Regulation of differentiation in trabecular bone-derived mesenchymal stem cells by T cell activation and inflammation

XINGHUO WU1*, WENJUAN WANG2*, CHUNQING MENG1, SHUHUA YANG1, DEYU DUAN1, WEIHUA XU1, XIANZHE LIU1, MING TANG1 and HONG WANG1

1Department of Orthopaedics, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Hubei, Wuhan 430022; 2Department of Plastic Surgery, Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology, Hubei, Wuhan 430030, P.R. China

Received April 29, 2013; Accepted June 4, 2013

DOI: 10.3892/or.2013.2687

Abstract. Mesenchymal stem cells (MSCs) are multipotent stem cells with the ability to migrate to sites of inflammation and injury, where they participate in tissue regeneration and repair. The present study aimed to investigate the effects of T cell activation and inflammation on the differentiation of MSCs. Human trabecular bone-derived MSCs were isolated from patients undergoing total hip replacement, and T cells were isolated and purified from peripheral blood mononuclear cells (PBMCs) using CD3 MicroBeads. MSCs were co-cultured with activated T cells to mimic the inflammatory microenvironment. MTS assay was used to detect cell proliferation. qRT-PCR, western blotting, histology and immunohistochemical staining were used to detect the adipo-/osteo-specific gene expression and the relative signaling pathway. The MTS results showed that higher concentrations of T cells significantly increased the proliferation of MSCs. Expression of the inflammatory gene IL-6 was upregulated, while expression of IL-10 and INFγ was downregulated in MSCs exposed to activated T cells. The results also showed that PHA-activated T cells significantly upregulated the expression of PPARγ and FABP4 (adipo-specific genes) in MSCs, but no difference was noted in the expression of RUNX2, osteocalcin and ALP (osteogenic-specific genes) at the protein level. T cell treatment and inflammation inhibited the protein expression of TGF-β1 and the phosphorylation of Smad3, resulting in the weakening of the TGF-β/Smad pathway and enhancing the adipogenic differentiation of MSCs. The results indicated that PHA-activated T cells and inflammation could promote adipogenesis without affecting the late stage of osteogenesis of MSCs, by increasing the expression of key adipogenic genes through TGF-β/Smad3 signaling.

Introduction

The most commonly used source of mesenchymal stem cells (MSCs) is bone marrow aspirate, but its disadvantages restrict clinical and laboratory practice, such as pain, morbidity and low cell number upon harvest. Recent studies have shown that MSCs can be obtained from trabecular bone fragments obtained during total hip/knee replacements, which can avoid the disadvantages of using bone marrow as a source. Human trabecular bone-derived cell populations are pluripotent stem cells, which have multilineage potential and can give rise not only to osteoblasts, but also to adipocytes, when subjected to appropriate treatment protocols (1). Sottile et al (2) compared the characteristics of mesenchymal cell cultures established either from trabecular bone or from bone marrow, and the results showed that both of the cell cultures actually had similar characteristics to bone marrow-derived MSCs, differentiating into osteoblasts, chondrocytes and adipocytes under appropriate differentiating conditions. Therefore, trabecular bone tissue is a good source of adult MSCs for in vitro investigation.

The fate and commitment of MSCs are regulated by microenvironmental conditions, such as injury, inflammation and tumors. For example, under conditions of chronic inflammation, MSCs may contribute to adverse manifestations, such as the accumulation of fat deposits in bone and muscles, impaired healing and fibrosis after severe injury, or altered hematopoiesis and autoimmunity (3). On the other hand, MSCs also exert anti-inflammatory effects that are important in maintaining homeostatic balance. To date, MSCs have been used as a treatment modality for several inflammation-related diseases, such as inflammatory bowel disease (IBD) (4), graft vs. host disease (5), rheumatoid arthritis (6) and multiple sclerosis (7). In addition, MSCs secrete large amounts of inflammatory cytokines which regulate the inflammatory process. Thus, these findings indicate a complex, functional interaction between MSCs and the inflammatory microenvironment.
T lymphocytes play a central role in the initiation and maintenance of inflammatory processes. Accumulation of T cells at inflammatory sites is one of the characteristic features of chronic inflammatory diseases (8). During the development of inflammation, T cells are activated by inflammatory messengers, and ‘activated’ T cells are a sign of ongoing inflammation. T cell activation depends on signals delivered from antigen-presenting cells (APCs) through triggering of their T cell receptor (TCR) complex and a co-stimulatory receptor such as CD28 (9). The T cell activation pathway is triggered when a T cell encounters its cognate antigen, such as plant mitogen phytohemagglutinin (PHA), which has a marked selectivity for T lymphocytes. Following short-term pre-incubation with PHA, T cell activation is maintained. T cells activated with PHA express Ia-like antigens (which play a role in the stimulation of T lymphocytes by autologous PHA-T lymphocytes) and acquire the ability to stimulate autologous T lymphocytes in mixed lymphocyte reaction (10).

