Treatment of CMV retinitis in an AIDS patient

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SUMMARY We present a case of cytomegalovirus (CMV) retinitis in an AIDS patient who survived for 10 months after the start of his ocular problems. The retinitis responded to dihydroxy propoxy methyl guanine (DHPG) but relapsed four to six weeks after each course of treatment with progressive retinal destruction. One relapse was therefore treated with trisodium phosphoformate hexahydrate (Foscarnet). There are few reports of the use of this drug in the treatment of CMV retinitis with AIDS, but it appeared to be less effective in our patient than DHPG, possibly because of poor penetration of the blood-ocular barrier. A final course of outpatient maintenance therapy with DHPG failed to prevent a preterminal relapse of the retinitis. Fundus photographs demonstrated the resolution and relapse of the retinitis associated with each course of treatment. Maintenance therapy with DHPG would appear to be necessary to prevent relapse, but the logistics of this are difficult, and the effective dosage of DHPG is as yet uncertain.

Patients with acquired immune deficiency syndrome are commonly found to have ocular lesions. Cotton-wool spots, isolated retinal haemorrhages, conjunctival Kaposi’s sarcoma, and cytomegalovirus (CMV) retinitis have been widely described.1–4 Other opportunistic infections occur less frequently and include cryptococcosis, toxoplasmosis, atypical mycobacterial infection, herpes simplex retinitis, and herpes zoster ophthalmicus.1,5 In a large series of eyes taken at necropsy from 35 consecutive AIDS patients CMV retinitis was the commonest infection, being found in 12 cases (18 eyes, 34%).6

The clinical features of CMV retinitis in AIDS are of progressive retinal destruction with fluffy white retinal infiltrates and intraretinal haemorrhages, often most marked along the major retinal vessels. Histologically there is full-thickness retinal necrosis with intracellular and intracytoplasmic inclusions. Virions may be identified on electron microscopy and virus antigen on indirect immunofluorescence. In AIDS it is not practical to confirm the diagnosis by obtaining intraocular specimens but the clinical diagnosis is supported by demonstrating cellular immune deficiency and human immunodeficiency virus (HIV, also known as HTLV III/LAV) antibody in association with positive CMV serology or virus isolation from blood, urine, and secretions.9

Pepose et al.4 have drawn attention to the frequent development of CMV retinitis in those AIDS patients with cotton-wool spots and propose that the cotton-wool spots are a manifestation of an immune complex mediated microvasculopathy which allows cell-free and cell-associated cytomegalovirus to infect the retina by traversing damaged retinal endothelial cells. They also distinguish between a non-inflammatory cytomegalovirus retinopathy seen in patients without AIDS and the cytomegalovirus retinitis seen in many patients with AIDS. In those with AIDS neutrophil function is preserved and may play a part in the progressive retinal necrosis of CMV retinitis. By contrast, infants, graft recipients, and patients with malignant disease often have defective neutrophil function, and when they develop disseminated CMV infection the retinopathy is less severe.

Experience of the specific treatment of disseminated CMV infection is limited, but over the past year remission of CMV pneumonitis, colitis, and retinitis in patients with immunodeficiency due to AIDS, allografts, and lymphoreticular neoplasms has been reported after administration of 9-(1,3-dihydroxy-2-propoxymethyl) guanine (DHPG, Syntex Research) and of trisodium phosphoformate hexahydrate (Foscarnet, Astra Pharmaceuticals). DHPG is also known as BW B759U (Burroughs Wellcome). We report our experience of the use of
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Figs. 1–7 Fundus photographs of the left macula and superotemporal retina demonstrate the retinopathy and effects of treatment and relapse.

Fig. 1 Fresh retinal infiltrates and haemorrhages at the macula before treatment.

both these agents in the treatment of one AIDS patient with relapsing CMV retinitis.

Case report

A 47-year-old male homosexual presented to the Eye Department in June 1985 with left visual loss. Seven weeks previously he had been treated with high-dose co-trimoxazole because of an atypical pneumonia presumed to be due to Pneumocystis carinii and at that time HIV antibody was demonstrated in the serum.

At presentation the right visual acuity was 6/5, N5, and the left was 6/36, N12. He read all the Ishihara colour plates correctly on the right and made four errors on the left. The eyes were white, and the anterior chambers were quiet. The right visual field, vitreous, and fundus was entirely normal. There was a dense central scotoma on the left side with a clearly demarcated 2-disc diameter area of intraretinal haemorrhages and white infiltrates at the macula. There were no cells in the left vitreous.

In July he developed oropharyngeal candidiasis and nystatin lozenges were prescribed. In late August the left visual acuity deteriorated to counting fingers, and fresh retinal lesions were seen (Fig. 1, 29 August 1985). A clinical diagnosis of CMV retinitis was made and dihydroxy propoxy methyl guanine (DHPG, Syntex Research) 10 mg/kg/day was administered for 21 days. The retinitis improved without change in the visual acuity (Figs. 2A, B, 9 October 1985). The patient felt better and his weight improved. No systemic side effects of treatment were observed. The white cell count and neutrophil count remained normal at 5.8x10⁹/l and 2.9x10⁹/l respectively, but there was a persistent absolute lymphopenia of below 1.0x10⁹/l. He developed oesophageal candidiasis,
which was treated with ketoconazole 200 mg once daily.

