LETTERS TO
THE EDITOR

Surgically removed submacular nematode

Editor,—Intraocular filariasis is an uncommon disease in civilised countries. In only a few cases has the worm been surgically removed from the posterior portion of the eye and identified.1 In this report we describe a patient who had a filaria-like worm in an epicentral and/or submacular lesion, which was successfully removed surgically.

CASE REPORT
A 38 year old woman complained of decreased visual acuity in her left eye of 5 days' duration. Her medical history was unremarkable. The patient had two pet dogs, and she had not travelled to foreign countries in recent years. On examination, her corrected visual acuity was 20/20 in the right eye and 20/100 in the left. No inflammation was present in the anterior segment or vitreous cavity bilaterally. Ophthalmoscopic examination disclosed a white worm, approximately 3 disc diameters in length, equivalent to about 4.5 mm, moving slowly in the macula at the epiretina. Epiretinal and intraretinal haemorrhages were observed in and around the macular region. The body of the worm was tapered at one end and slightly rounded at the other (Fig 1). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 2). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 1). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 2). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 1). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 2). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 1). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 2). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 1). A round, preretinal haemorrhage was observed on a slightly rounded at the other (Fig 2).

Numerous subretinal hypopigmented tracks with small haemorrhages were noted in the superior retina, and perivascular haemorrhages were observed around the inferior branch of the central retinal vein (Fig 2). Fluorescein angiography showed numerous hyperfluorescent tracks without dye leakage. The patient's blood test revealed a slightly elevated white blood cell count (10 200 ×103/l; normal <9000) and elevated IgE in the serum (680 U/ml; normal <250). On the following day pars plana vitrectomy was performed and the worm was found partially into the subretinal space of the macula. The worm was aspirated successfully through a 20 gauge silicone tipped needle and submitted for parasitological study; however, it was lost during transportation to a different laboratory. Parasitological study of the patient's serum detected an antibody to Dirofilaria Six months after surgery, the patient's visual acuity was still 20/50. Ophthalmoscopically the hypopigmented tracks had faded, and mildly irregular pigment was observed at the deep retina in the macular lesion. Cone and rod electroretinograms to full field stimuli were normal bilaterally.

COMMENT
Our patient owned two dogs, and her ocular findings could be differentiated from toxocariasis, in which only larvae of Toxocara canis can infect humans. The length of the Zosacara larva is about 400 μm, and grows no longer. Although no direct microscopic evidence of Dirofilaria infection was obtained in our patient, a positive antibody to Dirofilaria in the serum indicated its infection. There have been many reports of intraocular filariasis; however, the filariae were removed and identified in only six of 56 cases reviewed by Beaver in 1989.2 Direct parasitological microscopic examination is necessary for accurate diagnosis, but serological study can be helpful, as shown in the present case.

Various types of management for intraocular parasites have been reported. Direct photocoagulation to the worm body has been successfully reported in cases with filaria-like worms3 and in one case with insect parasites.4 It has been suggested that photocoagulation denatures the parasite proteins and mitigates the immune reaction.4 If the parasite is located in the posterior pole of the retina, however, photocoagulation may cause permanent visual impairment, and surgical removal should be selected. Furthermore, photocoagulation would make parasitological identification impossible. Preretinal or subretinal parasites were retrieved successfully by pars plana vitrectomy in several reported cases.5 In the present case, the visual acuity was still 20/50 6 months postoperatively. The migrating worm may have caused considerable damage to macular function, therefore, we believe that intraocular parasites should be removed as soon as possible.

The authors are grateful to Dr Keizo Yamaguchi for parasitological examination and Mr Shingo Yamazaki for fundus photographs.

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REFERENCES
5 Goodard RA, Riekhof FT, Beaver PC. Subretinal nematode: an unusual etiology for uveitis and retinal detachment. Retina 1985;5:87–90

Acanthamoeba keratitis occurring with daily disposable contact lens wear

Editor,—Up to 92% of cases of acanthamoeba keratitis occur in contact lens users of all types.1 Failure to disinfect soft contact lenses and the use of chlorine disinfection systems are major factors accounting for the increase in cases of acanthamoeba keratitis.2 It is thought that “daily disposable” contact lenses, which require no disinfection, will have a much lower risk for the development of all forms of infectious keratitis. We report a case of acanthamoeba keratitis occurring in a daily disposable contact lens wearer. We believe this is the first such reported case.

