Does Mycophenolate Mofetil Increase the Risk of Cytomegalovirus Infection in Solid Organ Transplant Recipients? – A Mini-Review

Alice Tung Wan Song¹, Edson Abdala¹², Patricia Rodrigues Bonazzi¹², Telésforo Bacchella² and Marcel Cerqueira César Machado²

¹Department of Infectious and Parasitic Diseases; ²Liver Transplantation Division, Department of Surgery, University of São Paulo Medical School; São Paulo, SP, Brazil

Mycophenolate mofetil (MMF) is currently used for prophylaxis of acute rejection in solid organ transplantation. There have been diverging reports regarding an association between MMF and the risk of cytomegalovirus (CMV) infection. We reviewed the main published studies in an attempt to clarify the association between the use of MMF and the risk, frequency and severity of CMV infections. In a search of the Medline database with the terms “mycophenolate” and “cytomegalovirus”, 42 articles were found to be relevant; among these, 29 articles were thoroughly analyzed. The first studies on MMF in renal transplantation already showed a tendency towards an association between this drug and the occurrence of CMV disease. Further studies were designed specifically to study this association; with the conclusion that an immunosuppressive regimen containing MMF increases the likelihood of CMV disease. Most studies were performed with kidney transplant recipients. We conclude that the use of MMF apparently increases the incidence of CMV disease in renal transplant patients; however, further studies are needed to confirm this association.

Key Words: Mycophenolate mofetil, cytomegalovirus, transplant.

This drug is currently used for prophylaxis of acute rejection in renal, heart, lung, pancreas and liver transplants, along with calcineurin inhibitors, such as cyclosporine and tacrolimus, and with corticosteroids. During the past few years, its use in clinical practice has increased as a consequence of its confirmed efficacy and the possibility of dose reduction or suspension of calcineurin inhibitors, thus diminishing the incidence of adverse effects, such as nephrotoxicity due to tacrolimus [4].

Cytomegalovirus (CMV) infection is a major cause of morbidity in patients undergoing solid organ transplants [5]. Additionally, CMV has been found to be an independent risk factor for the development of other infectious complications, such as bacteremias, invasive fungal diseases and Epstein Barr virus (EBV)-related post-transplant lymphoproliferative disease; it is also a cause of acute and chronic allograft injury [5]. There is a hypothesis that cytomegalovirus may cause endothelial damage in the transplanted organ, leading to chronic transplant dysfunction [6].

In solid organ transplantation, primary infection by cytomegalovirus is mainly acquired when the donor is CMV-seropositive and the recipient is CMV-seronegative (D+/R-), or else through blood products, though much less frequently [7]. Secondary infection is less frequent and occurs when there is reactivation of an endogenous virus, or through reinfection in a seropositive recipient [7]. CMV infection occurs mainly during the first three months following the transplant; but can be delayed in patients receiving CMV prophylaxis [5]. CMV infection is defined as evidence of CMV replication, regardless of symptoms [5]. CMV disease is a clinical expression of active infection, ranging from malaise, fever, myalgia, and arthralgia, to organ involvement, such as hepatitis, pneumonitis, gastroenteritis, colitis, and encephalitis [7].
Apart from the higher risk of CMV disease in the case of D+/R-, other risk factors for development of CMV disease include the type of transplant (lung, small intestine and pancreas transplant recipients are at the highest risk, while liver, heart and kidney recipients are at lower risk), and the recipient’s state of immunosuppression, which depends on the immunosuppressive regimen used (use of antilymphocyte antibody therapy for rejection treatment is associated with higher risk), and host factors, such as age, co-morbidity, and neutropenia [5].

Two strategies are commonly used for CMV prevention: universal prophylaxis and preemptive therapy. Universal prophylaxis involves antiviral therapy for all “at-risk” recipients, for a defined period of time, beginning immediately post-transplant. In preemptive therapy, patients are monitored at regular intervals for early evidence of CMV replication by use of a laboratory assay, and if positive, they receive antiviral therapy before the onset of symptoms [5].

Conflicting data has been published regarding an association between MMF and the risk of CMV infection. We reviewed the principal published articles in an attempt to clarify the association between the use of MMF and the risk, frequency and severity of CMV infections.

