular side effects is retinopathy. Interferon associated retinopathy is generally mild and resolves completely. We describe a severe retinopathy in a hypertensive patient treated with interferon for multiple myeloma.

Case report
A 56 year old man presented with a 3 week history of deterioration and distortion of right vision. Visual acuities (VA) were 6/60 right and 1/60 left. Funduscopy revealed bilateral extensive peripapillary cotton wool spots, retinal thickening, optic disc hyperaemia, and blot haemorrhages. Arteriolar changes were minimal.

He was anaemic (Hb 10.6 gdl/l) and slightly thrombocytopenic (platelets 93 × 10^9/l). Plasma viscosity was 1.59 (normal 1.5–1.7). Renal function was normal at presentation.

He underwent peripheral blood stem cell transplant for multiple myeloma 8 months previously after having melphalan 110 mg/m^2 and total body irradiation (including the head) in a total dose of 1200 cGy given in six fractions over 3 days. He then had interferon alfa therapy for 4 months, initially 3 mega units three times a week, later reduced to twice a week. It was stopped immediately after visual deterioration.

Five years previously, he had macular laser treatment following left inferotemporal branch retinal vein occlusion. On discharge 1 year later, VAs were right 6/5, left 6/36. He was a known hypertensive, taking lisinopril, but control was poor around the time his VA began to deteriorate, with readings up to 150/100 mm Hg. He was not diabetic. His myeloma status was stable. Cytomegalovirus (CMV) antigen checked by polymerase chain reaction (PCR) and pretransplant HIV status were negative.

One week after presentation at the eye clinic, VAs dropped to right 2/60, left finger counting and did not improve after a course of intravenous methylprednisolone (1 g/day for 3 days). He was registered blind.

A further 3 weeks later, bilateral cotton wool spots and haemorrhages were more numerous and both foveas showed gross thickening with exudates (Fig 1). Fundus fluorescein angiography revealed retinal ischaemia with capillary non-perfusion, pruning, and tortuosity of vessels, vessel wall staining, and leakage (Fig 2).

Five months later preproliferative retinopathy was noted and subsequent proliferative changes were treated with bilateral panretinal laser photocoagulation. At 9 months, VAs were 1/60 right and left.

Comment
In a review article on interferon retinopathy, initial interferon alfa doses ranged from 3–9 mega units three to six times per week for several weeks. In a prospective randomised placebo controlled trial of interferon alfa therapy for macular degeneration, retinopathy was noted with increasing frequency in the highest dose group (5% of the patients taking 6 mega units three times a week). The interferon doses in our patient were at the lower level of these regimens.

Severity of retinopathy was found to be related to the presence of the following risk factors: large initial dosages, long duration of treatment, and systemic diseases like diabetes mellitus or hypertension. Early onset of retinopathy was also a good indicator of severity.

Figure 1 Bilateral (right eye [A] and left eye [B]) extensive peripapillary cotton wool spots, retinal thickening with exudates, optic disc hyperaemia, and blot haemorrhages.

Figure 2 Fundus fluorescein angiography of the right eye: early frame (A) showing retinal ischaemia with capillary non-perfusion, pruning, and tortuosity of vessels; late frame (B) showing vessel wall staining and leakage.
References


Comment

The range of ocular findings associated with Bartonella henselae continues to expand. The classic follicular conjunctivitis described with lymphadenopathy and fever (Parinaud's ocular glandular syndrome) is due to direct inoculation of the conjunctiva. Neuroretinitis, a syndrome of acute visual loss associated with optic disc swelling and macular star, was the first established intraocular complication from disseminated Bartonella. Since then, other reported intraocular findings include inflammatory chorioretinal white spots, papillitis, serous detachment, vitritis, uveitis, vasculitis, retinal vaso-occlusive disease, and vitreous haemorrhage. A solitary macular lesion without other ocular inflammatory findings has also been reported. Such lesions in the posterior pole have been presumed to represent a mass lesion due to growth of Bartonella. We report a patient with a circumscribed, elevated lesion in the macula as well as other mass-like lesions at the optic nerve head and in the choroid. These lesions occurred in the absence of systemic or ocular inflammation and clinically resembled ocular metastases. This case highlights the importance of recognizing the wide spectrum of ocular bartonellosis. Furthermore, clinicians are reminded that cat exposure is not essential for bartonellosis, and casual contact with Bartonella should be obtained whenever there is a clinical index of suspicion, regardless of cat exposure.

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References


Optic neuritis with marked distension of the optic nerve sheath due to local fluid congestion

Distension of the subarachnoid space of the optic nerve is not a common feature of optic neuritis. We describe a patient with optic neuritis with swelling of the optic nerve head of the right eye. On magnetic resonance imaging (MRI) there was marked distension of the optic nerve sheath due to an increase of fluid in the subarachnoid space. The location of the lesion in the optic nerve and concurrent inflammatory changes of the arachnoid trabecula and septae may have had a role in the pathophysiology of this condition.

