Current Management Strategies for the Prevention and Treatment of Cytomegalovirus Infection in Pediatric Transplant Recipients

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Abstract

Cytomegalovirus (CMV) is a significant cause of morbidity and mortality following transplantation, especially in the pediatric population, who remain at high risk of primary infection. The availability of effective antiviral therapy has led to dramatic improvements in the outcome of CMV infection in patients undergoing transplantation. In recent years, three major strategies have been developed for the prevention of CMV disease in this population: (i) reduction of risk of viral acquisition or reactivation by management of risk factors; (ii) prophylaxis of all ‘at-risk’ patients using prophylactic strategies for a defined period of time, initiated at or near the time of transplant; and (iii) pre-emptive treatment with ganciclovir of selected ‘at-risk’ patients, guided by either laboratory markers indicative of subclinical infection or the presence of specific risk factors.

In general, well designed comparative studies of one or more antiviral agents for the prevention of CMV have not been carried out. While ganciclovir appears to be more effective than aciclovir, its tolerability profile is less optimal. The use of foscarin avoids myelosuppression, but is associated with significant nephrotoxicity. Its use should be reserved for patients unable to tolerate ganciclovir or with ganciclovir-resistant CMV disease. Similar to foscarin, the high frequency of nephrotoxicity associated with the use of cidofovir limits its use to clinical scenarios suggestive of ganciclovir resistance. Newer options, such as valaciclovir and valganciclovir, are currently under investigation and preliminary experience has been promising. Finally, passive immunoprophylaxis has been shown to prevent CMV disease after solid organ transplantation, but its use in bone marrow transplantation is controversial.
Essentially, pre-emptive strategies have relied on the quantitation in the peripheral blood of CMV phosphoprotein pp65 antigen and/or the polymerase chain reaction assay. Strict guidelines for the use of these assays as a guide to pre-emptive therapy have not been standardized. Prospective trials comparing pre-emptive therapy using either intravenous or oral ganciclovir, and now oral valganciclovir or valacyclovir, are necessary to determine the relative cost effectiveness and efficacy of these alternative strategies. Finally, it remains controversial as to whether prophylaxis or pre-emptive therapy is the optimal strategy for preventing CMV disease. While a growing body of literature describes these approaches in adult transplant recipients, published experience in children has been much more limited.

Infection with cytomegalovirus (CMV) has long been recognized as a major cause of morbidity and mortality after solid organ transplantation (SOT) and bone marrow transplantation (BMT).\(^1\) Pediatric recipients are a high risk group for this complication as a considerable proportion of them are CMV-seronegative and receive organs from CMV-seropositive donors. In the absence of preventive strategies, between 25 and 80% of SOT and allogeneic BMT recipients will develop CMV infection, with symptomatic disease developing in 8 to 41% of these patients.\(^2\) Further, recurrent episodes of CMV disease have been reported to occur in 6 to 59% of SOT recipients.\(^3\) Most of these studies have combined both children and adults. The incidence of CMV is much lower after autologous BMT and peripheral-blood stem cell transplantation than after allogeneic BMT.\(^4\) Although CMV infection continues to be an important cause of morbidity in transplant recipients, the use of ganciclovir for the treatment and prevention of CMV disease has led to a dramatic decline in CMV-associated mortality in these patients. This review discusses the current status of the treatment and prevention of CMV disease in children undergoing transplantation.

1. Epidemiology of Cytomegalovirus (CMV) Infection in Transplant Recipients

Children undergoing transplantation are at risk of developing either primary (first-time infection in a CMV-seronegative patient) or secondary (reactivation of latent virus or reinfection with a new strain) CMV infection. The major source of CMV in children experiencing primary infection after transplant is either the donor organ (including the possibility of transmission via passenger leukocytes that accompany the graft) or blood products. CMV infection is defined by the presence of viral replication (CMV-positive culture from any site or seroconversion in a previously seronegative patient) in the absence of any symptoms. CMV disease is defined by the presence of symptoms attributable to active CMV replication in a patient with a positive culture or histological evidence of CMV on a tissue biopsy in the absence of another pathogen to explain these symptoms.

