Infection with varicella-zoster virus (VZV) is common, occurring as primary varicella, usually during childhood, and as zoster, following reactivation of latent virus. Although serious VZV infections are infrequent, they can be life-threatening when they occur in susceptible immunosuppressed patients (1, 2). These patients have a high incidence of visceral involvement, including pneumonia, meningoencephalitis, and rarely, hepatitis. We report here 3 cases of VZV hepatitis in adult liver transplant recipients that occurred in our institution between 1984 and 1989. We describe their clinical presentations and use these cases to illustrate some aspects in diagnosis and prevention of VZV infection after solid organ Tx.

Case 1. A 19-year-old woman underwent orthotopic liver Tx for end-stage liver disease secondary to chronic active autoimmune hepatitis. She was receiving prednisone and azathioprine before the transplant operation. Her immunosuppression after Tx included cyclosporine (CSA) and steroids. At 28 days after her transplant operation, she started complaining of low back pain over the coccyx area, but the exam of this area did not reveal any skin lesions. Two days later she developed few skin vesicles on her face, abdomen, and hands. Laboratory examination showed elevation in LFTs with serum alanine aminotransferase (ALT) of 196 IU/L (normal value [nv] <40 IU/L), serum aspartate aminotransferase (AST) of 221 IU/L (nv <40 IU/L) and bilirubin of 3.5 mg/dl (nv <1.0 mg/dl), and gamma GTP of 137 (nv <40 IU/L). On day 33 after OLTx, her AST increased to 1057 IU/L and bilirubin increased to 4.7 mg/dl. Liver biopsy showed foci of coagulative necrosis with minimal mixed inflammatory cellular infiltrate and multinucleated cells, with negative histochemical stains for HSV 1 and HSV 2. The patient's skin examination revealed vesicular lesions on the palmar aspect of his hands and also on his neck. He was placed on i.v. acyclovir at 10 mg/kg after each dialysis, and was also given a dose of VZIG. Buffy coat and vesicular skin lesions samples yielded VZV.

He remained on a respirator, required hemodialysis, and developed bacterial and fungal abdominal sepsis secondary to leaks from the bowel. At 31 days after retransplantation he required distal pancreatectomy for pancreatic phlegmon. He was discharged 5 months after his original transplant operation. The patient's blood sample 32 days before Tx tested positive for VZV antibodies by IFA, enzyme-linked immunoassay (ELISA), and fluorescent-antibody-to-membrane antigen (FAMA). He denied having had chickenpox in the past. Interview of his family members revealed that the patient's young nephew was diagnosed with chickenpox 16 days before his first Tx, and later two of the patient's children also contracted this illness. The first child was diagnosed 3 days before Tx and the second child was diagnosed on the same day the patient was electively admitted to the hospital for Tx.

Case 2. A 38-year-old man with chronic active hepatitis B, underwent OLTx and was maintained on CSA, prednisone, and azathioprine. Nineteen months after his Tx he visited his friend's house, and two days later his friend's son, whom he met, was diagnosed with chickenpox. The patient was given an injection of varicella-zoster immune globulin (VZIG) intramuscularly the day the exposure was reported. Thirty-two days after exposure he noticed for the first time malaise and the onset of a rash on his chest. He was admitted to the hospital with high temperature and vesicular lesions on his face, chest, and back. He was started on i.v. acyclovir at 10 mg/kg every 8 hr and was given an intramuscular injection of VZIG. His laboratory tests showed ALT of 824 IU/L, AST of 2091 IU/L, and bilirubin of 1.0 mg/dl. No liver biopsy was performed. Auffy coat culture of blood drawn on admission was positive for VZV and IgG varicella antibody titer was positive by IFA. A sample of blood prior to his transplant was positive with a low titer by IFA, and negative by ELISA and FAMA test. The patient was discharged home 12 days after admission.

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immunostains for HSV 1 and HSV 2 in two patients, along with the recovery of VZV from the buffy coats in the three patients, suggests that VZV was the cause of the hepatitis. Patient 3 did not have liver biopsy and the diagnosis was based on liver function test abnormalities and the recovery of VZV from a buffy coat culture. Although without histology we can not really be sure that this patient had hepatitis, VZV viremia has been considered as a marker of visceral disease in immunocompromised patients (15).

