develop. Treatment of susceptible organisms early in the disease with a combination of intravitreal amphotericin and oral imida-
zoles, such as flucconazole or ketoconazole, may, however, be curative and reduce or pre-
vent the need for surgical intervention.7 Even in those eyes with a vigorous vitritis or anterior uveitis, the number of organisms is low2 and microbiological confirmation of infection by needle biopsy of the vitreous has a low success rate,8 in contrast with that in bacterial endophthalmitis. Thus, there is a tendency to treat these eyes empirically avoiding vitrectomy, often to good effect, but the patient described above illustrates the value of isolating the infecting organism and determining its drug sensitivity. Because of early and profound intravitreal feeding, oral imidazoles could not be given, and of the available intravenous preparations, the organism was sensitive only to micona-
zole.

Although intravenous miconazole can penetrate the eye and achieve adequate therapeutic levels, this was not effective in this patient as demonstrated by the clinical appearance of progression of the lesion in the right eye. In eyes that are not very inflamed, the blood-retinal barrier may remain intact and systemic drug penetration of the eye may not be adequate. Intravenous amphotericin B has poor ocular penetration even in inflamed eyes but may be effective if used in sufficiently high dosage,9 or after vitrectomy when the blood-
retinal barrier has been compromised.10 The overall effect of vitrectomy on the therapeutic efficacy of intracameral amphotericin B is difficult to gauge since ocular clearance of the drug is also enhanced.10 Alternative choices of therapy for this patient included repeated intravitreal injec-
tions of amphotericin B at weekly intervals or intravitreal injection of miconazole; in view of the limited therapeutic options, progressive retinal changes, and good outcome following surgery in the left eye, right pars plana vitrec-
tomy and repeat intravitreal injection of amphotericin B were undertaken. This allowed decompression of the lesion and resulted in an excellent clinical outcome. In Candida endophthalmitis, progression of lesions in spite of medical treatment is an indication for vitrectomy to ensure that appropriate drug therapy is given and to reduce the risk of sight threatening complica-
tions.

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Congenital stationary nightblindness in a patient with osteopetrosis

CASE REPORT
The patient was a 36-year-old woman whose parents were first cousins. She and two of her three siblings had had nightblindness and high myopia since childhood, although we had no chance to examine her siblings. A chest x ray showed increased bone density and diffusely sclerotic bone structures. Computed tomography scan showed diffuse thickening and increased density of the skull bones (Fig 1). Narrowing of the optic canals was not observed radiologically. She was diag-
nosed as having adult onset osteopetrosis.

Her visual acuity was 20/25 with a correc-
tion of –8.0 D in both eyes. No abnormality was observed in eye movement or in the pupils. Funduscopically, myopic changes were observed. The margins of the optic discs were clear and the colour was good, although the discs were tilted (Fig 2). The dark adapta-
tion curve showed a rod–dark adapta-
tion. The light peak/dark trough ratios on electro-oculogram were 2:2.

The electroretinogram (ERG) was recorded using contact lens electrodes (Kyoto-Control-Lens, AE, Kyoto, Japan) as active electrodes. The light source was a stro-


scopic light (Nihon-Kohden, SLS3100, Tokyo). After 30 minutes of dark adaptation, scotopic b-wave, photopic flickering, and a single white flash responses were recorded with a blue dim light, 30 Hz flickering light, and a white flash light of 40 J, respectively. Single flash ERGs showed negative-type ERG. Scotopic ERGs were extinguished (Fig 2C).

Figure 2A (A) Chest x ray showed increased bone density and diffusely sclerotic bone structures indicative of osteopetrosis. (B) Computed tomographic scan of the head showed diffuse thickening and increased density of the skull bones.

COMMENT
CSNB has been classified into CSNB with abnormal fundi (Oguchi's disease and fundus albipunctatus) and CSNB with normal fundi.6 In CSNB with normal fundi, the fund-

us appearance is essentially normal, although some patients show changes associ-
ated with high myopia5 or optic disc changes (tilted, pale, or dysplastic).6 CSNB with normal fundi, which is inherited as an autosomal recessive, autosomal dominant, or X linked recessive trait, is classified into two types. Type I shows a markedly reduced ERG. Type II shows a negative type ERG; a normal a-wave with a markedly reduced or
absent b-wave. Type II CSNB is further divided into complete and incomplete types. Complete type CSNB shows refractive error ranging from moderate to high myopia and corrected visual acuity ranging from 20/200 to 20/25. Dark adaptation shows absent rod-dark adaptation. On single flash ERG, oscillatory potentials are absent. The scotopic ERG is extinguished whereas the photopic flicker responses are normal or subnormal. Accordingly, this patient was diagnosed as complete type CSNB.

Patients having osteopetrosis with visual disturbance caused by compression of the optic canal have been recommended for optic nerve decompression. However, primary retinal degeneration has been reported as a cause of visual disturbance: three cases of reduced or extinguished ERGs, two cases of macular degeneration, and a case of marked atrophy of the rod and cone layer with degenerative outer nuclear layer. In our case, CSNB was suggested to be the cause of visual dysfunction. Electrophysiologically examination is important in detecting retinal dysfunction in patients with osteopetrosis.

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An unusual impediment to spectacle wear: pilomatrixoma

EDITOR—Pilomatrixoma or calcifying epithelioma of Malherbe is a relatively uncommon benign skin tumour arising from cells of the hair follicle. Sixty per cent present during the first two decades of life although pilomatrixoma can occur at any age. It is more common in females by a ratio of 3:2. Although pilomatrixoma has a predilection for the upper lid and eyebrow, each ophthalmologist will see a relatively small number. A case of pilomatrixoma is presented with discussion of its management.

CASE REPORT

A 48-year-old woman developed a 'thumb sized' lump on her left upper eyelid over the course of 1 year. Because of a pathological fear of doctors and hospitals she had delayed her presentation until the size of the lump interfered with her wearing of spectacles.

She attended the clinic with a gauze dressing attached to her spectacle frame with which she had been attempting to conceal the mass’s increasing bulk. On examination she was found to have a very large tumour, approximately 6 cm long and 3 cm in diameter, attached by a broad base to the left upper eyelid and anterior aspect of the temporal region. Granulation tissue was evident on the surface with a purulent discharge (Fig 1A). There was no evidence of local or systemic metastasis.

She underwent primary diagnostic excision of the tumour, and since the patient insisted upon local anaesthesia and day-case surgery, the tissue defect was left to heal by secondary intention (Fig 1B).

Figure 2 Fundus appearance (A) right eye, (B) left eye). Myopic changes were observed. Optic discs were tilted; however, the colour of the disc was good. (C) Electroretinogram (ERG). Single white flash ERG showed a negative-type ERG (that is, a normal a-wave with a markedly reduced b-wave). The oscillatory potentials were absent. The scotopic ERG were extinguished in both eyes. Photopic flicker responses were only slightly reduced compared with those of a normal subject.

Figure 1 (A) A large 6 cm by 3 cm tumour mass involving the left upper lid and temporal region. The mass demonstrates granulation tissue with a mild purulent discharge. The patient had disguised this mass by means of a gauze dressing attached to her spectacle frame. (B) Appearance of left upper lid and brow 6 months after primary excision of pilomatrixoma allowing large wound defect to heal by secondary intention.
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