Table I shows the principal risk factors associated with CMV infection and disease. Patients with primary CMV infection are at the greatest risk of developing CMV disease and experiencing CMV-associated morbidity; thus, CMV-seronegative recipients of organs from CMV-seropositive donors (D+/R−) are at the highest risk of developing symptomatic infection with CMV. Secondary CMV infection is more frequently associated with asymptomatic infection. When secondary infection is symptomatic, the risk of developing invasive disease is significantly lower than for patients experiencing primary CMV infection. The type and intensity of immunosuppression also has a strong influence on the risk of developing CMV disease; the higher the levels of immunosuppression, the higher the risk. Different forms of immunosuppression used in organ transplantation affect different aspects of CMV infection; antilymphocyte antibodies (e.g., antithymocyte globulin, muromonab CD3 [OKT3]) and cytotoxic drugs enhance viral activation from latency whereas cyclosporin, tacrolimus and corticosteroids promote the persistence and spread of the virus by suppressing the host’s antiviral immune responses.\(^5\) Also, a higher incidence of CMV disease has been described in recipients of kidney allografts receiving mycophenolate mofetil compared with those who received standard immunosuppressive regimens.\(^6\) Mycophenolate mofetil selectively suppresses the proliferation of B and T lymphocytes, and it may influence viral replication by altering the cytokine profile or modulating the down-regulation of adhesion molecules induced by CMV.\(^7\) Other factors, including rejection, systemic infection and inflammation have also been associated with an increased rate of CMV infection, presumably by reactivation of CMV from latency.\(^8\) Finally, the type of organ transplant has been associated with a variable risk of devel-

<table>
<thead>
<tr>
<th>Table I. Risk factors for CMV acquisition after transplantation</th>
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<tbody>
<tr>
<td>CMV-seronegative recipients who receive organs from CMV-seropositive donors</td>
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<tr>
<td>Use of CMV-seropositive blood transfusion</td>
</tr>
<tr>
<td>Use of intensive immunosuppression with antilymphocyte antibodies</td>
</tr>
<tr>
<td>Viral load of transplanted organ: more pronounced in lungs, intestine and pancreas</td>
</tr>
<tr>
<td>Situation with cytokine release: infections, surgery, stress situations</td>
</tr>
<tr>
<td>Acute graft-versus-host disease (after BMT)</td>
</tr>
<tr>
<td>Total body irradiation before transplantation (after BMT)</td>
</tr>
</tbody>
</table>

BMT = bone marrow transplantation; CMV = cytomegalovirus.
opining CMV disease. Recipients of intestinal and lung transplantation appear to be at a higher risk of developing CMV disease than patients undergoing other types of SOT. This enhanced risk is probably related to the higher viral load of the organ, and the higher levels of immunosuppression required to maintain these grafts compared with other types of SOT.

2. Clinical Manifestations of CMV

The peak incidence of CMV disease traditionally occurs 45 to 60 days after SOT and BMT.12-8,11 The range of clinically apparent effects of CMV in transplant recipients includes fever, leukopenia, thrombocytopenia and mild atypical lymphocytosis; invasive disease may also involve the liver, lungs, and gastrointestinal tract. CMV retinitis is uncommon after transplantation. Of interest, the allograft tends to be the most common site of tissue invasive disease. After BMT, pneumonia and gastrointestinal disease are the most common manifestations. In addition to these direct clinical effects, CMV has also been associated with a number of indirect effects. Active CMV infection is thought to increase the net state of immunosuppression, predisposing patients to opportunistic super-infections with bacteria, protozoa and fungi.13,16 It has also been suggested that CMV contributes to acute and chronic allograft rejection and injury, and to decreased long-term survival of SOT recipients.14,15-22 A history of CMV infection has been associated with an increased incidence of obliterative bronchiolitis after lung transplantation,20 progressive atherosclerosis after heart transplantation,21 and vanishing bile duct syndrome after liver transplantation.22 All of those entities are manifestations of chronic rejection. Whether CMV increases the risk for acute and chronic graft versus host disease after BMT remains controversial.23,24

3. Diagnosis of CMV

Diagnostic studies for CMV include serology, histology and virological tests, as well as antigenemia assays and polymerase chain reaction (PCR) techniques.

3.1 Serology

Serology is very important in defining the clinical risk from CMV at the time of transplantation; however, serology is a poor diagnostic tool after transplantation because of the immune response alteration.

3.2 Histopathology

Historically, CMV has been diagnosed by histology, with the visualization of cytomegalic inclusion bodies on tissue specimen. In addition, the use of immunohistochemistry has increased the sensitivity of histological diagnosis.26 An in situ DNA hybridization technique is less sensitive but highly specific; however, histological diagnosis is limited by the need to use invasive procedures to obtain samples.

3.3 Viral Isolation

Conventional detection of CMV in clinical specimens has been by direct viral culture in human fibroblasts, with follow-up visual examination for cytopathic effects over a period of 14 to 28 days. This lengthy period of time required for diagnostic confirmation limits its clinical usefulness. With the development of the shell vial assay using a monoclonal antibody directed at the immediate-early viral antigen, CMV can be detected earlier (1 to 2 days) with high accuracy.