Six weeks later, in November 1985, the right visual acuity fell to 6/36, and fresh lesions consistent with CMV retinitis were seen in both eyes (Figs. 3A, B, 25 November 1985). CMV was isolated on cell culture from blood, urine, and a throat swab. BW B759U (Burroughs Welcome) 7.5 mg/kg/day was administered for 21 days, with stabilisation and partial resolution of the retinitis (Figs. 4A, B, 18 December 1985). The right visual acuity improved to 6/18 and the left remained counting fingers. The peripheral neutrophil count was preserved at 2.3×10⁹/litre.

Four weeks later, in January 1986, fresh retinal
lesions were observed (Figs. 5A, B, 13 January 1986), and CMV was again isolated from urine. In view of the previous relapses after treatment with DHPG, Foscarnet (1.5 g loading dose followed by 0.15 mg/kg/min) was given intravenously for 19 days with only a marginal clinical response (Figs. 6A, B, 6 February 1986). The patient developed marked superficial thrombophlebitis at the infusion sites, which made peripheral venous access increasingly difficult. No other adverse responses to this treatment were observed.

BW B759U 7·5 mg/kg/day was then given for 14 days, with marked improvement (Figs. 7A, B, 5 March 1986), and this time the patient was dis-
charged on a maintenance regimen of 5 mg/kg three times weekly. The retinal lesions remained stable for six weeks, when the patient was readmitted because of fresh retinitis, weight loss, malaise, and salmonellosis. DHPG 7.5 mg/kg/day was given for 21 days. The patient died on 12 May. The course of the retinitis and treatment periods are summarised in Fig. 8.

At necropsy, the cause of death was found to be bronchopneumonia. Histology supported the clinical diagnosis of CMV retinitis, but viral cultures were negative.

Discussion

9-(1,3-Dihydroxy-2-propoxymethyl) guanine (DHPG, also known as BW B759U) is an acyclic purine analogue with antiviral activity against all herpes viruses, with a 30-fold greater activity against cytomegalovirus than acyclovir. The triphosphate metabolite of DHPG competitively inhibits viral DNA polymerase in a similar manner to acyclovir, affording relatively selective activity against cytomegalovirus infected cells. The principal adverse reactions are due to dose-related cytotoxicity causing neutropenia, inhibition of spermatogenesis, and gastrointestinal mucosal atrophy and necrosis. DHPG is administered by intravenous infusion over one hour, and a therapeutic effect is achieved in doses of 2.5 mg to 15 mg/kg/day given in two or three divided doses. Shepp et al. have shown that neutropenia developed when peak and trough plasma levels exceeded 50 and 10 μmol/l respectively, and that these levels were found in three out of five patients receiving 15 mg/kg/day and in none receiving 7.5 mg/kg/day. However, others have reported neutropenia in two out of six patients at doses of 7.5 mg/kg/day. Penetration of the central nervous system is high, with levels 38% of simultaneous blood levels, and 91% of the daily dose is excreted unchanged in the urine.

Trisodium phosphoformate hexahydrate (Foscarnet) is an agent with activity against reverse transcriptase from a large number of retroviruses and DNA polymerases from herpes simplex types 1 and 2 and CMV. At plasma concentrations of 32–55 μmol/l cellular DNA polymerase is inhibited. Cell cultures of herpes viruses and the animal retrovirus visna (which is related to HIV) are inhibited by 100 μmol/l of Foscarnet. In-vitro inhibition of HIV has also been described. Cellular toxicity is low, and the drug is eliminated rapidly in the urine. Foscarnet is given continuously as a 2% solution by intravenous infusion because of its short half-life, and only small amounts enter the central nervous system. A loading dose of 10–20 mg/kg followed by a continuous infusion of 0.5–0.15 mg/kg/min is calculated to achieve a steady state plasma level of 50–150 μg/l. Adjustments are made according to renal function.

Clinical cases of resolution of CMV pneumonitis and retinitis in AIDS patients after the administration of DHPG have recently been reported from North America. This year the Collaborative DHPG Study Group has reported resolution of disease activity and clearance of viraemia in nine out
of 11 patients with CMV retinitis. Palestine et al. reported the results of treatment with DHPG of 14 eyes in eight patients with CMV retinitis, eight of which demonstrated more than 90% resolution and four a partial response. Two out of these patients had chemotherapy induced immunosuppression and achieved remission for two and six months respectively. The remaining six patients had AIDS. Three were followed up and relapsed within 32 days of stopping treatment. One went on to further treatment, followed by maintenance therapy in reduced doses because of neutropenia and died four months after presentation. Rosencan et al. reported a response in all of a group of six patients with AIDS. Three relapsed within 21–28 days of cessation of treatment and three were lost to follow-up. Further treatment of one patient again was limited by drug induced neutropenia. These studies indicate that an initial response to DHPG is often obtained but that early clinical and virological relapse is common and that reversible neutropenia is the principal adverse reaction.

Reports of the clinical efficacy of Foscarnet in the treatment of disseminated CMV infection in the immunodeficient have appeared recently. However, there is only one report of the treatment of CMV retinitis in AIDS. Singer et al. describe satisfactory resolution in one patient who also received adjunctive cytomegalovirus hyperimmune globulin and emphasise the relatively long remission of five months which was obtained.

We report a more satisfactory response to DHPG than to Foscarnet without using hyperimmune globulin. This apparent superiority may relate to limited penetration of the blood-brain barrier by Foscarnet. Our experience was similar to others' in that, although the retinitis responded initially to treatment, relapse invariably occurred 4–8 weeks later. Maintenance therapy with three times weekly DHPG did not affect the relapse time in our patient. Although Bach et al. reported a remission of over eight weeks in one patient, the role of maintenance treatment, its frequency, and the dosage require further clarification, especially in view of the practical problems associated with intravenous therapy in AIDS patients.

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References

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