Management of CMV retinitis in HIV infected patients

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Cytomegalovirus (CMV) infection is a very common opportunistic manifestation of HIV disease ranging from 10% to 40% during AIDS. \(^{1,2}\) CMV retinitis is the most common manifestation.

From 1991 to 1996, CMV infection had been diagnosed in 632/1971 (32%) AIDS patients in our unit, 65% of them having retinitis. After AIDS diagnosis, the risk of CMV visceral manifestations was 16% at 1 year, 34% at 2 years, and 46% at 3 years. The risk of CMV has increased over time owing to both a longer survival induced by antiretroviral therapies and a better management of opportunistic infections in patients with a persistent low immune status. In a natural history study of 1002 patients the probability of CMV disease at 2 years was 21-4% for patients with levels of CD4 below 100 \(\times 10^3\)l compared with 10-3% for those with CD4 greater than 100 \(\times 10^3\)l. \(^3\) Interestingly "the protease inhibitors era" which started in the USA and some European countries, in spring 1996 has dramatically changed the clinical spectrum of HIV infection. Approximately 50-80% of patients receiving highly potent antiretroviral therapy including protease inhibitors exhibit a median decrease of 1 to 2 \(\log_{10}\) HIV RNA viral load with 60-80% of them going below detectable level, with a significant increase in CD4 lymphocytes even when initially they have a very low immune status. This had led to a dramatic decrease in different clinical syndromes related to HIV, such as wasting syndrome, cryptococcosis, microsporidiosis, or atypical mycobacterial infection. The consequences of highly active antiviral therapy (HAART) on CMV infection have not yet been fully established; however, in our experience, the incidence of CMV disease has decreased from 24/100 patients a year to 18/100 patients a year in 1996 (C Katlama, unpublished data). Therefore, it is reasonable to expect that better treatment of HIV infection, allowing control of HIV replication and thus an improved stability in immunological status with delay in the incidence and the time of occurrence of CMV infection.

**Diagnosis of CMV retinitis**

**CIRCUMSTANCES OF DIAGNOSIS OF CMV RETINITIS**

Diagnosis of CMV retinitis is based on clinical findings and may be made in different circumstances.

**Presence of visual symptoms**

CMV retinitis commonly induces symptoms such as visual blur, floaters, and loss of peripheral vision which should lead to a full fundus-copic examination. CMV retinitis is not associated with pain, redness of eye, or photophobia unless associated with anterior uveitis.

**Systematic funduscopic examination**

In patients with low immune status (CD4 lymphocytes below 100 or 50 \(\times 10^3\)l) it should be common practice to regularly examine the retina through funduscopic examination which may reveal anterior retinitis. CMV retinitis can be detected by systematic funduscopic performed in a context of other manifestations or disease—for example, colitis or CMV visceral localisations.

**CMV reactivation markers**

These markers of CMV reactivation are strong predictors of development of CMV infection including retinitis. In a prospective study, it has been shown that 50% of patients with a positive CMV viraemia at the time of AIDS diagnosis will develop a CMV disease within a period of 8 months compared with 11% of those with a negative viraemia within a mean period of 11 months. \(^4\) Furthermore, in a multivariate analysis, the presence of positive viraemia had a positive predictive value seven times higher in predicting the risk of further development of CMV focal disease compared with the absolute number of CD4 lymphocytes. Several studies indicate that approximately 50% of patients who were polymerase chain reaction (PCR) CMV DNA positive will develop CMV disease. \(^5,6\)

