Successful Haploidentical Transplantation with High Doses of CD34+-Selected Peripheral Blood Stem Cells in Wiskott-Aldrich Syndrome: A Case Report

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Abstract
Wiskott-Aldrich syndrome (WAS) is a rare X-linked recessive, primary immunodeficiency, characterized by eczema, microthrombocytopenia, recurrent infection, and increased susceptibility to autoimmune diseases and lymphoreticular malignancies. To date, HSCT is the only curative therapy. HSCT using HLA-matched related or unrelated donors is highly successful in treating WAS, but these donors may not always be available. We describe a 20 months-old-boy with WAS, who was transplanted by using highly-purified, 16.11x106/kg CD34+ stem cells from his HLA 2-loci mismatched mother, because of lacking donor and suffering life threatening, ongoing sinopulmonary infections. Conditioning consisted of busulfan (16 mg/kg), cyclophosphamide (200 mg/kg), rabbit anti-thymocyte globulin (10 mg/kg) and fludarabine (160 mg/m²). He received methylprednisolone for graft-versus-host disease (GvHD) prophylaxis. The time for neutrophil, platelet and erythrocyte recovery was 11 days, 14 days and 19 days, respectively. He showed immunologic reconstitution by day +180. He developed grade II aGVHD on day +12, and grade I cGVHD on day +240, resolution achieved in aGVHD with i.v. methylprednisolone and cyclosporine, in cGVHD with oral prednisolone. He is well and alive up to now six years after transplantation. He didn’t have any serious infection. Fluorescence in situ hybridization analysis revealed sustained full donor-type engraftment and flow cytometric analysis revealed sustained immunologic reconstitution. The convenient results in this patient can encourage further investigation of the role of using ‘high doses’ of related haploidentical CD34-selected peripheral blood stem cells in patients with WAS at early stages of disease, before the beginning of severe, life-threatening complications.

Keywords: Wiskott–Aldrich syndrome; Haploidentical blood stem cell transplantation; High dose CD34+ cells

Introduction
In 1937, Wiskott described a clinical entity characterized by thrombocytopenia, eczema, bloody diarrhea, and recurrent otitis media in male infants [1]. After rediscovery in 1954 by Aldrich as an X-linked recessive disorder, it was designated the Wiskott-Aldrich syndrome (XLT) [2]. X-linked thrombocytopenia (XLT), sometimes associated with mild eczema and/or infections, was recognized in the 1960s and was suspected to be a variant of WAS. This was confirmed when patients with XLT were shown to have mutations in the Wiskott-Aldrich syndrome protein gene (WAS). WAS mutations result in 3 distinct clinical phenotypes: classic WAS, XLT, and X-linked neutropenia, and a strong genotype phenotype correlation has been suggested. Mutations completely altering WAS protein (WASP) expression typically lead to the classic phenotype [3]. The classic WAS phenotype with microthrombocytopenia, severe eczema, increased susceptibility to pyogenic and opportunistic infections, and increased risk of autoimmune disease and cancer usually leads to death in early childhood or adolescence if left untreated.

Curative treatment by allogeneic hematopoietic stem cell transplantation (HSCT) should be offered to all such patients. The outcome is excellent if performed early in life from a human leukocyte antigen–matched related or unrelated donor [4]. The results of mismatched related or unrelated donor transplantations are associated with an increased risk of graft rejection and graft-versus-host disease (GvHD) [5]. To overcome these limitations, a ‘high dose’ of highly purified CD34+ peripheral blood stem cells (PBS) from his HLA 2-loci mismatched mother was used in this patient. This report details and draws attention to this kind of haploidentical peripheral blood stem cell transplantation in a patient with severe WAS.

Case Presentation
A four-month-old male infant was brought to our clinic with a history of itchy, scaly lesions on his neck and scalp, epistaxis and recurrent pulmonary infections since one month of age. He was the first child of non-consanguineous parents, born at term following an uncomplicated pregnancy. On examination, he was active and well hydrated. His weight, height and head circumference was within normal limits and his vital parameters were stable. There was no pallor, mucocutaneous bleeds or ear discharge. The dried and scaly lesions on his neck and scalp were consistent with eczema. Examination of other systems did not reveal any abnormality. Whole
blood count showed a hemoglobin concentration of 9.4 g/dL, platelet count of 16000/µL, total leukocyte count of 8800/µL, MCV 74.7 fL and MPV 5.4 fL. A blood smear revealed 612 neutrophils (ANC: 1056/µL), 62% lymphocytes (5456/µL), 12% monocytes, 14% eosinophils (1232/µL). There were also seen very small platelets (50% of normal size) on the blood smear, which was consistent with MPV value but there weren’t any big or giant platelets. Bone marrow aspirate smear findings were normal in terms of the presence, maturation, and differentiation of megakaryocytes, erythroid or granulocytic precursors. Wiskott-Aldrich syndrome was suspected in the patient because of microthrombocytopenia, recurrent infections and eczema. Molecular genetic analysis revealed WAS mutation. Liver and kidney functions, serum immunoglobulin levels, and urinalysis were all normal. The diagnosis was confirmed as severe WAS (score 5) by clinical findings and molecular genetic analysis.

