(Austin Publishing Group

Special Article - Hematopoietic Stem Cell Transplantation

The Use of Donor Mesenchymal Stem Cells in the Treatment of Steroid Refractory Graft Versus Host Disease. Ten Years of Single Center Experience

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Received: April 18, 2017; **Accepted:** May 08, 2017; **Published:** May 20, 2017

Short Communication

During the years 2003-2012 there were 94 sibling allogeneic stem cell transplants (alloTx) performed at the National Cancer Institute, Bratislava, Slovakia. From them 43 (46%) patients were treated with reduced intensity conditioning (RI) and 10 (11%) patients received a HLA haploidentical graft. Eight patients (8.5%) developed steroid refractory graft versus host disease (GVHD). After obtaining a general approval from Ethics Committee, all 8 patients had been eligible for the infusion of freshly prepared *ex vivo* expanded mesenchymal stem cells (MSC). The MSC for *ex vivo* expanded according to Koç, et

al. [1]. The patients' characteristics are shown in the Table 1. The mean number of the MSC infused was 0.4x10⁶ cells per kg of body weight. (Range 0.3-0.5x10⁶ cells/kg). Two patients (No 5 and 7) received the MSC twice. Four patients died because of refractory GVHD. From the other 4 patients, two died because of the disease progression, one because of the fungal infection and one is alive up to 10 years after alloTxin complete remission without any signs of cGVHD. It should be noted that all patients were heavily pretreated but the last one who is alive, with more than 5 lines of chemotherapy including autologous stem cell transplantation. Nevertheless, the steroid refractory GVHD resolved in half (4) of them. Moreover it seems to be plausible to have the MSC "in stock" for rapid *ex vivo* expansion in order to use the cells freshly prepared (not frozen, thawed and immediately applied) [2].

References

d +92, d+178

d +43

 Koç O, Gerson S, Cooper B, Dyhouse SM, Haynesworth SE, Caplan AI, et al. Rapid hematopoietic recovery after co-infusion of autologous cultureexpanded human mesenchymal stem cells (hMSCs) and PBPCs in breast cancer patients receiving high dose chemotherapy. J Clin Oncol. 2000; 18: 307-316.

infection

disease progression

Haplo Tx

RI Tx

2. Lakota J. Unpublished observation.

Patient No Gender Born Diagnosis Transplantation MSC infusion Death Reason of death Comment 1971 AMI 16 03 04 d +206 09 11 04 cGvHD 1 Μ 2 Μ 1976 HD 30.01.05 d +1 14.02.05 aGvHD Haplo Tx 3 Μ 1971 ALL 13.12.05 d +9 05.04.06 disease progression 4 F 1970 Ph- CML alive (31.12.16) 26.04.07 d +33 5 F 1978 HD 13.10.09 d +60, d+80 23.02.10 aGvHD **RI Tx** 6 Μ 1969 ALL 23.02.10 d +168 16.08.10 cGvHD

21.10.10

18.10.11

Table 1: Patients' characteristics, date of transplantation, MSC infusion, date of the patients' death and the reason of death (Haplo Tx: HLA Haploidentical Transplantation; RI Tx: Reduced Intensity Conditioning Transplantation).

F

Μ

1973

1987

HD

HD

7

8

22.09.11

09.04.13