Co-culture with lung cancer A549 cells promotes the proliferation and migration of mesenchymal stem cells derived from bone marrow

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Received April 11, 2016; Accepted March 17, 2017

DOI: 10.3892/etm.2017.4909

Abstract. The initiation and progression of various types of tumors, such as lung neoplasms, are driven by a population of cells with stem cell properties and their microenvironment. Bone marrow mesenchymal stem cells (BM-MSCs) in long-term in vitro culture may exhibit spontaneous changes in stem cell biological properties, including malignant transformations; however, the molecular mechanisms responsible for these have not been fully elucidated. In the present study, a BM-MSC and lung cancer A549 cell co-culture system was utilized to investigate how the tumor microenvironment may spontaneously change the proliferation, migration and differentiation of BM-MSCs. It was demonstrated that the lung cancer A549 microenvironment is able to induce changes in the cell morphology, proliferation, karyotype, cytoskeleton and migration ability of BM-MSCs in vitro. Compared with the control group BM-MSCs, the expression of Ras, phosphorylated-extra-cellular regulated protein kinases, nuclear factor-xB, P62 and B-cell lymphoma 2 (Bcl-2) proteins in groups of co-cultured BM-MSCs increased significantly (P<0.05) and the expression of P53, Bcl-2 associated X protein and caspase-3 protein decreased significantly (P<0.05). The mechanisms responsible for the changes observed in BM-MSCs may be related to abnormal expression of related genes in the ERK signaling pathway.

Introduction

Malignant tumors are a serious threat to human health and emphasis has been placed on developing novel therapies to treat such malignancies. Tumor biological therapy is the application of modern biological technology and products targeted at tumors to effectively ameliorate the side effects of chemotherapy (1). Bone marrow mesenchymal stem cells (BM-MSCs) cultured in vitro can be induced to proliferate, readily undergo transfection with liposomes carrying exogenous genes, have excellent mobility, are capable of undergoing multilineage differentiation (2) and have low immunogenicity (3). These properties make BM-MSCs effective carrier cells for biological treatments of tumors. Using stem cells as carriers to target drug delivery to malignant tumors alone may reduce the adverse reactions caused by systemic drug distribution (4). Furthermore, using genetically modified BM-MSCs as tumor target gene therapy vectors may enhance anti-tumor effects, providing a novel method for tumor therapy (5,6).

The stem cell niche is the microenvironment in which stem cells exist. The stem cell niche allows interaction between stem cells to regulate their function and fate, and it is a critical factor in stem cell homeostasis. The stem cell niche is able to tightly regulate stem cell self-renewal and proliferation by signal molecules (7). It has been reported that BM-MSCs undergoing long-term in vitro culture may undergo spontaneous changes in terms of their biological characteristics, and may even undergo malignant transformation (8-10). These results suggest that alterations to the cell microenvironment may affect the differentiation and proliferation of stem cells; however, the molecular mechanisms responsible for these

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Key words: bone marrow mesenchymal stem cells, lung cancer A549, proliferation, migration, co-culture
alterations have not been fully elucidated. It has not yet been reported whether changes to BM-MSC biological characteristics in the lung microenvironment are caused by cytokines, signaling molecules or cellular interactions.

To identify the risk of BM-MSCs undergoing malignant transformation when being used for biological therapies in the tumor microenvironment, the present study utilized a Transwell chamber to co-culture BM-MSCs and lung cancer A549 cells to simulate a tumor microenvironment. From this, it was possible to investigate whether BM-MSCs are able to spontaneously undergo changes in proliferation, migration and differentiation in the tumor microenvironment and whether it was possible to maintain BM-MSC genetic stability in these specific culture conditions. The results of the current study may provide an experimental basis for the clinical application of stem cell therapy.