Case report

A 38 year old man was admitted with pain on eye movements and loss of vision in the right eye. Best corrected visual acuities measured 20/40 on the right and 20/20 on the left. The patient identified 16 out of 18 Ishihara plates with the right eye and 18/18 with the left eye. There was a relative afferent pupillary defect (RAPD) on the right. Funduscopy demonstrated a swollen optic disc on the right. The left optic disc was normal. Spontaneous venous pulsations (SVP) were detectable bilaterally. Laboratory examinations, including red and white blood counts, C reactive protein, sedimentation rate, serologies for syphilis, HIV, herpes, toxoplasmosis, Lyme disease and cytomegalovirus, as well as collagen vascular disorders and coagulopathies were all in normal range. The right visual field (Octopus program G2) demonstrated a wedge-shaped defect in the inferior nasal and temporal visual field pointing towards the macula. The left visual field was normal. Neurological examination was normal. MRI of the brain was normal but showed enhancement of the right optic nerve in the T1 weighted axial and coronal (not shown) image and hyperintense fluid in the expanded optic nerve sheath on the T2 weighted image (Fig 1A and B). Two days after admission the visual acuity in the right eye decreased to 20/100 and only two out of 18 Ishihara plates were identified. SVP were no longer present on the right. The swelling of the right optic disc progressed and temporal peripapillary Patton folds appeared, suggesting the diagnosis of papilloedema (Fig 2). Within 2 weeks visual acuity improved to 20/25 right eye and colour vision returned to normal. A repeat MRI of the orbits 7 weeks later demonstrated normal diameters of both periophtalmic subarachnoid spaces (Fig 1C).

Comment

Distension of the periphtalmic subarachnoid space is a hallmark MRI feature of papilloedema due to an intracranial mass lesion, inflammatory disease, and pseudotumour cerebri. Unilateral distension of the optic nerve sheath due to increased fluid volume of the subarachnoid space of the optic nerve has previously been reported in some patients with optic hydrops, anterior ischaemic optic neuropathy, and anatomical anomalies such as arachnoid cysts. This patient with optic neuritis demonstrated marked distension of the subarachnoid space of the right optic nerve, presumed to be caused by an increase of total fluid following optic neuritis. As all cerebrospinal fluid compartments are thought to communicate, equalisation of fluid via the chiasmatic cistern would have been expected to occur. The MRI scan of the brain and orbits, however, demonstrated localised and isolated stasis of fluid in the right optic nerve subarachnoid space only. The reason for this fluid congestion causing a optic nerve sheath compartment syndrome could not be identified by neuroimaging. The site of inflammation of the optic nerve and local anatomical variations and alterations of the subarachnoid space—for example, the amount and number of trabecula and septae in the subarachnoid space—may have a crucial role in the pathophysiology of unilateral papilloedema.

References


Corneal opacification following keratoplasty in the rat model

I read with great interest the excellent perspective by Plsková et al., in which they raise the issue of transient corneal opacification following corneal transplantation in the mouse model and argue that it might be due to a sufficient number of endothelial cells regaining function.

What the authors describe for the mouse model also occurs in the rat model. In fact,
When discussing the other two papers it is of paramount importance to understand that given the long learning curve associated with deep sclerectomy, it is neither fair nor scientifically sound to compare a surgeon’s results to 20 cases of trabeculectomy with his first 20 cases of deep sclerectomy. As an example, one group reported 0% success rate in their first series of viscosanalostomy patients and then presented their second series with a success rate of 15%. The same group also analysed the depth of their dissection of the deep sclera to find that they dissected too superficially in 48% of their cases and too deeply in 17%; meaning that the proper depth of dissection, which should reflect transversely the Schlemm’s canal, was not achieved in the majority of their cases. The authors also failed to cite published long term results for deep sclerectomy with collagen implant. The study provided a qualified success rate of 94.8% and the complete success rate, 61.9%, after 60 months (survival analysis), with a mean IOP at 1 year of 11.8 (SD 3) mm Hg. Although the study reports a non-randomised consecutive series of patient, it should be taken as a proper indication of results achieved by experienced surgeons. It should be noted that no consideration that non-penetrating surgery is a broad genre of surgery, under which different surgeons perform fundamentally different procedures that include sinusotomy, ab externo trabeculectomy, deep sclerectomy with or without the use of an implant, viscosanalostomy, performance of postoperative gonipuncture, and the use of antimetabolites. The different techniques have one thing in common, the element of non-perforation.

What is true is that this type of surgery is continuously evolving, so it is unlikely that a proper judgment can be made yet. At the risk of sounding dramatic, it is valid to say that editors like the one by Khaw et al seem to indirectly sign a death certificate of non-penetrating surgery. It is much more useful to encourage research in non-penetrating surgery, including multi-centre follow-up of non-randomized studies, to see if deep sclerectomy will remain king.

