A p38 MAPK-Mediated Alteration of COX-2/PGE₂ Regulates Immunomodulatory Properties in Human Mesenchymal Stem Cell Aging

Kyung-Rok Yu¹,²,³, Jin Young Lee¹,²,³, Hyung-Sik Kim¹,²,³, In-Sun Hong¹,², Soon Won Choi¹,², Yoojin Seo¹,², In sung Kang¹,², Jae-Jun Kim¹,², Byung-Chul Lee¹,², SeungHee Lee¹,³, Andreas Kurtz¹,²,⁴, Kwang-Won Seo¹,²,³, Kyung-Sun Kang¹,²

¹Adult Stem Cell Research Center, College of Veterinary Medicine, Seoul National University, Seoul, South Korea, ²Research Institute for Veterinary Medicine, College of Veterinary Medicine, Seoul National University, Seoul, Korea, ³Institute for Stem Cell and Regenerative Medicine in Kangstem Biotech, Biotechnology Incubating Center, Seoul National University, Seoul, Korea, ⁴Berlin-Brandenburg Center for Regenerative Therapies, Berlin, Germany

Abstract

Because human mesenchymal stem cells (hMSC) have profound immunomodulatory effects, many attempts have been made to use hMSCs in preclinical and clinical trials. For hMSCs to be used in therapy, a large population of hMSCs must be generated by in vitro expansion. However, the immunomodulatory changes following the in vitro expansion of hMSCs have not been elucidated. In this study, we evaluated the effect of replicative senescence on the immunomodulatory ability of hMSCs in vitro and in vivo. Late-passage hMSCs showed impaired suppressive effect on mitogen-induced mononuclear cell proliferation. Strikingly, late-passage hMSCs had a significantly compromised protective effect against mouse experimental colitis, which was confirmed by gross and histologic examination. Among the anti-inflammatory cytokines, the production of prostaglandin E₂ (PGE₂) and the expression of its primary enzyme, cyclooxygenase-2 (COX-2), were profoundly increased by pre-stimulation with interferon gamma (IFN-γ) and tumor necrosis factor alpha (TNF-α), and this response was significantly decreased with consecutive passages. We demonstrated that the impaired phosphorylation activity of p38 MAP kinase (p38 MAPK) in late-passage hMSCs led to a compromised immunomodulatory ability through the regulation of COX-2. In conclusion, our data indicate that the immunomodulatory ability of hMSCs gradually declines with consecutive passages via a p38-mediated alteration of COX-2 and PGE₂ levels.

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* Email: kangpub@snu.ac.kr (KSK); kwang_wons@hotmail.com (KWS)

These authors contributed equally to this work.

Introduction

MSCs have been isolated from almost all tissues [1], and they exhibit a fibroblastic spindle shape and can be directed to differentiate into several different cell types, such as adipocytes, chondrocytes and osteoblasts [2]. It has been reported that MSCs play critical roles in many physiological functions, such as tissue homeostasis, regeneration and wound healing [3]. Together with their broad tissue distribution and ability to locate sites of injury, the immunomodulatory properties of MSCs hold great potential for therapeutic use [4,5]. The immunomodulatory properties of MSCs are elicited by proinflammatory cytokines, such as IFN-γ, TNF-α and IL-1, which produced during an immune response [6]. The combination of these proinflammatory cytokines provokes the production of several inducible soluble factors, specifically, transforming growth factor-β1 (TGF-β1), prostaglandin E₂ (PGE₂), nitric oxide (NO) and indoleamine 2, 3-dioxygenase (IDO), which in turn induce the immunosuppressive functions of MSCs [3,5]. Interestingly, proinflammatory cytokine-stimulated murine MSCs use NO as a major mediator to exert their immunosuppressive functions, whereas the immunosuppressive functions of proinflammatory cytokine-stimulated human MSCs are executed through IDO [7,8]. However, PGE₂ is secreted in both murine and human MSCs upon stimulation with inflammatory cytokines. PGE₂ induces macrophages to produce a higher level of IL-10 through the prostaglandin EP2 and EP4 receptors [9]. Furthermore, PGE₂ shows a strong inhibitory effect on monocyte-derived dendritic cells (DC) [10], natural killer (NK) cells and T cells [11,12]. Previous studies reported that transplantation of human MSCs into xenogeneic disease models, including mouse, rat, rabbit and dog, showed significant improvements, suggesting that human MSCs can regulate the immune/inflamm-
matory response in vivo with their immunomodulatory property [13]. We recently demonstrated that MSCs can suppress mononuclear cell proliferation and reduce the severity of colitis in mice by producing PGE2 via the nucleotide-binding oligomerization domain 2 (NOD2)-receptor-interacting serine/threonine-protein kinase 2 (RIP2) pathway [14]. Cyclooxygenase (COX) enzyme plays important roles in the biosynthesis of prostaglandins from arachidonic acid. There are two COX isoforms: COX-1 is constitutively expressed in a wide range of tissues and COX-2 is an inducible enzyme that produces PGE2 during inflammation [15]. p38 mitogen-activated protein kinase (MAPK) is preferentially activated by inflammatory stimuli and post-transcriptionally regulates COX-2 mRNA expression [16]. Treatment of SB203580, a specific inhibitor of p38 MAPK that acts by competing with ATP for the nucleotide binding site of p38, caused a rapid disappearance of COX-2 mRNA, suggesting that p38 MAPK is involved in the transcription and stabilization of COX-2 mRNA [17].