Methods

Articles that contained the terms “mycophenolate” and “cytomegalovirus” were identified in a search of the Medline database, without limitations, resulting in 222 articles, up till June 2005. One hundred and sixty-eight articles were excluded because they did not include a comparison of incidence/prevalence of CMV infection and use of MMF. Another five articles were excluded, as they were not published in English. Seven articles were excluded because the subjects were pediatric patients. Forty-two articles were considered relevant; among these, 29 original articles were thoroughly analyzed: three were the first studies to analyze the efficacy of MMF in renal transplant recipients; 14 were specifically designed to compare the association between MMF and CMV infections; 10 evaluated the efficacy of MMF, and two were renal transplant review articles.

The 29 articles were separated into three groups: the first group included the first articles whose main objective was to evaluate the efficacy of MMF; the second group included the articles which had as a primary goal an evaluation of the association between CMV infection and the use of MMF; the third group included the articles whose main objective was the analysis of the efficacy of the immunosuppressive drugs, and the occurrence of CMV was secondarily evaluated.

Results

First studies. Three major studies were performed to confirm the efficacy of mycophenolate mofetil as part of an immunosuppressive regimen for the prevention of acute rejection in renal transplants [8-10]. These studies were multicentric, prospective, randomized, controlled and double-blinded. Data from these three studies suggested an increase in cytomegalovirus invasive tissue disease, compared to azathioprine or a placebo. The European Mycophenolate Mofetil Cooperative Study comprised 491 recipients of cadaveric renal allografts divided into three groups: all received cyclosporine and corticosteroids, and the three arms received a placebo, or 2 or 3 g of MMF. The incidence of invasive CMV disease was 2.4%, 3%, and 6.9%, respectively [8]. The second study, done by the Tricontinental Mycophenolate Mofetil Transplantation Study Groups, was performed with 503 patients, all of whom received cyclosporine and corticosteroids, and each arm received either azathioprine, or 2 or 3 g of MMF. The incidence of invasive CMV disease was 6%, 7% and 11%, respectively, with predominance of gastrointestinal tract affection [9]. This study suggested that 2g MMF/day would be the most appropriate dosage, taking into account the immunosuppressive effect and the risk of tissue invasion by CMV. In both papers, the incidence of CMV syndrome or CMV viremia was similar in all groups [8,9].

In the third study, by the US Renal Transplant Mycophenolate Mofetil Study Group [10], 499 patients received cyclosporine, corticosteroids and antithymocyte globulin, and were randomized to receive azathioprine, 2 or 3g MMF. The donor/recipient cytomegalovirus serological status was similar among the three treatment groups. MMF treatment resulted in greater incidence of tissue-invasive CMV than in the azathioprine group (10.8% with 3g/day and 9.1% with 2g/day, versus 6.1% with azathioprine). None of the studies included statistical analysis. Their primary goal was to evaluate the outcome regarding acute rejection and not CMV infection. These were the initial studies that showed that there could be an association between the use of MMF and CMV disease, particularly in those patients receiving 3g MMF/d.

Studies designed to evaluate MMF x CMV. Fourteen articles evaluated the association between the use of MMF and the CMV infection/disease rates (Table 1).

Moreso et al. [11] compared three groups of renal transplant patients enrolled in the European Mycophenolate Mofetil Cooperative Study, treated with cyclosporine (target blood levels of 200-300 ng/mL) and prednisone (0.5mg/kg/d before surgery, gradually reduced to 0.1mg/kg/d in 3-6 months) and randomized to receive a placebo (n=27), 2g MMF/d (n=28), or 3g MMF/d (n=28), and a fourth group, which received 3g MMF/d, with low doses of cyclosporine (target blood levels of 25-175 ng/mL) and prednisone (0.25 mg/kg/d, gradually reduced to 0.1 mg/kg/d in 3 months). CMV disease was significantly increased in the 3g MMF/d plus conventional doses of cyclosporine and prednisone (35.7% versus 8% in the other groups) group, suggesting that the reduction of immunosuppression by diminishing the doses of other
Table 1. List of studies designed to evaluate the association between mycophenolate mofetil (MMF) and cytomegalovirus (CMV)