**DIAGNOSTIC CRITERIA FOR CMV RETINITIS**

Owing to the impracticality of obtaining retinal tissue for histopathological examination, the diagnosis of CMV retinitis is made on the appearance of the characteristic perivascular fluffy yellow-white retinal infiltrates often associated with retinal haemorrhages. In some circumstances, lesions may be more granular rather than fluffy in appearance. Progression of retinitis occurs from the periphery to the central part of retina with an agranular, white leading edge leaving behind an atrophic scar. The location of lesions has important consequences for vision. Retinitis localised in the immediate vicinity of the macula (zone 1) can be rapidly sight threatening and should lead to immediate initiation of treatment. In contrast, lesions outside major vascular vessels (zone 2 or 3), commonly referred to as "peripheral retinitis", are not immediately sight threatening even when the treatment is deferred for a few days. An important point to keep in mind is the fact that "destroyed retinal tissue" cannot be restored and remains as scarring. Furthermore, active CMV proliferation during maintenance therapy is only slightly impaired, and
frequently reactivates. Many factors may account for this including poor penetration into the eye, too low dosages of anti-CMV drugs, or reduced sensitivity of the virus to the drugs. Therefore, once settled, the CMV necrotic process is almost never arrested but is characterised by successive reactivations followed by residual necrotic retinitis after several nucleoside phases, with subsequent progressive loss of vision.

CMV retinitis is characterised by a minimal vitreal and anterior chamber inflammation. Retinal lesions, at an early stage, may be confused with other ocular lesions such as cotton wool spots which are common in HIV patients, but these are not progressive on serial funduscopy examination. Other agents directly causing retinal necrosis, and less common than CMV, include herpes zoster or herpes simplex infections, which have a rapid course, or toxoplasmic choroidoretinitis which is accompanied by a more pronounced inflammation.

Treatment of CMV retinitis

**ANTIVIRAL DRUGS ACTIVE AGAINST CMV**

**Foscarnet**

Foscarnet is a pyrophosphate analogue which directly inhibits viral DNA polymerase without the need for prior phosphorylation to be active. It has a broad antiviral spectrum and is active against ganciclovir resistant CMV strains, herpes simplex (HSV) including aciclovir resistant HSV strains, and varicella zoster virus (VZV). It has potential activity against HHV8, the putative agent for Kaposi’s sarcoma.

Foscarnet is given by intravenous daily infusion—90 mg/kg twice daily as induction therapy for 2 to 3 weeks, and 90–120 mg/kg/day as one daily infusion in maintenance therapy.

Concomitant hydration with isotonic saline has considerably reduced the incidence of renal toxicity which represents the major adverse event of this therapy. Evaluation of oral hydration is ongoing but preliminary results during induction therapy suggest that oral hydration can be safely used instead of intravenous hydration. Other side effects include hypocalcaemia, genital ulcersation, and nausea/vomiting.

Foscarnet offers the advantage of being active on ganciclovir resistant strains, and it has an anti-HIV activity with an average 0:3 to 0:5 log reduction in HIV RNA plasma viral load; it also has a synergistic activity with nucleoside analogues such as zidovudine which may account for the survival benefit in comparison with ganciclovir (12:6 months versus 8:5 months; p = 0:007) observed in the SOCA trial.

**Ganciclovir**

Ganciclovir is a nucleoside analogue which requires to be phosphorylated in CMV infected cells. Mutations of thymidine kinase as modifications of UL97 gene are main causes of ganciclovir resistance which develops over time during prolonged ganciclovir treatment. Mutation on DNA polymerase has also been reported with cross resistance with cidofovir.

Ganciclovir exists as two formulations: intravenous ganciclovir represents the standard treatment given as 5 mg/kg twice daily as induction therapy for 14 to 21 days, and 5 mg/kg one daily as maintenance treatment; oral ganciclovir is hampered by a low bioavailability (5%) leading to use of a high daily dosage (3 g/day given as a 1 g regimen three times daily). Its lesser efficacy than intravenous formulation gives a median time of progression to CMV retinitis measured by retinal photographs of about 55 days compared with 65 days with the intravenous formulation. This difference was not statistically significant, but what was significant was the progression rate when observed by funduscopy examination (75 days with oral ganciclovir and 100 days with intravenous ganciclovir). The use of oral ganciclovir is limited to maintenance therapy and to primary prophylaxis of CMV retinitis.

The most limiting adverse reactions for ganciclovir are haematotoxicity, mainly neutropenia and anaemia. Growth factors such as G-CSF/GM-CSF may be used in patients with severe neutropenia when a switch to foscarnet is not possible.