It is important to isolate and expand MSCs in vitro for therapeutic use. Unlike pluripotent stem cells, such as embryonic stem cells, MSCs undergo replicative senescence in vitro after 20–40 rounds of cell division, which is characterized by cell enlargement, changes in morphology, DNA damage response and growth arrest [18,19]. We and other groups have recently reported the molecular mechanisms are controlled by the hMSC aging process. During the progression of MSC senescence, the activity of histone deacetylases (HDACs), which regulates polycomb group genes (PcGs) and jumonji domain-containing 3 (JMJD3), is down-regulated [20]. ZMPSTE24, which is involved in the post-translational maturation of lamin A, is decreased during MSC senescence, leading to the accumulation of prelamin A in the nuclear envelope [21]. MSC properties, including multilineage differentiation, proliferation, homing and wound healing, gradually become compromised as MSCs undergo senescence [22]. However, the changes in immunomodulatory properties during MSC aging in the context of COX-2/PGE2 expression have not yet been elucidated. An understanding of the changes in the immunomodulatory properties of MSCs during the aging process is required for their use in clinical applications. In this study, we assessed the effect that consecutive MSC passages have on the immunomodulatory ability of hMSCs and the underlying mechanisms involved in these effects.

Results

Characterization of hMSCs

To determine the differentiation capacity of hMSCs, we induced the in vitro differentiation of the adipogenic, osteogenic and chondrogenic lineages. Under adipogenic induction, accumulated lipid droplets were visualized using Oil Red O staining after 2 weeks. Mineral deposition was visualized with Alizarin Red S staining after 2 weeks of osteogenic induction. Chondrogenic differentiation was confirmed by Toluidine Blue staining after 3 weeks (Figure 1A–C). The cell surface marker expression of hMSCs was examined by flow cytometric analysis. The hMSCs were positive for stromal markers (CD29, CD73, and CD105) but negative for hematopoietic cell and leukocyte markers (CD34, CD45, and HLA-DR) (Figure 1D).

Replicatively senescent hMSCs show senescence phenotypes with a compromised immunosuppressive ability

hMSCs must be expanded in vitro for maximum efficiency in clinical and pre-clinical applications. To determine the effects of consecutive hMSC cell divisions, we investigated the senescence phenotype and immunomodulatory properties in early-passage (5–10 passages, 10–20 population doublings) and late-passage (15–20 passages, 25–30 population doublings) hMSCs. Late-passage hMSCs exhibited a flattened shape and cell enlargement morphology as well as increased SA-β-gal activity compared to early-passage hMSCs (Figure 2A). Furthermore, late-passage hMSCs exhibited a significantly decreased proliferation rate (Figure 2B). To determine whether the immune modulatory functions of hMSCs were related to the physiological aging of the cells, we co-cultured human umbilical cord blood-derived mononuclear cells (hUCB-MNCs) with early- and late-passage hMSCs. Under cell-to-cell contact, the inhibitory effect of hMSCs on the proliferation of mitogen (Concanavalin A)-induced hUCB-MNCs was investigated. Remarkably, hUCB-MNC proliferation was suppressed more in the presence of early-passage hMSCs than late-passage hMSCs (Figure 2C, D and S1). Consistently, the decline in immunosuppressive function of late-passage hMSCs was observed when CD3/28 and IL-2 were used instead of ConA to induce the activation and proliferation of T cells specifically (Figure 2E, F). These data suggest that a decrease in the immunosuppressive properties of hMSCs correlates with the senescence induced by prolonged in vitro cell culture.

Early-passage hMSCs show enhanced protective effects against DSS-induced colitis in mice