Materials and methods

Cells and cell culture. BM-MSCs (Cyagen Biosciences, Inc., Santa Clara, CA, USA) and human lung cancer A549 cells (stored in the Provincial-Level Key Laboratory for Molecular Medicine of Major Diseases and The Prevention and Treatment with Traditional Chinese Medicine Research in Gansu Colleges and Universities, Lanzhou, China) were cultured in complete medium, consisting of Dulbecco's modified Eagle's medium/F12 supplemented with 10% fetal bovine serum (HyClone; GE Healthcare Life Sciences, Logan, UT, USA). The culture medium was replenished every 2-3 days. Cell aggregates were typically formed after 24 h incubation in a humidified chamber at 37°C (5% CO₂). Cell aggregates were grown in suspension for 3-5 days before they began to attach to the bottom of the culture bottle. When the cells covered 80-90% of the bottom of the bottle, they were digested with 0.25% trypsin to perform a co-culture experiment.

Establishment of co-culture system. A non-contact co-culture system of BM-MSCs and lung cancer A549 cells was established using a Transwell suspension culture chamber with polyethylene terephthalate film combined with a 6-pore plate (Corning 3450; Corning, Inc., Corning, NY, USA). The BM-MSC and A549 groups were groups in which BM-MSC cells and A549 cells were cultured respectively, in independent wells of a 6-well plate. The co-BM-MSC group, including BM-MSCs and A549 cells, co-cultured in the transwell system (BM-MSCs in the upper chamber and A549 cells in the lower chamber). The number of cells seeded per chamber for each group is 5x10⁶ cells. Cells were cultured in 6-well plates (Corning 3450) containing the aforementioned complete medium at 37°C (5% CO₂ incubator). Culture medium was replenished every 48 h and cell growth state was observed under an inverted microscope. On day 7 of culture, cell culture was terminated and three repeats per group and three repeats per replication. Briefly, cells were washed twice with 0.01 mol/l PBS and centrifuged for 3 mins at 167.7 x g at room temperature, then incubated with 70% precooled ethanol at 4°C overnight. The cells were then subjected to 50 µg/ml propidium iodide dye solution (containing 50 µg/ml RNase) and analyzed by EXPO™ 32 ADC version 1.1C software (Beckman Coulter, Inc.). Cell viability was analyzed using the 3-[4,5-dimethylthiazol-2-yl]-2,5 diphenyltetrazolium bromide (MTT) assay for three wells in each group every day, continuously for 9 days. The BM-MSCs group and A549 group served as the controls. The test was performed with 5 mg/ml MTT solution prepared by dissolving MTT powder (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) in warm PBS (37°C). Cells (~1x10⁶ cells/well) were incubated in 96-microwell plates (Nalge Nunc International, Penfield, NY, USA) at 37°C for 24 h. Following incubation, 20 µl MTT solution was added to each well and incubated for 4 h in a 5% CO₂ incubator at 37°C. Subsequently, the solution was removed and the construct was blotted with filter paper. Finally, 200 µl 99.5% dimethyl sulfoxide (Sigma-Aldrich; Merck KGaA) was added to each well and the plate was shaken for 5 min. For colorimetric readings, 100 µl of this solution was transferred to an ELISA plate and the absorbance values of the samples detected were read by an ELISA reader at 570 nm. Using cultivated time as the horizontal axis and the absorbance value as the vertical axis, a growth curve was constructed.