CASE REPORT
A healthy 21 year old woman, who had been wearing daily disposable contact lenses for 1 year, wore her lenses for 4 hours on 1 day and then stored the lenses in preserved saline solution overnight in a new contact lens case. She rewore the lenses for a further 3 hours the following day. She reports that this was the first time that she had done this. She then developed a painful left eye. This was initially treated as conjunctivitis by her general practitioner and local accident and emergency department. After 2 weeks her local eye unit insisted an epithelial scrape that “revealed amoeba species”. She was then referred to our unit for our opinion.

On examination her visual acuity was reduced to 6/18 in the left eye. There was conjunctival injection and a mild scleritis. The cornea showed diffuse punctate staining with linear epithelial infiltrates (Fig 1).

Figure 1 The cornea showing diffuse punctate staining with linear epithelial infiltrates.

5. Goodard RA, Riekhof FT, Beaver PC. Subretinal nematode: an unusual etiology for uveitis and retinal detachment. Retina 1985;5:87–90
cornea had diffuse punctate staining with linear epithelial and perineural infiltrates (Figs 1 and 2). She had a mild anterior uveitis. Acanthamoeba was strongly suspected. Corneal epithelium was removed for microscopy, culture, and histology. Her contact lens case and solutions were also sent for culture. Acanthamoeba was cultured from both corneal epithelium and lens case, but not the lens solutions. She was treated with topical polyhexamethylene biguanide 0.02% (PHMB) and prednisolone 0.3 % as well as oral flurbiprofen (Froben, Knoll Ltd, Nottingham). Six weeks after diagnosis she was asymptomatic with a visual acuity of 6/9. The eye was quiet, though some corneal infiltrates remained.

**COMMENT**

Acanthamoebaae are free living protozoa commonly found in soil and water, including bathroom tap water. Acanthamoeba keratitis is an uncommon but potentially devastating corneal infection. The number of cases diagnosed in the United Kingdom has steadily risen over the past 20 years owing to increased awareness of the condition and the rise in contact lens wear. Radford et al found that daily wear disposable contact lenses were associated with greatly increased risk of acanthamoeba keratitis compared with other lens types and wear systems.2,3 Multivariable analysis showed that this was largely attributable to a lack of disinfection, the use of non-sterile saline, and the use of chlorine based disinfection rather than alternative chemical systems. It was concluded that 80% of cases of acanthamoeba keratitis could be prevented by the adequate use of an effective disinfection system. It was felt that the “low care” philosophy of daily wear disposable lenses had become “no care” in practice.

Daily disposable contact lenses, in which the lens is discarded after 1 day’s wear only, were introduced in 1995. When used properly the lens is discarded after 1 day’s wear only, had become “no care” in practice.

**Letters**

**Acanthamoeba keratitis: multicentre survey in England 1989–92.**

**J K G DART**

BMJ 1995;310:1567–70.

**Acute zonal occult outer retinopathy (AZOOR) may present with venous infarcts in the eye.**

**J K G DART**


**Clinical course of acute zonal occult outer retinopathy in visual field and multifocal electroretinogram.**

**J K G DART**


**A previous report indicates its efficacy in the diagnosis of AZOOR.**

**J K G DART**


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**Figure 1** (Top) Ratio images of Humphrey 30–2 visual fields in the left eye. (Bottom) The relation between clinical course and mean deviation (MD) of Humphrey 30–2 visual field and visual acuity in the left eye. Asterisks indicate the day in which multifocal ERG was analysed. The roman numerals correspond with raw images in the upper part of the figure.
The findings of AZOOR that we observed in our patient suggest that the retinal recovery assessed by m-ERG was different for the lesion (main focus was related to the infero-temporal retina) and was delayed compared with visual acuity and HVF. Perhaps this delayed retinal recovery reflects a subtle microcirculatory disturbance that can not be clearly detected by FA or ICG. Slight leakage in FA and slight hypofluorescence on the late phase in ICG would suggest such a microcirculation disorder.

When better understanding of the aetiology and pathophysiology of AZOOR is available, the clinical response to appropriate therapy may perhaps best be followed by m-ERG.

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Hypotonic maculopathy following pneumatic retinopexy: a UBM study

Editor,—Pneumatic retinopexy is a promising surgical alternative in selected cases of retinal detachment.