To support the observation that the decrease in the hMSC immunosuppressive properties correlates with replicative senescence in vitro, we subsequently investigated the potential therapeutic efficacy of early- and late-hMSCs in the experimental model of acute colitis that was induced by oral dextran sulfate sodium (DSS) administration. It has been reported that DSS causes intestinal inflammation due to the exposure of the submucosa to external antigens (intestinal bacteria and food), leading to the recruitment or activation of inflammatory cells associated with innate immunity [23]. Mice were randomly assigned with similar body weight to investigate the therapeutic effect of early- and late-passage hMSC in intestinal inflammation. Oral administration of a 3% DSS solution induced acute colitis, characterized by clinical symptoms (diarrhea and bloody stool) with sustained weight loss resulting in 50% mortality (9/18). Intrapерitoneal injection of early-passage hMSCs reduced the loss of body-weight and decreased the mortality of mice compared to PBS- or late-passage hMSC injections (Figure 3A and B). Strikingly, the transplantation of early-passage hMSCs rescued 100% (n = 15) of the mice from colitis-induced lethality (Figure 3B). Treatment of late-passage hMSCs, however, did not exert these beneficial effects. On day 7, the disease activity index was significantly decreased by treatment with early-passage MSCs. In contrast, the administration of late-passage hMSCs did not show beneficial effects on the disease activity index (Figure 3C). The disease activity index was measured according to standards for the quantification of symptoms of patients with Crohn’s disease [24]. On day 10, the mice were sacrificed, and the entire colon from the caecum to the anus was acquired. The length, mass weight and histopathology of the colon were investigated to determine the inflammation status. The colon length decreased in mice treated with PBS or late-passage hMSCs. However, the colon length was restored by an injection of early-passage hMSCs (Figure 3D). The increased mass-to-length ratio revealed that both hyperemia and inflammation were exhibited in the colons of colitis-induced mice. Early-passage hMSCs reduced the mass-to-length ratio significantly compared to late-passage hMSCs (Figure 3E). Colon samples were processed and stained with H&E for histopatholog-
Figure 1. Characterization of hMSCs. (A–C) Images of differentiated hMSCs after induction into specific tissues. (A) The lipid droplet accumulation in differentiated cells was visualized using Oil Red O staining after 2 weeks of adipogenic induction. (B) Calcium deposition was stained with Alizarin Red S after 2 weeks of osteogenic induction. (C) Glycosaminoglycans in cell pellets were revealed by Toluidine blue staining after 2 weeks of chondrogenic induction. (D) hMSCs (1 x 10^6 cells/ml) were stained with FITC- or PE-conjugated antibodies specific for human CD29, CD34, CD45, CD73, CD105, and HLA-DR.

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Figure 2. Replicative senescent hMSCs show impaired immunosuppressive ability. (A) Phase contrast images of early- and late-passage hMSCs. β-gal staining was performed in the early-passage and late-passage hMSCs to confirm their senescence. (B) The cumulative population doubling levels of hMSCs were calculated after each repeated subculture to evaluate the loss of their proliferation potential, n = 3. (**P < 0.01). (C, D) hMNCs were treated with Concanavalin A to stimulate proliferation and colony formation. Stimulated hMNCs were co-cultured with early- or late-passage hMSCs at a 1:10 ratio (MSC:MNC). White arrows indicate colonies of MNCs (C). hMNC proliferation was assessed after 72 hours using a BrdU kit (D). The results show four representative results from experiments using three different hMSC lines. Scale bar = 100 μm. n = 3. (**P < 0.01). (E, F) Human primary T cell proliferation was induced by treatment of anti-CD3 monoclonal antibody, anti-CD28 monoclonal antibody and IL-2 to hMNCs. Activated human T cells were co-cultured with early- or late-passage hMSCs at a 1:10 ratio (MSC:MNC) (E). T cell proliferation was measured by BrdU assay after 72 hours (F). The graph shows two representative results from experiments using three different hMSC lines. Scale bar = 100 μm. n = 3. (**P < 0.01).

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ical evaluation. Histologic examination showed the destruction of the entire epithelium and submucosal edema and the infiltration of inflammatory cells into the lamina propria and submucosa in the colon of DSS-treated mice (Figure 3F). Importantly, the administration of early-passage hMSCs greatly reduced histologic damage and the histologic score in the colon. In contrast, late-passage hMSCs did not prevent histologic damage or decrease the histologic score (Figure 3F and G). These in vivo data suggest that replicative senescence significantly impairs the ability of hMSCs to deactivate the colonic inflammatory response.

Alteration of immunomodulatory activities during hMSC aging is dependent on PGE2 production

NO, PGE2, IDO and TGF-β1 have previously been identified as relevant mediators in the regulation of the immunomodulatory properties of MSCs [25]. To determine whether compromised immunosuppressive functions in hMSC aging are affected by these mediators, we measured the expression levels of each molecule from the culture media of early- and late-passage hMSCs. Because previous reports have shown evidence that the immunosuppressive functions of MSCs are elicited by proinflammatory cytokines [6],

![Figure 3. Administration of late-passage hMSCs reduces the protective effects against DSS-induced colitis in mice.](image-url)
early- and late-passage hMSCs were primed with IFN-γ and TNF-α for 24 hours. There was no significant difference in the NO and TGF-β1 levels in the presence or absence of proinflammatory cytokines in both early- and late-passage hMSCs (Figure S2A and B). However, the PGE2 concentration was dramatically increased in the presence of proinflammatory cytokines. Interestingly, the PGE2 concentration in early-passage hMSCs was higher than in late-passage hMSCs (Figure 4A). We next investigated the expression levels of immunomodulatory molecules after proinflammatory cytokine activation using Western blot analysis. Consistent with the results from an ELISA assay, only the expression of COX-2, a key enzyme in production of PGE2, was significantly decreased, while the expression of p16INK4A, a senescence marker, was increased in late-passage hMSCs (Figure 4B and S2C). Furthermore, COX-2 expression was significantly down-regulated in late-passage hMSCs compared to early-passage hMSCs, whereas Fibronectin expression was not changed, which was confirmed by immunocytochemistry (Figure 4C). To identify whether the expression of PGE2 and COX-2 decreased following hMSC passages, we investigated the secretion level of PGE2 and the expression of the COX-2 protein in consecutive passages of hMSCs. The PGE2 concentration and COX-2 expression decreased gradually with the increasing hMSC passage number (Figure 4D and E). To determine the effect of COX-2/PGE2 on hMSC immunosuppressive functions, we observed the inhibitory effect of hMSCs on hMNC proliferation in the presence or absence of celecoxib, a selective COX-2 inhibitor. Celecoxib-treated, early-passage hMSCs exhibited significantly compromised immunosuppressive abilities (Figure 4F). Taken together, these results indicate that the immunosuppressive properties of hMSCs gradually decrease upon consecutive passages via down-regulation of COX-2/PGE2 expression.