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Organ</th>
<th>Study design</th>
<th>N</th>
<th>Arms</th>
<th>CMV prophylaxis/preemptive therapy</th>
<th>CMV infection</th>
<th>CMV disease</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>kidney</td>
<td>prospective,</td>
<td>97</td>
<td>CYA + pred + placebo</td>
<td>NI</td>
<td>not evaluated</td>
<td>3.7%</td>
<td>0.002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>randomized</td>
<td></td>
<td>CYA + pred + MMF 2g</td>
<td></td>
<td></td>
<td>7.4%</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred + MMF 3g</td>
<td></td>
<td></td>
<td>35.7%</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred (low doses) + MMF 2g</td>
<td></td>
<td></td>
<td>6.7%</td>
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</tr>
<tr>
<td>12</td>
<td>kidney</td>
<td>prospective</td>
<td>445</td>
<td>MMF 2g + cort(+ anti-lymphocyte + CYA) AZA + cort + ATG + CYA</td>
<td>No</td>
<td>not evaluated</td>
<td>24.6%</td>
<td>NI</td>
</tr>
<tr>
<td>13</td>
<td>kidney</td>
<td>prospective</td>
<td>158</td>
<td>CYA + pred + MMF 2g</td>
<td>Yes</td>
<td>not evaluated</td>
<td>21.6%</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred + AZA + ATG</td>
<td></td>
<td></td>
<td>29%</td>
<td>0.005</td>
</tr>
<tr>
<td>14</td>
<td>kidney</td>
<td>case-control</td>
<td>741</td>
<td>CYA + cort CYA + AZA + cort CYA + AZA + cort + ATG CYA + MMF + cort</td>
<td>No</td>
<td>not evaluated</td>
<td>9.5%</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>tacrolimus + AZA + cort + MMF + cort taurolimus + AZA + cort</td>
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<td></td>
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<td></td>
<td>taurolimus + MMF + cort + cort</td>
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<td></td>
<td></td>
<td></td>
<td>CYA + rapamycin + cort + MMF + cort + cort</td>
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<tr>
<td>15</td>
<td>kidney</td>
<td>case-control</td>
<td>136</td>
<td>CYA + pred + MMF 2g</td>
<td>NI</td>
<td>not evaluated</td>
<td>no difference</td>
<td>0.958</td>
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<td></td>
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</tr>
<tr>
<td>17</td>
<td>kidney</td>
<td>retrospective</td>
<td>84</td>
<td>CYA + cort + MMF (median 2.6g)</td>
<td>no</td>
<td>analysis along</td>
<td>67%</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + cort</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>18</td>
<td>kidney</td>
<td>retrospective</td>
<td>1018</td>
<td>CYA + cort regimen with MMF regimen without MMF</td>
<td>NI</td>
<td>with CMV disease analysis along</td>
<td>30%</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>with CMV disease analysis along</td>
<td>8.4%</td>
<td></td>
</tr>
<tr>
<td>19</td>
<td>kidney</td>
<td>retrospective</td>
<td>280</td>
<td>CYA + pred + MMF</td>
<td>NI</td>
<td>not evaluated</td>
<td>1 episode/118 treatment mo</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred + AZA</td>
<td></td>
<td></td>
<td>1 episode/346 treatment mo</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>kidney</td>
<td>retrospective</td>
<td>66</td>
<td>tacrolimus + cort + MMF 2g</td>
<td>NI</td>
<td>11.4%</td>
<td>0.10</td>
<td>not evaluated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>tacrolimus + cort + AZA</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>21</td>
<td>kidney</td>
<td>retrospective</td>
<td>91</td>
<td>CYA + pred + MMF 1.5-2g</td>
<td>yes</td>
<td>none</td>
<td>22% (CMV/NI</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred + AZA</td>
<td></td>
<td>analysis along</td>
<td>fungal infection</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>kidney</td>
<td>retrospective</td>
<td>470</td>
<td>MMF + CYA + cort CYA (microemulsion) + cort CYA (standard) + cort</td>
<td>yes</td>
<td>with CMV disease</td>
<td>11%</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CYA + pred + AZA</td>
<td></td>
<td>64%</td>
<td>0.041</td>
<td>not evaluated</td>
</tr>
<tr>
<td>16</td>
<td>kidney</td>
<td>retrospective,</td>
<td>29</td>
<td>CYA + pred + MMF 2g</td>
<td>yes</td>
<td>not evaluated</td>
<td>58%</td>
<td>0.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td>descriptive</td>
<td></td>
<td>CYA + pred + AZA</td>
<td></td>
<td></td>
<td>18%</td>
<td></td>
</tr>
<tr>
<td>22</td>
<td>liver</td>
<td>prospective,</td>
<td>63</td>
<td>CYA + cort + anti-lymphocyte + MMF 1.5-2g CYA + cort + anti-lymphocyte</td>
<td>yes</td>
<td>no difference</td>
<td>NI</td>
<td>not evaluated</td>
</tr>
<tr>
<td></td>
<td></td>
<td>randomized</td>
<td></td>
<td>+ AZA</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>23</td>
<td>liver</td>
<td>prospective</td>
<td>157</td>
<td>CYA + cort + anti-lymphocyte + MMF 1.5-2g CYA + cort + anti-lymphocyte</td>
<td>yes</td>
<td>no difference</td>
<td>no difference</td>
<td>&gt;0.20</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>+ AZA</td>
<td></td>
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</tbody>
</table>