Western blot analysis. Using western blotting, the expression of proteins associated with the extracellular signal-related kinase (ERK) signaling pathway, cell proliferation and apoptosis were analyzed. Total protein was extracted from cells using Radio-Immunoprecipitation Assay cell lysis buffer (Qiagen GmbH, Hilden, Germany). Cells were washed with precooled PBS at 4°C, then scraped using a disposable cell scraper. RIPA cell lysis buffer (cat. no. R0101; Solarbio Science & Technology Co., Ltd., Beijing, China) was added to cells and the solution was mixed for 1 min. Cells were collected in 1.5 ml centrifuge tubes. All the aforementioned actions were completed on ice. Cells were stood for 10 min on ice. Then cells were centrifuged at 12,000 x g for 10 min at 4°C. Supernatant fluid is total protein of cells. The amount of protein was quantified with Bicinchoninic acid Protein quantitative kit (cat. no. P1511; Applygen Technologies Inc., Beijing, China). Briefly, the working reagent and standard solutions containing different concentrations of BSA were prepared according to manufacturer's instructions. The different protein concentrations of the standard solutions were determined using the enzyme marker. Then the standard curve was plotted, enabling the calculation of the protein concentration of the samples. A total of 30 µg of protein per lane was separated by 10% SDS-PAGE and transferred onto polyvinylidene difluoride membranes. The membranes were incubated with blocking buffer containing 5% skim milk powder and 0.1% Tween-20 (Bio-Rad Laboratories, Inc., Hercules, CA, 50 µg/ml; Beckman Coulter, Inc., Brea, CA, USA) and incubated at 37°C for 30 min. Following staining, the cells were washed once or twice with 0.01 mol/l phosphate-buffered saline (PBS) and cellular DNA content was measured using a Coulter® Epics® XL™ Flow Cytometer (Beckman Coulter, Inc.) to analyze the cell cycle, with three replications per group and three repeats per replication. Briefly, cells were washed twice with 0.01 mol/l PBS and centrifuged for 3 mins at 167.7 x g at room temperature, then incubated with 70% precooled ethanol at 4°C overnight. The cells were then subjected to 50 µg/ml propidium iodide dye solution (containing 50 µg/ml RNase) and analyzed by EXPO™ 32 ADC version 1.1C software (Beckman Coulter, Inc.). Cell viability was analyzed using the 3-[4,5-dimethylthiazol-2-yl]-2,5 diphenyltetrazolium bromide (MTT) assay for three wells in each group every day, continuously for 9 days. The BM-MSCs group and A549 group served as the controls. The test was performed with 5 mg/ml MTT solution prepared by dissolving MTT powder (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) in warm PBS (37°C). Cells (~1x10⁶ cells/well) were incubated in 96-microwell plates (Nalge Nunc International, Penfield, NY, USA) at 37°C for 24 h. Following incubation, 20 µl MTT solution was added to each well and incubated for 4 h in a 5% CO₂ incubator at 37°C. Subsequently, the solution was removed and the construct was blotted with filter paper. Finally, 200 µl 99.5% dimethyl sulfoxide (Sigma-Aldrich; Merck KGaA) was added to each well and the plate was shaken for 5 min. For colorimetric readings, 100 µl of this solution was transferred to an ELISA plate and the absorbance values of the samples detected were read by an ELISA reader at 570 nm. Using cultivated time as the horizontal axis and the absorbance value as the vertical axis, a growth curve was constructed.
USA) at 37°C for 3 h and then washed with PBS-Tween-20 three times. Subsequently, the membranes were incubated with primary antibodies P-ERK (cat. no. BS4759; 1:500; Bioworld Technology, Inc., St. Louis Park, MN, USA), nuclear factor (NF)-xB (cat. no. ab16502; 1:500; Abcam, Cambridge, UK), p53 (cat. no. ZM-0408; 1:500; Santa Cruz Biotechnology, Inc., Dallas, TX, USA), P62 (cat. no. ab91526; 1:500; Abcam), B-cell lymphoma-2 (Bcl-2; cat. no. ab194583; 1:500; Abcam), Bcl-2-associated X protein (Bax; cat. no. ab53154; 1:500; Abcam) and caspase-3 (cat. no. ab59388; 1:500; Abcam) at 4°C overnight. After the incubation of primary antibodies, membrane was washed four times with tris-buffered saline with Tween-20 for 10 mins. Following this, the membranes were incubated with horseradish peroxidase-conjugated secondary antibody (goat anti-rabbit immunoglobulin G; cat. no. ab150088; 1:5,000; Abcam). Proteins were quantified using enhanced chemiluminescence advanced western blotting detection solution (Abcam) and a gel imaging analysis system (Chemidoc; Bio-Rad Laboratories, Inc.). β-actin was used as a reference gene. The grey ratio value between the target protein and reference protein in the same sample was used as the relative expression value of the protein and analyzed using Image Lab™ software version 3.0 (Bio-Red Laboratories, Inc.) following three repeats.