p38 MAP kinase is responsible for the reduced expression of COX-2

p38 MAPK is one of the major intracellular signaling pathways activated by inflammatory stimuli, such as IFN-γ and TNF-α. It has been reported that p38 MAPK plays a significant role in activating the immune response by regulating PGE2/COX-2 synthesis [17,26]. To determine whether the p38 MAPK pathway is activated upon hMSC aging, we examined the phosphorylation level of p38 MAPK. Phosphorylation of p38 MAPK was determined by Western blot analysis after the treatment of hMSCs with IFN-γ and TNF-α. Early-passage hMSCs showed a higher level and duration of p38 MAPK activation without affecting the expression of IFN-γ and TNF-α receptors (Figure 5A and S3A). Our previous studies have demonstrated that COX-2/PGE2 expression is associated with the NOD2-RIP2 pathway, so we investigated the expression levels of NOD1 and NOD2 in early- and late-passage hMSCs. NOD1 and NOD2 expressions were not changed in early- and late-passage hMSCs (Figure S3B and C). Phosphorylation of p38 MAPK was also investigated immunocytochemically after exposure to IFN-γ and TNF-α and was found to decrease in late-passage hMSCs compared to early-passage hMSCs (Figure 5B and C). To confirm whether COX-2 expression was dependent on the phosphorylation of p38 MAPK, we investigated the expression of phosphorylated p38 MAPK and COX-2 after the treatment of hMSCs with SB203580, a p38 specific inhibitor. hMSCs were pretreated with SB203580 for 1 hour and were then stimulated with IFN-γ and TNF-α for 30 minutes to confirm the phosphorylation of p38 MAPK and for 24 hours for COX-2 expression. The expression of phosphorylated p38 MAPK and COX-2 was significantly decreased by treatment with SB203580 (Figure 5D and E). These results suggested that the impaired activation of p38 MAPK in late-passage hMSCs might lead to the down-regulation of COX-2/PGE2 expression.

Discussion

The results presented here are the demonstration that replicative senescence in hMSCs impairs the secretion of soluble factors in response to the inflammatory milieu, thereby resulting in the loss of the immunoregulatory functions of hMSCs. Several studies have reported that cellular senescence affects the proliferation, multilineage differentiation and soluble factor production of MSCs [27–30]. However, none of these studies investigated whether cellular senescence affects their immunomodulatory ability. In this study, serial passaging by long-term culture significantly decreased the inhibitory effect of hMSCs on the mitogen-induced proliferation of hMNCs. This finding led us to confirm in vivo anti-inflammatory effect of hMSCs using a chemically induced colitis model in which we previously verified the therapeutic effect of hMSCs [14]. Consistent with in vitro coculture experiments, the protective effect of early-passage hMSCs against DSS-induced colitis in mice was completely abrogated when late-passage hMSCs were administered. Recently, Scruggs et al. [31] reported that human adipose-derived MSCs from old donors failed to ameliorate mouse experimental autoimmune encephalomyelitis, a model for human multiple sclerosis. Although this result correlates with our study in the respect that donor age or long-term culture affected the immunoregulatory ability of hMSCs, a further study may be required to elucidate the precise senescence-related characteristics of old donor-derived MSCs and long-term cultured, late-passage MSCs.

