**AWARD NUMBER:**
W81XWH-15-1-0024

**TITLE:** Modulation of Ocular Inflammation by Mesenchymal Stem Cells

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**ABSTRACT:**
Ocular injury and inflammations are commonly treated with non-specific anti-inflammatory drugs such as corticosteroids. These non-specific treatments typically target both pathogenic and regulatory cells of the immune system, and are associated with side effects such as infection, cataract and glaucoma. Thus, there is a pressing need for new immunomodulatory strategies that not only inhibit pathogenic immune cells, but also promote critical regulatory immune cells that promote immune quiescence, such as myeloid-derived suppressor cells (MDSC). Bone marrow-derived mesenchymal stem cells (BM-MSC) have shown promise in exerting immunomodulatory functions, and present a novel and potentially effective alternative to the current non-specific immunosuppressive therapies. The goal of this project is to investigate the mechanisms by which BM-MSC modulate maturation and differentiation of immature myeloid progenitor cells into MDSC (away from pathogenic macrophages) to control ocular inflammation. We anticipate that these investigations will identify critical immunomodulatory factors which may be used to design novel, targeted strategies for treating eye injury and inflammation.
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1. INTRODUCTION

Bone marrow derived mesenchymal stem cells (BM-MSC) are a heterogeneous population of stromal cells that have considerable regenerative and immunomodulatory properties. The immunosuppressive effect of BM-MSC on pathogenic immune cells including effector T cells, macrophages and dendritic cells has been established. Studies have shown that BM-MSC inhibit generation, migration and function of macrophages. Preliminary data from our laboratory has shown that during ocular inflammation, BM-MSC are capable of promoting the differentiation of immature myeloid granulocyte-macrophage progenitors (GMP) into regulatory immune cells such as myeloid derived suppressor cells (MDSC) that could counter the effect of inflammatory macrophages. MDSC are a group of highly immunoregulatory cells that are categorized into monocytic MDSC (CD11b+ Ly6C^hi Ly6G^-) and granulocytic MDSC (CD11b+ Ly6C^lo Ly6G^+) based on their phenotype. Monocytic subpopulations of MDSC have significant immunosuppressive activity and their immunomodulatory role in tumors, transplant rejection and chronic inflammatory diseases of the eye has been established. The goal of this project is to investigate the mechanisms by which BM-MSC modulate maturation and differentiation of GMP into MDSC to control ocular inflammation.

2. KEYWORDS: ocular inflammation, mesenchymal stem cells, MDSC

3. ACCOMPLISHMENTS

● What were the major goals of the project?

**Task #1**: Determine mesenchymal stem cell (BM-MSC)-mediated immature myeloid progenitor (GMP) cell differentiation into monocytic versus granulocytic myeloid-derived suppressor cells (MDSC)  
(Jan 2015 – Aug 2015)

**Task #2**: Define critical factors expressed by BMMSC that promote GMP differentiation into MDSC  
(Sep 2015 – Apr 2016)

**Task #3**: Assess effect of in vivo administration of BMSC on MDSC frequency and ocular inflammation  
(May 2016 – Dec 2016)
What was accomplished under these goals?

**Task #1: Determine BM-MSC-mediated GMP differentiation into monocytic versus granulocytic MDSC**  
(Jan 2015 – Aug 2015)

Local IACUC Approval  (100% completion)  (Jan 2015 – Feb 2015)

1) **Major activities:** A postdoctoral research fellow (Dr. Afsaneh Amouzegar, MD) has been hired to perform experiments on this project. A personnel amendment for Dr. Amouzegar to animal protocol was submitted to IACUC and ACURO.

2) **Specific objectives:** ACURO approval, and Dr. Amouzegar’s animal handling training (both online and hands-on) by the Animal Facility Staff and experienced laboratory members.

3) **Significant results:** Both ACURO approval and Dr. Amouzegar’s training are completed.

**Subaim 1.A. Co-culture bone marrow-derived CD14⁺CD11b⁻/lo GMP with BM-MSC**  
(100% completion)  (Mar 2015 – May 2015)

1) **Major activities:** Determine BM-MSC-mediated GMP differentiation into MDSC

2) **Specific objectives:** Co-culture of flow cytometry (FACS) based sorted CD14⁺CD11b⁻/lo GMP cells from the bone marrow of C57BL/6 mice with in vitro cultured BM-MSC.

3) **Significant results:** Frequencies of CD14⁺CD11b⁻/lo GMP cells were determined in bone marrow, spleen, and cervical lymph node of C57BL/6 mouse (Figure 1). GMPs were characterized for the expression of progenitor markers, CD34, c-Kit, FcγRII/III and maturation markers, CD11b, Ly6G and Ly6C (Figure 2). Bone marrow (from femurs) of euthanized C57BL/6 mice was harvested, and single cell suspension was stained with CD14-FITC and CD11b-PE/Cy5 monoclonal antibodies in sterile conditions. Then, using FACS sorting (MoFlo cell sorter, Dako Cytomation), a purified population of immature CD14⁺CD11b⁻/lo GMP cells were freshly isolated. BM-MSC harvested from bone marrow from wild type C57BL6 mice were cultured in vitro. BM-MSC (5x10⁵ cells) were co-cultured with GMP (1x10⁶ cells) in the presence of pro-inflammatory cytokine IFNγ (100 ng/mL). After 72 hours, cultures were terminated and cells were immunostained with CD11b and Ly6G mAbs for multicolor flow cytometry to investigate the expression of CD11b (maturation marker expressed by macrophages) and Ly6G (maturation marker for granulocytes, such as neutrophils). As shown in Figure 3, with the addition of IFNγ, GMPs upregulate the expression of CD11b and Ly6G. However, in the presence of BM-MSCs in the inflammatory environment, there was a significant decrease in the expression levels of CD11b and Ly6G. These findings suggest that BM-MSC suppress GMP differentiation in the inflammatory environment. Furthermore, our results strongly suggest that in the presence of BM-MSCs, GMPs do not differentiate into MDCs and are rather kept in an undifferentiated state.
Figure 1. Frequencies of GMPs in different tissues.

Figure 2. Phenotypic characterization of GMPs. Surface expression of progenitor markers, CD34, c-Kit and FcγRII/III, and maturation markers, CD11b, Ly6G and Ly6C by GMPs.
Figure 3. BM-MSC prevent GMP acquisition of (a) CD11b and (b) Ly6G maturation markers in the inflammatory environment in vitro.

**Subaim 1.B. Co-culture corneal CD14⁺CD11b⁻/lo GMP with BM-MSC**

(100% completion) (June 2015 – August 2015)

1) **Major activities:** determine BM-MSC-mediated corneal GMP differentiation into MDSCs

2) **Specific objectives:** determination of frequencies of GMPs in normal cornea using flow cytometry and immunohistochemistry; co-culture of flow cytometry (FACS) based sorted CD14⁺CD11b⁻/lo GMP cells from the cornea of C57BL/6 mice with invitro cultured BM-MSC.

3) **Significant results:** Cornea of euthanized C57BL/6 mice was harvested and single cell suspension was stained with CD14-FITC, CD11b-PE/Cy5 and CD34-PE monoclonal antibodies for multicolor flow cytometry to determine the frequencies of CD34⁺CD14⁺CD11b⁻ GMPs in normal cornea. Corneal whole mounts were also stained with CD14-FITC and CD11b-PE monoclonal antibodies and visualized using a confocal microscope (Figure 4). In order to investigate the effect of BM-MSC on corneal GMP differentiation, corneas from C57BL/6 mice were enzymatically digested and after preparing single cell suspension, a purified population of immature CD14⁺CD11b⁻/lo GMP cells were freshly isolated using FACS sorting. BM-MSCs (2.5x10⁴ cells) were co-cultured with corneal GMPs (5x10⁴ cells) in the presence of pro-inflammatory cytokine IFNγ (100 ng/mL). After 72 hours, cultures were terminated and cells were immunostained for CD11b surface marker. As shown in Figure 5, in the presence of IFNγ approximately 12% of corneal GMPs acquired the CD11b maturation marker. However, with the addition of BM-MSC in the culture, there was a significant decrease in the frequencies of CD11b⁺ cells, with only 1.5% of cells acquiring the surface marker. Similar to our previous findings, these results
strongly suggest that in the presence of BM-MSCs, GMPs do not differentiate into MDSCs and are maintained in an undifferentiated state.

**Figure 4.** (a) Frequencies of CD34+ CD14+ CD11b- GMPs in the cornea. (b) Confocal microscopy image (×20) of corneal whole mount confirming the presence of CD14+ CD11b- cells in the peripheral corneal stroma.

**Figure 5.** Corneal GMP acquisition of CD11b surface marker. BM-MSCs prevent acquisition of CD11b maturation marker by corneal GMPs in the inflammatory environment.

**Task #2: Define critical factors expressed by BM-MSC that promote GMP differentiation into MDSC** (September 2015-April 2016)

Subaim 2.A. Define critical factors expressed by BM-MSC that promote GMP differentiation into MDSC (100% completion) (September 2015 – December 2015)

1) **Major activities:** determine mechanism of BM-MSC mediated suppression of GMP differentiation
2) **Specific objectives:** Determination of the mechanism by which BM-MSC suppress GMP differentiation and whether this effect is contact dependent or through secretion of soluble factors
3) **Significant results:** Bone marrow (from femurs) of euthanized C57BL/6 mice was harvested, and single cell suspension was stained with CD14-FITC and CD11b-PE/Cy5 monoclonal antibodies in sterile conditions. Then, using FACS sorting (MoFlo cell sorter, Dako Cytomation), a purified
population of immature CD14<sup>−</sup>CD11b<sup>−/lo</sup> GMP cells were freshly isolated. BM-MSC harvested from bone marrow from wild type C57BL6 mice were cultured in vitro. BM-MSC (5x10<sup>5</sup> cells) were cocultured with GMPs (1x10<sup>6</sup> cells) either directly or in the presence of Transwell polycarbonate membrane cell culture inserts (indirectly) in the presence of pro-inflammatory cytokine IFNγ (100 ng/mL). After 72 hours, cultures were terminated and cells were immunostained with CD11b mAb for multicolor flow cytometry to investigate the expression of CD11b differentiation marker. As shown in Figure 6, BM-MSC cultured in direct contact with GMPs suppressed GMP acquisition of CD11b, while in the presence of Transwell, BM-MSC did not have an inhibitory effect on differentiation of GMPs. This observation suggests that the immunomodulatory effect of BM-MSC on GMP is primarily contact-dependent and secretion of MSC-derived cytokines that were hypothesized to facilitate GMP differentiation into monocytic cells is not the primary mechanism by which BM-MSC modulate GMP differentiation.

