Influence of acrylamide on ROS, Hsp27 and NF-κB in bone marrow mesenchymal stem cells

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ABSTRACT

The bone marrow mesenchymal stem cells (BM-MSCs) treated with acrylamide (ACR) were used to make out the immune response to ROS, interleukin-8 and phosphorylated Hsp27 of ACR. ACR was reported as a probable human carcinogen, neurotoxic and mutagenic. BM-MSCs have the capability of immunoregulation, and participate in the process of multiple immune response. It has attracted the attention of researchers that these cells have priority to move to the damaged tissue, as a kind of potential therapeutic tool for tissue repair. ACR and BM-MSCs are related to immune reactions, especially those involving in tumours and cancers. However, the interaction between ACR and BM-MSCs is still poorly understood. In present study, we report the influence of ACR on BM-MSCs. At first, BM-MSCs were disposed with 0.5mM ACR for 72 h, and then the secretion of ROS, interleukin-8, phospho-Hsp27 and NF-κB activities, apoptosis and cell cycle, respectively, were determined. The results showed that the secretion of ROS, interleukin-8 and phosph-Hsp27 increased and NF-κB was activated, while the apoptosis and cell cycle have no obvious alteration. In conclusion, ACR probably activated the NF-κB pathway in BM-MSCs via oxidative stress, which may provide new insights to study the immune response and the influence mechanism of ACR.

Key words: Acrylamide, Bone marrow mesenchymal stromal cells, Hsp27, NF-κB, Porcine, ROS
that TNF-α dramatically increases human MSC (HMSC) migration via extracellular matrix interactions, which is partially regulated by the NF-κB pathway (Bocker et al. 2008).

Nuclear factor-kappa B (NF-κB) belongs to a family of inducible transcription factors (Dos Santos Nascimento et al. 2015), which involve cytokine- or pathogen-induced upregulation of inflammatory cytokines and subsequent immune responses. However, under certain circumstances, inflammatory cytokines may become immunosuppressive. For example, bone marrow stromal cells (BMSCs) from MM patients uniquely enhance the constitutive NF-κB activity in MM cells via a proteinaceous secreted factor in conjunction with interleukin-8 (IL-8) (Markovina et al. 2010).

In a word, we demonstrated that ACR activated the NF-κB pathway in BMSCs. It lays a foundation for further studies of the influence mechanism of ACR. At the same time, it also provides experimental and theoretical basis for better applying MSCs in the clinical treatment.

MATERIALS AND METHODS

Isolation and culture of porcine MSCs: Femur and tibia bones from 6-day to 6-month-old porcine (n>3) were isolated to extract the gelatinous bone marrow under sterile conditions, and had been evaluated by flowcytometric analysis, adipogenic and osteogenic differentiation as described previously (Huang et al. 2015). The extracted gelatinous bone marrow was resuspended in DMEM/F-12 (Dulbecco’s Modified Eagle Medium, Nutrient Mixture F-12) and dispersed gently. Samples were centrifuged (1,000 rpm, 5 min) and resuspended in 5 ml of DMEM/F-12 containing 15% fetal bovine serum (FBS) and 1% pen/strep. The cells were seeded in a 100-mm culture plate at a concentration of 1×10⁶/cm² and incubated in a humidified 5% CO₂ atmosphere at 37°C. Nonadherent cells were gently rinsed off by replacing half of the culture medium after plating cells for 48 h. The attached MSCs were confluent after 5–7 days (passage 0). For every passage, cells were trypsinised using a 0.25% Trypsin-EDTA solution. With the exception of passage 0, the FBS content in the complete culture medium was changed to 50% FBS solution. Then the culture medium was changed every 3 days to promote cell growth. The cells were split approximately 1:2 at each passage after the culture reached 80–90% confluency.

Measurement of reactive oxygen species: The content of the reactive oxygen species (ROS) that were produced by BMSCs treated with ACR was determined using the ROS Assay Kit. Cells were seeded in a 96-well cell culture plate at a concentration of 3×10⁴ cells/well in D-MEM/F-12, containing 10% FBS, 1% pen/strep and cultured at 37°C in a humidified atmosphere with 5% CO₂ for 12 h. The medium was changed before treating the cells with 0.5 mM (the optimal one according to the concentration gradient filter) ACR for 72 h. At the end of the treatment period, the medium in each well was removed, cells were washed with PBS, and the DCF of ROS assay was performed according to the kit instructions. The fluorescence was measured using a microplate fluorescence reader using a 488 nm excitation wavelength with a 525 nm emission filter.