3.4 CMV-pp65 Antigenemia Assay

The CMV-specific lower-matrix phosphoprotein pp65 is produced and secreted at the site of active infection, and subsequently phagocytosed by leukocytes; thus, CMV-pp65 antigen is identifiable in the nuclei of peripheral polymorphonuclear cells of patients with active CMV infection but is not detectable in patients where CMV is in a latent state.27 The CMV-pp65 antigen is visualized and quantitated under direct microscopy by either immunoperoxidase staining or an indirect immunofluorescence assay.27-29 In contrast to viral cultures, the antigenemia assay has a short processing time (3 to 5 hours) and a high diagnostic accuracy (being more sensitive than shell vial culture), and stays positive longer than culture after the institution of antiviral therapy.13,30,31 Elevated levels of the antigen are associated with the presence of, or progression to, CMV disease; the higher the level the greater the risk of disease or viremia. Hence, the ability to quantitate results of the pp65 assay allows the potential to stratify the risk of progression from asymptomatic infection to symptomatic disease based upon the height of the antigenemia assay. A positive assay result is the identification of anywhere from 1 to >1000 positively staining nuclei per 2 x 10⁴ leukocytes. In general it can be concluded that the greater the value of the antigenemia assay, the higher the risk of progression to symptomatic disease. Further experience in SOT patients has demonstrated that the assay becomes positive days to weeks in advance of clinical symptoms; therefore, this test should be able to serve as the basis for initiation of pre-emptive treatment of asymptomatic CMV infection.
3.5 CMV Polymerase Chain Reaction (PCR)

Nucleic acid amplification by PCR on DNA or RNA extracted from infected leukocytes allows the rapid diagnosis (same day) of CMV infection.\textsuperscript{13,32} Also, PCR techniques can detect CMV DNA in whole blood\textsuperscript{135} and cell-free body fluids such as serum and plasma.\textsuperscript{136,37} Unfortunately, although simple qualitative PCR techniques are very sensitive and become positive several weeks prior to the onset of symptoms, the positive predictive value of these assays for the development of disease is only around 50%.\textsuperscript{138,39} In an effort to improve specificity, quantitative PCR assays for the measurement of the CMV viral load in the peripheral blood leukocytes have been developed to be used as a guide for preemptive therapy.\textsuperscript{40} Additional amplification strategies to PCR have been developed and include the hybrid capture assay,\textsuperscript{41} the branched DNA assay,\textsuperscript{42} and nucleic acid sequence-based amplification.\textsuperscript{133,42} Each of these strategies serves as an alternative quantitative test to detect CMV DNA or RNA in whole blood or leukocytes.

4. Prevention of CMV Disease

Over the last decade, three major strategies have been developed for the prevention of CMV disease after transplantation: (i) reduction of risk of viral acquisition or reactivation by management of two risk factors (donor/recipient serological status, and blood transfusion); (ii) prophylaxis of all “at-risk” patients (using either an antiviral agent or an intravenous immunoglobulin product) for a defined period of time, initiated near the time of transplantation; and (iii) pre-emptive treatment with ganciclovir of selected patients guided by either laboratory markers indicative of subclinical infection or the presence of specific risk factors (e.g., exposure to augmented immunosuppression). While a growing amount of literature describes these approaches in adult transplant recipients, published experience in children has been much more limited. A consensus does not currently exist concerning whether prophylactic or pre-emptive therapy is the optimal strategy for preventing CMV disease. Further work is needed to identify whether specific preventive strategies will work equally well in recipients of different types of transplantation. This is particularly true for recipients of intestinal, lung and BMT in whom the consequences of CMV disease appear to be greater than other types of transplantation.

4.1 Reduction of Risk of Viral Acquisition

Many children undergoing transplantation are “immunologically naïve” to CMV and, therefore, are at high risk of acquiring a primary infection. An obvious strategy for the prevention of CMV disease in these children would be to only use organs from seronegative donors. In general, this approach is not practical because of the limited pool of seronegative donors; however, this policy has been recommended for CMV-seronegative candidates awaiting isolated intestine allografts in whom the transplant procedure may be deemed to be “elective”, and in whom the outcome of primary CMV infection continues to be associated with unacceptable morbidity and mortality.\textsuperscript{43} This approach has also been adopted by some pediatric lung transplant centers. One way that centers might maximize their chances of obtaining seronegative donors without necessarily requiring the use of a CMV-seronegative donor for CMV-seronegative candidates would be to consider a strategy of early listing of patients at high risk, allowing them the freedom to be “selective” when donor offers are made to their candidates.

