Chorioretinal biopsy in a patient with leukaemia

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SUMMARY The diagnosis of necrotising viral chorioretinitis in a child with leukaemia in remission was confirmed by the presence of viral particles on electron microscopy. Biopsy from the blind eye differentiated between leukaemic infiltrates and an infectious process after further investigations initially failed to support either diagnosis. Diagnostic chorioretinal biopsy may be indicated in a blind eye when the differential diagnosis includes conditions with radically different therapeutic alternatives.

The collection of chorioretinal tissue for diagnostic purposes has been considered unsafe until recently. Earlier reports in rabbits,1,2 monkeys,3 and man4 were associated with operative complications, particularly loss of vitreous humour. Recently a technique5 using systemic hypotensive anaesthesia has had no long-term sequelae in dogs6 or operative complications in human volunteers whose eyes were enucleated for malignant melanoma or glaucoma.7 We report a case in which chorioretinal biopsy was successfully used for diagnostic and therapeutic reasons in a child with T cell leukaemia in remission.

Case report

A 7-year-old boy was admitted to the Hospital for Sick Children, Great Ormond Street, on 18 February 1980 for routine reinvestigation following treatment of acute lymphatic leukaemia of the T cell type. He was originally diagnosed as having T cell leukaemia in September 1977 after presenting with fever and lymphadenopathy. Induction therapy consisted of 4 courses of cyclophosphamide, vincristine, cytosine arabinoside, and prednisolone (COAP). Central nervous system prophylaxis was given as 4 doses of intrathecal methotrexate and 2400 rads cranial irradiation. He received 2 years of maintenance therapy with alternating courses of COAP, 6 mercaptopurine, and methotrexate, and vincristine, adriamycin, and prednisolone every 3 weeks, remaining in continuous complete remission. The current admission was for testicular biopsy, bone marrow aspiration, and lumbar puncture before the anticipated discontinuation of chemotherapy.

The patient had had no visual symptoms, but the admitting haematologist noted numerous white retinal infiltrates in both eyes. Six weeks earlier the same doctor had found the fundi to be normal. Leukaemic infiltrates were suspected, and ophthalmic consultation was obtained.

The visual acuity was 6/24 in the right eye and 6/18 in the left eye. Both visual fields were constricted to approximately 15 degrees around fixation, tested by confrontation techniques. The patient could distinguish colours of objects in the room but formal testing was not done. Both retinas were deranged, right more than left, by multiple white lesions with a fluffy white border circumscribing a necrotic centre in which the inner retinal layers and much of the choroid and pigment epithelium were completely destroyed (Figs. 1 and 2). Many lesions were confluent. The peripheral retina was extensively involved in each eye, and the right parafoveal region was also affected. The retinal pigment epithelium was deranged with pigment dispersion and clumping. Very few vitreous cells were present posteriorly. Fluorescein angiography showed that the lesions were composed of relatively avascular centres surrounded by leaking retinal vessels.

It was felt that the differential diagnosis lay between leukaemic infiltration and opportunistic viral or fungal retinitis. Initial electron microscopy for cytomegalovirus particles in the urine was negative. Appropriate serum and urine samples for culture were submitted. The patient had a low white blood cell count on
area of the blind right eye was performed so that a definitive diagnosis might lead to appropriate treatment of the left eye. Cryotherapy to the biopsy site was not used. Vitreous fluid was cultured for virus particles, but no virus was grown.

Bone marrow aspiration, testicular biopsy, and lumbar puncture all confirmed continuing remission.

Because of the strong suspicion of opportunistic infection and the absence of other signs of leukaemic relapse, antiviral therapy was started on the day of biopsy. The patient received a 2-week course of intravenous adenine arabinoside, which was repeated after a 2-week intermission. Additionally, transfer factor was given daily for 1 week and then at weekly intervals.

admission to hospital (2.1 x 10⁹/l) with 30% lymphocytes. Within 3 days of admission the vision in the right eye deteriorated to no perception of light due to enlargement of the macular lesion. Trans-scleral chorioretinal biopsy (Fig. 3) of an apparently active

Fig. 1  The left fundus on admission. The macula was largely uninvolved and the acuity 6/18. There were white lesions nasal to the optic disc.