NA = not applicable; NS = non-significant; pred = prednisone; cort = corticosteroids.
NI = not informed; AZA = azathioprine; CYA = cyclosporine.
Immunosuppressive drugs along with 3g MMF/d reduces the incidence of CMV disease, without affecting rejection rates. Moreover, the incidence of CMV disease in patients treated with 3g MMF/d (7.4%), and reduced cyclosporine and prednisone doses was similar to the incidence in patients receiving 2g MMF/d and conventional cyclosporine and prednisone doses (6.7%). The CMV serological status was similar among the groups, but CMV prophylaxis was not reported.

A prospective trial by Giralt et al. [12] compared 445 renal transplant patients who were treated with either MMF 2g/d (n=126) or azathioprine (n=319) (along with corticosteroids and anti-thymocyte globulin (ATG) as induction therapy, followed by cyclosporine). None were treated with preemptive or prophylactic gancyclovir, and CMV seropositivity of donor/recipient was not informed. The incidence of CMV disease was similar in the two groups (24.6% in the MMF group versus 21.6% in the Az group); however, after treatment with gancyclovir for 14 days, graft survival increased significantly by up to one year in the patients in the MMF group, compared with those in the azathioprine group (90% versus 77%, respectively, p<0.02). Apparently, gancyclovir had better antivirus efficacy when associated with MMF, resulting in protection against the deleterious effects of CMV on allografts and against other consequences, such as graft dysfunction.

Another prospective study by Bernabeu-Wittel [13], made in Spain with renal transplant recipients, compared two groups that received cyclosporine and prednisone, combined with either 2g MMF/d (n=76) or azathioprine and antilymphocyte globulin (7-14 days, n=82), analyzed infectious complications during the first six months post-transplantation. Patients at high-risk for CMV infection (D+/R- and use of OKT3) received a course of anti-CMV immunoglobulin. Even though there were similar proportions of CMV seronegative recipient/CMV seropositive donor, the incidence of CMV disease was significantly higher in the MMF cohort than in the azathioprine-ATG cohort (29% versus 9.5%, respectively, p=0.005). The upper gastrointestinal tract was the most affected organ. In the multivariate analysis, with serostatus D+/R-, treatment with mycophenolate, and acute rejection episodes were independently associated with higher risk of developing CMV disease.

Immunosuppressive regimens were compared and analyzed in a case-control study on the development of tissue-invasive CMV infection in 741 renal transplant recipients [14]. There were 101 patients (13.6%) with CMV disease, with a total of 125 episodes. Seven basic drug regimens were identified; based on multivariate analysis, it was found that previous acute rejection treatment was a risk factor for developing CMV disease, as was an immunosuppressive drug regimen consisting of tacrolimus, MMF and steroids (odds ratio (OR) = 3.065, confidence interval (CI) = 1.817-5.169, p=0.0063). Also, the use of MMF did not influence the likelihood of developing gastrointestinal CMV disease. The protocols that used tacrolimus, azathioprine and steroids or cyclosporine, MMF and steroids were not found to be significant independent factors for the occurrence of CMV.

Sarmiento et al. designed a case-control study in 1998 [15] with renal transplant recipients, with three controls for each of the 34 cases of CMV infection/disease. After logistic regression, the significant risk factors for CMV were proof of past rejection episodes and positive CMV donor status. No association was found between MMF (2g/d) and CMV infection. In 2000, Sarmiento et al. analyzed 29 renal transplant recipients who developed CMV disease, and found that MMF was part of the immunosuppressive regimen used in 58% of the patients with organ involvement, versus azathioprine in 18% (p=0.03) [16]. The median number of organs involved was significantly greater in the MMF group than in the azathioprine group (2 versus 1, p=0.015). All patients received post-transplant prophylaxis with 200 mg acyclovir three times a day (tid) for 21 days. The frequency of CMV D+/R- was similar in the two groups. The small number of patients limits conclusions, as does the lack of controls.