Because of lacking related or unrelated matched donor and because of ongoing, recurrent, severe sinopulmonary and middle ear infections, he underwent haploidentical HSCT from his mother at 20 months of age. He received a myeloablative regimen consisting of busulfan (16 mg/kg), cyclophosphamide (200 mg/kg), rabbit anti-thymocyte globulin (10 mg/kg) and fludarabine (160 mg/m²). He also received methylprednisolone for graft-versus-host disease (GVHD) prophylaxis. T cell-depletion and positive selection of peripheral CD34+ stem cells from his G-CSF stimulated mother was performed using the method of immunomagnetic separation (CliniMACS, Miltenyi Biotec). The number of infused CD34+ cells was 16.11×10⁶/kg. He achieved full hematopoietic engraftment with the time for neutrophil, platelet, and erythrocyte recovery being 11 days, 14 days, and 19 days, respectively. He developed grade II aGVHD on day +12, neutrophil, platelet, and erythrocyte recovery being 11 days, 14 days, and 19 days, respectively. He achieved full hematopoietic engraftment with the time for neutrophil, platelet, and erythrocyte recovery being 11 days, 14 days, and 19 days, respectively. He developed grade II aGVHD on day +12, and grade 1 cGVHD on day +240, resolution achieved in aGVHD with i.v. methylprednisolone and cyclosporine A, in cGVHD with oral prednisolone. He was positive for cytomegalovirus which was detected by PCR in his blood, urine and stool samples on day +6 and resolved with ganciclovir. He is now 61/2 age, he has not experienced any serious infection and fluorescence-in-situ-hybridization analysis revealed sustained full donor-type chimerism and flow cytometric analysis revealed immunologic reconstitution.

Conclusion

Hematopoietic stem cell transplantation offers successful results for patients with WAS. Without HSCT 50% of these patients will die from infection, lymphoproliferative disease or autoimmune diseases by the third decade of their life. The WAS mutation affects hematopoietic and lymphoid compartments; both of these compartments are corrected by HSCT. To facilitate multilineage donor cell engraftment, most patients are conditioned with both immunosuppression and high dose chemotherapy. There is an almost 2-fold reduction in mortality among patients with full-chimerism compared to those with mixed or split chimerism. This showed the importance of achieving full donor chimerism in these patients who underwent HSCT. Mixed chimerism also is associated with a significantly higher risk for developing autoimmune manifestations after HSCT [6].

HSCT using HLA-identical sibling donors is highly successful in treating WAS, with approximately 88% event-free survival. The most commonly used conditioning regimen is the combination of busulfan (1 mg/kg × 16 doses) and cyclophosphamide (200 mg/kg total dose), with or without anti-thymocyte globulin. In younger patients, busulfan is metabolized faster. Therefore, it is prudent to monitor levels and target the dose to achieve a steady state concentration of greater than 200 ng/ml to assure engraftment. It is determined that patient age, disease severity, and splenectomy did not affect outcome in HSCT of WAS patients in which HLA-identical sibling marrow grafts were used [7,8].

Results of alternative donor HSCT for WAS having improved over time, particularly for unrelated marrow grafts. Most studies have shown approximately 70%-78% long-term survival. The International Bone Marrow Transplant Registry (IBMTR) which was facilitated in 2001 reported that, age of patients who are 5 years or older was associated with an increased risk of mortality after HSCT [6]. Survival for patients younger than 5 years was about the same as HLA-identical sibling HSCTs. In addition, it is found by several studies that there are comparable survival rates among recipients of HLA-matched related and unrelated HSCTs. The most commonly used regimen was a combination of busulfan, cyclophosphamide, and ATG. It is suggested that targeting the dose of busulfan to achieve a steady state concentration above 400 ng/ml facilitates engraftment.

Encouraging results have also been reported for umbilical cord blood transplants, but most of them are case reports. Among 15 patients reported by Ozsahin, et al. event-free survival was approximately 70%. It was similar to recipients of unrelated marrow grafts. Most patients in their reports were conditioned with the combination of busulfan, cyclophosphamide and antithymocyte globulin, which they found to be an important factor for improved survival when compared to other regimens. Reconstitution of immunity after cord blood approximates that observed with unrelated marrow grafts [9].

Although the historical results are less encouraging, there is more information about haploidentical transplants for WAS than umbilical cord blood transplants. John MJ, et al. reported literature review of eight studies between 1979 and 2014 about haploidentical transplants for WAS a case report about un-manipulated haploidentical transplant in WAS [10]. According to their review, between 1979 and 2014, there are 17 reported haploidentical transplants in the age group of 2-12 years with 8 survivors. Conditioning regimens used were cytotoxic arabinoside/total body irradiation (TBI) (2/17), busulfan/cyclophosphamide (5/17), cyclophosphamide/ TBI (3/17) and fludarabine based (4/17). Stem cell source was bone marrow in majority (13/17) and the rest used peripheral blood stem cell products (4/17). T cell depletion (TCD) was performed in most of the cases (12/17). Apart from their case, one more unmanipulated haplo-identical transplant was reported. There is a 4-5-fold increase in mortality after haploidentical compared to HLA-identical sibling grafts [7,11]. These reports show a high incidence of graft failure and poor immune reconstitution following T cell-depleted haploidentical HSCTs. Schwinger, et al. provided a promising case report of “mega-dose” purified CD34+ haploidentical grafts used to overcome the barrier to engraftment without endangering GVHD [12]. Our convenient results in this patient in a way that supports the results of Schwinger, et al. can encourage further investigation of the role of using ‘high doses’ of related haploidentical CD34-selected peripheral blood stem cells in patients with WAS at early stages of disease before the beginning of severe, life-threatening complications.
References


