Varicella Zoster Infection in Renal Transplant Recipients: Prevalence, Complications and Outcome

Z. Mustapic\textsuperscript{a} N. Basic-Jukic\textsuperscript{a} P. Kes\textsuperscript{a} V. Lovcic\textsuperscript{a} Lj. Bubic-Filipi\textsuperscript{a} I. Mokos\textsuperscript{b} Z. Kastelan\textsuperscript{b} S. Zekan\textsuperscript{c}

Departments of \textsuperscript{a}Nephrology, Arterial Hypertension and Dialysis, and \textsuperscript{b}Urology, Clinical Hospital Centre Zagreb and School of Medicine, University of Zagreb, and \textsuperscript{c}Clinical Hospital for Infectious Diseases ‘Fran Mihaljevic’, Zagreb, Croatia

Key Words
Renal transplantation • Immunosuppression • Varicella zoster • Mycophenolate mofetil • Varicella zoster virus infection

Abstract
Varicella zoster virus (VZV) is an important pathogen after renal transplantation. In the present study, we examined the prevalence, clinical presentation and outcome of VZV infections in renal transplant recipients. Charts and medical records of adult renal allotransplant recipients were investigated to find patients with VZV infection. From December 1972 until July 2010, 1,139 patients received kidney allograft at our institution. VZV infection was diagnosed in 40 patients (3.51%). 28 patients (70%) had intensified immunosuppression prior to VZV infection occurrence. Median time of onset was 2.13 years after transplantation (range 9 days to 19.2 years). 35 patients developed VZV during the first post-transplant year (median 0.61 years). Four patients developed VZV infection more than 12 years after transplantation. 33 patients (82.5%) had dermatomal distribution, 5 (12.5%) disseminated herpes zoster (HZ), and 2 patients (5%) who were VZV IgG-negative before transplantation, developed chickenpox. Immunosuppression was reduced and patients received acyclovir. Cutaneous scarring was recorded in 7 cases (17.5%). Two patients developed post-herpetic neuralgia, which was accompanied by scarring and skin depigmentation in 1 of them. Five patients (12.5%) experienced relapse of HZ. Timely initiation of therapy may prevent development of complications and the visceral form of disease. Based on our experience with development of chickenpox, we suggest active immunization for all seronegative patients before organ transplantation.

Introduction
An individual risk for development of infection after renal transplantation is determined by a relationship between the epidemiologic exposure of the individual and the state of immunosuppression which determines the individual’s susceptibility to infection [1].

Varicella zoster virus (VZV) is an important pathogen in organ transplant recipients [2, 3]. VZV infection causes two clinically different forms of disease: primary disease (varicella or chickenpox) is characterized by ve-
sicular lesions on the trunk, head or extremities, and herpes zoster (HZ) (shingles) is characterized by a painful unilateral vesicular eruption, which may rarely be disseminated.

In the present study, we examined the prevalence, clinical presentation and outcome of VZV infections in renal transplant recipients.

**Patients and Methods**

Charts and medical records of adult renal allotransplant recipients transplanted between December 1972 and July 2010 were investigated to find patients with VZV infection.

The immunosuppressive protocols included antiproliferative drugs, steroids and, from 1989, cyclosporin A (CyA). Before 2002 we used only azathioprine (AZA) as an antiproliferative drug, and since that time mycophenolate mofetil (MMF) has replaced AZA in immunosuppressive protocols in our center. Steroid dose at transplant was 0.5 mg/kg/day, which was tapered toward the maintenance dose of 0.05 mg/kg/day. From 2005, tacrolimus was introduced beside CyA. From the year 2004, patients with more than three mismatches received induction therapy with basiliximab or daclizumab, and until that time this group was treated with a steroid bolus (500 mg) prior to transplantation. While HLA matching was crucial for organ allocation until 2007 when we joined Eurotransplant, induction treatment was rarely used (18.4% of all patients).

