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References


To the editor:

Haploidentical hematopoietic stem cell transplantation for graft-versus-host disease after liver transplantation

A 74-year-old woman with hepatitis C and hepatocellular carcinoma underwent deceased donor liver transplant (LT) from a male donor in August 2008. Donor and recipient were blood group A+ and 2-loci HLA matched (DR01 and DR15). Initial immunosuppression included a corticosteroid taper and tacrolimus. On postoperative day (POD) 28, she developed a papular erythematous rash, diarrhea, and neutropenia. Skin biopsy showed perivascular lymphocytic infiltrate with necrotic keratinocytes and full thickness epidermal necrosis. FISH analysis demonstrated XY-chimerism in both peripheral blood (29.2%) and skin (18.5%, 37/200 nuclei) consistent with graft-versus-host disease (GVHD). Bone marrow biopsy showed < 10% cellularity with XY chimerism (2/10 nuclei) on karyotype analysis. Liver function tests were normal. High dose steroids and mycophenolate mofetil 500mg TID were added. The patient developed refractory neutropenia and thrombocytopenia. After failing a trial of IVIG (0.5mg/kg/d) and rabbit anti-thymocyte globulin (rATG; 1.5mg/kg/d), we sought a donor for rescue hematopoietic stem cell transplantation (HSCT) based on prior success with this strategy in GVHD following HSCT.1

On POD 42 she received a haploidentical CD34+ selected (Baxter Isolex 300i) HSCT (2.01 × 10^6 cells, CD3+ 0.3%) from her son (3-loci HLA match) after 7 days of rATG. A second

![Figure 1. Trend in XY chimerism of peripheral blood relative to bone marrow function. Peripheral blood FISH analysis demonstrating XY chimerism (number of XY nuclei/total nuclei counted) with corresponding absolute neutrophil count (ANC) representing bone marrow function. Recovery of ANC occurred with XY chimerism suggesting either transient engraftment of son’s cells or immunomodulatory effect of CD34+ infusion facilitating resolution of GVHD with subsequent stable marrow function and near complete restoration of patient XX chimerism on POD 113. Time of ANC recovery and platelet recovery are noted as defined by CIBMTR reporting standards. Timing of immunosuppression relative to clinical course is also noted.](image-url)
infusion of \(0.96 \times 10^6\) (CD3\(^+\) 0.4\%) cells was given on POD 45. HSCT was not on study protocol. Informed consent was obtained from both patient and donor. Immunosuppression regimens at HSCT included tacrolimus, mycophenolate mofetil 500mg TID, and prednisone (30mg BID). Neutropenia and thrombocytopenia resolved 10 (POD 53) and 46 (POD 88) days after HSCT, respectively (Figure 1). Chimerism studies demonstrated transient XY chimerism of peripheral blood (13\%) on POD 58 and near complete reconstitution of bone marrow with XX cells on POD 113 (98.85\%; Figure 1). Thirty-two months after LT, she has stable liver graft function and normal hematopoietic cell lines.

Acute GVHD is a rare complication following LT with an estimated incidence of <1\% and a high mortality rate (80\%).\(^1,2,3\) This is the first report of lymphodepletion followed by haploidentical HSCT for the management of GVHD following LT. After failure of traditional therapies involving augmentation of immunosuppression,\(^4,6\) we sought a novel intervention based on prior data demonstrating resolution of refractory GVHD following HSCT in 50\% of patients after secondary allogeneic HSCT.\(^1\)

The mechanism of action is unclear. Graft-versus-tumor effect has been demonstrated with higher doses of CD3\(^+\) infusions (\(\approx 1 \times 10^8\) cells). Our mean CD3\(^+\) yield post-CD3\(^+\) selection of 0.35\% (1 \(\times 10^5\) cells) measured by flow cytometry is below this reported threshold, making a direct cytotoxic graft-versus-graft effect possible but less likely.\(^7\) Transient engraftment has been reported as early as 10 days post-HSCT with subsequent restoration of recipient chimerism by as early as day 28.\(^7\) It is unclear if this occurred since STR analysis was not performed. CD34\(^+\) progenitor cell modulation of antigen presentation may have contributed to resolution of GVHD, similar to mechanisms proposed in mesenchymal stem cell infusions.\(^8,9\) We conclude that rescue HSCT should be considered for GVHD following LT, as most cases are refractory to current approaches.

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