Ter Meulen et al. [17] performed a retrospective analysis of 84 renal transplant recipients, in which all cases were donor CMV seropositive and recipient CMV seronegative. The objective was to determine if the addition of MMF (median 2.6g/day) to a regimen containing cyclosporine and prednisone would increase the frequency and/or severity of primary CMV infection in this high-risk population; no patients had received any prophylaxis for CMV. CMV serological status was similar among the groups, but CMV prophylaxis was not reported. Primary CMV infection, detected by IgG seroconversion, was similar in the two groups. However, CMV disease was more prevalent in the MMF group than in the control group (67% versus 30%, respectively, p<0.05). Although there was a tendency towards increased use of anti-T-cell therapy for treatment of acute rejections in the MMF group, the conclusions were not altered, when the patients treated with anti-T-cell therapy were excluded. In addition, the two groups had similar severity of disease, frequency of invasive tissue disease and post-transplantation time until the manifestation of symptoms (in 90% of the patients, the first symptoms developed within three months after the transplant).

A Spanish study analyzed the use of MMF and the incidence of CMV infection in 1,018 renal transplants: 8.4% of 381 patients receiving MMF had CMV infections, compared to 3.6% of 637 patients without MMF (p<0.01) [18]. The frequency of CMV D+/R- and treatment with OKT3 were similar in the two groups. Neither the MMF dosages nor CMV prophylaxis were reported.

In a Croatian study [19], 280 kidney transplant recipients were treated with azathioprine, cyclosporine and steroids, or azathioprine and steroids, while 219 transplant patients were treated with either MMF (dose not specified), cyclosporine and steroids, or MMF and steroids. There were no differences in donor-recipient CMV serological status. The AZA group...
had 51 CMV disease episodes (one episode per 346.5 treatment months), while the MMF group experienced 43 episodes (one episode per 118.1 treatment months) (p<0.01). The mean time till onset of disease was also different: median 4 months for the AZA group, and 1.8 months for MMF group. There were five cases of CMV pneumonitis in the AZA group, with a mortality rate of 80%; only one patient in the MMF group had CMV pneumonitis.

Satoh et al. [20] retrospectively compared two treatment regimens following renal transplant, consisting of tacrolimus and steroids, with either azathioprine (n=22) or 2g MMF/d (n=44). D+/R- cases were excluded from the study. The incidence of CMV infection in the MMF group was 11.4%, versus zero in the AZA group.

A study performed on geriatric (age > 60 years) renal transplant patients [21] retrospectively compared a cohort of 46 patients treated with AZA, prednisone and cyclosporine with a cohort of 45 patients treated with MMF (1.5-2g/d), prednisone and cyclosporine. Intravenous gancyclovir was administered to the patients during antibody administration, and acyclovir was given to all patients for the first six months after transplantation. Diagnosis of CMV was made with clinical presentation and one of the following: four-fold increase in IgG titers, new seroconversion, or direct immunohistochemical staining of tissue. The two groups were similar regarding antibody therapy and preoperative CMV serologic status. Infectious complications were evaluated during one year after transplantation. Fungal and CMV infections were analyzed together. The diagnostic methods for CMV had low sensitivity. The incidence was higher in the MMF group (22% versus 11% in the AZA group). MMF was demonstrated as the only independent risk factor for the development of CMV and fungal infection (RR 3.8, CI 1.5-9.8).

In a study by de Maar et al. [6], 470 renal transplant recipients were retrospectively evaluated according to the immunosuppressive regimen used: cyclosporine (standard formulation) and prednisolone, versus cyclosporine (microemulsion formulation) and prednisolone, versus MMF and cyclosporine and prednisolone. Patients who received induction therapy with OKT3 or ATG were excluded, and gancyclovir was given preemptively. The MMF dosage was not informed. The incidence of CMV infection was 35% in the first group, 53% in the second group, and 64% in the third group. There were no significant differences between the second and third groups, suggesting that the introduction of the microemulsion formulation of cyclosporine was mainly responsible for the increase in CMV infection, and not MMF. However, there was a significant difference in the duration of infection (prolonged viremia with MMF). No invasive disease was found, probably due to preemptive treatment. These results are difficult to evaluate, as the study comprises a large period (1989-1998), during which there were many different therapeutic approaches. In addition, there was no information regarding cyclosporine blood levels.