**Karyotype analysis.** Cells at the logarithmic growth phase were coated with 0.2 µg/ml colchicine (cat. no. c8190; Solarbio Science & Technology Co., Ltd.) and set aside for 3 h. The cells were fixed through low permeability of 0.075 mol/l KCl, methanol-acetic acid (3:1), dropped on an ice slide and treated with 0.25% trypsin for 1 min at room temperature. Karyotype analysis was performed using an ellipse 80i microscope (Nikon Corporation, Tokyo, Japan) to count 100 mitotic chromosomes using a chromosome image analyzer CytoKItype 6.2.2 (United Biotechnology Corporation, San Jose, CA, USA).

**Analysis of cytoskeleton and migration ability of cells.** Cell suspensions were placed on a slide, fixed with 4% paraformaldehyde for 15 min at room temperature, rinsed with PBS containing 5% bovine serum albumin (cat. no. A8020; Solarbio Science & Technology Co., Ltd.) and 0.1% Triton X-100 and stained with phallolidin-labeled fluorescein isothiocyanate (FITC; cat. no. P5282) and Hoechst (cat. no. H6024; both from Sigma Aldrich, Merck KGaA). The staining procedure was performed following the manufacturer’s protocol. Stained slides were placed under a laser scanning confocal microscope to observe the cell cytoskeleton.

Cell migration ability was then investigated using a Millicell μ-Migration Assay kit (EMD Millipore, Billerica, MA, USA), according to the manufacturer’s instructions. Briefly, following 7 days of culture, cells from the three groups (A549 group, BM-MSCs group, and co-BM-MSCs group) were transplanted into a Millicell (1x10^5 cells/Millicell, with a total volume of 100 µl) of the 24-well plate. A total of 500 µl A549 cell conditioned medium was added into the outer chamber of the Millicell. The aforementioned cells were cultured routinely for 8 h at 37°C, and cells that did not migrate were removed using a cotton swab. The 24-well plate was inverted to air-dry and 500 µl 0.1% crystal violet was added at 37°C for 30 min. The plate was placed under an inverted microscope to count cells in five fields selected at random. Levels of the cell migration-related gelatinases, matrix metalloproteinase (MMP) were detected by western blot analysis (identical to the aforementioned method, with primary antibodies: MMP-2 (cat. no. BS1236; 1:500; Bioworld Technology, Inc.), MMP-9, (cat. no. P14780; 1:500; Bioworld Technology, Inc.), β-actin (cat. no. SC-47778; 1:500; Santa Cruz Biotechnology, Inc.) and goat anti-rabbit immunoglobulin G secondary antibody (cat. no. 150088; 1:5,000; Abcam) to analyze the migration ability of the three groups of cells.

**Statistical analysis.** Data were expressed as the mean±standard deviation. SPSS v. 17.0 software (SPSS, Inc., Chicago, IL, USA) was used to determine statistical significance between groups. For statistical comparisons, parametric analysis of variance with the Turkey’s test. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**Changes in cell morphology.** BM-MSCs were uniform in morphology, fibroblast-like, fusiform or spindle-shaped, adherent on the plate, distributed evenly, arranged in an orderly manner with a clear boundary and exhibited swirl-like growth (Fig. 1A). Following co-culture with A549 cells for 7 days, BM-MSCs changed shape, cells became small and short, were arranged in a disorderly manner, exhibited irregular polygon overlapping growth, enlarged nuclei, an uneven nuclear shape and staining, and visible pathological mitotic figures. Similar biological cytomorphological features and growth characteristics were observed between A549 cells and co-BM-MSCs (Fig. 1B-D). These results indicate that co-BM-MSCs have a tendency to undergo malignant transformation (8-10).