A previous study by Liu et al. [32] showed that the immunosuppressive properties of MSCs decreased with increasing passage. The main finding of this study was that although mouse MSCs sustained the secretion of TGF-β during long-term cultivation, the immunosuppressive ability of MSCs was diminished as their immunogenicity concomitantly increased. Similarly, in our study, the basal secretion level of soluble factors from hMSCs, including NO, TGF-β1 and PGE2, was not affected by cellular senescence. However, when hMSCs were treated with IFN-γ and TNF-α, prominent cytokines in type 1 helper T cell-mediated autoimmune diseases, such as inflammatory bowel disease, multiple sclerosis and type I diabetes, early-passage hMSCs produced a much higher level of PGE2 in response to cytokine treatment compared to late-passage hMSCs. In addition, the elevation in PGE2 production and COX-2 expression of hMSCs by cytokine treatment gradually diminished as passage increased. Furthermore, we showed that celecoxib-induced COX-2 inhibition in early-passage hMSCs diminished the suppressive effect of hMSCs on hMNC proliferation to the same extent observed in late-passage hMSCs. These results imply that PGE2 is a critical soluble factor through which hMSCs exert their suppressive effect on MNC and that the responsiveness of hMSCs against inflammatory cytokines might be the main reason for the loss of immunomodulatory functions in late-passage hMSCs, regardless of their immunogenicity. Liu et al. [32] also demonstrated that TGF-β from MSCs and IL-10 from MNCs were pivotal regulatory factors for an in vitro immune-inhibitory effect but not for an in vivo effect, indicating that factors other than TGF-β or IL-10 are involved in the physiological function of MSCs. Our previous study showed that hMSCs alleviated the DSS-induced colitis model through the production of PGE2 and that COX-2 inhibition led to the loss of this therapeutic effect [14].
Figure 4. Declined immune-inhibitory effect of late-passage hMSCs is regulated by PGE2 and COX-2. (A–E) Early- and late-passage hMSCs were treated with or without IFN-γ and TNF-α for 24 hours. (A, D) The PGE2 concentration was measured from the culture supernatant by an ELISA after IFN-γ and TNF-α treatment for 24 hours. n = 3. (**P<0.01) (B, C, E) COX-2 expression was investigated using Western blot and immunocytochemistry after exposure to IFN-γ and TNF-α for 24 hours. GAPDH was used for normalization. n = 3. (**P<0.01) (F) Concanavalin A stimulated hMNCs were cultured alone or co-cultured with early-, late- passage and celecoxib-treated hMSCs at a 1:10 ratio (MSC:MNC). The proliferation of hMNCs was measured by a BrdU assay after 72 hours. n = 3. (**P<0.01). doi:10.1371/journal.pone.0102426.g004
Based on these findings, one can envision that hMSC-derived PGE2 may be the crucial factor for the regulation of both cellular and physiological inflammation and that impaired PGE2 production in senescent hMSCs upon an inflammatory trigger may result in the loss of their therapeutic effect against inflammatory diseases.

A recent study by Gu et al. [33] showed that human bone marrow-derived MSCs from systemic lupus erythematosus (SLE) patients were senescent and that the expression of p16INK4A, a major molecule that induces premature senescence, was significantly increased. Additionally, the inhibition of p16INK4A expression reversed the senescent characteristics of MSCs via the activation of the extracellular signal regulated kinase (ERK) pathway, resulting in the up-regulation of TGF-β production and regulatory T cell induction. Therefore, in this study, we explored the key signaling pathway responsible for the senescence-mediated loss of responsiveness to inflammatory stimuli. Interestingly, phosphorylation of p38 MAPK upon stimulation with IFN-γ and TNF-α was decreased in late-passage hMSCs, whereas ERK signaling upon cytokine stimulation was not affected by cellular senescence (data not shown). It has been reported that NOD2 stimulation induces ERK and p38 activation mediated by RIP2 [34]. We also recently showed that NOD2 regulates the functions of hMSCs [35] and is required to suppress mononuclear cell proliferation by inducing production of PGE2 [14]. However, we found that expression of the NOD2 gene was not significantly altered by serial in vitro passages, suggesting NOD2 signaling pathway is not implicated in decline of immunomodulatory property in late-passage hMSCs. Therefore, it is apparent that future work will require to elucidate the key signaling pathway responsible for the senescence-mediated decline of immunomodulatory property in late-passage hMSCs.

Figure 5. p38 MAP kinase is responsible for senescence-associated COX-2 expression. (A) Phosphorylation of p38 MAPK was investigated by Western blot analysis after treatment with IFN-γ and TNF-α, as indicated. (B, C) Expression of the phosphorylated form of p38 MAPK in early- and late-passage hMSCs was determined by immunocytochemistry after treatment with IFN-γ and TNF-α for 30 minutes. (D, E) After treating cells with SB203580, the expression levels of COX-2 and p-p38 were confirmed by Western blot analysis. Cells were treated with IFN-γ and TNF-α for 30 minutes to detect the expression of p-p38 and for 24 hours to detect COX-2 expression, n = 3. (*P<0.05).

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ulatory property to restore the therapeutic function of immuno-
senescent hMSCs.

We conclude that consecutive cell divisions for the ex vivo
expansion of hMSCs may alter their immunomodulatory prop-
erties. We show that PGE2 and COX-2, a key enzyme of PGE2
production, play positive roles in the immunomodulatory abilities
of hMSCs. In addition, the immunomodulatory ability of hMSCs
gradually declines with successive passages through a p38 MAP
kinase-dependent change of the PGE2 and COX-2 levels. This
study provides new insights into the mechanisms that regulate
hMSC aging and may have implications for prolonging the effect
of hMSCs in therapeutic applications.