Rejection episodes were treated with steroids (10 mg/kg), which had been given in 3–5 boluses. From the introduction of ganciclovir in 1999 in our center, all CMV IgG-negative patients who received an allograft from a CMV IgG-positive donor were treated with antiviral prophylaxis for 3 months (21 days intravenously, followed by oral ganciclovir).

Age, gender, time on dialysis, time of transplantation, immunosuppressive protocol, viral status before transplantation, episodes of acute graft rejection, other viral infections, and graft function were recorded. Detailed clinical characteristics of VZV infection were noted (localization, dissemination, complications and outcome).

The diagnosis was made on clinical grounds and/or VZV seroconversion. The study was approved by the Ethics Committee of the School of Medicine, University of Zagreb. We used descriptive statistics (Microsoft Office, Excel 2007).

**Results**

**Patients’ Characteristics**

From December 1972 until July 2010, 1,139 patients (58.7% male) received a kidney allograft at our institution. Mean age at transplantation was 52 years, with the dialysis vintage distribution from the preemptive transplantation to 25 years.

VZV infection was diagnosed in 40 patients (3.51%). There were 27 male patients and 13 female patients, with the mean age at diagnosis of 51.8 years (table 1). 39 of them received a renal transplant from a deceased donor. Average time on dialysis was 6.8 years.

At the time of onset, 36 patients had CyA, MMF and steroids, 2 had tacrolimus, MMF and steroids, and 2 AZA and CyA in therapy. The mean CyA concentration was 157.5 μmol/l (range 85–294) and the mean dose of MMF was 916 mg/m² (range 300–1,260) at the time of VZV infection. Mycophenolic acid concentration was not determined. Two patients receiving tacrolimus had a serum concentration of 12.9 and 13.6 ng/ml, and 7 patients had CyA C₀ >200 μmol/l. Eleven patients (27.5%) received induction therapy with basiliximab or daclizumab in their immunosuppressive protocol. Ten patients (25%), 2 of whom received induction therapy, had acute graft rejection and were treated with 3 (2 patients) to 5 (8 patients) boluses of intravenous methylprednisolone 3 weeks to 3 months prior to VZV reactivation. The average dose of methylprednisolone for rejection therapy was 2,875 mg. Thus, 28 patients (70%) had enhanced immunosuppression prior to VZV infection.

Six patients (15%) received antiviral treatment before development of VZV infection, due to CMV infection, and 3 patients received ganciclovir prophylaxis for CMV mismatches, 2–4 months before VZV infection. Three
patients had chronic hepatitis C infection. One patient had chronic hepatitis B infection, and 1 patient had positive BK virus in urine. We had no HIV-positive patients (table 1).

**Timing of VZV Infection**

Median time of onset was 2.13 years after transplantation (range 9 days to 19.2 years). 35 patients (85%) developed VZV during the first post-transplant year (median 0.61 years). Four patients (10%) developed VZV infection a long time after transplantation (10.3, 15.8, 17.2 and 19.2 years after transplantation, respectively). Two of these patients had received AZA since transplantation, while in 2 patients AZA was replaced with MMF after acute graft rejection, 3.7 and 7.8 years prior to VZV infection, respectively. None of the patients treated with AZA developed disseminated disease.

**Clinical Presentation of VZV Infection**

33 patients (82.5%) had dermatomal distribution, 5 (12.5%) disseminated HZ, and 2 patients (5%) who were VZV IgG-negative before transplantation developed chickenpox. Severe skin changes were recorded (fig. 1a). Two patients who received an organ from the same donor developed VZV infection 29 days after transplantation. The donor was VZV IgG-positive, IgM-negative. One recipient was VZV IgG-positive and developed disseminated vesicular rash (fig. 1b), and another was VZV IgG-negative (he developed chickenpox). There was no known exposure to the virus in the other patients.

Deterioration of graft function was recorded in 1 patient, and 2 had transient elevation of liver enzymes.