A second and uniformly available approach to reduce the risk of acquiring CMV is through the use of CMV-seronegative blood products during the transplant operation and throughout the posttransplant course. The use of CMV-seronegative blood products has been shown to be effective in preventing CMV infection in CMV-seronegative recipients of BMT and SOT.\textsuperscript{44} When CMV-negative blood products are not available, CMV-safe (e.g., filtered or leukocyte-poor) blood products should be used. While neither of these approaches is perfect, they are associated with a failure rate of only 1 to 4%.\textsuperscript{45}

4.2 Prophylaxis of All At-Risk Patients

4.2.1 Active Immunoprophylaxis

The ideal intervention for preventing CMV infection after transplantation would be active immunization of seronegative recipients with a well tolerated, immunogenic vaccine; however, efforts to date have not yet identified the appropriate candidate vaccine. Early efforts at vaccine development utilized the live, attenuated CMV Towne strain. This candidate vaccine was used in three randomized, placebo-controlled studies in adult renal transplant recipients.\textsuperscript{46-50} Although the vaccine appeared to be well tolerated, it did not alter the incidence of CMV infection in seronegative recipients of kidneys from seropositive donors; however, use of the vaccine did appear to modify the severity of CMV disease in these patients. Recent efforts have focused on subunit glycoprotein B and H candidate vaccines.\textsuperscript{51,52} The best studied is the glycoprotein B, which is responsible for at least one-half of neutralizing antibodies in the serum of naturally infected individuals.\textsuperscript{52} A glycoprotein B candidate vaccine has been engineered which is immunogenic in healthy adults and toddlers.\textsuperscript{53} Although a trial in dialysis patients yielded apparently promising results, it was discontinued prior to completion several
years ago. A canarypox vector-expressing CMV pp65 is another potential candidate vaccine which has been shown to induce long-lasting cytotoxic T cells in CMV-seronegative adult volunteers. While each of these candidates is promising, none of them is currently close to licensure nor have they been evaluated in children with end-stage organ disease.

4.2.2 Passive Immunoprophylaxis
CMV hyperimmune globulin (CMVIG) is standardized to provide an approximately 5-fold enrichment in anti-CMV titer as compared with unselected immune globulin (IVIG). While some data exist supporting a prophylactic effect of standard IVIG in the prevention of CMV in SOT recipients, the use of CMVIG is thought to provide superior levels of protection compared with standard preparations in this patient population. To date, however, no study has directly compared the effect of CMVIG with that of IVIG in the transplant population. Approved prophylactic regimens using CMVIG call for an initial infusion at the time of transplant, with subsequent doses at 2, 4, 6, 8, 12 and 16 weeks post-transplant. CMVIG has been shown to prevent CMV disease in both high and low risk liver transplant recipients and has been shown to reduce the rate of CMV disease by 250% in kidney transplant recipients at risk of primary CMV disease. A meta-analysis appeared to confirm the efficacy of CMVIG after SOT and its use has also been associated with increased survival after liver transplantation as well as the reduction of non-CMV opportunistic infection following SOT in general. In contrast to the demonstrated efficacy in SOT recipients, the use of CMVIG in BMT is controversial. Some authors have reported efficacy whereas others have not. Accordingly, recommendations for its routine use after BMT have no clear basis at the present time. Finally, the main disadvantage of CMVIG is the high cost.

4.2.3 Chemoprophylaxis
Chemoprophylactic strategies are based on the use of an antiviral agent initiated at or near the time of transplant in order to prevent CMV infection or suppress its reactivation. Seronegative recipients who receive organs from a seronegative donor should not receive chemoprophylaxis. In general, studies of chemoprophylaxis against the development of CMV disease have evaluated high dose oral aciclovir, valaciclovir, intravenous ganciclovir and oral ganciclovir. Table II summarizes general aspects of those drugs, as well as foscamet and cidofovir. In general, well designed studies providing direct comparisons of one or more of these agents have not been carried out and data regarding the use of prophylactic antiviral agents in SOT recipients are somewhat confusing. Coulson et al. performed a meta-analysis of randomized trials evaluating antiviral agents in the prevention of CMV infection and disease in adults and pediatric recipients of SOT. The studies compared prophylactic treatment with aciclovir and/or ganciclovir with control groups receiving either no treatment or placebo. The meta-analysis suggested that prophylaxis with ganciclovir resulted in a decreased rate of CMV infection and disease compared with placebo or no treatment. In contrast, the use of prophylactic aciclovir was associated with a decrease in CMV disease but not CMV infection. Similarly, a recent review by Payle suggested that while use of high dosage oral aciclovir was more effective than either no treatment or placebo, the results achieved using oral aciclovir were suboptimal, particularly when compared with those achievable with ganciclovir.

