Mesenchymal stem cells for inducing tolerance in organ transplantation

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INTRODUCTION

Organ transplantation is useful for treating the end stage of organ failure. The induction of tolerance to the transplanted organ is essential for its long-term survival. Immuneologic tolerance can be induced by immunosuppressive agents and mixed chimerism. Mixed chimerism is a state in which both recipient-and donor-derived blood cells remain in the hematopoietic system after allogeneic hematopoietic stem cells have been transplanted. Mesenchymal stem cells (MSCs), and immune cells such as dendritic cells and T-reg cells play an important role in the induction of tolerance. MSCs secrete cytokines, which modulate the immune response. In particular, they upregulate T-reg cell function and thereby induce tolerance. Intra-bone marrow-bone marrow transplantation recruits both donor-derived HSCs and MSCs, inducing persistent donor-specific tolerance without the use of immunosuppressants. In this review, we summarize the use of MSCs to induce tolerance in organ transplantation.

Keywords: tolerance, mesenchymal stem cells, organ transplantation, mixed chimerism, bone marrow transplantation

IMMUNE TOLERANCE

Immune tolerance includes central tolerance, which occurs in the thymus, and peripheral tolerance, which includes the deletion of effector T cells and the induction of expansion of active T-reg cells (Wood and Sakaguchi, 2003). In central tolerance, T-cell precursors from the bone marrow (BM) enter the thymus and are selected by thymic epithelial cells via positive and negative selection. These selected T cells show tolerance to autoantigens when they enter peripheral lymphoid tissues. Peripheral tolerance includes the induction of anergy, and the active regulation of effector T cells. Immature DCs, T reg cells and MSCs play important roles in the induction of peripheral tolerance. BM-derived immature DCs express low MHC class II and co-stimulatory molecules to promote tolerance to solid organ allografts, and the injection of donor-derived DCs may prevent acute graft-versus-host disease (GVHD) during a follow-up for of more than 2 years (Scandling et al., 2008). Another review summarizes the induction of long-term allograft tolerance through mixed chimerism in small and large animal models, and includes clinical studies (Sachs et al., 2011). Allogeneic chimerism and tolerance have been induced by HSC transplantation in human leukocyte antigen-mismatched patients when kidneys were transplanted—without any evidence of acute GVHD in the recipients. And stable renal function was maintained as a result of the persistent donor chimerism without the use of immunosuppressive agents (Leventhal et al., 2012). Immune cells such as regulatory T (T-reg) cells, immature dendritic cells (DCs), and mesenchymal stem cells (MSCs) play important roles in the induction of immune tolerance in organ transplantation (Wood et al., 2012). Recent reports have suggested that MSC-derived-exosomes, which are released by exocytosis with the plasma membrane, may benefit therapy-refractory GvHD patients. The use of MSC exomes, being-the therapeutically active component of MSCs, provides a number of advantages compared to just MSCs (Kordelas et al., 2014).
The markers of T-reg cells, CD4, CD25, and FoxP3, mainly mature in the thymus. The presence of T-reg cells has been reported to reduce the need for conditioning regimens in the generation of mixed chimerism (Raimondi et al., 2010), being the presence of both donor- and recipient-derived hematopoietic cells in the recipients after BM transplantation (BMT) (Pilat and Wekerle, 2010). It has been shown that mixed chimerism is more effective than full chimerism in combating infectious risk when allogeneic kidney transplants were performed in humans, with non-myeloablative conditioning promoting the mixed chimerism and helping the renal allografts to survive (Buhler et al., 2002; Kawai et al., 2011). MSCs can be isolated from many tissues, and not only support the growth of hematopoietic stem cells (HSCs), but also secrete cytokines to regulate the immune response. Human MSCs promote the generation of CD4+CD25+FoxP3+T-reg cells, and induce tolerance to allografts (Casiraghi et al., 2008; English et al., 2009). Moreover, MSCs secrete matrix metalloproteinases (MMP), protect allogeneic islets, and maintain long-term normoglycemia through MMP-2 and -9 in vitro (Ding et al., 2009).

