Novel Induced-Mesenchymal Stem Cells (i-MSCs) Attenuate Severity of ARDS in Septic Sheep
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Abstract or Introduction

Acute respiratory distress syndrome (ARDS) is a respiratory failure resulting from noncardiogenic pulmonary edema. ARDS is commonly caused by bacterial or viral pneumonia. ARDS is associated with very high mortality (36-45%) (1). Currently there is no FDA approved drug therapy for ARDS. A growing body of evidence supports mesenchymal stem cell (MSC)-based therapy as a potential treatment for ARDS. MSCs target a large number of dysregulated inflammatory cytokines and provide tissue repair and pathogen clearing capabilities via multimodal mechanisms-of-action. Previously, we have demonstrated beneficial effects of bone marrow (BM)-derived MSCs in an ovine model of lung injury (2). Although BM-MSCs have shown excellent clinical safety, they have struggled to demonstrate therapeutic efficacy in many diseases of interest. High efficiency and lack of vector persistence make miRNA reprogramming an ideal platform for the development of new autologous and allogeneic MSC therapies. Previously, we have shown that i-MSCs exhibit superior growth potential, secretome, and therapeutic efficacy in a demyelinating disease model compared to BM-MSCs (3), suggesting that i-MSCs may be an ideal cell therapy for devastating inflammatory diseases, such as sepsis induced ARDS.

Objectives

Objective: To test the safety and efficacy of a novel potent iPSC-induced MSCs (i-MSCs) in a clinically relevant sheep model of sepsis-induced ARDS.

Hypothesis: i-MSCs reduce the severity of lung injury and will be well tolerated in sheep model of ARDS induced by instillation of P. aeruginosa into lungs.

Methods

Preparation of induced pluripotent stem cell-derived MSCs (i-MSCs):
First, iPSCs were generated from adult human fibroblasts derived from a dermal punch biopsy through a one-step, high efficiency, immature pluripotent, self-renewing reprogramming protocol using miRNA encoding Oct4, Sox2, c-Myc, and Lin28 (A). These cells expressed pluripotency factors (B). Second, i-MSCs were generated using the STEmAd Mesenchymal Progenitor Kit (STEMCELL Technologies) (A). i-MSCs were differentiated to adipocytes, osteoblasts and chondrocytes (B). i-MSCs underwent 74.8 population doublings before senescence compared to 18.2 population doublings for the BM-MSCs (C).

Results

Figure 1. Effects of i-MSCs on pulmonary gas exchange (A) and pulmonary edema shunt fraction (B). i-MSCs reduced pulmonary shunt and improved oxygenation, thus preventing onset of ARDS. Data are presented as nS.E.M.

Figure 2. i-MSCs markedly reduced pulmonary edema assessed by measuring lung lymph flow (A) and lung wet-to-dry weight ratio (B). The results indicate that i-MSCs reduced pulmonary microvascular hyperpermeability to water. Data are expressed as nS.E.M.

Figure 3. i-MSCs attenuated atelectasis in the lung (A) and better preserved alveolar epithelial integrity (B). Figures C and D illustrate macroscopic pictures of the lungs and chest X-ray at 48 hrs after the injury. Treatment and grouping: Two groups of sheep (n=3 in each) were randomized either to i-MSCS treatment (10x106 cells/kg) intravenously infused at 1 and 24hrs after the injury, or vehicle (Plasma-Lyte A in same manner).

Measured variables:
Pulmonary gas exchange and Pulmonary edema Cardiopulmonary hemodynamics Fluid balance Bacterial clearance

Figure 5. i-MSCS markedly reduced excess fluid requirement (A) and did not augment increases in pulmonary artery pressure (B), indicating that i-MSCs are safe to use.

Figure 6. i-MSCS reduced pulmonary microvascular hyperpermeability to protein as evidenced by higher plasma (A) and lower lung lymph (B) proteins. Data are expressed as nS.E.M.

Figure 8. i-MSCs significantly reduced bacterial burden in lungs, bronchoalveolar lavage fluid (BALF), and spleen (A). Data are expressed as nS.E.M. Number of samples in each group is 3. **p<0.001, *p<0.05.

Conclusions

• i-MSCS displayed characteristics of donor MSCs by differentiating to adipocytes, osteoblasts and chondrocytes
• i-MSCS population doubling was ~4-fold higher vs. that of bone marrow-derived MSCs
• i-MSCS improved oxygenation and prevented onset of ARDS
• i-MSCS reduced pulmonary microvascular hyperpermeability to water and protein and ameliorated severity of pulmonary edema
• i-MSCS reduced fluid requirement
• i-MSCS reduced vasopressor requirement to maintain arterial blood pressure
• i-MSCS significantly improved bacterial clearance
• i-MSCS had no hemodynamic adverse effects

In summary, use of i-MSCs is safe and effective in ameliorating severity of sepsis induced acute lung injury. Further studies increasing sample sizes are warranted. Data from this sheep ARDS study will inform the design and dosing of planned future human clinical trials in ARDS using these i-MSCs.

References

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