Figure 6. Effect of BM-MSC on GMP differentiation is contact dependent. BM-MSCs cultured indirectly with GMPs in the presence of Transwell fail to suppress CD11b surface marker acquisition and GMP differentiation into mature inflammatory cells.

Subaim 2.B. Define specific function of BM-MSC expressed factors promoting GMP differentiation into MDSC

(100% completion) (January 2016- April 2016)

1) Major activities: To characterize the receptor involved in BM-MSC’s contact-dependent immunoregulatory function

2) Specific objectives: To determine the mechanism by which in vitro expanded BM-MSC suppress GMP differentiation through cell-cell contact and the receptor involved in such contact dependency.

3) Significant results: Single cell suspension was prepared from in vitro expanded BM-MSC from C57BL/6 mice. In order to investigate the receptor involved in immunoregulatory function of BM-MSC, cells were screened for cell surface expressed inhibitory molecules including Programmed death-ligand 1 (PD-L1), V-set domain-containing T-cell activation inhibitor 1 (VTCN-1), CD200 and Carcinoembryonic antigen-related cell adhesion molecule 1 (CEACAM-1) receptors. mRNA expression levels of these candidate receptors was evaluated using real-time PCR. Our data demonstrated relatively higher expression levels of CD200 on BM-MSCs compared to other receptors (Figure 7a). Furthermore, in order to assess the effect of IFNγ stimulation on upregulation of CD200, in vitro expanded BM-MSC
were stimulated with IFNγ for 18 hours and then using real-time PCR mRNA expression level of CD200 was investigated. These results further demonstrated that BM-MSCs significantly upregulate their expression of CD200 in the inflammatory environment (Figure 7b). Many studies have suggested that the interaction between CD200 and its receptor (CD200R), which is expressed by myeloid cells, is involved in regulation of myeloid cell function. These observations and our PCR data demonstrating high mRNA expression of CD200 on MSCs, suggested to us that CD200 could be the critical receptor mediating the immunoregulatory effect of BM-MSC on GMPs. We then performed flow cytometry to confirm the expression of CD200 and its counter receptor, CD200R on BM-MSCs and GMPs, respectively. In vitro expanded BM-MSC and FACS sorted splenic GMPs were immunostained for CD200 and CD200R antibodies, respectively. As shown in Figure 7c & d, our FACS data confirmed the expression of CD200 and CD200R by MSCs and GMPs, respectively. Next, to investigate the role of CD200 expression by BM-MSC in mediating their suppressive effect on GMP differentiation, using CD200-shRNA, expression of CD200 was silenced on BM-MSCs. Then, either control-shRNA or CD200-shRNA BM-MSCs (5x10^5 cells) were co-cultured with GMP (1x10^6 cells) in the presence of pro-inflammatory cytokine IFNγ (100 ng/mL). After 72 hours, cultures were terminated and cells were immunostained for CD11b surface marker. As shown in Figure 8, while wild-type BM-MSCs (control-shRNA) suppressed GMP acquisition of CD11b differentiation marker, CD200-silenced BM-MSCs (CD200-shRNA) demonstrated significantly abrogated ability in suppressing GMP differentiation, further confirming the critical role of CD200 in the immunoregulatory function of BM-MSCs.

**Figure 7.** (a) Real-time PCR analysis of mRNA expression levels of PDL-1, VTCN-1, CD200 and Ceacam-1 by BM-MSCs. (b) Real-time PCR analysis of CD200 expression levels on resting versus IFNγ-stimulated BM-MSCs. (c & d) FACS plots demonstrating the expression of CD200 and CD200R1 by BM-MSCs and GMPs, respectively.
Figure 8. Regulatory effect of BM-MSCs on GMP differentiation depends on their expression of CD200. Unlike control-shRNA (wild-type) BM-MSCs, CD200-shRNA BM-MSCs fail to suppress acquisition of CD11b maturation marker by GMPs.

Task #3: Evaluate effect of in vivo administration of BM-MSC on MDSC frequency and ocular inflammation (May 2016- December 2016)

Subaim 3.A. Assess effect of BM-MSC on frequencies of MDSC versus macrophages and on expression of inflammatory cytokines (IL-1β and TNF-α) in the eye (100% completion) (May 2016- August 2016)

1) Major activities: To investigate the in vivo effect of BM-MSCs on ocular inflammation

2) Specific objectives: To investigate the in vivo effect of wild-type and CD200-silenced BM-MSC administration on frequencies of GMPs and mature inflammatory cells and expression of pro-inflammatory cytokines in corneal injury

3) Significant results: Corneal injury was induced in C57BL/6 mice by mechanical removal of the corneal epithelium and anterior stroma. Functional expression of CD200 was silenced in in vitro expanded MSCs using shRNA. Control shRNA (wild-type) or CD200 shRNA MSCs (0.5 × 10^6 cells per mouse) were intravenously injected through the tail vein to mice 1 hour after injury. Corneas were harvested 48 hour post-injury. Corneal single cell suspensions were made and immunostained with CD14, CD11b and CD45 mAbs to determine the frequencies of CD14⁺CD11b⁻ GMPs and CD45⁺ inflammatory cells. Our flow cytometry results demonstrated an approximate 5-fold increase in the frequencies of corneal CD14⁺ CD11b⁻ GMPs in mice treated with control shRNA MSCs compared with non-treated mice (Fig. 9a). Interestingly, silencing of CD200 in MSCs abrogated their ability to expand corneal GMP frequencies (Fig. 9a). Next, we investigated the effect of administration of control shRNA versus CD200-shRNA treated MSCs on infiltration of inflammatory cells into the cornea after injury by assessing the frequencies of corneal CD45⁺ cells. Flow cytometry data demonstrated that systemic administration of control shRNA MSCs resulted in significant suppression of inflammatory cell infiltration in the injured cornea. CD200-silenced MSCs were, however, unable to suppress the corneal infiltration of CD45⁺ inflammatory cells (Fig. 9b). Next, real time PCR was performed for the expression of inflammatory cytokine IL-1β in the cornea. Similarly, PCR data demonstrated substantially lower transcript levels of IL-1β in conjunctiva of mice treated with control shRNA MSCs compared to untreated mice. Similar to flow cytometry results, CD200 silenced MSCs lost their ability
to suppress expression of inflammatory cytokine IL-1β in the injured cornea (Fig. 9c). Collectively, our data suggest that expression of CD200 by MSCs is essential for expansion of ocular GMPs and suppression of ocular inflammation.

**Figure 9.** Systemic BM-MSC treatment suppresses ocular inflammation and expands corneal GMP frequencies. (a) Systemic treatment of mice with corneal injury with BM-MSC (shCON) leads to an approximate 5-fold increase in frequencies of corneal GMPs compared to mice treated with CD200-silenced MSCs (shCD200) or untreated mice. (b) Corneal injury leads to a significant increase in the frequencies of inflammatory CD45+ cells. Systemic treatment with BM-MSC (shCON) dramatically suppresses corneal CD45+ cell frequencies, while CD200-silenced BM-MSC (shCD200) show abrogated ability in suppressing CD45+ cell frequencies. (c) Systemic treatment with BM-MSC (shCON) results in a significant decrease in the transcript levels of inflammatory cytokine IL-1β in the eye compared to mice treated with CD200-silenced MSCs (shCD200) and untreated mice.

**Subaim 3.B. Assess effect of BM-MSC on repair of corneal injury**

(100% completion)  (September 2016- December 2016)

1) **Major activities:** To investigate the effect of BM-MSC treatment on corneal wound repair
2) **Specific objectives:** To determine the in vivo effect of wild-type and CD200 silenced MSCs on restoration of corneal structure after injury
3) **Significant results:** Corneal injury was induced in C57BL/6 mice by mechanical removal of the corneal epithelium and anterior stroma. Functional expression of CD200 was silenced in in vitro
expanded MSCs using shRNA. Control shRNA (wild-type) or CD200 shRNA MSCs (0.5 × 10^6 cells per mouse) were intravenously injected through the tail vein to mice 1 hour after injury. Corneas were harvested 48 hour post-injury. Corneal cross sections were prepared and fixed in 4% paraformaldehyde and stained with hematoxylin and eosin (H & E). Images were obtained using a bright field microscope at ×20 magnification. Our H&E staining results demonstrated significant cell infiltration, thinning of corneal epithelium and increased stromal thickness in injured corneas. However, systemic treatment with wild-type or control shRNA-treated MSCs resulted in less cell infiltration and significant restoration of normal corneal structure. Silencing of CD200 expression in MSCs abrogated their regenerative abilities and resulted in significant inflammatory cell infiltration and disruption of corneal structure (Fig. 10).

![Figure 10](image)

**Figure 10.** H&E staining of corneal cross-sections (×20) from normal, untreated, control-shRNA BM-MSC-treated and CD200-shRNA BM-MSC-treated mice demonstrating epithelial and stromal layers and inflammatory cell infiltration. Systemic treatment with wild-type BM-MSC (shCON) results in less cell infiltration and significant restoration of corneal structure. Silencing of CD200 expression in MSCs (shCD200) abrogates their ability to suppress immune cell infiltration and restore normal corneal structure.

**Additional findings:**
We also found that BM-MSCs secrete high levels of hepatocyte growth factor (HGF) in the inflammatory environment of the injured cornea. Our results demonstrated that BM-MSCs restore corneal transparency after injury by inhibiting the expression of opacity inducing α-SMA (α-smooth muscle actin) and TGF-β (transforming growth factor β) through secretion of HGF. In addition, we found that administration of HGF alone can suppress corneal opacity and inflammation.