While ganciclovir appears to be more effective than aciclovir, its tolerability profile is less optimal and its use may require central line placement to provide long-term therapy. Bone marrow and renal toxicity are the most frequent adverse effects. This is a particular concern in recipients of allogeneic BMT, in whom ganciclovir-related neutropenia has been associated with bacte-

Table II. Characteristics of antiviral drugs

<table>
<thead>
<tr>
<th>Antiviral drug</th>
<th>In vitro activity</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Principal adverse effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aciclovir</td>
<td>+</td>
<td>Oral</td>
<td>Low bioavailability</td>
<td>Neuro- and nephrotoxicity</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Inexpensive and well tolerated</td>
<td>Gastrointestinal symptoms</td>
<td></td>
</tr>
<tr>
<td>Valaciclovir</td>
<td>++</td>
<td>Oral</td>
<td>Greater bioavailability than aciclovir</td>
<td>Neurotoxicity (hallucinations)</td>
</tr>
<tr>
<td>Intravenous ganciclovir</td>
<td>+++</td>
<td>Proven efficacy</td>
<td>Long-term use</td>
<td>Bone marrow suppression</td>
</tr>
<tr>
<td>Oral ganciclovir</td>
<td>++</td>
<td>Oral</td>
<td>Requires central venous access</td>
<td>Nephrotoxicity</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Poor availability</td>
<td>Same as intravenous ganciclovir</td>
</tr>
<tr>
<td>Foscarnet</td>
<td>+++</td>
<td>No bone marrow suppression</td>
<td>Requires probenecid and prehydration</td>
<td>Nephrotoxicity</td>
</tr>
<tr>
<td>Cidofovir</td>
<td>++++</td>
<td>Weekly administration</td>
<td>Requires probenecid and prehydration</td>
<td>Nephrotoxicity</td>
</tr>
</tbody>
</table>

+ indicates intensity of activity.
rial and fungal infection and late CMV disease secondary to recovery delay of CMV-specific cytotoxic T-cell function.\(^{69,71}\) Finally, although antiviral resistance among strains of CMV in transplant recipients has been described,\(^{12-11}\) increasing use of these agents may result in diminished benefits of these prophylactic strategies. Hence, continued vigilance for changes in the prevalence of resistance to CMV will be necessary in order to assure the ongoing benefits of any chemoprophylactic approach. A brief summary of results achieved with individual agents follows below.

Aciclovir

The use of high dosage, oral aciclovir was the first widely accepted chemoprophylactic strategy for the prevention of CMV. Although several studies did demonstrate the effectiveness of this agent,\(^{67,78,79}\) subsequent studies were not consistently able to reproduce these findings, and enthusiasm for its use has waned.\(^{83,84}\) One major reason for the lack of sustained enthusiasm has been the availability of intravenous, and subsequently oral, ganciclovir. Finally, aciclovir prophylaxis has not been shown to be effective in CMV-seropositive patients receiving muromonab CD3 or antilymphocyte preparations, or high risk patients after SOT.\(^{81,82}\) Although limited data evaluating the use of intravenous aciclovir suggest a potential role in the prevention of CMV in the allogeneic BMT population,\(^{83}\) it is not effective in autologous BMT.\(^{84}\) This approach has not been widely adopted and is not recommended at this time.

Valaciclovir

This ester valine prodrug of aciclovir has greater bioavailability and hence, results in higher serum levels than oral aciclovir.\(^{85}\) Consequently, valaciclovir should offer more effective oral prophylaxis compared with oral aciclovir. Use of valaciclovir for 90 days significantly reduced the risk and delayed the onset of CMV disease, herpes simplex virus infection, and acute graft rejection compared with a placebo control in renal transplant recipients.\(^{86}\) In contrast, other investigators found that the use of valaciclovir was not cost effective in CMV D+/R- recipients of renal transplant compared with placebo.\(^{87}\) The use of valaciclovir for patients who had undergone allogeneic BMT was equivalent to results achieved with high dosage aciclovir prophylaxis.\(^{88}\)

Intravenous Ganciclovir

Two meta-analyses suggest that ganciclovir is an efficacious prophylactic agent against CMV disease in SOT.\(^{67,89}\) Furthermore, the use of intravenous ganciclovir protected against the development of CMV disease in adult liver transplant recipients receiving muromonab CD3.\(^{90}\) What remains unclear is the optimal duration and route of ganciclovir prophylaxis in this population. Published experience suggests that different regimens will likely be necessary for different types of patients with varying levels of risk of developing CMV disease (e.g. seropositive liver transplant recipients versus CMV D+/R- intestinal transplant recipients). Intravenous ganciclovir administered for 100 days has been shown to be effective in preventing CMV disease after both liver transplantation and allogeneic BMT.\(^{150,151}\) Nevertheless, its use does not impact on overall survival after BMT, maybe because of myelosuppression and reconstitution delays of immune function. In contrast, Patel et al.\(^{160}\) concluded that shorter courses (2 to 4 weeks) of intravenous ganciclovir should be beneficial in non-high risk heart and liver transplant recipients (and probably in high risk renal transplant recipients); however, longer courses of intravenous therapy may be necessary to prevent CMV disease in high risk populations, including at-risk lung (e.g. all but D-/R-) and D+/R- heart and liver transplant recipients.\(^{151,152}\) In some scenarios, data are contradictory as to what the appropriate duration of intravenous therapy should be. For example, while some studies have suggested that short course therapy is not adequate for high risk liver transplant recipients,\(^{160}\) use of 14 days of intravenous ganciclovir prophylaxis resulted in good outcomes in pediatric liver transplant recipients.\(^{161}\) Head-to-head studies comparing short versus long courses of intravenous ganciclovir prophylaxis will be necessary to resolve these contradictions. In addition, consensus must be achieved on what the appropriate endpoints (e.g. prevention of all CMV infection versus preventing CMV disease) should be prior to the initiation of such studies.