**Analysis of cell cycle and viability.** Flow cytometric analyses demonstrated that, compared with BM-MSCs, the number of cells in the G1 (stationary) phase in the co-BM-MSCs group was lower and the number of cells in the S-(proliferating) phase were significantly higher ($P<0.05$). There was no difference in the percentage of cells in the G0/G1 between the BM-MSC group and the co-BM-MSCs group. The percentage of cells in the G0/G1 in BM-MSCs group is 77.8±5.04% and in co-BM-MSCs group is 62.57±6.65%. The percentage of cells in S in the BM-MSCs group is 9.05±0.8% and in co-BM-MSCs group is 23.54±0.6% (Fig. 2). Using cultivation time as the horizontal axis and the absorbance value as the vertical axis to draw the growth curve, it was determined that the cell growth curves of the three groups were S-shaped. co-BM-MSCs and BM-MSCs exhibited a slow initial increase in viability from days 1-3 followed by more rapid increases in viability from day 4. The co-BM-MSC group underwent a more rapid increase in viability than the BM-MSC group; however, this increase leveled off after day 8. The viability of cells in the A549 group was higher than that of cells in the co-BM-MSCs and BM-MSCs groups (Fig. 3). These results demonstrate that the tumor microenvironment created by the A549 cells is able to promote the viability of BM-MSCs.

**Changes in the expression of proteins associated with the ERK signaling pathway, proliferation and apoptosis.** To investigate...
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the progression of proliferation and morphological changes of BM-MSCs growing in the tumor microenvironment formed by A549 cells, total proteins were extracted from the three groups of cells following 7 days culture, in order to measure their expression by western blotting. Proteins associated with the ERK signaling pathway were selected to analyze the difference in levels of expression among the three groups of cells. Compared with the BM-MSC group, levels of Ras, P-ERK, NF-κB, P62 and Bcl-2 expression in the co-BM-MSC group were significantly increased (P<0.05) and levels of P53, Bax and caspase-3 expression decreased significantly (P<0.05; Fig. 4). Consequently, the changes to the biological properties of BM-MSCs following co-culture with A549 cells may be related to abnormal expression of genes associated with the ERK signaling pathway.

Changes in karyotype. Analyzing chromosome karyotypes, which is used to determine cellular genetic structure and type, chromosome number and genetic aberrance, is a basic method of studying cell chromosomes. In the present experiment, the results of the chromosome karyotype analysis determined that BM-MSCs were 46, XX, diploid. By contrast, the co-BM-MSC karyotype was between 46 and 70, hypotriploid, triploid. Part

Figure 1. Cell morphology of different groups on day 7 of culture. (A) BM-MSCs were uniform in morphology, fibroblast-like, fusiform or spindle-shaped and adherent on the plate. They were distributed evenly, arranged in an orderly manner with a clear boundary and demonstrated swirl-like growth. Magnification, x100. (B) A549 cells were long fusiform or irregular polygon and arranged in a disorderly manner. Magnification, x100. (C) co-BM-MSCs were short and small, irregular polygon, arranged in a disorderly manner and exhibited overlapping growth. Magnification, x100. (D) co-BM-MSCs stained with hematoxylin staining exhibited enlarged nuclei, an uneven nuclear shape and stain, and exhibited visible pathological mitotic figures. Magnification, x1,000. BM-MSC, bone marrow mesenchymal stem cells; Co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

Figure 2. Cell cycles of the different groups were measured using a flow cytometer following 7 days culture. (A) BM-MSC cell cycle analysis demonstrated that 77.8% of cells were in the G0/G1 phase and 9.05% in the S-phase. (B) Analysis of the co-BM-MSC group demonstrated that 62.57% of cells were in the G0/G1 and 23.54% in the S-phase. (C) A549 cell cycle analysis demonstrated that 59.95% of cells were in the G0/G1 phase and 21.41% in the S-phase. There was a significant difference between the cell cycle of BM-MSCs and that of co-BM-MSCs (P<0.05). BM-MSC, bone marrow mesenchymal stem cells; Co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

Figure 3. Growth curve profiling of the different cell groups during 9 days culture. MTT analysis demonstrated that the viability of co-BM-MSCs increased more rapidly than that of the BM-MSC group. Data are presented as the mean ± standard deviation. BM-MSC, bone marrow mesenchymal stem cells; Co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

Figure 4. Changes in karyotype. Analyzing chromosome karyotypes, which is used to determine cellular genetic structure and type, chromosome number and genetic aberrance, is a basic method of studying cell chromosomes. In the present experiment, the results of the chromosome karyotype analysis determined that BM-MSCs were 46, XX, diploid. By contrast, the co-BM-MSC karyotype was between 46 and 70, hypotriploid, triploid. Part
of the co-BM-MSCs cells exhibited an aneuploid chromosome number following co-culture, with obvious abnormal chromosomes (Fig. 5; Table I). These results suggest that, in a tumor microenvironment, BM-MSCs had a malignant tendency.