● **What opportunities for training and professional development has the project provided?**

Abstract of the current findings was presented as a poster presentation in ARVO conference 2016 in Seattle, Washington. The poster was also selected as a finalist and was entered into the members-in-training (MIT) outstanding poster award competition and won the best poster award in the cornea study section.
How were the results disseminated to communities of interest?

Abstract of the current findings was presented as a poster presentation in ARVO conference 2016 in Seattle, Washington. Results of the current study have been submitted as an original research article to the Stem Cells journal and have been accepted for publication.

What do you plan to do during the next reporting period to accomplish the goals?

Nothing to report.

4. IMPACT

- What was the impact on the development of the principal discipline(s) of the project?
  Our novel findings have revealed the critical role of expression of CD200 by BM-MSC in mediating their regulatory function on the differentiation of myeloid progenitor cells. This observation has high translational value as it can serve as the background for the development of potential CD200-based therapies that could modulate the differentiation of progenitor cells to inflammatory cells at the very early stages of inflammation.

- What was the impact on other disciplines?
  CD200-mediated suppression of tissue inflammation can also be investigated in other organ/tissue inflammatory diseases.

- What was the impact on technology transfer?
  Nothing to report

- What was the impact on society beyond science and technology?
  Findings from our study could significantly improve the public knowledge, as BM-MSCs have long been recognized for their substantial regenerative abilities and much less was known about the regulatory function of these cells on immune cells. Our results demonstrate that in addition to their regenerative capacity, BM-MSCs show immunoregulatory properties and have the ability to regulate the function of other immune cells during inflammation.

5. CHANGES/PROBLEMS

- Changes in approach and reasons for change
  Nothing to report

- Actual or anticipated problems or delays and actions or plans to resolve them
  Nothing to report
- **Changes that had a significant impact on expenditures**
  Nothing to report

- **Significant changes in use or care of human subjects, vertebrate animals, biohazards, and/or select agents**
  Nothing to report

### 6. PRODUCTS

- **Publications, conference papers, and presentations**

  **Abstract** of the current findings was presented at the ARVO conference 2016 in Seattle, Washington.


  An original article on the **major findings** of the current project has been accepted for publication in the *Stem Cells* journal.


  An original article from the **additional findings** of this project has been published in the Stem Cell Reports.


- **Website(s) or other Internet site(s):** Nothing to report
- **Technologies or techniques:** Nothing to report
- **Inventions, patent applications, and/or licenses:** Nothing to report
- **Other products:** Nothing to report
7. PARTICIPANTS & OTHER COLLABORATING ORGANIZATIONS

● What individuals have worked on the project?

Name : Sunil Chauhan, DVM, PhD
Project Role : PI
Researcher Identifier : eRA
Nearest person month worked : 2 months
Contribution to Project : PI supervises all aspects of the project, including technical training of postdoctoral fellow, experimental design, troubleshooting, and data analysis. This also includes holding meetings with fellow (twice/week) to discuss progress and direction of the project.

Name : Afsaneh Amouzegar, MD
Project Role : Postdoctoral research Fellow
Researcher Identifier : eRA ID: 12617641
Nearest person month worked : 24 months
Contribution to Project : Under direct supervision of PI, research fellow is responsible for performing the experiments described in this project, collaborating on experimental design and analysis and presentation of data.

● Has there been a change in the active other support of the PD/PI(s) or senior/ley personnel since the last reporting period?
Nothing to report

● What other organizations were involved as partners?
Nothing to report

8. SPECIAL REPORTING REQUIREMENTS
None

9. APPENDICES
i). Quad Chart
Modulation of Ocular Inflammation by Mesenchymal Stem Cells
MR130457 and Vision Research Program - Hypothesis Development Award
W81XWH-15-1-0024

PI: Sunil Chauhan, DVM, PhD  Org: Schepens Eye Research Institute  Award Amount: $249,828

Study/Product Aim(s)
1. Determine mesenchymal stem cell (BM-MSC)-mediated differentiation of immature myeloid progenitors (GMP) into myeloid-derived suppressor cells (MDSC).
2. Define critical factors expressed by BM-MSC that promote GMP differentiation into MDSC.
3. Evaluate effect of in vivo administration of BM-MSC on MDSC frequency and ocular inflammation.

Approach
We first plan to investigate BM-MSC-mediated differentiation of both bone marrow- and cornea-derived immature CD14+CD11b-/-lo progenitors into MDSC. Secondly, we will identify and functionally characterize the BM-MSC-expressed factors that promote MDSC generation. Finally, using a mouse model of corneal injury, we will determine the effect of systemic administration of in vitro expanded BM-MSC on MDSC frequency and ocular inflammation.

Goals/Milestones
✓ IACUC/ACURO Approvals
CY14 Goals –
Define mechanisms by which BM-MSC promote immature CD14+CD11b-/-lo GMP differentiation into MDSC
✓ Culture bone marrow-derived GMP with MSC
✓ Co-culture corneal GMP with BM-MSC
✓ Evaluate BM-MSC expression of factors promoting MDSC

CY15 Goals –
✓ Assess function of BM-MSC factors promoting GMP into MDSC
✓ Effect of in vivo administered BM-MSC on MDSC frequency
✓ Effect of in vivo administered BM-MSC on ocular inflammation

Comments/Challenges/Issues/Concerns: None
Budget Expenditure to Date
Projected Expenditure: $249,828
Actual Expenditure: $249,828

Updated: (6th March 2017)
Restoration of Corneal Transparency by Mesenchymal Stem Cells

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SUMMARY

Transparency of the cornea is indispensable for optimal vision. Ocular trauma is a leading cause of corneal opacity, leading to 25 million cases of blindness annually. Recently, mesenchymal stem cells (MSCs) have gained prominence due to their inflammation-suppressing and tissue repair functions. Here, we investigate the potential of MSCs to restore corneal transparency following ocular injury. Using an in vivo mouse model of ocular injury, we report that MSCs have the capacity to restore corneal transparency by secreting high levels of hepatocyte growth factor (HGF). Interestingly, our data also show that HGF alone can restore corneal transparency, an observation that has translational implications for the development of HGF-based therapy.

INTRODUCTION

A transparent cornea is crucial for optimal vision. Ocular trauma, a leading cause of loss of corneal transparency, accounts for approximately 25 million cases of blindness annually (Resnikoff et al., 2008; Whitcher et al., 2001). During ocular injury, inflammation-induced transforming growth factor β (TGF-β), particularly TGF-β1 and TGF-β2, drive the differentiation of corneal fibroblasts (activated keratocytes) into α-smooth muscle actin (αSMA)-expressing myofibroblasts (Jester et al., 1997; Torricelli et al., 2016), which are themselves opaque and produce disorganized extracellular matrix, leading to the development of corneal opacity and scarring (Jester, 2008; Jester et al., 2012; Ljubimov and Saghizadeh, 2015). Recently, mesenchymal stem cells (MSCs) have been linked to a variety of anti-inflammatory and repair functions in both ocular and non-ocular tissue injuries (Basu et al., 2014; Jiang et al., 2002; Lan et al., 2012; Lee et al., 2014; Uccelli et al., 2008; Wang et al., 2011). However, ocular injuries involving the cornea undergo a wound-healing process that often results in scar formation and loss of corneal transparency. Here, we report that bone marrow-derived MSCs are capable of restoring corneal transparency after injury involving corneal stroma. Specifically, we show that MSCs secrete high levels of hepatocyte growth factor (HGF), which inhibits the generation of opacity-inducing myofibroblasts. Furthermore, we show that HGF alone can restore corneal transparency in an in vivo model of eye injury, a finding that offers an HGF-based therapeutic approach that could potentially eliminate the need for cell-based and conventional therapies.

RESULTS AND DISCUSSION

Inflammatory Milieu Drives MSCs to Secrete Elevated Levels of HGF

The cornea is the most anterior tissue of the eye that comprises the epithelium, stroma, and endothelium (Nishida and Saika, 2011). Ocular injuries involving the stroma (Figure 1A) lead to corneal scarring and compromised vision (Jester, 2008; Whitcher et al., 2001). The aim of this study was to determine whether MSCs have the potential to restore corneal transparency following injury. To investigate this, we first screened MSCs for expression of potential anti-inflammatory and growth factors under both homeostatic and inflammatory conditions. In vitro expanded and functionally characterized bone marrow-derived MSCs (Figures 1B and 1C) were cultured in the absence (medium alone) or presence of interleukin-1β (IL-1β) (to mimic injury-induced inflammatory milieu) for 24 hr, followed by quantification of tumor necrosis factor-stimulated gene 6 (Tsg-6), IL-10, Tgf-β1, and Hgf transcripts using real-time qPCR (Figure 1D). Strikingly, IL-1β stimulation greatly enhanced the expression of Hgf in MSCs compared with unstimulated cells. In contrast, Tgf-β1 expression was significantly reduced in IL-1β-stimulated MSCs. The steady-state expression of Tsg-6 was moderately increased, and IL-10 remained unchanged upon IL-1β stimulation. In addition, ELISA performed on culture supernatants corroborated the qPCR data and showed a 2.5-fold increase in HGF secretion by IL-1β-stimulated MSCs (Figure 1E). These in vitro data demonstrate that MSCs express high levels of HGF in an inflamed environment. We also confirmed these findings using human MSCs. Our data showed that human...
bone marrow-derived MSCs constitutively expressed high levels of HGF, which was significantly upregulated upon stimulation with recombinant human IL-1β (Figure S1A).

To determine whether in vivo administration of MSCs leads to high levels of HGF at inflamed injury site, we utilized a well-characterized sterile injury model of mouse cornea.
Injury was induced by mechanical removal of corneal epithelium and anterior stroma (Figure 1A); 1 hr after injury, MSCs (5 × 10^5/0.1 mL/mouse) were intravenously injected in mice. Using GFP-expressing MSCs (Figure S2), we additionally confirmed that MSCs specifically home to the injured eye (Lan et al., 2012; Omoto et al., 2014). Normal corneas without injury and corneas with injury alone (without MSC administration) served as controls. On day 3 after injury, corneas were harvested, and qPCR and ELISA were performed to measure HGF levels. Indeed, injured corneas from MSC-injected mice showed significantly higher levels of HGF at both transcript (Figure 1F) and protein (Figure 1G) levels compared with injured and normal corneas.