We report a rare complication, to our knowledge the first ever reported, of this procedure—namely, hypotonic maculopathy, caused by a wound dehiscence on a patient who had previously undergone a standard extracapsular extraction with posterior chamber implant. Ultrasound biomicroscopy (UBM) proved to be a valuable adjunct in both the diagnosis and management of this complicated case.

CASE REPORT

A 63 year old white man, complaining of a shadow in his right visual field, was referred to our department in April 1997 for further management. Ocular history was notable for bilateral extracapsular cataract extraction with posterior chamber lens implantation (PC/ IOL); right eye in January 1997 and left eye in June 1996. Visual acuities were 20/25 in both eyes. IOPs were within normal limits. Anterior segment examination was unremarkable with well positioned IOLs. Fundal examination of his right eye revealed a supertemporal macula on rhegmatogenous retinal detachment, produced by a single horseshoe retinal tear and after additional cryoprobe manipulation caused the

Figure 1 UBM picture of right eye showing the external part of the dehiscent cataract wound, as a narrow slit, before repair (area between arrows). There is a full blown hypotonous maculopathy in this eye, with VA 20/200.

Figure 2 UBM picture of right eye showing the same area after repair. Note that the external part of the wound is totally closed (area between arrows) while internal wound gap appears to be more pronounced postoperatively. However, at this point the leak is rectified. IOP and macula are normal with VA of 20/50.

Between October 1997 and November 1997 he was presented to our unit on three occasions complaining of further deterioration of vision to 20/200, IOPs between 3–5 mm Hg, and a full blown hypotonous maculopathy. After 3 months of hypotony of “unknown origin”, a diagnosis was made by a glaucoma specialist using indentation gonioscopy revealing a dehiscence of the cataract wound. A preoperative UBM study confirmed the presence of an internal wound gap, behind the limbus, supertemporally, 3 mm in circumference. Ultrasonically the wound dehiscence was depicted as a narrow slit (Fig 1) with a flat inadvertent bleb above which was not apparent clinically. A surgical repair of the wound was decided upon. Intraoperatively no definite dehiscence could be clinically identified. Balanced salt solution through a 30 gauge needle was repeatedly injected under pressure to the anterior chamber but this failed to localise any suspicious area. At this point, based on the ultrasonic study, two 10-0 nylon interrupted sutures were placed through the sclera parallel to the limbus in the suspicious area. These bites were moderately deep in an attempt to engage the internal flap of the cataract wound.

Two weeks postoperatively, the pressure normalised (IOP 12 mm Hg), maculopathy was reversed, and visual acuity improved to 20/50.

Hypotonic maculopathy is an unusual and, to our knowledge, the first reported complication of this kind after pneumatic retinopexy. We hypothesise that the original cataract wound did not heal properly and the additional cryoprobe manipulation caused the

Figure 2 (Top left) In the multifocal ERG, the fundus was divided into four foci. (Top right) Sum of the amplitudes in each of four foci of the m-ERG were indicated.

Figure 2 (Bottom) The three dimensional topography and sum of the amplitudes in each of four foci of the m-ERG were indicated.
wound to leak. The UBM study provided us with an interesting insight into how the scleral suture repair may have worked to correct the wound leak as shown in Figure 2. The fact that hypotony resolved after suturing the wound, indicated that the external part of the wound was secure (no slit is apparent) despite the fact that the internal part of the wound was gaping even more postoperatively. This finding implies, therefore, that only minimal overall alteration of the wound architecture postoperatively, sufficient to rectify the leak.

We believe that UBM is a valuable adjunct in the management of similar cases by clearly identifying both the presence and exact location of leak. Finally, pneumatic retinopexy should be performed with caution, especially in cases of previously operated eyes with large incision wounds.

**COMMENT**

Involvement of the lens exclusively, sparing other ocular structures is rare. This case documents such a possibility and also highlights the salient features involving electric trauma to the lens. The scalp burn in this case represents the entrance wound for the electrical energy but the lack of an exit wound makes this case particularly peculiar. Both entry and exit sites for the electric current have been reported by all previous authors. The excellent surgical results noted in both eyes of this patient are in keeping with the similar result reported by Portellos et al. This observation should encourage the ophthalmologist to undertake surgery for electric cataract, where necessary, without any undue concern.

**Figure 1** Cross section through temporal artery showing disrupted internal elastic lamina (arrow) and large calcific plaques (C) in the tunica media (haematoxylin and eosin; original magnification ×40).