In another trial [22], 63 recipients of a complete or right lobe split liver graft were prospectively randomized to receive AZA or 1.5-2g MMF/d, as part of an immunosuppressive regimen containing lymphocyte antibodies, corticosteroids, and cyclosporine. Gancyclovir was given preemptively, and information on CMV serological was not given. No significant differences were found in the incidence of CMV infection.

In a study performed by Paterson et al. [23], liver transplant recipients who developed neurotoxicity or nephrotoxicity, supposedly due to tacrolimus, had their doses of this drug lowered (5-10) and they started using 2g MMF/d. Preemptive CMV therapy was also implemented. Out of 157 patients, 46 had their immunosuppressive regimen altered by the time of observation (16 in the first month, five from 1-5 months after transplant, and 25 >6 months after transplant). After six months of observation, no significant differences were found in the occurrence of cytomegalovirus infection or disease, when compared to patients not treated with MMF. Patients who had the regimen changed more than six months post-transplant were grouped along with those without change, as the analysis was made during the six months after transplant.

Finally, renal transplant recipients in the United States Renal Data System were analyzed in a historical cohort study of patients with a primary discharge diagnosis of CMV disease during a three-year period [24]. Of 33,479 recipients of renal transplants, 695 patients were hospitalized for CMV disease. Controlled for potential confounders (such as CMV serology and rejection), the use of mycophenolate mofetil was found to be a risk factor, based on univariate and multivariate analysis. A systematic review evaluated the safety of MMF versus azathioprine in renal transplantation, and identified 20 trials, including a total of 6,387 patients [25]. The incidence of CMV infection was higher with 3g MMF compared to azathioprine; there were no significant differences between 2g MMF and azathioprine or between 2g and 3g MMF.

Studies designed to evaluate the efficacy of MMF. Seven articles compared immunosuppressive regimens, with or without MMF, for renal allograft recipients for the prevention of acute rejection. Higher rates of CMV infection or disease were found in four of them (three retrospective and one prospective) [26-29]. In two studies, there was no differences in CMV infection frequencies; although in one study there was a non-significant increase [30,31]. Another study [32] was conducted to examine the association between MMF and chronic allograft nephropathy; CMV infection and disease was more frequent in the azathioprine group. However, this group received more antirejection therapy.

A randomized trial of cardiac recipients was performed [33], comparing 3g mycophenolate versus azathioprine; CMV disease was more invasive in MMF patients, with similar rates of CMV infection. Eckhoff et al. [34] performed a study of liver transplantation, in which there were similar rates of CMV infection among patients receiving tacrolimus versus...
tacrolimus plus 2g MMF. Lastly, a randomized multicentric trial of 2g MMF versus azathioprine treatment of lung transplant recipients [35] revealed similar rates of CMV infection.

Conclusions

The first studies of MMF had already shown a tendency towards an association between this drug and CMV disease. Based on further studies especially designed to evaluate this association, an immunosuppressive regimen containing MMF apparently increases the likelihood of CMV disease. However, it is not clear whether it is the drug itself or the global immunosuppression caused by an association of multiple immunosuppressive drugs that causes such an increase. Data from the study by Moreso et al. [11] suggest that it is the degree of immunosuppression that determines the increased risk of CMV infection, and not the drug itself. However, Ter Meulen et al. [17] argue that the use of MMF is not accompanied by an increase in bacterial or fungal infections, which goes against the idea that general attenuation of the immune response is the sole factor responsible for the increased incidence of CMV disease. It is then suggested that MMF induces a specific change in the primary immune response to CMV infections, which more frequently leads to symptomatic CMV disease.

We found that most of the studies published on this subject were performed with renal transplant recipients, making it difficult to reach conclusions regarding other solid organ transplants. The two prospective studies on liver recipients that we reviewed did not show any difference in the incidence of CMV infection. Furthermore, unfortunately, most studies were done with patients with a cyclosporine-based immunosuppressive regimen. However, tacrolimus is currently the most commonly used calcineurin inhibitor.

We conclude that the use of MMF increases the incidence of CMV disease in renal transplant patients, though further studies are needed to confirm this association.

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