Corneal transparency recovery following MSC transplantation was also assessed in a sterile injury model of mouse cornea (Figures 1A and 2A). HGF expression in MSCs was knocked down using small interfering RNA (siRNA) (Abed et al., 2015), which led to nearly 80% reduction in Hgf expression compared with control siRNA (Figure 2B). MSCs transfected with Hgf siRNA or control siRNA were pre-stimulated with IL-1β for 6 hr, then intravenously administered 1 hr post injury and followed for 5 days. At days 1, 3, and 5 post injury, photographs of injured cornea with or without green fluorescein stain were captured using slit-lamp biomicroscopy. Corneal fluorescein staining was used to indicate epithelial defects and bright-field micrographs were used to evaluate corneal opacity.

The values shown are mean ± SD and each corneal injury group consists of n = 6 mice. *p < 0.02, **p < 0.005.

Capacity of MSCs to Restore Corneal Transparency Is Dependent upon Their HGF Expression

Based on our in vivo data and because previous reports have ascribed an anti-fibrotic function for HGF (Herrero-Fresneda et al., 2006), we hypothesized that HGF could be a putative MSC-expressed factor that could contribute to the restoration of transparency in injured corneas. We therefore determined whether altering HGF expression within MSCs influenced opacity in a sterile injury model of mouse cornea (Figures 1A and 2A). HGF expression in MSCs was knocked down using small interfering RNA (siRNA) (Abed et al., 2015), which led to nearly 80% reduction of Hgf expression compared with control siRNA (Figure 2B). MSCs transfected with Hgf siRNA or control siRNA were pre-stimulated with IL-1β for 6 hr, then intravenously administered 1 hr post injury and followed for 5 days. At days 1, 3, and 5 post injury, photographs of injured cornea with or without green fluorescein stain were captured using slit-lamp biomicroscopy. Corneal fluorescein staining was used to indicate epithelial defects and bright-field micrographs were used to evaluate corneal opacity.

The values shown are mean ± SD and each corneal injury group consists of n = 6 mice. *p < 0.02, **p < 0.005.
Figure 3. HGF Alone Is Sufficient to Inhibit Corneal Opacity and Inflammation
(A and B) A corneal fibroblast cell line (MK/T1) was stimulated with TGF-β1 in the presence or absence of HGF for 24 hr. α-SMA expression was assessed (A) at mRNA level using real-time PCR and (B) at protein level by immunohistochemistry. The values shown are the mean ± SD of three independent experiments.
administered to the mice 1 hr post injury. Injured corneas with no MSC administration served as untreated controls. Slit-lamp biomicroscopy was used to monitor the extent of corneal opacity and wound healing for 5 days. Corneas of mice injected with control siRNA-treated MSCs showed a significant reduction in corneal opacity at days 3 and 5 post injury compared with corneas from Hgf siRNA-treated MSCs and untreated mice (Figures 2C and 2D). To determine the extent of wound repair, we used corneal fluorescein staining to assess the epithelial defect (Figures 2E and 2F). A smaller area of fluorescein (green) represents a faster rate of wound healing. A complete and significantly more rapid wound repair was seen in mice injected with control siRNA-treated MSCs compared with corneas from Hgf siRNA-treated MSCs and untreated control mice. Previous reports have shown similar effects of wild-type MSCs on wound repair (Lan et al., 2012; Lee et al., 2014). After 5 days of injury, corneas were harvested to assess expression levels of α-Sma and Tgf-β1 using qPCR. Data showed a markedly decreased expression of α-Sma and its inducer cytokine Tgf-β (Yi et al., 2014) in the corneas of mice injected with control siRNA-treated MSCs compared with the corneas of Hgf siRNA-treated MSCs and untreated mice (Figures 2G and 2H). These data clearly demonstrate that HGF expression by MSCs is crucial for inhibiting the expression of opacity-inducing α-SMA and TGF-β, and restoring corneal transparency in the injured eye.

**Topical Administration of HGF Alone Is Sufficient to Restore Corneal Transparency in Ocular Injury**

Finally, the functional and translational relevance of HGF in restoring corneal transparency was confirmed by investigating the effect of HGF alone (without MSC administration) using both in vitro and in vivo model systems. First, to experimentally address whether HGF can inhibit expression of α-SMA in corneal fibroblasts, we stimulated a well-characterized corneal fibroblast cell line (MK/T1) (Gendron et al., 2001) with TGF-β1 in the absence or presence of recombinant mouse HGF for 24 hr. Unstimulated cultures served as a control. HGF treatment showed a dose-dependent suppression of TGF-β-induced α-Sma expression in corneal fibroblasts (Figure 3A). Consistent with our data in mice, we also observed that human recombinant HGF completely suppressed TGF-β1-induced α-SMA expression in human corneal fibroblasts (Figure S1B).

We also confirmed the effect of HGF on TGF-β-induced α-SMA protein expression using immunohistochemistry. HGF completely suppressed TGF-β-stimulated α-SMA protein expression in corneal fibroblasts and prevented their conversion to myofibroblasts (α-SMA+ cells; green) (Figure 3B), which are the primary cause of corneal opacity (Jester, 2008; Jiang et al., 2002). Interestingly, HGF treatment (Figures 3A and 3B; media versus HGF) also significantly reduced the baseline expression of α-SMA in corneal fibroblasts, suggesting that HGF alone could be effective in reversing pre-formed myofibroblasts into α-SMA-negative fibroblasts. Using this information, we sought to investigate whether in vivo administration of HGF can suppress corneal opacity. Corneal injury was induced as described above (Figure 1A), 5 μL of 0.1% recombinant mouse HGF or mouse serum albumin (control) was applied topically to the injured eye twice daily for up to 7 days after injury, and slit-lamp biomicroscopy was used to monitor corneal opacity (Figure 3C). At day 3 post injury, both groups showed a significant development of corneal opacity. However, the corneas of HGF-treated mice exhibited a significant reduction in opacity on day 5 and a near complete restoration of transparency on day 7 compared with mouse albumin-treated control corneas (Figure 3D). After 7 days post injury, corneas were harvested to confirm the effect of HGF on injury-induced opacity at cellular and molecular levels. H&E staining of corneal cross-sections revealed normalization of corneal tissue structures only in HGF-treated mice (Figure 3E), whereas albumin-treated control corneas showed a significant increase in tissue thickness accompanied by infiltration of inflammatory cells (Figures 3E and 3F). Moreover, HGF-treated corneas showed increased stratification of the epithelial cell layer (Figures S3A and S3B). Both confocal micrographs of immunostained corneas (Figure 3G) and qPCR (Figure 3H) showed a significant reduction in the expression of α-SMA in HGF-treated corneas compared with control corneas. Moreover, mRNA expression levels of α-SMA-inducer cytokines Il-1β (Figure 3J) and Tnf-α (Figure 3K) were significantly reduced in HGF-treated corneas compared with
albumin-treated corneas. The fact that HGF-treated corneas showed high expression of Hgf-R (c-Met) compared with control corneas (Figure S3C) further supports our finding that HGF signaling inhibits α-SMA expression. Collectively, these findings indicate that HGF administration alone is sufficient to restore transparency in corneal injury by suppressing conversion of corneal fibroblasts into αSMA+ myofibroblasts and by inhibiting tissue infiltration of inflammatory cells, which secrete inflammatory cytokines and proteolytic enzymes, leading to degradation and remodeling of the extracellular matrix (Ljubimov and Saghizadeh, 2015).

Conventional treatments for ocular injuries involving corneal scarring vary from topical immunosuppressive steroids to corneal transplantation. However, (1) the increased risk of infection and delayed wound healing, (2) immune rejection of the transplant, and (3) shortage of cornea donors remain major limitations to such treatment (Hamil, 2011). Recently, due to their unique immunomodulatory property, MSCs have been used in experimental and clinical settings to treat a variety of tissue injuries and inflammatory diseases (Basu et al., 2014; Lan et al., 2012; Lee et al., 2014; Uccelli et al., 2008; Wang et al., 2011). Here, we ascribe a hitherto unknown function of MSCs in restoring corneal transparency following ocular injury. We report that MSCs inhibit the expression of opacity-inducing α-SMA and its inducer TGF-β in the injured cornea by secreting HGF. Furthermore, we show that administration of HGF alone can suppress corneal opacity and inflammation. Given that clinical-grade production of cell-based therapies is cost prohibitive, our findings offer the promise of HGF-based modalities for treating ocular conditions that compromise corneal transparency and vision.

EXPERIMENTAL PROCEDURES

Animals
Six- to 8-week-old male C57BL/6 wild-type mice (Charles River Laboratories) were used in these experiments. The protocol was approved by the Schepens Eye Research Institute Animal Care and Use Committee, and all animals were treated according to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Corneal Injury
Mice were anesthetized and a 3-mm superficial keratectomy was performed as previously described (Basu et al., 2014; Hutcheon et al., 2007). In brief, under a dissecting microscope the central area of the cornea was demarcated with a 3-mm trephine and rotated gently to cut into the stroma. The circular area was traced with a sharp pair of surgical forceps, and the corneal epithelium and basement membrane, including the anterior portion of the stroma, were removed using a hand-held Algerbrush II (Alger Equipment). Following injury, corneas were flushed with sterile saline and subsequently covered with Vetropolyvin (bacitracin-neomycin-polymyxin) ophthalmic ointment.

Conical opacity was determined by taking bright-field images using a biomicroscope. Corneal wounds were monitored by placing 1 μL of 2.5% sodium fluorescein (vital staining) on the ocular surface. After 3 min, the ocular surface was visualized by slit-lamp biomicroscope under cobalt blue light, and digital pictures of corneal defects were captured. Degree of opacity and area of injury (fluorescein-stained green color) were calculated using the NIH ImageJ (version 1.34s) software.