Therefore, T cell activation using PHA stimulation is a good pathway with which to curb inflammation.

To better understand the effects of T cell activation and inflammation on the adipogenic and osteogenic differentiation of MSCs, PHA-activated T cells were used for co-culturing with MSCs in the present study. The inflammatory gene expression was detected by quantitative RT-PCR, and the adipo- and osteo-specific proteins were determined by western blotting. TGF-β/Smad signaling was described by western blotting; and ALP activity was detected by pNPP method; and the adipo-/osteo-differentiation of MSCs was further verified using histological and immunohistochemical staining.

Materials and methods

All the procedures were approved by the Ethics Committee of the Union Hospital and the Tongji Medical College. Informed consent was obtained from the patients included in the study.

Isolation and culture of MSCs. Human trabecular bone-derived MSCs were isolated from patients undergoing total hip replacement. One piece of bone was collected from the removed femoral neck under sterilization. The bone was rinsed several times (3-5 times) with PBS to remove excess blood. Subsequently, the bone was placed onto a sterile Petri dish and cancellous bone was broken into 2-3 mm² x 1 mm fragments (the cortical bone was discarded). Subsequently, the bone fragments were placed into 15-ml centrifuge tubes in collagenase solution (3 mg/ml), and the tubes were incubated on a rotator at 37°C for 3 h. After 3 h of digestion, the same volume of complete culture medium (CM) was added to neutralize the reaction, and the supernatant was filtered through a 70-µm cell strainer. Finally, we obtained MSC solution, the cell density was adjusted and cells were seeded in T-75 cell culture flasks. The flask was placed in a humidified incubator at 37°C with 5% CO₂. After a 24-h culture, the culture medium was refreshed; the culture medium was replaced every 2-3 days.

In vitro osteogenic, adipogenic and chondrogenic differentiation of MSCs. MSCs of passage 3-5 were used in the experiments. MSCs were plated in a 24-well plate in coverslips at a density of 1.5x10⁵ cells/ml. For osteogenic differentiation, after becoming confluent, the cells were incubated in an osteogenic medium (OS) containing dexamethasone, ascorbate-phosphate and β-glycerophosphate in complete medium. OS was replaced every 2-3 days for 3 weeks. The cells were fixed and subjected to Alizarin Red S staining and immunostaining (mouse anti-human osteocalcin monoclonal antibody, osteogenic marker). For adipogenic differentiation, MSCs were seeded at a density of 7.4x10⁵ cells/ml. The subconfluent cultured cells were cultured in adipogenic differentiation medium, containing hydrocortisone, isobutylmethylxanthine and indomethacin in complete medium. After 7-21 days of stimulation, the cells were fixed and then detected using Oil Red staining and immunostaining (goat anti-mouse FABP-4 polyclonal antibody, adipogenic marker). As previously described, a pellet culture system was used for chondrogenic differentiation. A total of 2.5x10⁵ cells were transferred into a 15-ml tube and pelleted by spin-down. The pellet was cultured in 0.5 ml of chondrogenic differentiation medium, containing dexamethasone, ascorbate-phosphate, proline, pyruvate, TGF-β3, insulin, transferrin and selenious acid in complete medium. The chondrogenic culture medium was changed every 2-3 days for 3 weeks. The pellets were then fixed and subjected to frozen sections. Immunostaining was used to detect the expression of aggrecan protein (goat anti-human aggrecan polyclonal antibody, chondrogenic marker).