**Treatment of VZV infection**

In the treatment of VZV infection, 33 patients (82.5%) received acyclovir orally, 6 (15%) acyclovir intravenously and 1 (2.5%) had no antiviral treatment because he failed to visit a doctor. Antiviral therapy was introduced 0–3 days after vesicular eruption. Patients received 5 × 800 mg of acyclovir for 7–10 days (doses were adjusted for graft function if necessary). Seven (17.5%) patients had no changes in immunosuppressive treatment (they were treated in local hospitals). MMF was reduced in 28, and temporarily switched off in 5 patients. CyA and tacrolimus were reduced in patients with elevated C0 concentrations. Steroid dose was decreased in 6 patients. Steroids were tapered off in 2 patients (those with the relapsing form of disease, previous CMV and/or BK infection).

**Complications and Outcome**

All patients and grafts survived. Complications occurred in 9 patients (22.5%). Cutaneous scarring was recorded in 7 cases (17.5%). Two patients (5%) developed severe skin changes.
improved graft survival and immunosuppressive treatment in the following studies [1]. According to our results, introduction of MMF in an immunosuppressive protocol resulted in a higher incidence of different viral infections was recorded [9, 10]. However, an increased incidence of different viral infections was recorded [11–16]. According to our results, introduction of MMF in an immunosuppressive protocol resulted in a higher incidence and more severe VZV disease. All our patients with disseminated disease were in the MMF era. Early therapy with acyclovir orally with a reduction of MMF dose is a therapy of choice. We believe that dose adjustment and finding an upper limit of the therapeutic range of mycophenolic acid, above which the risk of different viral infection is increased, needs to be determined for MMF therapy in the following studies [17], at least in patients who received enhanced immunosuppression early in the post-transplant period. Seventy percent of patients were exposed to intensive immunosuppressive treatment before VZV infection, while they received either an induction or steroid bolus therapy for treatment of acute rejection, or had a high calcineurin inhibitor concentration. This is in line with the previous observations that intensive immunosuppression presents a risk factor for development of VZV infection [1, 8].

The majority of our patients developed VZV infection during the first post-transplant year. Only 4 patients experienced disease a long time after transplantation, thus prolonging the median time of onset (2.13 years). Previous studies reported the onset of VZV infection after solid organ transplantation to be between 2 and 92 months [3, 5, 18]. Switch from AZA to MMF resulted in more intensive immunosuppression and subsequently in development of VZV infection in 2 of our patients (up to 228 months after transplantation).

Five cases of disseminated HZ were recorded, which is a relatively high proportion in comparison with other studies (fewer than 40 cases described in the literature). A mortality rate of 34% was described in patients with disseminated HZ [2]. None of our patients died, probably because of the fast recognition and initiation of antiviral therapy. However, in immunocompromised patients, visceral dissemination of VZV may occur without skin eruptions [19, 20] rendering diagnosis difficult and prolonging the time to initiation of antiviral therapy. A lethal case of VZV infection has been reported without skin lesions [21, 22]. Keeping in mind the retrospective design of our study, it may be possible that such cases have been missed and the risk of VZV infection might be underestimated.

We report a low rate of PHN (5%) in our patients, but a relatively high rate of cutaneous scarring (17.5%). Other authors reported PHN in up to 42.7% of solid organ recipients [5, 23].

Active immunization for VZV-seronegative patients before transplantation should be performed. Taking into account that cellular immunity in patients with end-stage renal disease is impaired and results of vaccination against hepatitis B are suboptimal [24, 25], it may be suspected that vaccination against VZV in patients awaiting renal transplantation may be of limited efficacy. However, vaccination should be recommended to all seronegative patients in order to try to decrease the incidence of this potentially fatal post-transplant complication.

High-dose acyclovir therapy together with reduction of immunosuppression is a cornerstone of VZV infection treatment. Timely initiation of therapy may prevent development of complications and the visceral form of disease. Based on our experience with the development of chickenpox, we suggest active immunization for all seronegative patients before organ transplantation.
References