Isolation, Expansion, and Characterization of MSCs
Bone marrow was harvested from femurs of euthanized C57BL/6 mice. MSCs were phenotypically and functionally characterized as per criteria defined by The International Society for Cellular Therapy (Dominici et al., 2006), using the previously described plastic adherence method of MSC cultivation (Lan et al., 2012; Lee et al., 2014), and bone marrow cells were cultured in murine MSC-specific MesenCult medium with supplement (STEMCELL Technologies). Non-adherent cells were removed by changing medium every 2 days, and at passage 2 the MSCs were harvested to be used in experiments. Before using MSCs in indicated experiments, cells were characterized phenotypically for the expression of MSC markers (CD45-SMA+ myofibroblasts and by inhibiting tissue infiltration of inflammatory cells, which secrete inflammatory cytokines and proteolytic enzymes, leading to degradation and remodeling of the extracellular matrix (Ljubimov and Saghizadeh, 2015).

siRNA Transfection
MSCs (1.5 × 10^6 cells) were plated in a 75-cm² flask and incubated for 18–24 hr to reach to 60%–70% confluency. The cells were then washed and transfected with 4.8 μg of Hgf-specific or non-specific control siRNA duplex using transfection reagent in siRNA transfection medium according to the protocol suggested by the manufacturer (Santa Cruz Biotechnology). After overnight incubation, transfection medium was replaced with normal MSC growth culture medium and cells were cultured for an additional 2 days. Knockdown efficiency of siRNA was validated by real-time PCR using Hgf-specific primers after 2 and 5 days of transfection.

MSC or HGF Administration
In vitro expanded wild-type or Hgf-silenced MSCs were pre-stimulated with IL-1β for 6 hr, and 5 × 10^5 MSCs in 100 μL of normal saline per mouse were injected to mice 1 hr after corneal injury. Mice were placed in a restraining tube without anesthesia and the tail cleaned with 70% ethanol. The tail was pulled gently and 5 μL of PBS were injected into the tail vein. Five microliters of 0.1% murine recombinant HGF protein (R&D Systems) or mouse serum albumin (Sigma-Aldrich) staining was used to confirm the differentiation of MSCs into the adipocytes.

In Vitro MK/T1 Cell Stimulation
The mouse corneal fibroblast cell line MK/T1 (Gendron et al., 2001) was seeded at 1 × 10^5 cells per well in 24-well plates and
cultured in medium alone or stimulated with 100 ng/mL murine recombinant TGF-β1 (R&D Systems) in the presence or absence of murine recombinant HGF (R&D Systems) at indicated doses for 24 hr. Cells were then used for evaluation of α-Sma expression by real-time PCR and immunohistochemistry.

RNA Isolation and Real-Time qPCR
Total RNA was isolated using the RNeasy Micro Kit (Qiagen). Isolated RNA was reverse transcribed into cDNA using oligo(dT) primer and SuperScript III (Invitrogen). Real-time qPCR was then performed using Taqman Universal PCR Mastermix and pre-formulated Taqman primers for murine glyceraldehyde-3-phosphate dehydrogenase (Gapdh), Hgf, II-10, Tsg6, II-1β, Tgf-β1, Tnf-α, and α-Sma (Life Technologies). The results were analyzed by the comparative threshold cycle method and normalized to Gapdh as an internal control.

Immunohistochemistry and Histology
Cryosections of the whole eyeball and fibroblast culture on 8-chamber slides were fixed in acetone and blocked with 2% BSA and anti-FcR antibodies (catalog #14-0161-86, Affymetrix eBioscience). The sections were immunostained with Alexa Fluor 488-conjugated anti-α-SMA or isotype-matched control antibodies (#53-6496-80, Affymetrix) overnight at 4°C. Slides were then mounted using Vector Shield mounting medium (Vector Laboratories) and examined under a confocal microscope. For histological evaluation, corneal sections were stained with H&E and examined using bright-field microscopy.

Flow Cytometry
A single-cell suspension of MSCs was prepared and stained with fluorochrome-conjugated monoclonal antibodies and appropriate isotype controls. Antibodies (Biolegend) against CD45 (catalog #103133), CD34 (#119310), SCA-1 (#108105), CD29 (#102207), CD105 (#120407) were used for the phenotypic characterization of MSCs. Stained cells were analyzed on an LSR-II flow cytometer (BD Biosciences).

ELISA
Levels of TGF-β1 and HGF in supernatants of MSC cultures or corneal lysates were analyzed using commercially available murine ELISA kits (R&D Systems) as per the manufacturer’s instructions.

Statistical Analysis
Mann-Whitney U tests or Student’s t tests were performed to determine significance, which was set at p < 0.05. Results are presented as the mean ± SD of three independent experiments. In vivo evaluations and quantification of images of corneal injury and opacity were performed in a masked fashion. Samples sizes were estimated on the basis of previous experimental studies on corneal injury and inflammation (Lan et al., 2012; Basu et al., 2014).

SUPPLEMENTAL INFORMATION
Supplemental Information includes three figures and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2016.09.001.

AUTHOR CONTRIBUTIONS
S.K.M. and M.O. performed experiments, and contributed to data analysis and manuscript writing. A.A., A.S., A.R., and K.R.K. assisted in performing experiments and data analysis. S.K.S. contributed to manuscript revision and data analysis. D.I.S. assisted in GFP-MSC homing experiments. S.K.C. contributed to the underlying hypothesis, designed the experiments, analyzed data, and wrote the manuscript.

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Mesenchymal stem cells modulate differentiation of myeloid progenitor cells during inflammation

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Mesenchymal stem cells modulate differentiation of myeloid progenitor cells during inflammation

Running Title: MSCs modulate myeloid progenitor differentiation

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\textbf{Key Words}: mesenchymal stem cells, myeloid progenitor cells, immunoregulation, differentiation
Abstract

Mesenchymal stem cells (MSCs) possess distinct immunomodulatory properties and have tremendous potential for use in therapeutic applications in various inflammatory diseases. MSCs have been shown to regulate pathogenic functions of mature myeloid inflammatory cells, such as macrophages and neutrophils. Intriguingly, the capacity of MSCs to modulate differentiation of myeloid progenitors to mature inflammatory cells remains unknown to date. Here, we report the novel finding that MSCs inhibit the expression of differentiation markers on myeloid progenitors under inflammatory conditions. We demonstrate that the inhibitory effect of MSCs is dependent on direct cell-cell contact and that this intercellular contact is mediated through interaction of CD200 expressed by MSCs and CD200R1 expressed by myeloid progenitors. Further, using an injury model of sterile inflammation, we show that MSCs promote myeloid progenitor frequencies and suppress infiltration of inflammatory cells in the inflamed tissue. We also find that downregulation of CD200 in MSCs correlates with abrogation of their immunoregulatory function. Collectively, our study provides unequivocal evidence that MSCs inhibit differentiation of myeloid progenitors in the inflammatory environment via CD200-CD200R1 interaction.
Introduction

During hematopoiesis, myeloid lineage-committed progenitors derived from hematopoietic stem cells (HSCs) in the bone marrow give rise to mature myeloid cells such as macrophages and neutrophils [1]. The bone marrow is also home to non-hematopoietic stromal cells such as mesenchymal stem cells (MSCs), which, in addition to providing a niche and trophic support for HSCs, maintain hematopoiesis by sustaining a part of the HSC population in an undifferentiated quiescent state through release of soluble factors and intercellular interactions [2, 3].

Acute inflammatory stresses lead to deviation of hematopoiesis toward preferential induction of committed myeloid progenitors and their subsequent differentiation into mature macrophages and neutrophils [4]. The highly proliferative capacity of myeloid progenitors plays a central role in inflammation-induced myelopoiesis, restoring consumed macrophages and neutrophils at the site of inflammation [5]. Despite the critical role of mature myeloid cells in host defense and resolution of inflammation, excessive innate immune response can have deleterious effects on tissue homeostasis and lead to undesired tissue damage.

In addition to supporting hematopoiesis, MSCs are characterized by their self-renewal and multilineage differentiation potential and unique immunoregulatory properties [6]. Studies on the interaction between MSCs and immune cells have shown that MSCs can regulate functions of mature innate immune cells, including polarization of inflammatory macrophages into an anti-inflammatory phenotype and enhancement of the phagocytic capacity of neutrophils [7, 8].

Although much is known about the regulatory role of MSCs on function of mature myeloid cells, information regarding potential regulatory interactions between MSCs and myeloid progenitor cells is lacking. Given the central role of myeloid progenitors in...
inflammation, regulating differentiation of these precursors into pathogenic myeloid cells could effectively inhibit inflammatory response at an earlier stage. In this study, we sought to determine whether MSCs can inhibit the differentiation of myeloid progenitors into mature inflammatory cells during inflammation. Specifically, we demonstrate that MSCs inhibit differentiation of myeloid progenitors and maintain these cells in an immature state. Using both in vitro co-culture assays and an in vivo model of injury-induced sterile inflammation, we show that MSCs exert immunoregulatory effects on myeloid progenitors in a cell-cell contact dependent manner – a process mediated through CD200-CD200R1 interaction.
Materials and Methods

Animals

Six- to eight-week-old male C57BL/6 mice (Charles River Laboratories, Wilmington, MA, USA) were used in the experiments. Mice were kept in a pathogen-free environment at the Schepens Eye Research Institute Animal Facility. The protocol was approved by the Schepens Eye Research Institute Animal Care and Use Committee, and all animals were treated in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Isolation, expansion, and characterization of mesenchymal stem cells

Bone marrow was harvested from femur bones of euthanized C57BL/6 mice. Using the previously described plastic adherence method of MSC cultivation [9, 10], bone marrow cells were cultured at 37°C in murine MesenCult basal medium and supplement (Stem Cell Technologies, Vancouver, BC, Canada). Cells were passaged at every three to five days intervals. Before using in experiments, MSCs from third passage were characterized phenotypically for the expression of MSC markers (CD45−CD34−SCA1+CD29+) using flow cytometry, and functionally by in vitro differentiation using adipocytes using MesenCult adipogenic stimulatory supplements (Stem Cell Technologies). Oil-red-O (Sigma-Aldrich, St. Louis, MO) staining was used to confirm the differentiation of MSCs into the adipocytes. MSCs from third passage were used in both in vitro and in vivo experiments.