Materials and Methods

Isolation and culture of hMSCs

Umbilical cord blood samples were obtained from the umbilical
vein with the written informed consent of the mother and the
approval of the Boramae Hospital Institutional Review Board
(IRB) and the Seoul National University IRB (IRB no. 1109/001-
006). The UCB samples were mixed with the HetaSep solution
(StemCell Technologies, Vancouver, Canada) at a ratio of 5:1 and
were incubated at room temperature to deplete the erythrocytes.
The supernatant was carefully collected, and the mononuclear
cells were isolated using Ficoll density-gradient centrifugation at
1200 g for 20 minutes. The cells were washed twice with PBS.
Cells were seeded at a density of 2 x 10^5 to 2 x 10^6 cells/cm^2 on
plates in growth medium consisting of D-medium (formula no. 78-
5470EF, Gibco BRL, Grand Island, NY) containing EGM-2
SingleQuot and 10% fetal bovine serum (Gibco BRL). After 3
weeks, the cells were subcultured upon reaching 80–90% confluency.
The adherent cells were removed. For long-term culture, the cells were
subcultured upon reaching ~80-90% confluency.

Flow cytometric analysis

hMSCs were tritured into single cells and labeled with
monocolonal mouse anti-human fluorescein-conjugated antibodi-
esties; CD29-PE, CD34-FITC, CD45-FITC, CD105-FITC, CD73-
PE, and HLA-DR (BD Bioscience, San Jose, CA). The labeled

cells were analyzed by flow cytometry using a FACS calibur (BD
Biosciences).

In vitro differentiation assay

The in vitro differentiation assay was performed as previously
described [36]. For differentiation into osteoblasts and adipocytes,
1 x 10^5 cells were plated in 6-well plates. After the cells reached
70–80% confluency, they were treated with an osteogenic differen-
tation induction medium (DMEM containing 10% FBS, 100 nM
dexamethasone, 50 μM ascorbic acid 2-phosphate, and 10 mM β-
glycerophosphate) or an adipogenic differentiation induction
medium (DMEM supplemented with 10% FBS and 200 μM
indomethacin, 1 μM dexamethasone, 0.5 mM isobutyl methyl-
exanthine, and 0.5 μg/ml insulin) (all materials from Sigma-
Aldrich, St. Louis, MO). The medium was changed every 3 days.
After 2 weeks of induction, the cells were stained to confirm
osteogenic or adipogenic differentiation. To confirm osteogenic
differentiation, Alizarin Red S staining, which is specific for
calcium, was performed to detect alkaline phosphate activity.
Briefly, cells were rinsed with PBS and fixed with 70% ice-cold
ethanol for 1 hour at 4°C. After 3 washes with distilled water, the
cells were stained using 40 mM Alizarin Red S (Sigma-Aldrich) for
10 minutes at room temperature. To confirm the adipogenic
differentiation, Oil Red O staining was conducted to detect fat
droplets in differentiated cells. Briefly, cells were fixed with 10%
formalin for 1 hour and were rinsed with 60% isopropanol before
incubation in fresh diluted Oil Red O for 10 minutes. For
chondrogenic differentiation, 5 x 10^5 cells were placed in a 15-mL
polypropylene tube and maintained with 1 mL of chondrocyte
differentiation medium (Lonza, Walkersville, MD) for 3 weeks. The
round pellets were embedded in paraffin and cut into 3-μm
sections. The sections were stained with toluidine blue to detect
glycosaminoglycans.

Isolation and culture of hUCB-derived MNCs

Human mononuclear cells were isolated from UCB samples.
The UCB samples were mixed with the HetaSep solution
(StemCell Technologies) at a ratio of 5:1 and then were incu-
batated at room temperature to deplete erythrocyte counts. The super-
nant was carefully collected and mononuclear cells were obtained
using Ficoll density-gradient centrifugation at 2500 rpm for 20
minutes. The cells were washed twice in PBS and seeded in growth
media consisted of RPMI 1640 (Gibco BRL) containing 10% fetal
bovine serum.

Senescence-associated betagalactosidase (SA β-gal)
staining

SA β-gal staining was performed as previously described [21].
The hMSCs were seeded on 6-well plates at a density of 1 x 10^5
cells/well for late-passage cells and 5 x 10^4 cells/well for early-
passage cells. The cells were incubated for 3 days until they
reached the appropriate confluency. The cells were then washed
twice with PBS and fixed with 0.5% glutaraldehyde in PBS
(pH 7.2) for 5 minutes at room temperature. The cells were then
washed with PBS containing MgCl2 (pH 7.2, 1 mM MgCl2) and
stained with X-gal solution [1 mg/ml X-gal, 0.12 mM
K3Fe(CN)6, 1 mM MgCl2 in PBS, pH 6.0] overnight at 37°C.
The cells were washed twice with PBS and the images were
captured using a microscope (IX70, Olympus, Japan).

Cumulative population doubling level

hMSCs were maintained in a medium containing 10% FBS and
were subcultured every 3 day. The proliferation potential of early-
passage and late-passage hMSCs was determined by calculating
the cumulative population doubling level in continual subculture
and growth from a known number of cells. At each subculture,
the CPDL was calculated from the cell count using the equation:
ln(Nf/Ni)/ln2, where Nf and Ni are the initial and final cell count
numbers, respectively, and ln is the natural log. A total number of
5 x 10^4 cells were initially plated in a 6-well culture plate (Nunc,
Rochester, NY) and were counted at each subculture. Calculated
CPDL rates were added serially and represented as a broken line
graph.