**MSCs INDUCE TOLERANCE IN ORGAN TRANSPLANTATION**

The functions of MSCs and their effects on immune cells have been summarized in two reviews (Uccelli et al., 2008; Li and Ikehara, 2013). MSCs suppress allogeneic T cell responses by secreting soluble factors such as PGE2, IL-10, and IL-6 (English, 2013), and modulate DC function, indirectly regulate T and B cell activity, delay and prevent the development of GVHD and suppress DC function (Zhang et al., 2009; Aldinucci et al., 2010). MSCs have been shown to alter the NK cell phenotype and suppress proliferation, decrease cytokine levels such as those of TNF, IFNγ and IL-12, and increase IL-10. One report has shown that porcine MSCs inhibit alloreactive T cells through the induction of PGE2 and indoleamine 2,3-dioxygenase (IDO) (Hsu et al., 2013).

The immunoregulatory properties of MSCs have been reported in vitro and in vivo. MSCs suppress T-cell responses, inducing tolerance to transplanted kidney via the expression of IDO (Ge et al., 2010). An infusion of MSCs and rapamycin has been shown to induce heart allograft-specific tolerance, supporting the idea that MSCs might be used for inducing tolerance in a clinical setting (Ge et al., 2009). The MSC infusion leads to an expansion of T-reg cells and prolongs allograft survival in a MHC matched heterotopic heart transplantation model. Moreover, MSC infusion is characterized by reduced numbers of Th1 effector cells (Casiraghi et al., 2008). However, there is one report indicating that donor-specific MSC pre-treatment resulted in a higher degree of kidney cortex tissue damage and elevated creatinine levels in a rat kidney transplantation model (Seifert et al., 2012). Another report showed that MSCs suppressed allogeneic T-cell responses and prolonged the survival of transplanted hearts by improving the Th1/Th2 balance when allogeneic heart transplantation was combined with the intravenous infusion of MSCs (Zhou et al., 2006). Transplanted MSCs may promote revascularization and improve islet graft function after the co-transplantation of islets with MSCs in streptozotocin-induced diabetic rats (Ito et al., 2010).

In clinical trials, intravenous infusions of autologous BM-derived MSCs were given to kidney allograft recipients. Although immunosuppression remained unaltered, there was a resolution of tubulitis without interstitial fibrosis/tubular atrophy in one third of patients. Additionally, five of the six patients displayed a donor-specific downregulation of peripheral blood mononuclear cell proliferation, which was not reported in patients that did not have the MSC treatment. These results suggest that autologous BM-derived MSC treatment provides systemic immunosuppression in allograft transplantation (Reinders et al., 2013). Calcineurin inhibitors (CNIs) have been reported to reduce acute rejection rates in kidney recipients. The infusion of MSCs combined with CNIs improved renal function such as the estimated glomerular filtration rate at the first month after treatment. This therapy decreased the incidence of acute rejection, and also decreased the risk of opportunistic infection at 1 year after treatment (Tan et al., 2012). Infusion of MSCs was used for the treatment of patients who received kidney transplants, and these infused MSCs increased the percentage of CD4+CD25+FoxP3+CD127- T-reg cells and decreased memory T cells, and CD8+ T cell activity. The infusion of MSCs thus appears to be a safe and clinically feasible method for patients receiving organ transplants (Perico et al., 2011). BM-derived MSCs have been used clinically to treat GVHD, decrease the risk of infection, and help induce tolerance in organ transplantation, and one review indicates that MSCs promote tolerance in the case of kidney transplants (Casiraghi et al., 2014).

When MSCs are infused by intravenous injection, they become trapped in the lungs and other tissues, and it is therefore preferable for the MSCs to be directly injected into the bone cavity. Intra-bone marrow-BMT (IBM-BMT) has been shown to efficiently recruit not only donor-derived HSCs but also MSCs in animal experiments (Fukui et al., 2007; Guo et al., 2008; Song et al., 2008). Furthermore, IBM-BMT induced tolerance to adult allogeneic liver in mice (Okazaki et al., 2008). IBM-BMT is a feasible strategy for the induction of persistent donor-specific tolerance, enabling the use of reduced radiation doses as conditioning.
regimens, and obviates the need for immunosuppressants (Guo et al., 2008). IBM-BMT induced tolerance in the case of allogeneic lung transplants, while intravenous BMT failed to do so (Kaneda et al., 2005). HSCs can normally proliferate in major histocompatibility complex (MHC)-compatible MSCs even in allogeneic microenvironments.

In conclusion, MSCs have been shown to prevent GVHD, and to induce tolerance in organ transplantation in both animal and clinical studies (Figure 1). MSCs can be easily isolated from bone marrow and adipose tissue, and their use thus represents a feasible approach in the clinical setting for inducing tolerance in organ transplantation.

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