Development of resistance of CMV to ganciclovir is correlated with the duration of drug exposure. In a study of 16 viral isolates from patients treated for more than a month, the percentages of both high level and low level ganciclovir resistance were respectively 19% and 81% when treatment was less than 9 months and 64% and 36% when treatment was longer than a year. Most of this resistance was correlated with the appearance of a mutation within the UL97 gene which encodes for the CMV specific transferase. Furthermore, high level ganciclovir resistance has a strong correlation with cidofovir resistance; some isolates of highly ganciclovir resistant strains might develop foscarnet resistance more quickly than expected. In brief, these data mean that the development of viral resistance is correlated with antiviral treatment which allows persistent viral growth; ganciclovir resistance is common over long term treatment and may induce cross resistance to other drugs.

**Cidofovir**

Cidofovir (Vistide HPMPC) is a nucleotide analogue of cytosine with potent in vitro and in vivo activity of prolonged duration against a broad spectrum of herpes viruses including CMV and herpes simplex type 1 and 2, VZV, and Epstein-Barr virus (EBV). Unlike aciclovir or ganciclovir which require intracellular activation by viral encoded enzyme, conversion of cidofovir to cidofovir diphosphate, which is the active cellular metabolite, is performed by host cellular enzymes and is therefore independent of viral infection. The diphosphate has a long intracellular half life (17–65 hours) which results in prolonged antiviral effects of cidofovir. The major dose
limiting toxicity is dose dependent nephrotoxicity leading to degeneration and necrosis of renal proximal tubule cells. Concomitant administration of probenecid is thought to reduce the toxicity by competing with cidofovir for uptake at the surface of the tubule thereby minimising cidofovir toxicity, although the pharmacokinetics is complicated and probably irrelevant at the lower dose of 3 mg/kg.

Cidofovir is administered in acute therapy as 5 mg/kg once a week for 2 weeks, followed by 5 mg/kg every 2 weeks as maintenance therapy, and prehydration with a 1 litre intravenous infusion of normal saline. Oral probenecid is given as a 2 g dose in the 3 hours before infusion, and 1 g, 2 and 8 hours after infusion.

When there is concomitant use of probenecid, zidovudine clearance decreases by 50%, so patients should be advised to decrease by 50% or interrupt zidovudine use on each day of cidofovir injection.

In a prospective controlled trial\(^20\) comparing immediate treatment with cidofovir (5 mg/kg weekly) with other treatments in non-sight threatening retinitis, the median time to progression of retinitis was 22 days compared with 120 days for the immediate treatment group (p < 0.0001). In this study, proteinuria occurred in 12% of patients, elevation of serum creatinine in 5% of patients; overall, 24% of patients discontinued treatment because of nephrotoxicity. Renal toxicity may not always be reversible. Early immunological symptoms of renal tubular necrosis such as hypophosphorhaemia, hypoprotinaemia, and hypoglycaemia should be looked for. Patients with pre-existing renal dysfunction or mild proteinuria should avoid the use of cidofovir. After two or three cidofovir treatments 50–60% of patients had evidence of mild to moderate probenecid reactions, most commonly consisting of fever, chills, and rash.\(^{20,21}\)

Cidofovir is currently licensed for the treatment of CMV retinitis and in Europe for patients failing standard therapy.

Although cidofovir is not an easy drug to manage, because of its potential side effects, the major advantage is the possibility for maintenance therapy to be given as one injection every 2 weeks. The impact on quality of life, especially as the patient does not need an indwelling central catheter, compared with daily administrations of foscarnet or ganciclovir is of importance.

**Lobucavir**

Lobucavir is a guanine nucleoside analogue with a broad spectrum antiviral activity against CMV, herpes simplex, and VZV; interestingly, the compound has also in vitro inhibiting activity on HIV replication in monocyte-macrophage cell line, probably through inhibition of reverse transcriptase activity and it appears also to have in vitro activity against hepatitis B. Bioavailability following oral administration of lobucavir varies around 50%.\(^{22,23}\)

Phase I/II studies of lobucavir have demonstrated an in vivo activity on CMV treatment in urine or semen. Phase III trials in treatment and prophylaxis of retinitis are planned.