Myeloid progenitor cell characterization and isolation

Single cell suspensions from spleen, bone marrow, and draining submandibular lymph nodes harvested from C57BL/6 mice were stained with fluorochrome-conjugated monoclonal
antibodies to CD14 (#123308), CD11b (#101210), CD34 (#119307), c-kit (#105817), and 
FcγRII/III (#101327) (Biolegend, San Diego, CA, USA) for characterization of myeloid 
progenitors. Due to higher frequencies of myeloid progenitors in the spleen, CD14⁺CD11b⁺ 
progenitors were then isolated from the spleen by flow sorting (MoFlo XDP, Beckman Coulter).
Purity of isolated myeloid progenitors (>95%) was determined by flow cytometry. Isolated 
spleen-derived progenitors were characterized for the expression of progenitor and mature 
myeloid cell markers before being used in in vitro experiments, as described later in Flow 
cytometry method.

**Co-culture and transwell assays**

Isolated spleen-derived myeloid progenitors (2×10⁵ cells) were cultured with or without in-vitro 
expanded MSCs (4×10⁴ cells) for 72 hours in the presence of 100ng/mL IFNγ, 100 ng/mL IL-1β, 
or 10 ng/mL GMCSF (Biolegend, San Diego, CA, USA) as inflammatory or hematopoietic 
growth factor stimuli. For the indirect co-culture, MSCs were first cultured in a monolayer on 6.5 
mm transwell inserts with 0.4 µm pore size (Corning, NY, USA) and then co-cultured with 
isolated myeloid progenitors at the ratio of 1:5 MSCs to myeloid progenitors in the presence of 
100ng/mL IFNγ for 72 hours.

**shRNA transfection**

MSCs (1.5×10⁶ cells) were plated in a 75 cm² flask and incubated for 18-24 hours to reach to 60-
70% confluency. The cells were then washed and transfected with CD200-specific or non-
specific control shRNA using transfection reagent in shRNA transfection media according to the 
protocol suggested by the manufacturer (Santa Cruz Biotechnology, Dallas, TX). After overnight 
incubation, transfection media was replaced with normal MSC growth culture media and cells
were cultured for additional 2 days. Knockdown efficiency of shRNA was validated by real-time PCR using CD200-specific primers 48 hours after transfection (Supplement Fig. S4).

Corneal injury model

Corneal injury was induced in mice as described previously [11, 12]. Briefly, Mice were anesthetized by intraperitoneal injection of Ketamine and Xylazine. Central cornea of deeply anesthetized mice was marked by a 2mm trephine. Using the tip of a hand-held motor brush (AlgerBrush II, Alger Company Inc., Lago Vista, TX), total corneal epithelium and anterior stroma were removed mechanically to create corneal injury. Upon completion of the procedure, triple antibiotic ointment was applied to the injured eyes, and a subcutaneous injection of Buprenorphine was given to mice to minimize injury-induced pain. To study the therapeutic effect of MSCs on corneal inflammation, mice were randomly divided into injury only or MSC (wild-type or CD200 shRNA)-recipient groups, with n=5 in each group. In vitro expanded and characterized MSCs or CD200 shRNA-treated MSCs (0.5×10⁶ cells suspended in 100µL sterile saline) were injected into the tail veins of mice 1-hour post injury. Mice were euthanized 48 hours post injury to collect corneas for flow cytometry, real-time PCR, and fluorescence microscopy analyses as described later.

Flow cytometry

Flow cytometry was performed to characterize the phenotype of in vitro expanded MSCs and myeloid progenitors, to evaluate in vitro differentiation of myeloid progenitors, and to quantify the frequencies of CD45⁺ and myeloid progenitors in the cornea. Cultured MSCs in single cell suspension were stained with conjugated monoclonal antibodies to CD45 (#103115), CD34 (#119307), Sca-1 (#108107), CD29 (#102207), CD11b (#101210), c-Kit (#105817), CD105 (#120407), CD31 (#102407), and CD200 (#123807). Single cell suspensions were prepared from...
bone marrow, spleen, and draining submandibular lymph nodes and were stained with conjugated monoclonal antibodies to CD14 (#123308), CD11b (#101235), CD34 (#119307), c-kit (#105817), FcγRII/III (#101327), Ly6G (#127627), Ly6C (#128007), and CD200R1 (#123907). Single cell suspensions of cultured myeloid progenitors were stained with conjugated monoclonal antibodies to CD11b (#101210), Ly6G (#127627) and CD11c (#117329). Corneas were harvested 48 hours post injury and were digested in RPMI media (Lonza, Walkersville, MD) containing 2 mg/mL collagenase type IV (Sigma-Aldrich, St. Louis, MO) and 2 mg/mL DNase I (Roche, Basel, Switzerland) for 45 minutes at 37 °C and then filtered through a 70-μm cell strainer. Corneal single cell suspensions were then stained with conjugated monoclonal antibodies to CD45 (#103133), CD34 (#119307), CD14 (#123308) and CD11b (#101210). All the antibodies with their matched isotype controls were purchased from Biolegend (San Diego, CA, USA). Stained cells were analyzed using an LSR II flow cytometer (BD Biosciences, San Jose, CA, USA).

RNA isolation, RT-PCR, and quantitative real-time PCR

Corneas were harvested at 48 hours post injury from each group, and mRNA was isolated using the RNeasy Micro Kit (Qiagen, Germantown, MD, USA). Isolated RNA was reverse transcribed into cDNA using oligo (dT) primer and Superscript TM III (Invitrogen, Grand Island, NY, USA). Real-time PCR was performed using Taqman Universal PCR Mastermix and preformulated primers for PDL-1 (Mm00452054_m1), VTCN-1 (Mm00628552_m1), Ceacam-1 (Mm04204476_m1), CD200 (Mm00487740_m1), IL-1β (Mm00434228_m1), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH, Mm99999915_g1) (Thermofisher Scientific, Waltham, MA, USA). The results were analyzed by the comparative threshold cycle method and normalized to GAPDH as an internal control.
Immunofluorescence and histopathology

Freshly excised corneas were washed in PBS and fixed with 4% paraformaldehyde for 20 minutes and permeabilized with 0.5% Triton X-100 for 10 minutes. Whole corneas were then immunostained with FITC-conjugated anti-CD14 (#123308) and PE-conjugated anti-CD11b (#101207) (Biolegend (San Diego, CA, USA) overnight at 4°C to detect myeloid progenitors and mounted onto slides with mounting medium (Vector Laboratories, Burlingame, CA, USA) and visualized using a confocal microscope (Leica TCS-SP5; Buffalo Grove, IL, USA) at ×20 magnification. Corneal sections fixed in 4% paraformaldehyde were stained with hematoxylin and eosin. Images were obtained using a bright field microscope (Nikon Eclipse E800; Melville, NY, USA) at ×20 magnification.

Statistical analysis

A two-tailed Student's t-test was performed and P values <0.05 were regarded as statistically significant. Results are presented as the mean ± standard error of the mean (SEM) of at least three independent experiments. Samples sizes were estimated on the basis of previous experimental studies on corneal injury and inflammation [10, 13].
Results

Characterization of myeloid progenitor cells

As immune cells are primarily developed in lymphoid organs, single cell suspensions from bone
marrow, spleen, and submandibular lymph nodes were immunostained for flow cytometry
analysis as per the gating strategy shown in Supplement Figure S1. First, a population of
CD14^+CD11b^- cells was identified (Fig. 1A) and gated for further characterization and for
examining the expression of progenitor cell markers, including CD34, c-Kit, and FcγRII/III, as
well as the mature myeloid cell markers, Ly6G granulocytic marker and Ly6C monocytic marker
(Fig. 1B). Majority of CD14^+CD11b^- cells (~80%) were positive for the expression of CD34, c-
Kit and FcγRII/III progenitor markers, and all (~99%) were negative for Ly6G and Ly6C mature
myeloid cell markers (Fig. 1B). Based on our results, we estimate that myeloid progenitors
(CD34^+c-Kit^+FcγRII/III^+CD14^+CD11b^-Ly6G^-Ly6C^-) constitute 4.8±1.09% of bone marrow
cells, 8.7±0.29% of splenocytes, and 4.1±0.31% of lymph node cells (Fig. 1C).

MSCs inhibit differentiation of myeloid progenitor cells in vitro

Bone marrow-derived MSCs were cultured and characterized as per criteria defined by The
International Society for Cellular Therapy [9, 10]. MSCs were expanded using the plastic
adherence method, and were characterized phenotypically for positive expression of SCA1 and
CD29 and negative expression of CD45 and CD34 surface markers, and functionally by their
ability to differentiate into adipocytes (Fig. 2A). Next, sorted myeloid progenitor cells were
cultured with or without MSCs in the presence or absence of inflammatory or hematopoietic
growth stimuli such as IFNγ, IL-1β or GMCSF, which have been implicated in myeloid cell
differentiation[14-16]. After stimulation with IFNγ, IL-1β or GMCSF, expression of mature
myeloid cell markers, including CD11b (marker for macrophages; also known as macrophage-1 antigen [Mac-1]) and Ly6G (marker for granulocytes) was investigated using flow cytometry to assess progenitor cell differentiation. Our data showed that upon stimulation with IFNγ, myeloid progenitors acquire high expression of both CD11b and Ly6G. Further analysis demonstrated a significant reduction (55%) in expression of CD11b by myeloid progenitor cells cultured with MSCs in contrast to those cultured without MSCs (MFI 4.15 ±1.04 vs. 9.2 ±1.6; p= 0.000065) (Fig. 2B), and a significant 58% suppression in expression of Ly6G in progenitors cultured with MSCs compared to progenitors cultured without MSCs (MFI 9.9 ± 0.75 vs. 23.02 ±1.14, p=0.0000013) (Fig. 2C). Strikingly, our data that myeloid progenitors fail to express CD11c in the steady state or upon stimulation, suggest that these myeloid progenitors do not differentiate into dendritic cells (Supplement Fig. S2). Similar to effects of IFNγ on myeloid progenitors, stimulation with IL-1β and GMCSF also resulted in selective expression of CD11b by myeloid progenitors, which was significantly suppressed in myeloid progenitors co-cultured with MSCs (Supplement Fig. S3). Taken together, these data suggest that MSCs suppress acquisition of differentiation markers by myeloid progenitors and maintain these cells in an immature state in an inflammatory environment.