**Monckeberg’s sclerosis in temporal artery biopsy specimens**

**EDITOR,—**Temporal artery biopsies are performed routinely on patients suspected of having giant cell arteritis. Of 131 pathology specimens examined at University of Illinois at Chicago Eye Center from 1975 to 1998, the most common diagnosis was atherosclerosis with moyointimal fibrosis (63%) followed by giant cell arteritis (13%). In about 6% of cases we encountered calcific sclerosis confined to the tunica media which was associated with mild tissue disorganisation surrounding the calcific plaque and disruption of the internal elastic lamina (Fig 1).

Monckeberg’s sclerosis as seen in these specimens was first described by Monckeberg in 1903. It commonly affects medium size muscular arteries and is described in femoral, tibial, radial, coronary, cerebral, and visceral arteries. However, its association with the temporal artery is uncommon. The infrequent occurrence of this condition in the temporal artery and the presence of a fragmented inter-

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**Figure 2** The characteristic anterior subcapsular lens opacities. (A) Right eye, (B) left eye.

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**Figure 2** The characteristic anterior subcapsular lens opacities. (A) Right eye, (B) left eye.
nal elastic lamina should not be erroneously interpreted as sequelae of previous arterial inflammation. The pathophysiology of Monckeberg’s arteriosclerosis is still unclear, but it can be induced in animal models by injecting adrenalin, nicotine, parathyrionid hormone, and vitamin D. In addition, lumbar sympathectomy has been shown to promote occurrence of Monckeberg’s arteriosclerosis of the lower extremities in humans. Automatic dysfunction from diabetic neuropathy is thought to be responsible for the occurrence of Monckeberg’s in diabetic patients. Unlike atherosclerosis, Monckeberg’s arteriosclerosis is a benign condition and does not cause vascular thrombosis. In conclusion, Monckeberg’s arteriosclerosis of the temporal artery may be seen occasionally in the temporal artery. It is an interesting histological diagnosis that has little clinical significance but can be recognised in temporal biopsy material.

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Retinal vascular abnormality in Poland’s syndrome

EDITOR.—Poland’s syndrome is a congenital abnormality first described in 1841 consisting of unilateral hypoplasia or aplasia of the pectoralis major muscle and ipsilateral upper extremity abnormalities which often include ipsilateral syndactyly. Since then a number of associated anomalies have been reported. These include absence of the pectoralis minor muscle, absence or atrophy of ipsilateral ribs two to five, aplasia of the ipsilateral breast or nipple, and simian crease of the affected extremity. Although vascular alterations associated with this syndrome have been described, no involvement of eye vasculature has been reported so far.

CASE REPORT
We examined a 39 year old man previously diagnosed with Poland’s syndrome who came to our clinic because he had experienced blurred vision in the right eye for 2 months. Computed axial tomography, arteriography, and abdominal echography revealed absence of the left kidney. There was no history of diabetes or hypertension. The ophthalmological examination of the right eye revealed a visual acuity of 20/50 and a paracentral relative scotoma. The right eye fundus showed perimacular capillary telangiectasis, retinal vascular distortion, moderate perimacular hard exudates, and retinal swelling, all of them more prominent in the temporal perimacular area. Fluorescein angiography clearly showed the vascular abnormalities (Fig 1). The left eye fundus was normal. A diagnosis of juxtapfoveal retinal telangiectasis was made.

COMMENT
Juxtapfoveal telangiectasis is difficult to detect ophthalmoscopically and therefore can be neglected by routine eye fundus examinations made before the first visual symptoms appear. Unilateral renal agenesis occurs in approximately 1 per 1000 births and can be associated with vascular anomalies.


Bilateral optic disc oedema associated with latanoprost

EDITOR.—Latanoprost is a recent addition to the medical management of raised intraocular pressure in chronic open angle glaucoma and ocular hypertension. It is a potent ocular hypotensive agent with few ocular or systemic side effects.

We report a case of bilateral optic disc oedema developing soon after commencing treatment with latanoprost which resolved once therapy was stopped.