T cell isolation and activation. Peripheral blood mononuclear cells (PBMCs) were collected from healthy donors and separated by Ficoll-Hypaque density gradient centrifugation. The T cells were then isolated and purified from PBMCs by magnetic-activated cell sorting using the CD3 MicroBead-based isolation kit according to the manufacturer's instructions (MACS; Miltenyi Biotec). To achieve higher purities, the positively selected CD3+ cell fractions were separated again over a new, freshly prepared LS column. The viability of CD3+ cell fractions was measured by trypan blue, with cells generally being >95%. For activation, CD3+ cell fractions (T cells) were resuspended in α-MEM medium containing PHA (2.5 μg/ml), 10% FCS, penicillin (100 U/ml), streptomycin (100 μg/ml), and L-glutamine (2 mM) and incubated at 37°C for 2 days in tissue culture tubes with filtered caps.

MTS assay for cell proliferation. MTS assay was used to detect the cell proliferation of MSCs after treatment with different concentrations of PHA-activated T cells. MSCs were plated into each well of 96-well plates at a density of 6x10⁵ cells/ml. At 24 h after plating for attachment, cells were incubated with different concentrations of activated T cells (6x10⁵ to 6x10⁶ cells/ml). On day 4, suspended cells were removed by gentle washing with phosphate-buffered saline (PBS). The number of adherent cells remaining in each well was then quantified using a coupled enzymatic assay, which resulted in the conversion of a tetrazolium salt into a red formazan product (MTS assay). Recording of the absorbance at 490 nm in the MTS assay was carried out.

MSC differentiation under T cell activation and inflammation. MSCs were co-cultured with activated T cells to mimic the inflammatory microenvironment. In the experiment, the cells were divided into the control group and inflammatory
group (inflammatory). In the inflammatory group, MSCs were seeded onto plates at a concentration of 1.2x10^5 cells/ml in CM, containing 10% FCS, 100 µg PSN (penicillin, streptomycin and neomycin) and 100 µM L-ascorbic acid phosphate in α-MEM. Activated T cells were then added at a concentration of 0.6x10^5 cells/ml, after the cultures became 90% confluent. In the control groups, MSCs were cultured without T cell exposure. The cultures were refreshed with complete medium (CM) every 2-3 days for the following measurements.

**RNA isolation and quantitative RT-PCR.** To investigate the effects of the inflammatory microenvironment induced by activated T cells, quantitative RT-PCR was used to detect the expression of inflammatory genes and osteo/adipo-specific genes. After a 1-day co-culture with activated T cells, MSCs were collected in TRIzol reagent (Invitrogen, Carlsbad, CA, USA), followed by RNA isolation according to the manufacturer's instructions. The RNA samples were subjected to cDNA synthesis, followed by quantitative PCR assays. The reverse transcriptase reaction was carried out using ThermoScript™ reverse transcription reagents (Roche Applied Science). PCRs were performed according to the real-time PCR machine manufacturer's instructions (MJ Research, Inc., Watertown, MA, USA), which allow real-time quantitative detection of the PCR product by measuring the increase in SYBR-Green fluorescence caused by binding of SYBR-Green (Bio-Rad Laboratories, Quebec, Canada) to double-stranded DNA. To amplify specific gene products, the following primers were used: IL-6 (forward, CCTCGAGCGGCATCTCAGCCC and reverse, TGCCCCAGTGCAAGTTCTGGA); IL-10 (forward, CAAGGCCTGGAGCAGTTGAAA and reverse, GTTTTCTAAGGGGCTTGGTCA); INFγ (forward, TCG CCAGCGACTTAAAAACAGGGA and reverse, GCTGCTA GTTGCCCTGTGA); PPAR-γ (forward, GCTGTTATGGGT AAACTCT and reverse, ATGGAATGTCTTCGTAATGT); RUNX2 (forward, GTGCTACTGGCAGTTTCTC and reverse: GCTCTTCTTACTGAGAGTGGAAGG); and 18S rRNA (forward, CCGAGCTAGAATATGCAGTA and reverse, TCTAGCCGGCGCAATACGGA), which served as an internal control. Amplification was performed using a profile at 94°C for 1 min (denaturation), 60°C for 30 sec (annealing), 72°C for 45 sec (elongation) for a total of 38 cycles, followed at the end by 72°C for 5 min (extension). Negative controls without RT were carried out in parallel for every PCR reaction to exclude amplification of contaminating DNA.