We have previously reported 12 patients who developed herpes simplex hepatitis after solid organ Tx (16). Five of these patients developed disseminated intravascular coagulation and expired. This fulminant disease was more common in primary infection compared with reactivation, similar to the clinical picture of patient 1 in this report. The main dissimilarity is that all three patients with varicella hepatitis had skin lesions. In our report of herpes simplex hepatitis, skin lesions were not common, even in cases with autopsy-proven disseminated disease to multiple organs (16). This is similar to bone marrow Tx, where higher rates of VZV visceral dissemination without skin manifestations have been reported (17).

In a surveillance performed among our adult candidates for liver Tx, only 5% were seronegative for VZV and therefore at risk for infection (unpublished data). Because seronegative patients are more susceptible to VZV infection, prophylaxis with varicella zoster immunoglobulin (VZIG) is recommended for them after virus exposure. However, there are conflicting reports regarding its beneficial effect (18). In fact some patients given VZIG after exposure to VZV nevertheless acquired severe infection (19, 20). On the other hand, passive immunity may modify and "mitigate" the disease even if not giving an absolute protection. Patient 3 was most likely seronegative for VZV before Tx, since the ELISA and FAMA tests were negative. The low-positive antibody titer detected by IFA may have been related to passive acquisition of antibodies through blood transfusion. This patient received VZIG one day after exposure; he did contract varicella 32 days later, with visceral involvement, but survived. Patient 1 was seronegative for VZV, since serum from day 33 after Tx tested negative. The exposure in this case was not identified and the rash was seen for the first time 28 days after the transplant operation. The patient did not receive prophylactic treatment and developed a fulminant disease with DIC and expired.

It is assumed that seropositive individuals are not susceptible to varicella. However, there are some reports of immunosuppressed patients who developed varicella more than one time (20, 21). Patients may be reinfected with a different strain of virus and develop a second episode of varicella (22). This is what probably occurred to patient 2, who was seropositive before Tx but was exposed to a new virus in the family from his two children and developed varicella with hepatitis. The patient might have had disseminated zoster secondary to VZV reactivation and not varicella. However, his vesicular rash did not follow any dermatomal distribution at any time. Moreover, visceral dissemination is not a common event in dermatomal zoster infection, compared with varicella infection (23). The three cases that we reported here suggest that transplant recipients in close contact with VZV may benefit from VZV prophylaxis, even when the patients are seropositive. The question of whether this prophylaxis...
should be done with VZIG or with oral acyclovir, as recently reported in the pediatric population (24, 25) will require further investigation. Recently a second oral antiviral agent with activity against VZV, famciclovir, was introduced for clinical use, and it may have a role in prophylaxis (26). There are some data from leukemic children showing that varicella vaccine is both safe and immunogenic in immunocompromised patients (27). Seronegative transplant candidates would be good candidates for the use of this vaccine, which has recently been released for clinical use.

Since VZV infection may be life-threatening in solid organ transplant patients, and after our experience with these three cases, we suggest a tentative management plan for prevention of VZV infection in adult transplant recipients. These suggestions should be used as a working plan until more data are available and conclusive recommendations are made regarding the use of passive immunity, antivirals, and vaccination in this patient population:

1. A history of varicella (chickenpox) and zoster (shingles) is obtained from all patients during pre-Tx evaluation, and VZV IgG titer is checked. (Our laboratory uses ELISA.) Seronegative patients and possibly those with low titer may be candidates for varicella vaccine.

2. Transplant candidates and recipients are educated to avoid exposure to varicella (chickenpox) or zoster (shingles), and to immediately report to their nurse coordinators any accidental exposure.

3. After an accidental exposure to VZV (chickenpox or shingles), VZIG is offered to seronegative recipients and those with low-positive titer. Varicella-zoster immune globulin (VZIG) is administered intramuscularly as a dose of 125 mg/10 kg of weight (maximum dose 625 mg).

4. After any exposure to VZV, susceptible patients should be considered for placement on oral acyclovir at 500 mg five times a day (or possibly famciclovir) for the duration of the incubation period (2–3 weeks).

5. Development of vesicular skin lesions after exposure should prompt viral cultures for herpesvirus (skin lesion and buffy coat), and immediate institution of high-dose acyclovir therapy (10 mg/kg every 8 hr, i.v.) while awaiting confirmation of diagnosis.

6. Liver function tests abnormalities in a patient who has had a recent exposure to VZV should prompt liver biopsy.

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REFERENCES