Proliferation assay

hMSCs were treated with 25 μg/ml of mitomycin C (Mitomy-
cin C from Streptomyces caepitosus) at 37°C for 1 hour. After
washing twice in PBS, the cells were seeded in 96-well plates at
1 x 10^5/well and were incubated for 24 hours. hMNCs were
prepared as previously described, treated with Concanavalin A
(Con A from Canavalia ensiformis) in RPMI medium and added
cultured at 1 x 10^5 cells/well. After 3 days of co-culture, MNC
proliferation was determined using a cell proliferation ELISA kit,
BrdU kit (Roche, Indianapolis, IN). For the activation of T cells,
CD3/CD28 beads (Invitrogen Life Technologies, Carlsbad, CA)
were added according to manufacturer’s instruction followed by
additional treatment of recombinant human interleukin-2 (30 U/
ml, R&D systems, Minneapolis, MN). Changes of absorbance
(optical density, OD), as the level of BrdU incorporation, were measured spectrophotometrically at 450 nm in a microplate reader.

Mice and Ethical Statement
C57BL/6j mice (male; aged 8–10 weeks; 18–25 g) were obtained from Jackson Laboratory (Bar Harbor, ME). Mice were group-housed in the animal facility of Seoul National University. All experiments were conducted at the Seoul National University. All of the experiments were approved by and followed the regulations of the Institute of Laboratory Animals Resources (SNU-131209-1 Seoul National University). The experimental protocol was approved by the Seoul National University Institutional Animal Care and Use Committee (SNU IACUC). Institutional guidelines for animal care and use were followed throughout the experiments. Mice were monitored until they had reached criteria for humane endpoints. Mice losing above 30% of their original body weight or showing signs of distress (unphysiological bodily posture, shaggy fur, and breathlessness/panting etc) were sacrificed using CO₂ asphyxiation. At the designated time points for tissue harvesting, mice were sacrificed by CO₂-asphyxiation. Every effort was made to minimize suffering.

Colitis induction
This study was performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. Colitis was induced by administration of dextran sulfate sodium (DSS, MP Biochemicals, Solon, OH) in drinking water as previously reported [37]. Briefly, colitis was induced in mice from specific pathogen-free (SPF) facilities by the administration of 3% (w/v) DSS in drinking water for 7 days followed by normal drinking water until the end of the experiment. Early- and late-passage hUCB-MSCs resuspended in PBS (2×10⁶ cells in 200 μl) were injected intraperitoneally into mice 1 day after the administration of DSS. Colitis-induced mice were randomly assigned to the following groups in experiment. The mice were checked each day for morbidity (n = 12 for negative control mice, n = 12 for positive control mice, n = 12 for early-passage hMSC injected mice and n = 12 for late-passage hMSC injected mice) and body weight was recorded over 14 days at a specific time of a day (n = 6 for normal control mice, n = 18 for positive control mice, n = 15 for early-passage hMSC injected mice and n = 15 for late-passage hMSC injected mice). At day 9, colitis severity was measured by evaluating the disease activity index through the scoring of weight loss (0–4), stool consistency (0–4), bleeding (0–4), coat roughness (0–4), mouse activity (0–2), and bedding contamination by stool and blood (0–2). At the peak of the disease (day 10), the mice were sacrificed by CO₂-asphyxiation and their colon length and weight were measured (n = 7 for negative control mice, n = 15 for positive control mice, n = 16 for early-passage hMSC injected mice and n = 16 for late-passage hMSC injected mice). Histopathological evaluation was performed with the tissue acquired.

Histopathologic Evaluation
Colon samples were collected, fixed in 10% formalin, subjected to consecutive steps of alcohol-xylene changes, and embedded in paraffin. Sections that were 5-mm thick were prepared and stained with H&E. Scattered infiltration of inflammatory cells in the lamina propria and submucosa (0–4) and intestinal damage based on destruction of entire epithelium and severe submucosal edema (0–4) were graded blindly.

Cytokine production
Cells were treated with recombinant human IFN-γ and TNF-α (Peprotech. Rocky Hill, NJ) for 24 hours, and PGE₂ production was determined from the culture supernatant using a commercial ELISA kit (R&D Systems). For the NO measurement, hMSCs were treated with IFN-γ and TNF-α for 24 hours, and the concentration was measured using an ELISA kit (Promega, Madison, WI). TGF-β1 production was measured from the MSC culture supernatant after treatment of IFN-γ and TNF-α for 24 hours using a commercial ELISA kit (eBioscience, San Diego, CA).