**Western blot analysis.** Under the inflammatory microenvironment induced by activated T cells, cell lysates of MSCs were collected at 0, 1, 3, 6, 12 and 18 h and stored at -20°C for western blot analysis. The protein concentrations were measured using Bio-Rad protein assay. The proteins were then electro-transferred to 0.45 µm-pore diameter polyvinylidene difluoride (PVDF) membranes (Invitrogen). PVDF membranes were blocked in binding agent (5% skim milk, 0.1% Tween-20 in basic buffer) overnight at 4°C. After blocking of non-specific immunoglobulin (IgG) binding, the membranes were incubated in primary antibodies at a 1:400 dilution for 2 h at 4°C under rocking. The membranes were then incubated for 2 h at 4°C with a 1:2,000 dilution of goat secondary antibodies in antibody diluents. Finally, the ECL-Plus western blotting system was used, and immunoreactive bands were revealed and quantified using ImageQuant LAS 400 software (GE Healthcare Life Sciences). The primary antibodies were anti-PPAR-γ, anti-RUNX2, anti-TGF-β1, anti-Smad3 and anti-P-Smad3 produced in rabbit (Sigma-Aldrich).

**Immunohistochemical staining.** After 2-3 weeks of culture, the cells were fixed and saved for immunostaining, to assess the possible adipogenic or osteogenic differentiation of MSCs. The culture medium was aspirated and the cells were washed twice PBS. The cells were fixed with 4% paraformaldehyde for 20 min at room temperature. After fixation, the cultures were washed with 1% BSA in PBS and blocked with 0.3% Triton X-100, 1% BSA, 10% normal donkey serum in PBS for 45 min. Subsequently, the primary antibody was added and incubated overnight at 4°C. After washing, the secondary antibody was added and incubation was carried out in the dark for 60 min at room temperature. The primary antibodies used in the experiment were goat anti-human FABP4 and mouse anti-human osteocalcin antibodies (R&D Systems); and the secondary antibodies were fluorochrome-conjugated donkey anti-goat and donkey anti-mouse (R&D Systems). Staining was examined using fluorescence microscopy immediately.

**Oil Red O staining and Alizarin Red staining.** To determine whether the inflammatory microenvironment induced by activated T cells affects the differentiation of MSCs, the adipogenic and osteogenic differentiation of the cells was evaluated after 2-3 weeks of culture. Briefly, the cultures were washed with PBS. 2.5% glutaraldehyde was added (0.5 ml/well) (24-well plate), and the reaction was carried out at room temperature for 20 min. For adipogenic differentiation, Oil Red O solution was added to the fixed cells; for osteogenic differentiation, 2% Alizarin Red solution was added. The plate was kept at 37°C in an incubator for 10-20 min. The staining was monitored under a microscope every 2-5 min and images were captured under the microscope immediately for analysis.

**In vitro osteogenic differentiation and pNPP assay.** For osteogenic differentiation, the cells were cultured in OS, containing dexamethasone, ascorbate-phosphate and β-glycerophosphate in CM. After 2-3 weeks of culture, the cells were subjected to pNPP alkaline phosphatase assay to detect alkaline phosphatase (ALP) activity. Assay buffer 1X was prepared and the cells were gently washed twice with the buffer. Then, 300 µl of 1X assay buffer in Triton X-100 per well (6-well plate) was added to the cells, and the adherent cells were scraped off and transferred into a microcentrifuge tube. The cell suspension was incubated at 4°C for 10 min under agitation. After centrifugation, the supernatant was collected for alkaline phosphatase assay, according to the instructions for the SensoLyte® pNPP alkaline phosphatase assay kit. Meanwhile, the concentrations of the proteins were determined using Bio-Rad protein assay. The results of ALP activity were determined using the standard curve and normalized by the total protein content. ALP activity = ALP (µg/ml)/protein (mg/ml).