**Adefovir**

Adefovir (PMEA) is a nucleotide analogue of adenosine with activity on a broad spectrum of retro and herpes viruses including HIV and CMV.\(^{24,25}\) Phosphorylation of PMEA in cells to PMEApp is independent of virus infection; thus, PMEA may prime uninfected cells to resist viral infection when subsequently infected. Because of low oral bioavailability, several compounds have been synthesised to circumvent this problem; among them adefovir dipivoxil (bis POM PMEA) has an advantageous pharmacokinetic profile. Preliminary data suggest that adefovir leads to a 0·6 log\(_{10}\) reduction in HIV viral load and 0·3 to 0·7 log\(_{10}\) decrease in CMV viral load when studied in a small number of patients. A large phase III placebo controlled trial with adefovir both as anti-HIV and prophylactic drug is being planned.

**Prodrug of ganciclovir**

Valganciclovir is the valine ester of ganciclovir, a prodrug that is de-esterified to produce ganciclovir in the peripheral blood A dose ranging study\(^26\) in 32 HIV positive patients—460 mg, 875 mg, 1750 mg, 2625 mg once daily—has shown that the bioavailability of the drug is approximately 60% and increased by 20% with food absorption. A single daily dose of 900 mg would achieve concentrations that are equivalent to a dosage of 5 mg/kg/day intravenous ganciclovir.\(^{26}\) Further studies to evaluate clinical efficacy of valganciclovir in induction and maintenance therapy of CMV retinitis are in progress.

**LOCAL THERAPY**

The rationale for using local therapy is to administer a specific treatment at the site of the infectious process, thus avoiding all the complications of systemic therapy. Several local treatment strategies have been used over time.

**Intravitreal injections**

Ganciclovir has been the drug used most often. Intravitreal injections with ganciclovir include one injection twice weekly as induction therapy followed by one injection weekly as maintenance therapy (400 µg ganciclovir per injection). Although no randomised studies have compared intravitreal injections with intravenous ganciclovir, results demonstrate a similar efficacy of the two treatments for progression in the affected eye.

**Intravitreal foscarnet**

There have been fewer experiences with intravitreal injections of foscarnet which is not easy to infuse given the large volume needed.

**Intravitreal implants**

An important approach has been the concept of permanent intravitreal devices to deliver ganciclovir continuously over several months.
(up to 6–8 months). These intraocular ganciclovir implants have been evaluated in two randomised studies in comparison with intravenous ganciclovir. Better control of CMV retinitis was obtained with ganciclovir implants (2 μg/h) with a median time of progression of 180–200 days; a higher risk of extracocular disease (15%) was usually observed and the development of CMV retinitis in the fellow eye, with 50% of cases within 6 months.27 28

The main advantages of local therapy, particularly with intraocular ganciclovir implants, is a better control of CMV retinitis, compared with systemic intravenous infusion and a major improvement in quality of life avoiding any systemic potentially toxic therapy. However, the major drawback of this local therapeutic strategy is to treat only the eye affected by CMV and not the whole CMV pathological process, with the consequence of development of retinitis in the fellow eye or occurrence of extracocular CMV complications. Other specific complications of local therapy include conjunctival bleeding and infection; retinal detachment, a common complication in the long term CMV retinal process, might be facilitated by regular injections.

MANAGEMENT OF CMV RETINITIS THERAPY

One particular feature of CMV retinitis is its high tendency to relapse. Because of an overall longer survival, patients may experience several acute CMV retinitis episodes. Therefore, it is important to adapt to each individual clinical situation and give the optimal therapy. The acute treatment of CMV retinitis necrosis optimally consists of 2 or 3 weeks of intravenous treatment. The choice between foscarnet and ganciclovir will depend on various factors such as concurrent or previous anti-CMV treatment, renal function, the polymorphonuclear cell count and haemoglobin level, the current clinical and social status of the patient, and any concomitant antineoplastic chemotherapy.

Reasons to choose ganciclovir are ease of administration, and a good clinical tolerance, whereas reasons to prefer foscarnet are a proved or possible resistant CMV strain to ganciclovir, neutropenia or low haemoglobin level, a specific anti-HIV effect that may complement antiretroviral therapy, and the absence of need for expensive leucocyte growth factors.