Oral Ganciclovir

Oral ganciclovir has a relatively low mean bioavailability in humans (6 to 9%), which is more pronounced in children. A multicenter study in adult recipients of liver allografts found that the incidence of CMV disease at 6 months was 18.9% with placebo versus 4.8% with oral ganciclovir.\(^{93}\) Disease was also prevented in this study in the high risk (D+/R-) population. Oral ganciclovir has also been found to be more effective than oral aciclovir in D+/R- kidney transplant recipients.\(^{94}\) Unfortunately, the pharmacokinetics, tolerability, tolerance and antiviral effects of oral ganciclovir in children have not been well defined. Three studies of the pharmacokinetics of oral ganciclovir in children have been reported; two of them after transplantation (table III).\(^{150,152}\) Although minimal toxicity was reported, intolerance of the large volume of suspension or numerous capsules was the primary reason for discontinuation of ganciclovir. The results from these studies suggest that a dosage of oral ganciclovir 230 mg/kg every 8 hours should provide effective prophylaxis of CMV disease in children. One study also presented a method to calculate the ganciclovir dose for children with an impaired renal allograft.
Table III. Pharmacokinetic studies of oral ganciclovir in children

<table>
<thead>
<tr>
<th>Recommended dosage</th>
<th>No. of patients and study population</th>
<th>CMV effective prophylaxis</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 mg/kg/6h</td>
<td>9 high-risk CMV kidney and liver transplant recipients</td>
<td>Yes</td>
<td>Frenkel et al. [37]</td>
</tr>
<tr>
<td>20-40 mg/kg/6h</td>
<td>36 children infected with HIV</td>
<td>Yes</td>
<td>Pescevititz et al. [38]</td>
</tr>
<tr>
<td>100 mg/kg/day in 3 doses</td>
<td>14 CMV D+/R− kidney transplant recipients</td>
<td>Yes</td>
<td>Filler et al. [39]</td>
</tr>
</tbody>
</table>

CMV = cytomegalovirus; D+/R− = seropositive donor/seronegative recipient.

Volganciclovir
Volganciclovir is a valine ester of ganciclovir that has recently been approved for use in patients with AIDS. As in the case of valaciclovir, the addition of the valine moiety results in a dramatic improvement in the bioavailability of the parent drug. Preliminary experience in liver transplant recipients has been promising.[100]

Foscarnet
This pyrophosphate analog does not cause myelosuppression but is associated with significant nephrotoxicity. Use of foscarnet should be reserved for ganciclovir-resistant CMV disease. Although clinicians must be aware that cross-resistance between foscarnet and ganciclovir (and rarely even cidofovir) has been reported.[102,103] Because of its lack of bone marrow suppression, foscarnet has been evaluated and found effective for CMV prophylaxis following allogeneic BMT in adult patients who were unable to receive ganciclovir.[103] It has also been used for success for pre-emptive therapy in allogeneic BMT and hemopoietic stem cell recipients, alone or in combination with ganciclovir, respectively.[103,104]

Cidofovir
Cidofovir is a nucleotide analog, 10-fold more potent in vitro than ganciclovir.[105] The drug has a long intracellular half-life (>48 hours) and may inhibit CMV for 5 to 10 days, which allows for weekly administration.[106] It has efficacy in CMV retinitis in patients with AIDS.[107] Occasionally cidofovir has been used in the pediatric population, and it may hold promise for improving the treatment of pediatric viral infection.[108] Like foscarnet, the high frequency of nephrotoxicity associated with the use of cidofovir limits its use to clinical scenarios suggestive of ganciclovir resistance; however, cidofovir resistance has been described among allogeneic BMT in children.[109] Ophthalmological adverse effects have also been reported with cidofovir.[109]

4.3 Pre-emptive Therapy
The aim of pre-emptive therapy is to identify and treat patients with subclinical CMV infection before they progress to symptomatic disease.[110] The strategy depends upon the use of laboratory markers or clinical characteristics to identify patients at high risk of developing disease prior to the onset of symptoms. The first laboratory marker used for pre-emptive therapy was virus with viral cultures using the shell vial assay. Unfortunately, positive cultures in CMV-seropositive transplant recipients have not always accurately predicted progression from asymptomatic infection to CMV disease.[112,113] More recently, pre-emptive strategies have relied on quantitation in the peripheral blood of CMV-pp65 antigen and/or PCR assay.[128,129]