**Statistical analysis.** All experiments were repeated three times, and the results are expressed as the means ± SEM. Data
Results

Characterization of MSCs isolated from human trabecular bone. One of the defining characteristics of MSCs is their multilineage differentiation potential. Under certain inductive conditions, MSCs are able to acquire the characteristics of cells derived from the embryonic mesoderm, such as osteoblasts, chondrocytes and adipocytes. To define the characteristics of the trabecular bone-derived MSCs, cell type-specific histological and immunochemical staining were used. To identify adipogenic and osteogenic differentiation in vitro, the cultures were stained with Oil Red O and Alizarin Red S, respectively. Trabecular bone-derived MSC differentiation was also verified by analyzing the gene expression of adipogenic, chondrogenic and osteogenic markers, such as osteocalcin, FABP-4 and aggrecan (Fig. 1). The results showed that the MSCs were successfully differentiated into adipocytes, osteocytes and chondrocytes.

MSC proliferation with PHA-activated T cell supplements. MTS is a common and useful method for monitoring cell proliferation, for the detection of viable cells. To quantify the proliferation of trabecular bone-derived MSCs following exposure to activated T cells of different concentrations, an MTS assay was performed (Fig. 2). The results showed that lower concentrations (ratios of 1:100 and 1:10) of T cells had no effect on the proliferation of MSCs. However, high concentrations of T cells significantly increased the proliferation of MSCs, particularly at ratios of 10:1 and 100:1 (P<0.01).

Expression of inflammatory factors induced by T-cell exposure. T cells induce the expression of genes mostly related to inflammatory cytokines, which are thought to play an important role in the process of chronic inflammation. The present study was undertaken to detect the gene expression profile of MSCs exposed to activated T cells using qRT-PCR. The results showed that the expression of the proinflammatory gene IL-6 was significantly upregulated (6.89-fold compared with the control) and acted to promote excessive inflammation. On the other hand, the expression of anti-inflammatory genes, such as IL-10 (-44.24-fold) and INFγ (-5.96-fold), was significantly downregulated when compared to the control group (Fig. 3). Therefore, T-cell exposure increased the expression of proinflammatory genes but inhibited the expression of anti-inflammatory genes and led to an excessive inflammatory microenvironment.

Adipo-/osteo-specific gene expression is influenced by activated T cells and inflammation. In order to investigate the effects of an inflammatory microenvironment induced by activated T cells on the adipogenic and osteogenic differentiation of MSCs, qRT-PCR and western blotting were performed. The
results of qRT-PCR showed that the adipo-specific gene expression of PPARγ was significantly upregulated (2.58-fold compared with the control), while the osteogenic-specific gene expression of RUNX2 was significantly downregulated (-3.63-fold) (Fig. 4A and B). Western blot analysis revealed that the expression level of PPARγ was upregulated in MSCs after inflammatory stimulation with activated T cells, reaching a peak value at 6 h, which was subsequently decreased (Fig. 4C). In contrast, MSCs expressed barely detectable levels of RUNX2 protein for the duration of the experiment; thus, we could not compare the expression levels of RUNX2 protein between the groups. However, the results still indicated that the alterations resulted in increased adipogenic differentiation and decreased osteogenic differentiation of MSCs at an early stage.