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References

The role of optometry in vision 2000
The latest issue of Community Eye Health (No 43) discusses the mobilisation of optometry to deal with uncorrected refractive error, which is now a major cause of functional blindness. For further information please contact: Journal of Community Eye Health, International Centre for Eye Health, Institute of Ophthalmology, 11–43 Bath Street, London EC1V 9EL, UK (tel: +44 (0)20 7608 6910; fax: +44 (0)20 7250 3207; email: eyeresource@ucc.ac.uk; web site: www.jch.co.uk). Annual subscription (4 issues) UK£25/US$40. Free to workers in developing countries.

International Centre for Eye Health
The International Centre for Eye Health has published a new edition of the Standard List of Medicines, Equipment, Instruments and Optical Supplies (2001) for eye care services in developing countries. It is compiled by the Task Force of the International Agency for the Prevention of Blindness. Further details: Sue Stevens, International Centre for Eye Health, 1–43 Bath Street, London EC1V 9EL, UK (tel: +44 (0)20 7608 6910; email: eyeresource@ucc.ac.uk).
welfare service, help members and their families cope with the everyday concerns caused by retinitis pigmentosa. Part of the welfare service is the telephone helpline (+44 (0)1280 860 363), which is a useful resource for any queries or worries relating to the problems retinitis pigmentosa can bring. This service is especially valuable for those recently diagnosed with retinitis pigmentosa, and all calls are taken in the strictest confidence. Many people with retinitis pigmentosa have found the Society helpful, providing encouragement, and support through the Helpline, the welfare network and the BRPS branches throughout the UK. (tel: +44 (0)1280 821 334; email: lynda@brps.demon.co.uk; website: www.brps.demon.co.uk)

Detachment Course with international faculty on: Retinal and Vitreous Surgery with Case Presentations preceding Retina Meeting

The detachment course with international faculty on: Retinal and Vitreous Surgery with Case Presentations and the Retina Meeting will be held 14–15 March 2003 and 16 March 2003 respectively, in Mexico City, Mexico. Further details: Scientific programme: Prof Ingrid Kreissig, University of Tuebingen, Schleichstr. 12, Breuningerbau, 72076 Tuebingen, Germany (tel: +49 7071 295209; email: ingrid.kreissig@med.uni-tuebingen.de). Local organisation: Prof. Quiroz-Mercado, Prof. Munoz, and Prof. Gonzalez “Hospital Luis Sanchez Bulnes”, Asociacion para Evitar la Ceguera en Mexico Vicente Garcia Torres #46, Coyoacan, Mexico DF 04330 (fax: +525 5639 5928; email: retinamex@yahoo.com).

16th Annual Meeting of German Ophthalmic Surgeons

The 16th Annual Meeting of German Ophthalmic Surgeons will be held 8–11 May 2003 in Nürnberg, Germany, Messezentrum. Organised by the Professional Association of German Ophthalmologists Ophthalmic Surgery Group the conference will cover cataract surgery, refractive surgery, glaucoma surgery, vitreoretinal surgery, corneal surgery, eye surgery in developing countries, and orbita, lacrimal and lid surgery. Further details: MCN Medizinische Congress organisation Nürnberg AG, Zerzabelshofstr 29, 90478 Nürnberg, Germany (tel: +49 911 3931621; fax: +49 911 3931620; email: doc@mcnag.info; website: www.doc-nuernberg.de).

3rd British Oculoplastic Surgery Society Meeting

The 3rd British Oculoplastic Surgery Society Meeting will be held 18–19 May 2003 in Birmingham, UK. For further details please contact the Secretary of the British Oculoplastic Surgery Society Jane Olver (tel: +44 (0)121 424 5464; fax: +44 (0)121 424 4464; email: MartiDI@heartsol.wmids.nhs.uk; website: www.bopss.org).

13th Meeting of the EASD Eye Complication Study Group

The 13th Meeting of the EASD Eye Complication Study Group will be held on the 23–25 May 2003, in Prague, Czech Republic. The scientific programme includes keynote lectures from Professor John H Fuller (UK) on The epidemiology of diabetic retinopathy; Dr P Martin van Hagen (The Netherlands) on Growth factors and diabetic retinopathy; Professor Jerecic Pelikanova (Czech Republic) on Pathophysiology of diabetic microvascular complications; Dr Tomas Sosna (Czech Republic) on Risk and protective factors of diabetic retinopathy.

Three travel grants of €1000 each, sponsored by GlaxoSmithKline for young scientists (under 35 years at the time of the meeting). Applications should be made with the submission of abstracts. The deadline for abstracts is 14 February 2003.

Further details: Ortopedické Centrum, s.r.o., Střeňovské nábřeží 51, 400 03 Usti nad Labem, Czech Republic (tel: +420 47 521 6588; fax: +420 47 533 40 77; email: ortcentrum-ul@volnv.cz; website: www.ortopedick-centrum.cz).