4.3.1 CMV-pp65 Antigenemia Assay
Despite the fact that there is extensive literature about the CMV-pp65 assay, strict guidelines for its use as a guide to pre-emptive therapy have not been agreed upon. Suggested thresholds for initiation of pre-emptive treatment have not been standardized and have varied among centers and different types of transplant recipients. For SOT, most clinicians believe that for CMV-seronegative pre-transplant recipients, any positive result (pp65 antigenemia ≥1) indicates primary infection and is an indication for pre-emptive therapy. In contrast, there is no “gold standard” for patients who were CMV-seropositive prior to SOT. Studies in heart transplant recipients suggested that patients with an assay of ≥100/2 x 10^6 leukocytes might benefit from pre-emptive therapy.[123] In contrast, Kusnet et al. [124] have suggested that pre-emptive therapy should be initiated for seropositive SOT recipients when the antigenemia assay is >10/2 x 10^5 leukocytes, because that threshold was associated with a 7-fold increase in the risk of developing CMV disease in a retrospective analysis of seropositive adult liver transplant recipients. For BMT recipients, in addition to the CMV-serostatus of the recipient and the donor, whether the patient underwent an autologous or allogeneic BMT also has to be taken into consideration. One recent study reported that only 10 to 15% of CMV-seropositive recipients of autologous BMT with antigenemia will progress to CMV disease compared with 50 to 60% of seropositive recipients of allogeneic BMT with antigenemia.[113] It was suggested that recipients of autologous transplants with >5 positive cells/1.5 x 10^5 leukocytes should receive pre-emptive therapy.

Several disadvantages of the pp65 antigenemia assay are noted in table IV. We recommend its use for patients at high risk of developing CMV disease after small bowel and lung transplan-
Table IV. Disadvantages of the pp65 antigenemia assay

Staining with immunoperoxidase may lead to different results than indirect immunofluorescence.
Results of tests performed on shipped specimens may not be accurate.
Limited utility in patients with neutropenia (insufficient neutrophils to count).

4.3.2 CMV PCR

As is the case with the pp65 antigenemia assay, consensus on appropriate CMV PCR thresholds for initiation of pre-emptive therapy has not yet been achieved. One set of investigators has suggested using 100 genome copies/10^6 leukocytes as the threshold to initiate pre-emptive therapy in allogeneic BMT patients, as well as in SOT recipients experiencing primary CMV infection;[10,114] however, they recommend a higher cutoff of 1000 genome copies/10^6 leukocytes when monitoring CMV-seropositive SOT recipients.[115] Prospective studies confirming the appropriateness of these proposed cutoffs have not been carried out. Additional limitations to the use of quantitative CMV PCR as a guide to initiation of pre-emptive therapy include the fact that standardization of tests between transplant centers has not been accomplished; a variety of different primer sets and quantitative strategies have been used. However, one potentially important advantage of PCR compared with the antigenemia assay is the fact the specimen can be stored safely and shipped without significantly affecting the results of the assay.[116,117]

Theoretical advantages and disadvantages of pre-emptive therapy compared with chemoprophylaxis are shown in Table V. Routine prophylactic strategies result in up to 65% of patients receiving antiviral therapy unnecessarily. It is of particular interest after BMT, where the use of pre-emptive therapy is associated with a reduction in ganciclovir use, less neutropenia, earlier recovery of CMV-specific immunity and less invasive fungal infection.[115,118] In contrast, the use of pre-emptive therapy has been associated with an increase of late CMV disease and CMV pneumonia after discontinuation of ganciclovir when antigenemia was no longer detected.[115,118] Screening for CMV infection with antigenemia or PCR beyond day 100 seems reasonable.

5. Treatment of CMV Infection and Disease

Ganciclovir is the drug of choice for the treatment of CMV disease after SOT and BMT in children. Its long record of success and relatively infrequently encountered adverse effects support its use over either foscamet or cidofovir. Bone marrow suppression may limit its use in BMT recipients. Unresolved issues regarding the treatment of CMV disease include the duration of therapy, the potential utility of combination therapy using CMV-IVIG, and the value of following the pp65 antigenemia assay or quantitative CMV PCR as a guide to the duration of therapy. Because of the so-called indirect effects of CMV (e.g., chronic rejection), some investigators have felt that even asymptomatic CMV infection should be avoided. However, data do not exist to support the treatment of asymptomatic CMV infection as a means of preventing chronic rejection or late graft loss after transplantation. An overview of proposed guidelines for the treatment of CMV infection and disease is provided in Table VI. Finally, while mycophenolate mofetil enhances the severity and morbidity of CMV disease after kidney transplantation,[119] a recent study has suggested that the drug provides a protective effect against CMV-mediated injury and long-term graft loss.[120] In fact, Neyts et al.[121] observed that mycophenolate mofetil could strongly potentiate the anti-CMV activity of aciclovir and ganciclovir, both in vitro and in vivo, in a murine model.