**Adipogenic differentiation of MSCs is regulated by activated T cells.** MSCs were cultured in CM supplied with activated T cells. After a 2-week culture, the cultures were fixed; half of them were subjected to Oil Red O staining, and the other half were used for the detection of the expression of FABP4 using fluorescence immunohistochemistry. The fluorescence staining showed a very weak diffuse fluorescence pattern in the control group (Fig. 5A) but strong FABP4 expression of MSCs in the inflammatory group (Fig. 5B). The results of Oil Red O staining showed accumulated intracellular lipid droplets in the inflammatory group (Fig. 5D), while no lipid droplet was found in the control group (Fig. 5C). Both types of staining indicated that an inflammatory microenvironment induced by activated T cells promoted the adipogenic differentiation of MSCs.
Osteogenic differentiation of MSCs in the presence of activated T cells. In order to investigate the effect of activated T cells and inflammation on the osteogenic differentiation of MSCs, fluorescence immunohistochemical staining and Alizarin Red staining were used. The fluorescence staining showed that MSCs expressed lower levels of osteocalcin in both the inflammatory group and the control group. Alizarin Red staining showed similar results; only mild mineralization was observed in both of the groups (Fig. 5E-H). These results confirmed the finding that the inflammatory microenvironment induced by activated T cells had no significant effect on the osteogenic differentiation of MSCs.

ALP activity as determined by pNPP method. The activity of ALP was detected to evaluate the effect of activated T cells on the osteogenic differentiation of MSCs in OS medium by pNPP method. The ALP levels expressed by MSCs increased with time in both the control and inflammatory groups in the presence of OS medium. However, at the time-points studied (week 2 and 3), no significant difference was noted in ALP activity due to the inflammatory microenvironment induced by the activated T cells, when compared to the control group (Fig. 6). Therefore, the results confirmed that the activated T cells had no obvious effects on the osteogenic differentiation of MSCs at the late stage.

TGF-β/Smad3 signaling. To evaluate the effect of T cell treatment on the TGF-β/Smad pathway, the TGF-β1 level was detected by western blotting. The expression of the TGF-β1 gene in MSCs decreased significantly after inflammatory stimulation with activated T cells at 6 h, compared to the control group. In addition, the gene expression of Smad3 and the phosphorylation levels of the MSC culture in the absence or presence of T cells were investigated by western blotting. At this time-point, the phosphorylation level of Smad3 protein declined in the presence of the T cells, when compared to control group. Therefore, T cell treatment inhibited the expression of TGF-β1, resulting in the weakening of the TGF-β/Smad3 pathway and enhancing the adipogenic differentiation of MSCs (Fig. 7).

Discussion

Mesenchymal stem cells (MSCs) have chemotactic ability and appear to migrate to sites of inflammation (11), where they appear to play an active role in tissue remodeling. MSCs are generally isolated from an aspirate of bone marrow; in addition, MSCs have been isolated from bone marrow, periosteum, trabecular bone, adipose tissue, synovium, skeletal muscle and deciduous teeth (12). Trabecular bone-derived mesenchymal stem cells (MSCs) are multipotent cells, which can differentiate into a number of different types of cells, including osteocytes, adipocytes, chondrocytes, myocytes and neurocytes (1,2). Differentiation of MSCs into different lineages of cells is strictly regulated by various instructive signals, and alteration or malfunction of this regulation results in pathological consequences, such as osteoporosis or a high bone mass phenotype (13,14). In the present study,
and immunohistochemistry methods were used to confirm the multipotential characteristics of the trabecular bone-derived cells. Oil Red O staining and anti-FABP-4 staining showed trabecular bone-derived MSCs had adipogenic differentiation potential; Alizarin Red S staining and anti-osteocalcin staining showed osteogenic differentiation capability; and chondrogenesis of MSCs was evidenced by anti-aggrecan fluorescent staining. Therefore, the results showed that MSCs derived from trabecular bone were multipotent stem cells; that is, they can give rise to diverse cell types.

Another property of MSCs is the ability to reduce the proliferation of lymphocytes of various types (15,16). For example, BM-MSCs have been reported to impair the proliferation of activated T cells (17). In contrast, activated T cells do affect the proliferation of in vitro cultured MSCs, but this depends on the concentration of the cells. In the present study, we examined the effects of various concentrations of activated T cells on the proliferation of MSCs by MST assay. The results showed that activated T cells had no effect on the proliferation of MSCs at low concentrations, but higher concentrations of activated T cells significantly increased the proliferation of MSCs, particularly at the ratios of 10:1 and 100:1 (T cells/MSCs; P<0.01). Therefore, in order to exclude effects of an increase in the proliferation of precursors on the differential outcome of MSCs; in subsequent experiments, T cells at a lower concentration were added to the cultures after MSCs reached 90% confluency.