Western blot analysis
After treatment with recombinant human IFN-γ and TNF-α (Peprotech) for 24 hours or 15 to 60 minutes, whole cell protein lysates were extracted in a solution containing 1% Triton X-100, 20 mM Tris HCl (pH 8), 137 mM NaCl, 10% glycerol and 2 mM EDTA (Sigma-Aldrich), and the protein concentrations were determined using a DC assay kit (Bio-Rad, Hercules, CA). The proteins were separated by 10% sodium dodeyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), transferred to nitrocellulose membranes at 100 V and 330 mA for 2 hours and probed with primary antibodies. The primary antibodies used to detect each protein were as follows: COX-2 (polyclonal, Abcam, Cambridge, MA, 1:1000), p-p8 (polyclonal, Cell Signaling, Danvers, MA, 1:1000), p38 (monoclonal, Abcam, 1:1000), IFNγR1 (polyclonal, Abcam, 1:1000), TNF (monoclonal, Millipore, Billerica, MA 1:1000), p16INK4A (monoclonal, Abcam, 1:1000), Glycerald ldehyde 3-Phosphate Dehydrogenase (monoclonal, Millipore, 1:3,000), and β-actin (monoclonal, Cell-signaling, 1:5000). The secondary antibodies were used according to the manufacturer’s specifications and binding was detected using an enhanced chemiluminescence (ECL) detection kit (Amersham, Piscataway, NJ) according to the manufacturer’s instructions.

Immunocytochemistry
Cells were fixed in 4% paraformaldehyde for 10 min at room temperature after treatment with recombinant human IFN-γ and TNF-α (Peprotech) for 30 min or 24 hours. The cells were permeabilized by exposure to 0.5% Triton X-100 in PBS for 10 min and were then blocked for 2 h with 10% normal goat serum (Zymed, San Francisco, CA) at room temperature. The cells were then stained with antibodies against COX-2 (polyclonal, Abcam, 1:200) or p-p8 (polyclonal, Cell Signaling, 1:200), followed by incubation for 1 hour with an Alexa-Fluor-488- or Alexa-Fluor-549-labeled secondary antibody (1:1000; Molecular Probes, Oregon). The nuclei were stained with Hoechst 33258 (1 μg/ml for 10 minutes), and the images were captured using a confocal microscope (Eclipse TE200, Nikon, Japan).

Reverse transcription polymerase chain reaction (RT-PCR) and real-time PCR
The total cellular RNA was extracted from cells using the TRIzol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. Purified RNA was transcribed into complementary DNA using Super-script III First-Strand Synthesis System (Invitrogen) followed by qualitative PCR using the SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA) with each primer. According to the manufacturer’s instructions, AB 7500 Software Version 2.0.5 (Applied Biosystems) was used to quantify gene expression. Each gene expression level was normalized with GAPDH as housekeeping controls. At least three independent analyses were carried out for each gene. The
following primers were used: for NOD1 forward, 5'-CCACTT-CACAGCTGGAGACA-3' and reverse, 5'-TGAGTAGAAG-CAGGATTGTGG-3'; for NOD2 forward, 5'-CCTGATA- TTTGGGCACTC-3' and reverse, 5'-GGTCAGACCCA-AAGGCAGG-3'; for GAPDH forward, 5'-TGATGACATTCA-GAGTTGTTG-3' and reverse, 5'-ACCCTGCTTGCTGTACG-CAAT-3'.

Statistical analysis

All of the experiments were conducted at least three times (n = 3) and the results are expressed as the mean ± SD. All of the statistical comparisons were made by one-way analysis of variance followed by a Bonferroni post hoc test for multigroup comparisons using GraphPad Prism software (version 5.01; GraphPad Software, San Diego, CA). Statistical significance designated is indicated in the Figure Legends.

Supporting Information

Figure S1 Early-passage hMSCs and late-passage hMSCs show different inhibition rates of MNC proliferation. Inhibition rates of hMSC proliferation in the presence of early- and late-passage hMSCs were calculated. Ten tests were performed. (*P<0.05).

Figure S2 Replicative senescence results in no change in NO and TGF-β1 secretion. (A) NO secretion levels were investigated in early- and late-passage hMSCs after treatment with IFN-γ and TNF-α. (B) Secretion of TGF-β1 was measured in early- and late-passage hMSC culture media after the concomitant addition of IFN-γ and TNF-α. (C) Expression of IDO, Galactin 3 and inducible Nitric Oxide Synthase (iNOS) showed no difference in early- and late-passage hMSCs. (TIF)

Figure S3 Expression of receptors shows no difference between early- and late-passage hMSCs. (A) After exposure to IFN-γ and TNF-α for 30 minutes, Western blot analysis was performed to confirm the expression levels of each cytokine receptor. (B, C) Early- and late-passage hMSCs were analyzed for mRNA expression of NOD1 (B) and NOD2 (C) using real-time RT-PCR. Error bars denote standard deviation of triplicate reactions. (TIF)

Author Contributions

Conceived and designed the experiments: KRY JYL HSK KSK KWS. Performed the experiments: KRY JYL HSK. Analyzed the data: KRY JYL HSK ISH SWC YS KSK. Contributed reagents/materials/analysis tools: KRY JYL HSK IK JJK BCL SHL AK KSK. Contributed to the writing of the manuscript: KRY JYL HSK KSK. Designed the software used in analysis: KRY JYL HSK KSK.

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