Fig. 2  The left fundus on admission. In the periphery there were several lesions with avascular centres, surrounded by a white ring which itself was surrounded by a small amount of haemorrhage.

Fig. 3  Chorioretinal biopsy. The suture retracts the right lateral rectus muscle, and a few fibres of the inferior oblique muscle are visible just under the lower arm of the lid speculum. A Desmarre’s retractor provides exposure for the 4 x 4 mm scleral trapdoor through which the choroid may be seen to bulge. Excision of a 1 x 3 mm triangle was accompanied by a gush of non-viscous clear fluid, but there was no haemorrhage. Profound hypotensive anaesthesia was not used.
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Fig. 4 Ultrastructural studies of the chorioretinal biopsy showed, in one area only, aggregatous of electron-dense bodies, irregularly circular in outline, and smaller lamellated virions. Bar marker = 1 μm.

Sections stained with toluidine blue revealed necrotic retinal tissue without either tumour cells or viral inclusion bodies on light microscopy. Electron microscopy (Fig. 4) showed intracytoplasmic particles within choroidal cells; the average measured size was 150 to 200 nm. These were identical to the dense bodies described by Craighead et al. in cytomegalovirus infected cells. These dense bodies were not virions but were found to have had similar antigenic determinants to the cytomegalovirus envelope and were subsequently clearly separated by special centrifugation techniques. Subsequently urine and serological studies supported the diagnosis of cytomegalovirus. Four to 6 urine cultures were positive for cytomegalovirus, though electron microscopy of urine repeatedly failed to show virus particles. Additionally the VK strain of papova virus was isolated on urine cultures. The cytomegalovirus complement fixation test antibody titre was positive at 1:32 dilutions on 25 February and rose to 1:256 by 11 March. Other studies included a rubella HA1 antibody titre of 256, a toxoplasma dye test titre of less than 1:16, and a herpes simplex complement fixation test antibody titre less than 8.

For 3 months after biopsy the patient's vision in the left eye remained 6/9 to 6/12 despite constriction of visual fields and an ill-defined difficulty in detecting one half of the Ishihara test plates. The averaged skin-electrode electroretinogram showed no definite response recognisable from either eye. Despite anti-viral therapy, however, the necrotising retinitis slowly worsened in the left eye (Figs. 5 and 6) to the point of total blindness in early June. The biopsy site in the right eye has healed without complication.

Discussion

The advent of multiple agent chemotherapy, and central nervous system radiotherapy prophylaxis, has markedly improved the prognosis for children with leukaemia. This treatment, however, may rarely be associated with significant morbidity. Isolated reports have attributed visual loss to toxic drug
effects,12 13 vascular changes secondary to high dosage radiotherapy,14 or opportunistic infection in the immunosuppressed patient.15–17 The clinician must consider these side effects of treatment when evaluating leukaemia patients who either complain of visual disturbance or who have atypical findings on physical examination. It may be very difficult to differentiate infectious infiltrates from leukaemia recurrent within the retina.16 18 A positive diagnosis dictates whether further treatment should consist of more antileukaemic chemotherapy and radiotherapy, or of antiviral, antifungal, or other therapy. Clinical judgment as well as serological and urine studies have thus far been the only guides to diagnosis.

Chorioretinal biopsy can provide a tissue diagnosis within days of the operation. Although the risks involved are not clearly defined, we suspect on the basis of animal experiments that they are low.9 We feel that at present chorioretinal biopsy may be indicated only in a blind eye and then only if a definitive diagnosis would lead to treatment which would either prevent blindness in the fellow eye or arrest a life-threatening systemic process. We do not believe, for instance, that chorioretinal biopsy is indicated in the retinal dystrophies. So far antiviral therapy has been of limited effectiveness against cytomegalovirus19–22 but has been shown to be effective against herpes simplex. Although our patient did indeed become completely blind, it is not known whether the treatment he was given delayed this outcome. It is hoped that further advances in antiviral therapy might provide a more specific and effective treatment for cytomegalovirus.

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