Furthermore, one of the undisputed features of MSCs is their ability to produce a wide variety of chemokines and cytokine receptors, including those for tumor necrosis factor α (TNF-α), interleukin (IL)-10, IL-6 and interferon γ (INFγ). The cytokine IL-10, which is produced by various immune cells, in particular monocytes/macrophages and T cell subsets, has a crucial role in limiting the inflammatory response caused by activated T cells (18). MSCs are also known to produce high levels of IL-10 (19). What is more, MSCs constitutively expressed mRNA of IL-6, which acts as a predominantly pro-inflammatory cytokine. IL6 is produced by undifferentiated dividing MSCs, and the secretory pathway of MSCs is mediated through the activation of p38 MAPK (20).

Both MSCs and T cells express inflammatory cytokines, thus ELISA methods cannot be used directly to measure the expression levels of cytokines secreted by MSCs. Alternatively, the detection of mRNA expression was used. The expression of inflammatory genes was regulated in MSCs exposed to PHA-activated T cells. According to the results, IL-6 (pro-inflammatory gene) was significantly upregulated, while IL-10 and INFγ (anti-inflammatory genes) were significantly downregulated. This indicated that the inflammatory microenvironment induced by activated T cells could increase the expression of proinflammatory genes but inhibited the expression of anti-inflammatory genes, leading to an excessive inflammatory microenvironment. Actually, these inflammatory factors not only attribute to inflammatory response but also play an important role in the differentiation of MSCs. Previous studies have shown that MSCs cultured in OS differentiated along the osteogenic lineage and downregulated...
protein and mRNA levels of IL-6 (22,23). Our results are consistent with the findings of the adipogenic differentiation of MSCs following upregulated mRNA levels of IL-6.

The balance between peroxisome proliferator-activated receptor-γ (PPARγ) and Runx-related transcription factor 2 (RUNX2) leads to MSC differentiating into adipocytes or osteoblasts, respectively. PPARγ is a member of the nuclear receptor superfamily of ligand-activated transcriptional factors, which act as a key regulator of adipogenesis. PPARγ is commonly termed the master regulator of adipogenesis; no factor has yet been identified that can induce normal adipogenesis in the absence of PPARγ (24). In contrast, RUNX2 promotes osteogenesis but inhibits adipogenesis of MSCs. These master regulators of different lineages are expressed at low levels in undifferentiated cells, maintaining the differentiation potential of MSCs (25,26). In the experiment, expression of PPARγ and RUNX2 genes was detected by qRT-PCR and western blot analyses. The results showed that PHA-activated T cells significantly upregulated the expression of PPARγ (adipo-specific gene) in MSCs at both the RNA and protein levels. However, the expression of RUNX2 was altered at the RNA level but not at the protein level. qRT-PCR showed that the expression of RUNX2 was significantly downregulated in MSCs treated with activated T cells. Whether treated or not with activated T cells and inflammation, RUNX2 protein expressed by MSCs remained at or below the limits of detection for the duration of the experiment. Thus, it was difficult to compare the expression levels between the groups. According to the results, activated T cells and inflammation promoted adipogenesis in MSCs and may play a role in the early stage of MSC osteogenesis.

To carry out the specialized functions of T cell activation and inflammation, the adipogenic differentiation of MSCs was further verified using histological and immunohistochemical staining. Fatty acid binding protein 4 (FABP4), also called aP2 (adipocyte protein 2), has a high affinity for a variety of fatty acids and facilitates their storage, trafficking and solubilization (27). FABP4 has been used as a marker (specific gene) to follow the differentiation of adipocytes. The expression of FABP4 can be induced by fatty acids, likely through changes in PPARγ expression or activity, both at the transcriptional and post-transcriptional level (28,29). In accordance with its stimulatory effect on PPARγ expression, we found that PHA-activated T cells upregulated the expression of FABP4 in MSCs, as detected by fluorescence immunohistochemistry. In addition, the increase in FABP4 expression by activated T cells was consistent with the observed increase in Oil Red O staining. Therefore, these studies confirmed once again that PHA-activated T cells promoted MSC adipogenesis, by increasing the expression of key adipogenic genes.