**Study of the cytoskeleton and cell migration.** The three groups of cells stained with Hoechst and phalloidin-labeled FITC were placed under a laser scanning confocal microscope to observe the cytoskeleton microfilament. The results demonstrated that the microfilament of BM-MSCs was bundle-like or radial-gathered in the projection and the cytoplasm, whereas that of the co-BM-MSC group was irregular and resembled that of the A549 cells. The microfilament was observed under the microscope (Fig. 6). Compared with the BM-MSCs group,
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Figure 6. Analysis of cytoskeleton and cell migration. Cells were stained with Hoechst and phalloidin-labeled fluorescein isothiocyanate. The microfilament of (A) BM-MSCs was bundle-like or radial gathered in the projection and the cytoplasm, while that of the (B) Co-BM-MSCs was irregular, similar to that of the (C) A549 cells. Microfilament could be seen under microscope at magnification of x1,000. Scale bar, 50 μm. BM-MSC, bone marrow mesenchymal stem cells; co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

Table I. Chromosome number analysis of the different groups of cells.

| Group         | Chromosome number | Rate of chromosome aberration, % |
|---------------|-------------------|----------------------------------|
| BM-MSC        | 46.0              | 0.0                              |
| Co-BM-MSC     | 54.4±8.2          | 66.7                             |
| A549          | 62.1±4.6          | 100.0                            |

Data are presented as the mean ± standard deviation where appropriate. P<0.05 vs. BM‑MSC chromosome number. BM‑MSC, bone marrow mesenchymal stem cells; Co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

Table II. Analysis of the number of cells in each group that migrated to the Millicell chamber of the plate.

| Group         | Number of migrating cells |
|---------------|---------------------------|
| BM-MSC        | 89.00±7.55                |
| Co-BM-MSCs    | 18.67±3.21                |
| A549          | 12.67±2.08                |

Data are presented as the mean ± standard deviation. P<0.05 vs. the BM-MSC group. BM-MSC, bone marrow mesenchymal stem cells; Co-BM-MSCs, BM-MSCs co-cultured with lung cancer A549 cells.

The number of cells in the co-BM-MSC group that migrated to the Millicell chamber significantly decreased (P<0.05; Table II). There was no significant difference in the number of migrated cells in the Millicell chamber between the co-BM-MSC group and A549 group (P>0.05; Table II). Furthermore, compared with the BM-MSC group, levels of MMP-2 and -9 expression were significantly lower compared with the co-BM-MSCs group (P<0.05; Fig. 7).

Discussion

The cellular microenvironment serves an important role in maintaining normal proliferation, differentiation, metabolism and function. Abnormal changes to microenvironment components may result in the formation of cell lesions (11). The results...
of the present study demonstrate that the viability and number of chromosomes of the BM-MSCs changed abnormally when in the lung cancer A549 cell microenvironment.

Proliferation and differentiation are two important aspects of cellular activities. The proliferation of cells prepares them for differentiation and there is a very close relationship between them. Cell differentiation is inhibited during the proliferative phase, whereas the proliferation ability of differentiated cells is decreased (12). In the present study, the growth curve of the cells of the three groups was identified to be S type. The co-BM-MSC and BM-MSC groups exhibited a slow increase in viability from days 1-3 and a rapid increase in viability starting from day 4. The increase in viability was faster in the co-BM-MSC group than the BM-MSC group. After day 8, increases in cellular viability ceased. The viability of the A549 group was higher than that of the co-BM-MSC and BM-MSC groups.

The cell cycle is the basic process of cell life and refers to the entire life cycle of a cell from the end of one division to the end of the next one (13). Using flow cytometry and PI staining to detect the content of DNA in cells, the cell cycle can be divided into the G1/G0, S- and G2/M phases. G1 phase initiation is the key to cell cycle progression. Whether cells may enter the division cycle and whether the cycle may be completed predominantly depends on its ability to smoothly pass through a series of checkpoints, of which the most important are the G1/S and G2/M transformations (14).