**MSCs interact with myeloid progenitors in a cell-cell contact-dependent manner**

To delineate whether the inhibitory effect of MSCs on myeloid progenitor differentiation was through direct cell-cell contact or by MSC-secreted soluble factors, MSCs were either cultured in direct contact with isolated myeloid progenitors, or were first plated into transwell inserts and then cultured with myeloid progenitor cells with IFNγ stimulation. Expression of CD11b surface marker was assessed using flow cytometry. As shown in Figure 3A, MSCs that were cultured
directly with myeloid progenitors significantly suppressed acquisition of CD11b differentiation marker by these cells. However, MSCs in the transwell chamber system failed to suppress CD11b expression by myeloid progenitors, suggesting that the suppressive function of MSCs is dependent on direct cell-cell contact rather than secretion of soluble factors by MSCs. To further explore the molecular mechanism underlying such contact-dependency, we investigated the expression of following cell membrane-bound inhibitory molecules by MSCs using real time PCR: programmed death-ligand 1 (PD-L1), a transmembrane protein, which delivers inhibitory signals to immune cells upon binding with PD-1 expressed by T cells and activated monocyte; v-set domain containing T cell activation inhibitor 1 (VTCN-1) or B7-H4, a transmembrane protein that negatively regulates the function of T cells and neutrophils [17]; CD200 (OX2), a transmembrane glycoprotein that inhibits function of myeloid immune cells [18]; and carcinoembryonic antigen-related cell adhesion molecule 1 (Ceacam-1), a cell adhesion molecule involved in contact-dependent regulation of the innate and adaptive immune responses [19] (Fig. 3B). Significantly higher mRNA expression of CD200 compared to other molecules prompted us to speculate that CD200 may be the critical ligand mediating the immunoregulatory function of MSCs. Our data further demonstrated that MSCs significantly upregulate their expression of CD200 in the inflammatory environment (Fig. 3C). Using flow cytometry, we also confirmed protein expressions of CD200 on MSCs and its receptor, CD200R1 on myeloid progenitor cells (Fig. 3D & E).

**MSCs inhibit differentiation of myeloid progenitors via CD200-CD200R1 interaction**

Next, to investigate the role of CD200 in mediating the immunoregulatory function of MSCs in vitro, functional expression of CD200 on MSCs was silenced using CD200-shRNA (Supplement
Control-shRNA or CD200-shRNA treated MSCs were then cultured with myeloid progenitors in the presence of IFNγ. Our data regarding expression of CD11b demonstrated that CD200-shRNA-treated MSCs had 22% less ability in suppressing myeloid progenitor differentiation compared to control-shRNA-treated MSCs (p=0.008) (Fig. 3F). Compromised ability of CD200-shRNA MSCs to suppress myeloid progenitor acquisition of CD11b suggests that expression of CD200 by MSCs is critical for their inhibitory function on myeloid progenitor differentiation.

**CD200 expression in MSCs is indispensable for suppression of inflammation and accumulation of undifferentiated myeloid progenitors in the inflamed tissue**

Lastly, we chose a sterile inflammation in vivo model of mouse eye injury – a well-established system to study inflammation [13, 20] – to confirm the immunoregulatory effect of MSCs on myeloid progenitors. This well-characterized model provides an excellent system to study inflammation. Simple anatomy of the eye and its paucity of resident immune cells facilitate study of recruited immune cells and their contribution to the inflammatory response [21]. As demonstrated previously [10, 22], we show that MSCs administered systemically home specifically to the injured cornea (Supplement Fig. S5). Interestingly, similar to the bone marrow, spleen and lymph node, we identified a population of CD34^+CD14^-CD11b^- myeloid progenitors in the cornea (Fig. 4A). Our immunofluorescence microscopy results also confirmed the presence of CD14^-CD11b^- progenitors primarily in the stromal layer of cornea (Fig. 4B). Similar to lymphoid tissue-derived progenitors, upon stimulation with IFNγ, sorted corneal myeloid progenitors expressed CD11b, and MSCs suppressed their acquisition of CD11b in vitro (Supplement Fig. S6). To determine the effect of systemic administration of MSCs on myeloid
progenitor cell frequencies and tissue inflammation, mice were intravenously injected with
control-shRNA or CD200-shRNA-treated MSCs 1 hour after corneal injury induction, followed
by harvesting of corneas 48 hours post-injury (Fig. 4C). Our flow cytometry data demonstrated
that normal (control-shRNA-treated) MSCs led to a 5-fold increase in the frequencies of corneal
myeloid progenitors, while CD200-shRNA-treated MSCs failed to do so, suggesting that MSC
expression of CD200 is important for expansion of myeloid progenitor cell frequencies in the
inflamed tissue (Fig. 4E). Similar to our previous findings on the anti-inflammatory effect of
MSCs in the inflamed tissue, our data demonstrated that normal (control-shRNA-treated) MSCs,
but not CD200-silenced MSCs, have a significant suppressive effect on tissue inflammation as
evidenced by reduced frequencies of CD45^+ cells (Fig. 4D), decreased expression of
inflammatory cytokine IL-1β (Fig. 4G) and less inflammatory cell infiltration in the corneal
stroma (Fig. 4F) compared to untreated mice with corneal injury. These findings strongly suggest
that MSCs suppress tissue inflammation by reducing inflammatory cell infiltration and by
expanding frequencies of myeloid progenitor cells through a CD200-dependent mechanism.
Discussion

The current study ascribes a novel immunoregulatory function for MSCs on myeloid progenitor cell differentiation. Our data indicate that MSCs inhibit differentiation of myeloid progenitor cells in an inflammatory environment through direct cell-cell contact. Furthermore, we demonstrate that this intercellular contact is mediated by CD200-CD200R1 interaction, and that CD200 expression by MSCs is indispensable for inhibition of myeloid progenitor differentiation and suppression of tissue inflammation.

Myeloid progenitors are precursors of mature myeloid cells, critical effector cells in innate immune response. Upregulation of pro-inflammatory cytokines such as IFNγ, IL1β and TNFα during inflammation activates steady state progenitors in the bone marrow to differentiate into myeloid effector cells [4, 23]. Myeloid progenitors are primarily found in the bone marrow and cord blood [24]. Some studies have demonstrated the presence of undifferentiated monocytes and DC precursors in non-bone marrow tissues such as spleen [25, 26]. Our findings demonstrate a population of myeloid progenitors, which in addition to the bone marrow are also present in peripheral lymphoid tissues, including spleen and lymph nodes. These progenitors express high levels of CD34, CD14, c-Kit and FcγRII/III progenitor markers, which makes them phenotypically similar to early myeloid progenitors such as common myeloid progenitors (CMPs) and granulocyte/macrophage progenitors (GMPs) [27, 28]. CMPs are thought to be precursors of common DC progenitors, which eventually give rise to DCs [27]. Our data, however, demonstrate that the myeloid progenitors identified do not express the DC marker CD11c in the steady or activated states, suggesting that these cells are not DC precursors. Rather, these progenitors acquire high levels of CD11b and Ly6G myeloid markers in the inflammatory
milleu, suggesting these myeloid progenitors are phenotypically closer to GMPs that give rise to macrophages and granulocytes [1, 27, 29].

Interestingly, we find that MSCs inhibit acquisition of CD11b and Ly6G differentiation markers on myeloid progenitors. MSCs have been shown to interact with cells of both innate and adaptive immunity [30]. Recent reports on the interaction of MSCs with DC precursors have demonstrated that MSCs inhibit differentiation of peripheral blood-derived CD14+ monocytes to mature DCs [31, 32]. Here, our data show that MSCs negatively regulate both bone marrow- and peripheral lymphoid tissue-resident myeloid progenitors. MSCs maintain these cells in an undifferentiated quiescent state and further prevent their differentiation into inflammatory cells.

MSCs primarily exert their immunoregulatory effects through secretion of paracrine factors such as IDO, IL-10, TGF-β and TSG6 [33, 34]. In contrast, we find that MSCs inhibit differentiation of myeloid progenitors mainly through direct cell-cell contact. The results of our study demonstrate that silencing of CD200 expression in MSCs abrogates their ability to suppress myeloid progenitor cell differentiation, suggesting that CD200-CD200R1 interaction is critical for MSCs to exert their immunoregulatory effect. CD200 or Ox-2 is a transmembrane glycoprotein, which binds to its receptor CD200R1 [35]. The CD200R family of receptors consists of 4 isoforms [36], among which CD200R1 is mainly expressed by myeloid cells and T cells [35, 37]. CD200-CD200R1 pathway plays a central role in regulation of innate immune system by inhibiting myeloid cell activation [38, 39]. We show that bone marrow-derived MSCs constitutively express CD200, and significantly upregulate CD200 expression in response to inflammatory stimuli. These results are consistent with previous studies which demonstrated that IFNγ in particular induces CD200 expression in bone marrow-derived stromal cells [40].
Finally, the functional relevance of MSC regulation of myeloid progenitor cell differentiation during inflammation was tested using a standardized mouse cornea model of sterile injury [10, 41]. Similar to the bone marrow, spleen and lymph nodes, we have identified myeloid progenitors residing in the cornea. MSCs have been shown to migrate to the sites of inflammation and promote wound repair [42, 43]. Previously, we showed that systemically administered MSCs home to the inflamed eye, and accelerate wound healing [10, 22]. Here, we show that MSCs suppress infiltration of inflammatory cells and increase the frequencies of corneal myeloid progenitors. Consistent with our in vitro findings, systemically administered CD200-shRNA-treated MSCs lose their ability to suppress differentiation of myeloid progenitors and tissue inflammation. The increase in myeloid progenitor frequencies at the inflamed site could be the result of MSC-mediated expansion of corneal resident myeloid progenitors, or due to MSCs inhibiting differentiation of recruited myeloid progenitors from the bone marrow. Early myeloid progenitors have recently been identified as immunosuppressive cells that are capable of inhibiting T cell proliferation [44]. If MSCs promote recruitment of myeloid progenitors to the inflamed tissue, suppression of inflammation could be the cumulative result of MSC-and myeloid progenitor-mediated regulation of the immune response. However, we acknowledge that further experiments will be needed to elucidate the exact mechanism by which MSCs promote myeloid progenitor frequencies at the site of inflammation.
Conclusion

In conclusion, our findings provide new insight into the immunoregulatory effect of MSCs on myeloid progenitor cell differentiation. Herein, we show that MSCs suppress inflammation not only by regulating inflammatory cell infiltration, but also by preventing differentiation of early myeloid precursors into inflammatory cells. Our data further supports a critical role for CD200 expressed by MSCs in regulating function of myeloid progenitors and thus inhibiting inflammatory response. These observations could provide a framework for the development of potential CD200-based therapeutics that could effectively modulate the generation of innate immune cells and inhibit inflammation at early stage.
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Disclosure of Potential Conflicts of Interest: The authors declare no potential conflicts of interest.


