pigment epithelium or the choroid. A diagnosis of cryptococcal meningitis was eventually made and both the visual symptoms and the fundal abnormality resolved quickly on systemic treatment.

This is strong circumstantial evidence that the visual symptoms were due to the observed fundal abnormality and that Cryptococcus was the offending agent even in the absence of pathological proof.

The ophthalmologist plays a valuable role in the management of patients with AIDS since 70% of these patients have ocular disease.5-6 Cryptococcus neoformans is the most commonest ocular opportunistic infection and occurs in 30-40% of patients. The incidence of opportunistic infections which metastasise to the choroid is much lower and includes Cryptococcus neoformans, Mycobacterium avium, and Pneumocystis carinii.7 The prognosis for these patients is very poor by this stage but, nevertheless, the diagnosis may elude the physician until the choroidal involvement develops. It is, therefore, important for ophthalmologists to ‘recognise’ the pattern of choroidal involvement produced by opportunistic infections in AIDS as prompt treatment will prolong life.

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COMMENT
This case report describes a patient with visual symptoms and a rare fundal picture which, on fluorescein angiography, was consistent with pathology of either the retinal pigment epithelium or the choroid. Examination revealed a striking blotchy appearance of the retinal pigment epithelium and choroid, which looked like clouds beneath the retinas (Fig 1A). The optic discs and retinal vessels were normal. A fluorescein angiogram confirmed the presence of lesions which were underneath the neuroretina. These lesions masked fluorescence in an irregular pattern and there was no significant leakage in the late stages of the angiogram. The retinal component to the angiogram was normal (Fig 1B).

Although an opportunistic infection was suspected, this fundal picture was not recognised by us or other specialists in HIV and no treatment was given. During the next 2 weeks he became unwell with headaches and general lethargy. The differential diagnosis included cryptococcal meningitis, toxoplasmosa encephalitis, or cerebral lymphoma. Investigation showed a cryptococcic meningitis with an antigen level in both the CSF and serum of $1 \times 10^{10}$. He was treated with intravenous amphotericin for 10 days followed by a maintenance oral dose of fluconazole (400 mg daily).4 This resulted in regression not only of his systemic symptoms but also of his visual symptoms and dramatic resolution of the fundal changes over a 1 month period (Fig 2). A repeat lumbar puncture, 3 weeks later, showed mild improvement with reduction of the cryptococcal antigen level of $1 \times 10^{10}$.

Diagnosis and management of an occult cyclodialysis cleft

EDITOR—Non-intentional cyclodialysis may occur up to 6 months1 after anterior segment surgery or following trauma.3,4 The hypotony is due to aqeous outflow through the cleft to the uveoscleral pathway.5,6 In these patients, there is normal aqueous production dynamics,3,5,7,8 an abnormal outflow facility,2 the cleft size is unrelated to the degree of the hypotony and maybe microscopic and hence occult.9

CASE REPORT
Preoperatively Mrs AB, a 53-year-old white woman, had visual acuity of 6/18 in the left eye owing to cataract and perception of light in the right eye following retinal detachment. The left eye was otherwise normal and the IOP had varied from 11 to 15 mm Hg over the preceding 5 years. A routine extracapsular cataract extraction with a limbal section and posterior chamber lens implantation was performed. The eye maintained an IOP of 11 mm Hg until she banded her head 1 month later and the vision worsened. Examination revealed an IOP of 2 mm Hg and a visual acuity of 6/9. Gonioscopy did not reveal any cyclodialysis cleft. Four months postoperatively, the IOP remained at 2 mm Hg but the refraction was dramatically unstable as a result of blinking and eye movements. An 8-0 mm diameter hard contact lens stabilised the corneal topography and reduced the IOP to 6/6. There was no evidence of uveitis, no cleft was seen on gonioscopy, and ultrasound showed no evidence of choroidal or ciliary body detachment.

Laser flare studies revealed an anterior chamber flare count of 11 photon counts, which was within normal limits for her age. Topical timolol increases the anterior chamber protein concentration in normal eyes by reducing aqueous production. Two hours after administration of timolol drops, the aqueous protein concentration decreased by 35% and by 4 hours it had risen to 63%, compared with 75% in a normal eye. Following intravenous injection of fluorescein, an area of increased scleral fluorescence was demonstrated adjacent to the site of cataract section. These observations suggested an occult cyclodialysis cleft at the site of the previous surgical wound.

Gonioscopy with viscoelastic and surgical exploration of the wound failed to reveal a cleft. The wound was closed and the viscoelastic was removed. The IOP rose to 46 mm Hg at 12 hours and this required acetazolamide, mannitol, and levobunolol drops. By 10 days the IOP was 14 mm Hg with the patient receiving levobunolol and dexamethasone, the choroidal folds had resolved, and the visual acuity was 6/9.

For the next 10 months the IOP was 10 mm Hg with no medication and the visual acuity was stable at 6/9. At 11 months, hypotony and macula oedema suddenly redeveloped. Argon laser trabeculoplasty to the wound region was unsuccessful on two occasions. Surgical treatment was unhelpful as exploration of the original wound failed to find a cleft. The scleral flaps were closed and 12 hours later the IOP had risen to 55 mm Hg. The pressure fell slowly to 14 mm Hg over 14 days with medical treatment. One month later, the visual acuity was stable at 6/9 and the IOP has remained at 14 mm Hg on no treatment for 6 months.

COMMENT
The diagnosis of cyclodialysis cleft requires an assessment of aqueous production and the facility of outflow but the latter is not possible because the eye is hypotonic. Therefore the cleft requires visualising with gonioscopy, sometimes with perioperative chamber deepening. With an occult cleft the diagnosis depends on proving both normal aqueous production and an abnormal outflow pathway.

Aqueous production must fall to less than 10% of normal to produce hypotony4 so that any test which shows approximately normal aqueous dynamics excludes ciliary body dysfunction as the cause of hypotony. Laser flare measurements can quantify the amount of protein in the anterior chamber aqueous.

Figure 1B
Figure 1 (A) Right fundus showing diffusely abnormal pigment epithelium but normal retinal vessels. (B) Anterior ciliary phase of fluorescein angiogram of same eye confirms the presence of abnormal pigment epithelium with patchy mashing of the lesions.

Figure 2
Figure 2 Right fundus 4 weeks later showing considerable resolution of the lesions.
The presence of a normal flare count and the increased protein concentration by 63% after timolol was highly suggestive that the ciliary body was producing normal amounts of aqueous in this case. Similar observations can be made with fluorophotometry.

The anatomical site of the cleft was suggested by the scleral staining after an intravenous injection of fluorescein. An alternative method would be to inject the fluorescein directly into the anterior chamber but this was considered too hazardous in an only eye. However, the technique may be useful during surgery as drainage of fluorescein stained fluid from a sclerotomy site would indicate a patent cleft and that the sclerotomy is close to the cleft.

This case illustrates that occult clefts can be confirmed using laser flare measurements, provocative testing with timolol, and intravenous fluorescein. Hypotony induces a dynamically unstable refraction which can be stabilised with a hard contact lens. Closure of the cleft can then be attempted with argon laser, diathermy, cryotherapy, or surgery. If the distance across the cleft is small, the tissues may appose solely from the inflammatory swelling but if the distance is large, sutures or plombage may be required.

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