CASE REPORT
An asymptomatic 64 year old woman presented with raised intraocular pressure. She maintained good general health, had no significant medical history and was not on any medication. Snellen visual acuities were 6/5 in both eyes. The intraocular pressures were 28 mm Hg right eye and 26 mm Hg left eye. Ocular examination was otherwise unremarkable with open angles, normal optic nerves, and full Humphrey 24-2 visual fields. She was thus diagnosed as having ocular hypertension and consented to enter a prospective double masked trial comparing some of the intraocular pressure lowering drops. Therapy was commenced with one of the drugs involved in the study and at a 1 month review she reported no problems with the drops. The intraocular pressures had lowered to 16 mm Hg in both eyes and the examination was otherwise unchanged. At her third visit 2 months later, she was again asymptomatic with visual acuities of 6/5 in both eyes and intraocular pressures of 15 mm Hg. However, examination of the optic nerves revealed bilateral oedema which was more prominent in the left eye. There were no signs of uveitis in either eye, pupillary reflexes were normal, colour vision and Amsler testing were not affected, and the visual fields were full. At this point the code for the trial drug was broken and it was seen that she had been using latanoprost 0.005% eye drops at night to both eyes over the 3 month period. A neurological consultation failed to find any neurological abnormality and all haematological and biochemical analyses were normal. A computed tomography scan with and without contrast showed no abnormality and she was discharged from neurological review. Follow up in the eye clinic revealed no change after 72 hours. The latanoprost was stopped and the disc swelling had largely resolved at 1 week. By 10 weeks

Figure 1 Early venous phase fluorescein angiogram of the right eye, macular area. Note the easily visible telangiectatic retinal capillaries (arrows). These abnormal vessels showed significant leakage of fluorescein in the late phase of the angiogram.

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both optic nerves looked normal. Visual acuities were still 6/5 in both eyes and there was no loss of colour vision or visual field. The intraocular pressures had increased to 22 mm Hg in both eyes.

**COMMENT**

Latanoprost is a prostaglandin F₂α analogue which acts by increasing uveoscleral outflow. Side effects include increased iris pigmentation, hypertrichosis and increased eyelash pigmentation, anterior uveitis in patients with complicated glaucoma or in those having had previous incisional surgery, and cysoid macular oedema occurring soon after beginning latanoprost in pseudophakic or aphakic eyes. Ocular hypotony with choroidal effusions and facial rash have also been attributed to latanoprost. The best of our knowledge, optic disc oedema associated with latanoprost has not previously been described. The mechanism behind this association is unclear. One may not be surprised to see optic nerve swelling in association with signs of posterior uveitis or hypotony but in this case it occurred without any sign of ocular inflammation and the lowest recorded intraocular pressure was 15 mm Hg. It may be possible that the perfusion to the optic nerve heads via the short posterior ciliary arteries was compromised by a prostaglandin-like action manifesting as disc oedema and that latanoprost acid and prostaglandin F₂α, at high concentrations could cause vasoconstriction of bovine ciliary arteries and a similar action cannot be discounted in this case. The rapid resolution of the swelling with seemingly no long term sequelae once latanoprost was stopped would perhaps support this hypothesis.

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**Somatostatin scan positive gastrinoma ocular metastasis**

**EDITOR.—** Symptomatic ocular metastases are uncommon despite the 4% prevalence in patients dying of all types of malignancy in postmortem series. We report a case of ocular metastasis from a gastrinoma, which was part of the Wermer’s syndrome (multiple endocrine neoplasia (MEN) type 1), diagnosed by indium labelled octreotide scanning.

**CASE REPORT**

A 57 year old man presented with a 1 week history of blurring in his peripheral vision in his right eye and severe loss of visual acuity, worse early morning. He had been diagnosed with MEN type 1, 8 years previously after two perforated jejunal ulcers (1978, 1990) led to a diagnosis of Zollinger–Ellison syndrome, and a hyperplastic parathyroid gland had been removed for hypercalcaemia (1990). His mother had MEN type 1.

Ophthalmic examination revealed 6/12 acuity in the right eye and 6/5 in the left. His anterior segments were unremarkable. His right fundus showed a solid amelanotic lesion about one disc diameter above the right disc. Ultrasonography demonstrated a base of 13 mm and a height of 8 mm. He also had bilateral inferior retinoschisis. One month later the tumour base measured 14.5 mm and the thickness measured 7.9 mm. The posterior edge now practically abutted the optic disc (Fig 1). There was subretinal fluid accumulation.

