Blastic Epstein-Barr virus associated post-transplant lymphoproliferative disorder after allogeneic stem cell transplantation for severe aplastic anemia

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Abstract

Post-transplant lymphoproliferative disorder (PTLD) is a well-recognized complication of organ transplantation. Progress has recently been made in the pathological classification of PTLD. However, the clinical course has not been clarified because of the rarity of this disease. We experienced a case of PTLD with a fulminant clinical course. The patient had been under long-term immunosuppressive treatment for aplastic anemia. He received related allogeneic hematopoietic stem cell transplantation. Soon after transplantation, he developed post-transplant lymphoproliferative disorder (PTLD). His immunosuppression and prednisolone were stopped. However, high fever persisted and he died of multi-organ failure. There may be various clinical types of PTLD, which may include some fulminant cases. In such a case, it is not sufficient to reduce immunosuppression. The patient should be carefully observed and an appropriate individual treatment should be chosen.

Case Report

A 31-year-old man was admitted to our hospital to receive related allogeneic hematopoietic stem cell transplantation (HSCT). He was diagnosed with mild aplastic anemia (AA) when he was 9 years old. Subsequently, his condition repeatedly oscillated from better to worse and he was given immunosuppression intermittently. He had previously been treated with antithymocyte globulin (ATG) and cyclosporine (CSA). More recently, his condition deteriorated; allogeneic HSCT was therefore planned. His older brother proved to be an 8/8 HLA-matched donor. Before transplantation, laboratory examination showed that his white blood cell count was 1700/µL, hemoglobin was 5.9 g/dL, platelet count was 10,000/µL, and reticulocyte count was 36,960/µL. Bone marrow examination showed a total nucleated cell count of 27,000/µL. Expression of CD55 and CD59 on granulocytes was within the reference range. He was diagnosed as stage 3 aplastic anemia. A conditioning regimen was administered consisting of 120 mg/m² fludarabine, 9 mg/m² cyclophosphamide with continuous CSA infusion, and 10 mg/m² methotrexate on days +1, 3, 6 and 11 as graft-versus-host disease (GVHD) prophylaxis. While receiving the conditioning regimen, there were no grade ≥3 adverse events other than mucosal erosion and hematological toxicities. Neutrophil engraftment was achieved on day 17. From day 30, he developed high-grade fever and grade 2 skin GVHD. We started prednisolone (1g/kg) as a GVHD treatment. However, his fever was uncontrolled. At the same time, abnormal lymphocytes appeared in the peripheral blood examination. On day 35, he developed sudden hepatomegaly and cervical lymphadenopathy. Liver biopsy was performed. Histopathological analysis showed diffuse large B cell lymphoma with Epstein-Barr Virus (EBV)-encoded small nuclear RNA positivity. The EBV-DNA level was 8.2×10⁵ copies. We considered that he had developed post-transplant lymphoproliferative disorder (PTLD). His immunosuppression and prednisolone were stopped. However, high fever persisted and he died of multi-organ failure on day 40 (see Figure 1 for the course of HSTC).

Discussion and Conclusions

PTLD is a well-recognized complication of both solid organ transplantation and HSCT. In most cases, PTLD is associated with EBV infection, either as a consequence of post-transplantation viral reactivation or of primary EBV infection. Approximately 70% of PTLD cases are associated with EBV, however, the impact of EBV on survival is not clear.¹ EBV positivity has been reported to be a poor prognostic factor, while non-EBV PTLD has also been seen to have a poor prognosis.² ³ In the updated 2016 World Health Organization lymphoma classification, PTLD is subclassified as plasmacytic hyperplasia PTLD, infectious mononucleosis PTLD, florid follicular hyperplasia PTLD, polymorphic PTLD, monomorphic PTLD (B- and T-NK-cell types) or classical Hodgkin lymphoma PTLD.⁴ Diffuse large B-cell lymphoma accounts for the majority of PTLD cases.

Risk factors for PTLD are HLA-mismatch between donor and recipient, conditioning regimens that include ATG, and T-cell-depleted transplants.⁵ ⁶ EBV-DNA load is elevated exponentially in patients with PTLD, and weekly screening of the EBV-DNA level is recommended for at least 3 months in high-risk patients who have undergone HSCT.⁷ ⁸ ⁹

The current case was at high risk of developing PTLD because of his long-term use of immunosuppression and previous use of ATG. The one point we regret was that we did not check his EBV-DNA level before transplantation. However, even if we had followed EBV-DNA levels during the treatment course, we might not have been able to predict the PTLD. The current case showed rapid onset and fulminant blastic disease. The fulminant form of PTLD is very uncommon and occurs in approximately 1% of PTLD patients.¹⁰

Historically, the initial treatment for PTLD is reduction of immunosuppression...
Case Report

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Figure 1. The course of hematopoietic stem cell transplantation. From day 30, he developed high-grade fever and grade 2 skin GVHD, and abnormal lymphocytes appeared in the peripheral blood examination. On day 35, he developed sudden hepatomegaly and cervical lymphadenopathy, and he died on day 40. CY: cyclophosphamide; FLU: fludarabine; ATG: antithymocyte globulin; CSA: cyclosporine; MTX: methotrexate; SCT: stem cell transplantation; PSL: prednisolone; WBC: white cell blood.

(RIS). Thus, we stopped immunosuppression immediately; however, the disease course was too rapid for this to provide a therapeutic effect. Rituximab has recently been used for the treatment of PTLD, with an overall response rate (ORR) of 44-100%. Furthermore, R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine and prednisolone) is currently the preferred regimen for B-cell PTLD, with an ORR of up to 90%. In our case, the reduction of immunosuppression was not sufficient. We should have given chemotherapy as soon as we suspected PTLD. The standard treatment for PTLD has not been established. Current guidelines recommend RIS to the lowest tolerated levels, based on PTLD type, stage, critical illness and allograft risk, followed by close monitoring for response and escalation of therapy in the event that RIS alone is not sufficient. Classification of the pathology associated with PTLD has progressed recently. However, the clinical course has not been clarified because of the rarity of this disease. There has been no report on incidence according to original disease. We assume that, even if the primary disease is benign (such as AA), but the patient is under long-term immunosuppressive therapy, then there might be a higher risk of developing PTLD rather than malignant disease. There may be various clinical types of PTLD and these may include some fulminant cases. In such a case, it is not sufficient to reduce immunosuppression alone. Each case should be carefully observed and an appropriate individual treatment should be chosen.

References


[page 44] [Hematology Reports 2018; 10:7527]