In the present study, analysis of the cell cycle by flow cytometry demonstrated that, compared with the BM-MSC group, the ratio of G1 phase cells in the co-culture group significantly decreased and those in the S-phase increased markedly. There was a significant difference between the cell cycles of the two groups. The G0 (stationary phase) ratio in the co-culture group was lower than that in the BM-MSC group and the proliferating cell ratio in the S-phase increased markedly compared with that in the BM-MSC group. This demonstrates that the co-culture group had greater proliferation ability than the BM-MSCs group. Compared with the A549 group, there was no significant difference in the G1 and S-phase ratio in the co-culture group, suggesting that the cell cycle of the co-culture group was similar to that of the A549 group. The lung cancer A549 cell microenvironment promoted bone marrow mesenchymal stem cells from the G1 phase to the S-phase and drove the cell cycle transition from G2 phase to M phase (15). This change may be associated with the increased number of MSCs, which were no longer able to undergo multi-directional differentiation (16).

Chromosomal changes include alterations to chromosome number and structure. Compared with gene mutations, aneuploidy is more relative to cancer. Cytogenetic data have demonstrated that almost all tumor cells are represented as aneuploid karyotypes (17,18). Li et al (19) proposed the aneuploid hypothesis of cancer, where the normal gene expression imbalance caused by aneuploidy is the root of tumor formation, and aneuploidy rather than other mutations is able to provide a more reasonable explanation of tumor cells in a variety of phenotypes (19,20). In the experiment, regular karyotype analysis demonstrated that total cell chromosome number in the control group was 46, while in the experimental group, cell chromosome numbers ranged from 46-70 (19,20). The results illustrate that the tumor microenvironment may stimulate aneuploidy in BM-MSC chromosomes.

A key problem with stem cell transplantation is how to regulate stem cell migration, proliferation and differentiation following transplantation. Previous studies have demonstrated that signals for the survival, proliferation and apoptosis of various cells are transduced through the activation of mitogen-activated protein kinase (MAPK) and the phosphoinositide 3-kinase/protein kinase B signaling pathways (21-23). MAPK is a widely distributed intracellular protein kinase that contains serine/threonine residues (21-23). MAPKs are involved in various physiological activities, including cell growth, division, differentiation and death, which regulate a series of complex biological functions. The activation of the ERK1/2 pathway is triggered by cell surface receptors. In cultured cells, the ERK pathway is associated with cell proliferation, and whether it promotes or inhibits proliferation depends on the cells, as the ERK pathway activates different genes in different cellular environments. The ERK pathway has different effects due to different activation times and strengths (22). In certain cells, the ERK pathway may also mediate differentiation signals. The ERK pathway is involved in the regulation of the cell cycle and its activation may be anti-apoptotic (24), leading to unlimited cell proliferation. The P38 MAPK pathway is associated with the proliferative response induced by cytokines and inhibiting the P38 MAPK pathway may inhibit the induction of cell proliferation by interleukin (IL)-2 and IL-7 (25). Overall, pathways related to apoptosis, such as the P38 MAPK pathway, and catabolic signals together determine whether caspase activation leads to irreversible apoptotic processes (26). Research has demonstrated that the P38 MAPK pathway is related to BM-MSC proliferation (27). In the present study, P-ERK in the P38 MAPK pathway was analyzed by western blot analysis. These results indicate that BM-MSCs co-cultured with lung cancer A549 cells may spontaneously change their cell proliferation ability, possibly via the P38 MAPK pathway.