A liver ultrasound scan showed multiple lesions and a tumour biopsy was composed of small solid islands of polygonal cells with granular cytoplasm, diagnostic of metastatic neuroendocrine carcinoma (immunostaining positive for chromogranin, neuron specific enolase, and NCAM, negative for S-100 and HMB45 (melanoma markers)). His urinary 5HIAA was marginally raised at 133 pmol/24 hours (normal range 0–40 pmol/l) but normal levels of other polypeptides.

No primary tumour or further metastases were seen using body computed tomograph and magnetic resonance imaging scans and the I-123 MIBG scan (meta-iodobenzyl guanidine) was also negative. An indium (In-111) labelled octreotide scan at 1 and 4 hours +SPET showed focal areas of increased uptake in the right orbit (Fig 2), the nasal region, mediastinum, multiple sites in the liver, and possibly other abdomen sites. However, a positive octreotide scan is not exclusively seen with neuroendocrine tumours, since other tissues have somatostatin receptors. These include high grade lymphoma, some small cell lung cancers, occasional tumours of the breast, and in chronic inflammatory conditions where there is T cell activation including endocrine ophthalmopathy with orbital involvement. Thus, although an absolute positive diagnosis of metastatic gastrinoma cannot be made definitively in the absence of histology, in the context of this clinical case it is highly probable that the choroidal tumour is due to ocular metastasis from gastrinoma.

**COMMENT**

Gastrinoma may occur sporadically or as part of multiple endocrine neoplasia. MEN type 1 is a rare disorder, usually inherited in an autosomal dominant fashion with high penetrance which affects multiple endocrine glands (hyperparathyroid hyperplasia in 80–90%, pancreatic islet tumours in 50%, and pituitary adenomas in 40–50%).

Ocular metastasis from gastrinoma has not previously been reported, although it has been reported in other neuroendocrine tumours—for example, carcinoid. Multiple or metastatic gastrinomas (especially if actively secreting) are best localised by a new radioisotope scan using radioactively labelled somatostatin analogues. Somatostatin is a peptide elaborated by the delta cells of the islets of Langerhans and the hypothalamus. It inhibits the secretion of gastrin by gastric mucosa and many other hormones such as insulin, thyrotropin, and corticotropin. Somatostatin receptors are found on neuroendocrine tumours (80% of gastrinomas) and indium labelled octreotide visualises all somatostatin receptor positive gastrinomas. Our scans revealed somatostatin avid liver metastases. Interestingly, the scan also very clearly confirmed the ocular/orbital disease. Orbital radiotherapy and chemotherapy were recommended.

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Retinal neovascularisation in Goltz syndrome (focal dermal hypoplasia)

EDITOR.—This is the first reported case of Goltz syndrome with documented peripheral retinal non-perfusion with subsequent retinal neovascularisation and vitreous haemorrhage.

In the eye this represents solely a mesodermal artefact. In-111 ocreotide scintigraphy in newly diagnosed endocrine gastroenteropancreatic tumour has been shown to be a valuable tool in the management of patients with Goltz syndrome. However, in our case, there was peripheral retinal non-perfusion and right temporal neovascularisation with subsequent neovascularisation within weeks.

COMMENT

Focal dermal hypoplasia is a rare disorder of ectodermal and mesodermal dysplasia originally described by Goltz et al. It is characterised by congenital atrophic skin changes often associated with herniation of the subcutaneous fat; skeletal anomalies, in particular syndactyly, polydactyly, or adactyly as well as coloboma, strabismus, cataract, and dental anomalies, especially hypodontia.

Ocular anomalies occur in 40% of cases. Coloboma have been reported in one third of cases, then less frequently microphthalmia, aniridia, heterochromia, and optic atrophy. Rarely ectropion, epiblepharon, miosis, and cataracts have been reported. The choroidal and retinal neovascularisation is a result of the retinal vessels appearing normal till the transitional zone.

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Retinitis sclopetaria associated with airbag inflation

EDITOR.—Chorioretinitis sclopetaria is a severe form of blunt trauma, caused by a high velocity object grazing the globe but not penetrating it. It is a concussion injury, which usually manifests as severe choroidal and retinal rupture associated with haemorrhage but an intact sclera. We report a case of chorioretinitis sclopetaria resulting from deployment of an airbag in a stationary motor vehicle.