**Figure legends**

**Figure 1. Frequencies and phenotypic characterization of myeloid progenitor cells.**

A. Representative flow cytometric dot plots showing gating strategy for selecting CD14+CD11b- cells in the bone marrow, spleen, and submandibular lymph nodes. 


C. Bar chart comparing the frequencies of myeloid progenitors in the bone marrow, spleen and lymph node as analyzed by flow cytometry. Representative data from 3 independent experiments are shown and each experiment consisted of 5 animals. Data is represented as mean ± SEM.
Figure 2. MSCs inhibit differentiation of myeloid progenitor cells in vitro. 

A. Left Panel: Expansion and characterization of MSCs. Microscopic images of MSCs cultured in MSC or adipogenic media. Oil-Red-O staining after 2 weeks showed red colored fat vacuoles (black arrows) in the cytoplasm of MSCs cultured in adipogenic media, confirming their differentiation into adipocytes. 

B. Right Panel: Representative flow cytometry plots demonstrating the phenotype of bone marrow derived MSCs as CD45-CD34-CD11b-Kit-CD31-Sca-1-CD29-CD105+ cells. 

C. B. Representative flow cytometric histograms and bar chart demonstrating CD11b expression by myeloid progenitors (MPs) cultured with or without MSCs with IFNγ stimulation for 72 hours. 

C. C. Representative histograms of flow cytometric data and bar chart showing Ly6G expression by myeloid progenitors (MPs) cultured with or without MSCs with IFNγ stimulation for 72 hours. 

Results are representative of 3 independent experiments. Myeloid progenitors were isolated from a pool of 5-6 animals in each experiment. P values are calculated using student’s t-test and data is represented as mean ± SEM. *p< 0.0001.
Figure 3. MSCs inhibit differentiation of myeloid progenitors in a contact-dependent manner. A. Representative flow cytometric histograms and bar chart demonstrating CD11b expression by myeloid progenitors (MPs) cultured with MSCs either in direct contact or using a transwell chamber system, which separated MSCs from MPs, in the presence of IFNγ for 72 hours. Results are representative of 3 independent experiments. Myeloid progenitors were isolated from a pool of 5-6 animals in each experiment. B. Real-time PCR analysis of PD-L1, VTCN-1, CD200 and Ceacam-1 mRNA expression levels by resting MSCs. C. Real-time PCR analysis of CD200 expression on resting and IFNγ-stimulated MSCs. Representative flow cytometric histograms demonstrating expression of D. CD200 on MSCs, and E. CD200R1 on myeloid progenitor cells. F. Representative flow cytometric histograms and bar chart showing CD11b expression levels in myeloid progenitors cultured with control-shRNA (shCON) or CD200-shRNA (shCD200) MSCs with IFNγ stimulation for 72 hours. Results are representative of 3 independent experiments. Myeloid progenitors were isolated from a pool of 5-6 animals in each experiment. P values are calculated using student’s t-test and data is represented as mean ± SEM. *p< 0.01, ** p< 0.001
Figure 4. MSCs suppress ocular inflammation through expansion of myeloid progenitor cells in a CD200-dependent manner. A. Representative flow cytometric plots demonstrating CD34^+CD14^+CD11b^+ myeloid progenitor in the cornea. B. Confocal microscopy image (×20) of corneal whole mount confirming the presence of CD14^+ CD11b^- cells in the peripheral corneal stroma (Green: CD14, Red: CD11b). C. Schematic representation of sterile injury induction in mouse and experiment timeline. Corneal epithelium and anterior stroma are mechanically removed using Algerbrush II. D. Bar chart demonstrating the frequencies of infiltrating corneal CD45^+ cells in naïve mouse, injured mice without systemic MSC treatment, control-shRNA (shCON) MSC-treated and CD200-shRNA (shCD200) MSC-treated mice. E. Representative flow cytometric plots and bar chart demonstrating the frequencies of myeloid progenitors in naïve cornea, injured cornea, injured cornea with IV administration of control-shRNA-treated MSCs, and injured cornea with IV administration of CD200-shRNA-treated MSCs 48 hours after injury induction. F. H&E staining of corneal cross-sections (×20) from naïve, untreated, control-shRNA MSC-treated and CD200-shRNA MSC-treated mice demonstrating epithelial and stromal layers and inflammatory cell infiltration. G. Real-time PCR analysis of relative expression of IL-1β mRNA in naïve mice, injured mice without systemic MSC treatment, control-shRNA MSC-treated and CD200-shRNA MSC-treated mice. Results are representative of 2 independent experiments. Each group consisted of 4-5 animals in each experiment. P values are calculated using student’s *t*-test and data is represented as mean ± SEM. * p< 0.05, ** p< 0.01, *** p< 0.001
Figure 1.

(a) Isotype, Bone Marrow, Spleen, Lymph Node.

CD14

<table>
<thead>
<tr>
<th>Isotype</th>
<th>Bone Marrow</th>
<th>Spleen</th>
<th>Lymph Node</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1%</td>
<td>4.9%</td>
<td>8.4%</td>
<td>3.1%</td>
</tr>
</tbody>
</table>

CD11b

(b) Gated on CD14^+CD11b^-.

Counts

- CD34
- c-Kit
- FcγRI/II

MFI: 29
MFI: 17
MFI: 14

Counts

- CD11b
- Ly6G
- Ly6C

<1%
<1%
<1%

(c) Myeloid Progenitors

Frequency (%)

- BM
- Spleen
- LN

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Figure 2.

(a) Culture morphology and differentiation to adipocytes.

(b) MFI analysis of CD11b expression under different conditions.

(c) MFI analysis of Ly6G expression under different conditions.
Figure 3.

a) MPS+IFNγ, MPS+IFNγ+MSC (Contact), MPS+IFNγ+MSC (Transwell) with CD11b expression.

b) mRNA expression of CD200, Fold change.

c) CD200 expression with IFNγ.

d) MSCs with CD200 and CD200R1 expression, 80% MFI: 21.1, 60% MFI: 12.4.

e) MPs with CD200R1 expression.

f) MPS+IFNγ, MPS+IFNγ+shCON-MSC, MPS+IFNγ+shCD200-MSC with CD11b expression.

Protein expression (Fold change)
Figure 4.

a) Isotype

b) Count

Figure 4.

c) Epi.

Figure 4.

d) Total CD45+ cells

Figure 4.

e) CD14

Figure 4.

f) Normal

Figure 4.

g) IL-1β

Figure 4.

h) CD14

Figure 4.

i) CD11b

Figure 4.

j) CD34

Figure 4.

k) CD14 | CD11b

Figure 4.

l) CD14 | CD11b

Figure 4.

m) CD14 | CD11b

Figure 4.

n) CD14 | CD11b

Figure 4.

o) CD14 | CD11b

Figure 4.

p) CD14 | CD11b

Figure 4.

q) CD14 | CD11b

Figure 4.

r) CD14 | CD11b

Figure 4.

s) CD14 | CD11b

Figure 4.

Paper dimensions: 612.0x792.0

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SUPPLEMENTAL DATA

Figure S1. Myeloid progenitor cell characterization. Representative flow cytometry plots demonstrating the gating strategy for characterization of myeloid progenitors as CD34⁺CD14⁺c-Kit⁺FcγRII/III⁺CD11b⁻ cells.

Figure S2. Myeloid progenitor cells do not differentiate into dendritic cells in the inflammatory environment. Isolated myeloid progenitors were stimulated with 100 ng/mL of IFNγ for 72 hours and acquisition of dendritic cell marker CD11c was evaluated using flow cytometry. Flow cytometric histograms demonstrate that myeloid progenitors do not express CD11c in the steady state or after stimulation with IFNγ.
Figure S3. MSCs inhibit differentiation of myeloid progenitors stimulated with IL-1β or GMCSF. Myeloid progenitors were cultured with or without in vitro-expanded MSCs in the presence of (a,b) 10 ng/mL GMCSF or (c) 100 ng/mL IL-1β. As demonstrated in flow cytometric histograms and bar charts, myeloid progenitors express CD11b once stimulated with IL-1β or GMCSF, and MSCs suppress acquisition of CD11b by both IL-1β-treated and GMCSF-treated myeloid progenitors.

Figure S4. Knockdown efficiency of CD200-specific shRNA. MSCs were transfected with CD200-specific or non-specific control shRNA. CD200-specific shRNA suppressed endogenous expression of CD200 in MSCs by 70% 48 hours after transfection.
Figure S5. MSCs home specifically to the injured cornea. Confocal microscopy image (×40) of corneal whole mounts demonstrating the presence of systematically administered green fluorescent protein (GFP)-positive MSCs in the injured cornea, but not in the contralateral cornea 48 hours after induction of injury.

Figure S6. MSCs inhibit acquisition of CD11b by corneal myeloid progenitors in vitro. Isolated corneal myeloid progenitors (corMPs) were cultured with or without in vitro-expanded MSCs in the presence of 100 ng/mL IFNγ for 72 hours. As demonstrated in flow cytometric histograms, once stimulated with IFNγ, corneal myeloid progenitors express CD11b. However, MSCs inhibit acquisition of CD11b differentiation marker by IFNγ-treated corneal myeloid progenitors.