NF-κB is a fast response transcription factor readily detected in eukaryotic cells and also serves an important role in proliferation, differentiation, apoptosis and autophagic cells. Research suggests that the damage factor of NF-κB activation may produce pro-apoptotic factors, including p53, c-Myc and cyclin D1 in neural cells, leading to the induction of apoptosis (4,28,29). The tumor suppressor p53 is a key mediator of apoptosis. P53 is the target gene of NF-κB. Nuclear translocation of NF-κB may upregulate p53 expression and inhibition of NF-κB nuclear translocation may downregulate p53 expression; thus NF-κB may serve an anti-toxicity role (30-32). P53 regulates a variety of genes and some of them may affect mitochondrial genes, thereby affecting cellular apoptosis. P53 may also be transferred to the mitochondria, affecting depolarization of the mitochondrial membrane, releasing pro-apoptotic factors and inducing apoptosis. Furthermore, p53 is an important transcription regulator of cell proliferation, not only in tumor cells, but also in stem cells (33). Research has demonstrated that there are regulatory pathways in which p53 is not only the cell genome supervisor but also the protector for the proliferation and differentiation of cell precursors (32).

Apoptosis, as in cell proliferation and differentiation, serves an important role in maintaining cellular homeostasis as it is
the process of programmed cell death induced by physical and chemical factors (34). Apoptosis is regulated by a series of genes, such as those from the Bcl-2 gene family. High expression of Bcl-2 protein may protect cells from apoptosis (35), whereas Bax induces cellular apoptosis. Bax and Bcl-2 therefore regulate cell apoptosis positively and negatively, respectively, and their ratio determines whether cells undergo apoptosis or not.

In the present study, the viability of BM-MSCs co-cultured with human lung adenocarcinoma A549 cells was greater than that of the control group BM-MSCs. Western blotting demonstrated that levels of NF-κB, Bcl-2, Bax, p53, caspase-3 and Ras expression in co-cultured BM-MSCs were between the levels of those in the A549 and BM-MSCs groups. These results indicate that the co-culture tumor microenvironment may alter gene expression via molecular pathways, resulting in the cell evading physiological apoptosis and leading to abnormal cellular proliferation. The tumor suppressor gene p53 and the oncogene Ras are key genes involved in cell signal transduction pathways, which determine cellular fates (36). P53 and Ras regulate key cellular functions, including cell proliferation, programmed cell death, cell movement, inflammation and angiogenesis (37). At present, it has been demonstrated that hundreds of proteins interact with p53. Similarly, Ras has a complex regulatory network. Although p53 and Ras have different molecular pathways, together they control key cellular procedures (38). A study by Buganim et al (39) demonstrated that activation of Ras and inactivation of p53 were able to induce expression of the NF-κB gene and a large number of studies have demonstrated that the regulation of NF-κB by p53 and Ras is important in carcinogenesis (40,41). Such findings suggest that there are synergistic effects between p53 and Ras, and both direct and indirect regulatory interactions and networks may exist between them. Judging from the results of the present experiment, the tumor microenvironment may result in the abnormal proliferation of MSCs due to the synergism between p53 and Ras.

The key to successful stem cell transplantation is the successful migration, proliferation and the genetic stability of stem cells following transplantation (42). The response of cells to outside stimulation is through interactions between different signaling pathways. No path exists independently in cell reactions and each pathway may integrate with many other pathways. The intensity of the regulation and the interplay between signaling pathways decides the fate of the associated cells (1). The results of the present experiment indicate that the tumor microenvironment is able to induce spontaneous changes in the proliferation and migration ability of BM-MSCs. However, the exact molecular mechanism has not been fully elucidated and the application of BM-MSCs in biological treatment requires further investigation.

The tumor microenvironment is a complex system that consists of numerous stromal cells. Tumor cells, through the production of growth factors and proteases, regulate the tumor stromal environment and may also activate various types of cells in tumor stroma at the same time (43). Following activation, stromal cells secrete insulin-like growth factor-1, hepatocyte growth factor and other cytokines, which promote the malignant transformation of cells (8).

In conclusion, the in vitro lung cancer A549 microenvironment may induce BM-MSCs to undergo changes in cell morphology, proliferation, karyotype, cytoskeleton and migration ability. The underlying mechanisms for these changes may be related to abnormal expression of related genes in the ERK signaling pathway and apoptosis.

Acknowledgements

The present study was supported by the Outstanding Youth Fund Projects of Gansu Province (grant no. 1308RJDA008) and the National Natural Science Foundation of China (grant nos. 81360588 and 81473457).

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