In cases of relapsing retinitis, it might be wise to change the treatment, especially if time to relapse has been short, suggesting a lesser sensitivity of the virus. It has been demonstrated that a combination of foscarnet and ganciclovir was more effective in halting progression of CMV retinitis in patients with multiple CMV retinitis episodes.

Local therapy, mainly with intravitreal implants, may be used in the absence of systemic CMV disease.

CMV prophylaxis

The ideal management of any infectious disease is a preventive strategy, particularly in infections with severe morbidity. Therefore, unsurprisingly, there have been different approaches to study the possibility of primary prophylaxis of CMV infection.

ORAL GANCICLOVIR

Two randomised placebo controlled phase III studies have been completed,30 31 with controversial results. In study 1654 from Roche, oral ganciclovir (3 g per day) led to a 50% reduction in the risk of CMV disease compared with placebo with a occurrence rate of CMV retinitis in the ganciclovir group of 24% compared with 49% after 18 months.3

The CPCR study (994 patients) had planned a different management strategy, where retinal lesions were not systematically looked for and funduscopy examination performed only after occurrence of visual symptoms; after a 12 month median time of follow up, there was no significant difference regarding the occurrence of CMV disease in the oral ganciclovir group (11%) compared with the placebo group (12%); one suggested hypothesis for the difference observed has been the protective role of DDI on CMV while combination of DDI and oral ganciclovir might have a negative effect.31

Overall these data suggest that the prophylactic role of CMV is far from being optimal and, given the benefit-risk ratio, it appears necessary to better define the population of HIV infected patients who might benefit the most from prophylaxis. Interestingly, in study 1654, the analysis of CMV viral load measured by PCR in correlation with the occurrence of CMV visceral disease shows that:

- a positive viral load was correlated to the occurrence of CMV disease;
- patients who were CMV PCR positive at baseline had a CMV disease rate of 26% in the ganciclovir group compared with 43% in the placebo group (p = 0.017). However, in patients with CMV viral load greater than 50 000 copies, there was no efficacy of ganciclovir compared with placebo (75–100% CMV events in each group).
- the best prophylactic index was obtained in patients with negative CMV PCR at entry, where the rate of CMV disease was 1% in the ganciclovir group compared with 14% in the placebo (p < 0.001); however, the low risk of CMV in this group decreases the interest in any prophylactic regimen which is potentially toxic, represents a high number of pills (12/day), and is expensive. Oral ganciclovir is the only drug approved for prophylaxis of CMV retinitis in patients with less than 100 CD4 lymphocytes × 109/l.

VALACICLOVIR

Valaciclovir is the prodrug of aciclovir and allows plasma concentrations equivalent to that observed with aciclovir. A large phase III study32 had evaluated valaciclovir (2 g × 4 per day) versus two dosages of aciclovir (high dose 800 mg × 4 per day/low dose 400 mg × 4 per day). Overall, the valaciclovir group experienced less retinitis and less CMV organ dis-
eases. The time to confirm CMV disease was significantly longer in the valaciclovir arm that in the two aciclovir groups. However, there was a trend (p = 0.06) towards earlier mortality in the valaciclovir group compared with low dose aciclovir but not compared with high dose aciclovir for which, to this date, the reason remains unknown. Furthermore, an unusual number of patients with haemolytic uraemic syndrome/thrombotic thrombocytopenic purpura was observed with valaciclovir. Further work is required to identify a dose of valaciclovir which is better elevated while maintaining a CMV protective effect and which provides a survival rate similar to that of aciclovir. For all these reasons, valaciclovir is not, in practice, used as prophylactic agent in HIV.

Conclusion
The control of CMV infection is a major challenge given the morbidity/mortality of this viral disease in HIV infection. In the past few years, important progress has been made regarding diagnosis, predictive virological markers, and quantification of CMV viral load for a better understanding of CMV disease; investigation of new drugs, with broad viral spectrum of activity, will be of importance for a better clinical management of patients with immune suppression.

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