Table V. Theoretical advantages and disadvantages of pre-emptive versus prophylactic therapy

<table>
<thead>
<tr>
<th>Pre-emptive therapy</th>
<th>Prophylactic therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Advantages</strong></td>
<td></td>
</tr>
<tr>
<td>Avoidance of universal prophylaxis</td>
<td>Decrease incidence of opportunistic infections</td>
</tr>
<tr>
<td>Cost reduction</td>
<td>Prophylaxis for other herpes virus</td>
</tr>
<tr>
<td>Reduction of adverse effects attributable to the use of antiviral agents</td>
<td>Proven efficacy</td>
</tr>
<tr>
<td>Decreased likelihood of emergence of drug-resistant strains of cytomegalovirus</td>
<td>Universal prophylaxis</td>
</tr>
<tr>
<td><strong>Disadvantages</strong></td>
<td></td>
</tr>
<tr>
<td>No well-proven advantages</td>
<td>Development of resistance</td>
</tr>
<tr>
<td>Intensive surveillance</td>
<td></td>
</tr>
<tr>
<td>Logistical difficulties</td>
<td></td>
</tr>
<tr>
<td>Patient compliance</td>
<td></td>
</tr>
<tr>
<td>Ambiguity concerning its effect on patient care cost</td>
<td></td>
</tr>
<tr>
<td>Standardization of testing procedures has not been accomplished</td>
<td></td>
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</tbody>
</table>
Table VI. Proposed guidelines for the treatment of CMV infection and disease

<table>
<thead>
<tr>
<th>Treatment of CMV infection and disease after solid organ transplantation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug of choice: intravenous ganciclovir (5 mg/kg/12h) for 14 days (with dosage adjustment according to renal function)</td>
</tr>
<tr>
<td>If resistant to ganciclovir: intravenous ganciclovir (5 mg/kg/12h) + foscarnet (90 mg/kg/12h) + CMVIG 100 mg/kg weekly (with dosage adjustment according to renal function)</td>
</tr>
<tr>
<td>Follow up with antigenemia or PCR</td>
</tr>
<tr>
<td>If resistant to ganciclovir and foscarnet: intravenous ganciclovir + cidofovir (5 mg/kg once a week for two doses, and then once every other week) + CMVIG</td>
</tr>
<tr>
<td>Follow up with antigenemia or PCR</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Treatment of CMV disease after bone marrow transplantation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drug of choice: intravenous ganciclovir (5 mg/kg/12h) for 14-21 days, followed by 5 mg/kg/day for at least 3-4 weeks</td>
</tr>
<tr>
<td>Marrow failure: foscarnet (60 mg/kg/12h) for 14 days followed by 90 mg/kg/day for 2 weeks, plus G-CSF</td>
</tr>
<tr>
<td>CMV pneumonitis: intravenous ganciclovir (5 mg/kg/12h) for 14-21 days, followed for at least 3-4 weeks by 5 mg/kg/day plus CMVIG</td>
</tr>
<tr>
<td>Follow up with antigenemia or PCR</td>
</tr>
</tbody>
</table>

CMV = cytomegalovirus; CMVIG = cytomegalovirus hyperimmune globulin; G-CSF = granulocyte colony-stimulating factor; PCR = polymerase chain reaction.

6. Conclusions

Current efforts have gone beyond the treatment of CMV disease to its prevention. The major focus of these efforts have centered on prophylaxis (using either immunoprophylactic or chemoprophylactic strategies) or on preemptive therapy. Comparative studies are needed to investigate which strategy is the most cost effective. While some authors advocate prophylactic therapy as the preferred approach to preventing CMV disease,[122] data directly comparing the outcomes of the two different approaches in various transplant populations are needed. The comparative efficacy of preemptive treatment strategies, based on either CMV-pp65 antigenemia assay or quantitative CMV PCR, has not been evaluated in well designed clinical trials in either adults or children. Trials carried out in the pediatric population would be of particular interest because there is a greater proportion of children undergoing primary CMV infection (and therefore at risk of greater morbidity) than adults. Prospective trials comparing preemptive therapy using either intravenous or oral ganciclovir, and now oral valganciclovir, are necessary to determine the relative cost effectiveness and efficacy of these alternative strategies with these related agents. The use of a combined approach may be necessary for very high risk patients, e.g. D+/R- intestinal transplant recipients. Pending the availability of these proposed studies, individual centers will need to evaluate the incidence and impact of CMV disease in their transplant population to determine whether or not their current prophylactic strategies are appropriate.

References

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