Whether or not T cell activation and inflammation have an effect at later stages of osteogenic differentiation from MSCs was investigated. The osteogenic differentiation of MSCs was evaluated by the expression of osteocalcin, mineralization and alkaline phosphatase (ALP) activity. According to the results, it had no obvious effect on osteogenic differentiation from MSCs. Osteocalcin (bone gla-protein), secreted by osteoblasts, is generally regarded as an osteoblast-specific marker (30). The cells showed low-level expression of osteocalcin and mild mineralization in both groups, as evidenced by Alizarin Red S staining and fluorescence immunohistochemical staining. The findings were consistent with the protein expression of RUNX2. No significance was observed between the groups. Alkaline phosphatase (ALP) activity is another marker for osteoblast differentiation (31). Under osteogenic conditions (OS), pNPP assay revealed that activated T cells had no significant effect on ALP activity on the 14th and 21st day of osteogenic differentiation. Totally, these results indicated that PHA-activated T cells promote adipogenesis without affecting the osteogenesis of MSCs.

Growth factors such as transforming growth factor-β1 (TGF-β1) are known to inhibit adipocyte differentiation in vitro. MSCs can produce significant levels of transforming growth factor-β1 (TGF-β1) which can be further enhanced by anti-inflammatory cytokines (32). Moreover, Smad proteins play a key role in the regulation of TGF-β signaling and Smad3 is the central intracellular mediator of TGFβ signaling. After activation of Smads, the effectors of TGFβ signaling result in Smad translocation from the cytoplasm into the nucleus where they act as transcriptional modulators to regulate target gene expression (33). The TGF-β1/Smad3 signaling pathway was found to play a key role in adipogenesis, which inhibited adipogenesis independent from the Wnt and β-catenin pathway (34). TGF-β inhibits adipocyte differentiation. Previous results indicate that endogenous TGF-β signaling regulates the rate of adipogenesis, and that Smad3 has distinct functions in this endogenous control of differentiation (35). TGF-β targets the transcription factor cascade upstream of PPARγ, and the inhibition of adipogenesis by TGF-β was accompanied by reduced mRNA and protein levels of PPARγ. In the present study, western blot analysis confirmed that T cell treatment inhibited the protein expression of TGF-β1 and the phosphorylation of Smad3, resulting in a weakening of the TGF-β1/Smad pathway which enhanced the adipogenic differentiation of MSCs (Fig. 7).

In conclusion, we demonstrated the effects of T cell activation and inflammation on osteoblast and adipocyte differentiation of MSCs. The results showed that PHA-activated T cells upregulated the expression of adipocyte-specific genes and led to adipogenic differentiation, possibly due to gene expression of PPARγ which plays a more decisive role in the differentiation of MSCs exposed to activated T cells through TGF-β1/Smad3 signaling. This may explain why we observed no obvious effect of PHA-activated T cells on osteogenic differentiation of MSCs, but it is complicated. Additionally, there are some limitations to the current research that merit further study. First, T cells, even activated T cells, normally have a short lifespan, which may have partially influenced the experimental outcomes. Thus, in future experiments, a constant supply of T cells will be necessary to observe the effects on MSC differentiation. Second, the present study mainly focused on the downstream regulatory events, and the precise molecular mechanisms and signaling pathways are still not clear. Thus, further study is needed to investigate the upstream regulatory mechanisms, particularly using knockdown techniques to elucidate the mechanism involved in the differential gene expression. In spite of these limitations of the present study, further investigation of the individual effects of activated T cells and inflammation on MSC differentiation in vivo is warranted.
Acknowledgements

The authors appreciate the help of the Osteonecrosis Research Team of Union Hospital, Tongji Medical College, Wuhan, China. The present study was supported by the National Natural Science Foundation of China (no. 81201393).

References