CASE REPORT

A 32 year old pregnant woman was the driver in a car accident in which the airbag failed to deploy at the time of impact but inflated when she returned to sit in the driver’s seat approximately 5 minutes after the accident occurred. The patient was not wearing spectacles and there was no evidence of injury from other sources.

At presentation the visual acuity was hand movements in the left eye and 6/6 in the right eye. Examination revealed evidence of swollen eyelids and marked chemosis and subconjunctival haemorrhages on the left, with no bony injury and a full range of eye movements. Anterior segment examination revealed a clear left cornea and a quiet anterior chamber with a microhyphaema. The lens was clear and there was no angle recession and the intraocular pressure was normal.

The differential diagnosis of retinal vascular anomalies associated with skin lesions includes incontinentia pigmenti (IP) and Cockayne’s syndrome. Although focal skin atrophy may occur in IP, the initial skin lesions are vesicles and bullae which may later become pigmented maculae. The skin lesions in Cockayne’s syndrome are pigmented scars due to light sensitivity and trauma. Neither syndrome is associated with digital anomalies.

The genetic anomaly in Goltz syndrome remains to be determined. Most cases are sporadic. It is generally thought to be X-linked dominant with lethality in males, like IP; however, 9% of cases are male. These are proposed to be the result of half chromatid mutations. Deletions in the region of the chromosome Xp22 are a suggested site, though these must be differentiated from the deletions seen in microphthalmia with linear skin defects (MLS) and that of microphthalmia, dermal aplasia, and sclerocornea (MIDAS), which are now considered to be distinct entities. The wide variation in severity of expression is thought to be due to mosaicism.

Letters
pressure was recorded at less than 4 mm Hg. Fundal examination revealed evidence of a retinal tear although details were obscured by a diffuse vitreous haemorrhage. An ultrasound examination showed a vitreous haemorrhage and large retinal tear but no evidence of a scleral perforation.

An examination under anaesthesia performed the following day confirmed that there was no scleral rupture and indirect ophthalmoscopy confirmed the findings of diffuse vitreous haemorrhage and a retinal tear.

At the 2-week postoperative clinic visit, visual acuity had improved to 6/18. There was 2+ cells in the anterior chamber and the intraocular pressure was 10 mm Hg. Posterior segment findings were an intragel haemorrhage and retini s sclopeteria with no evidence of retinal detachment.

Four months later, the best corrected visual acuity was 6/36. The anterior chamber activity had settled; the intraocular pressure was recorded at less than 4 mm Hg. More crystalline deposits were found in the patients with more advanced retinopathy. It is supposed that corneal deposits and fundus deposits are essentially the same and it is suggested that crystalline retinopathy is caused by systemic abnormality. Although the exact pathogenesis of crystalline deposits is still uncertain, it is possible that destroyed fibroblasts appear to glitter or fibroblasts with crystalline-like deposits look glittering during break down. Further biochemical or cellular biological studies are needed to clarify these possibilities.

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Specular microscopic findings of corneal deposits in patients with Bietti’s crystalline corneal retinal dystrophy

Editor.—In 1937, Bietti first described three cases of tapetoretinal degeneration characterised by yellowish glistening retinal crystals, tapetoretinal degeneration with choroidal sclerosis, and marginal crystalline deposits of the cornea. Although more than 100 cases of crystalline retinopathy have been reported, crystalline deposits of the corneal limbus have been observed in only four out of 52 Japanese patients with crystalline retinopathy. Recently, observation using specular microscopy has been reported to be useful in detecting crystalline deposits at the limbus of patients with crystalline corneal retinal dystrophy. Therefore, in this study, we examined four patients with crystalline retinopathy using specular microscopy under a “con-surface” mode, which is used for the observation of the corneal surface, and we detected the deposits at the limbus.

Twelve months after the initial specular microscopic examination, we re inspected the crystalline deposits of two cases. Interestingly, the changes in the locations and forms of the crystalline deposits in the corneal limbus were exposed over time (Fig 1). More crystalline deposits were found in the patients with more advanced retinopathy. It is supposed that corneal deposits and fundus deposits are essentially the same and it is suggested that crystalline retinopathy is caused by systemic abnormality. Although the exact pathogenesis of crystalline deposits is still uncertain, it is possible that destroyed fibroblasts appear to glitter or fibroblasts with crystalline-like deposits look glittering during breakdown. Further biochemical or cellular biological studies are needed to clarify these possibilities.

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Retinal vascular abnormality in Poland's syndrome

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