Measurement of phosphorylated Hsp27 by ELISA: Phospho-Hsp27 in control and ACR-treated BMSCs was measured using ELISA. BMSCs were seeded in 96-well plates at a concentration of 2×10⁴ cells/well and treated with 0.5 mM ACR. Control cells were treated with complete culture medium. After 72 h, cells from the 2 groups were collected and prepared for the detection of phospho-Hsp27 according to the manufacturer’s protocol. The plate was read using a fluorescence plate reader with an excitation wavelength at 540 nm and an emission wavelength at 600 nm, followed by an excitation wavelength at 360 nm and an emission wavelength at 450 nm. The readings at 600 nm represented the amount of phosphorylated Hsp27 in the cells, and the readings at 450 nm represented the amount of total cytochrome c in the cells.

Measurement of IL-8 secretion by ELISA: Secretion of IL-8 in the spent media of control and ACR-treated BMSCs was measured with the Quantikin® Porcine IL-8/CXCL8 Immunoassay. Cells were treated with ACR as described above for 72 h. The supernatants of the ACR-treated cells and control cells were collected according to the manufacturer’s protocol. The optical density was read using a fluorescence plate reader with the emission wavelength set to 450 nm and the correction wavelength at 540 nm. IL-8 concentrations of the 2 groups were extrapolated from a standard curve that was derived using known concentrations of IL-8. Sample values were normalised to the total cell protein concentrations, which were determined using the BCA protein assay.

Relative quantification of IL-8 gene expression by real-time PCR: Total RNA of BMSCs cultured for 72 h in culture medium containing 0.5 mM ACR was extracted. One microgram of total RNA was reverse-transcribed into cDNA with oligo (dt) primers. For real-time PCR, the following primers were designed using the Primer 5.0 software IL-8 F: 5'-cttgagctcattccggt-3', IL-8 R: 5'-atagaggaagttcggagaggg-3'. GAPDH F: 5'-gccgaactgtgatggtg-3', GAPDH R: 5'-ctcgctcgggatgtggt-3'.

Real-time PCR was performed using the ABI 7500 real-time PCR system with the Real Master Mix (SYBR Green). The RT-PCR conditions were as follows: 50°C for 2 min and denaturation at 95°C for 30 sec followed by 40 cycles of denaturation at 95°C for 20 sec, reannealing at 60°C for 30 sec, and extension at 72°C for 30 sec. The gene GAPDH was used as an internal control to normalise cDNA quantity. The expression level of target gene was quantified using the comparative threshold (Ct) method. Each real-time PCR reaction was performed in triplicate, and the data were shown as mean ± SE (n=3). The significant differences were determined using the one-way analysis of variance (ANOVA) test using SPSS software 13.0, and the significance levels were set to P<0.05.
Transfection, luciferase reporter assays: BM-MSCs were seeded in 96-well plates at a density of 2×10^4 cells/well. Once cells reached approximately 60% confluence, the cells were transfected using Lipofectamine 2000 using a procedure of Li et al. (2011). The samples were divided into 3 groups, as follows: positive, ACR-treated and control groups. Samples were transfected with 70 ng/well of the reporter plasmid pNF-κB-Luc and 70 ng/well of pRL-TK plasmid for the control and ACR-treated groups, 70 ng/well of the expression plasmid pcDNA3.1 (+)-p65 (Li et al. 2011) to induce p65 overexpression as the positive control or 70 ng/well of the empty control plasmid pcDNA3.1 (+) as the negative. After 6 h of the transfection, the culture medium of the treatment group was replaced with complete culture medium containing 0.5 mM ACR. The media of the other groups were placed with normal culture media. Luciferase assays were performed 72 h after transfection using the dual-specific luciferase assay kit according to the manufacturer’s directions. All of the reporter assays were repeated at least three times. The samples were collected according to the manufacturer’s protocol.

Cell cycle analysis: Cells (Passage 3–6) were harvested using 0.25% trypsin, washed in PBS and fixed in absolute ethanol at −20°C. The cells were washed once with PBS, incubated with DNA staining solution containing propidium iodide and RNase A for 30 min at room temperature and analysed using FACS Calibur flow cytometry.

Analysis of cell apoptosis: Apoptotic cells (Passage 3–6) were detected using the Annexin V-FITC apoptosis detection kit using the manufacturer’s protocol. Cells were collected after ACR treatment and resuspended in PBS. Approximately 1×10^6 cells were incubated with Annexin V-FITC and analysed using FACS Calibur flow cytometry.

Statistical analysis: Data were expressed as mean ± SE. The statistical significance was determined in all of the experiments using the one-way ANOVA test by SPSS software 13.0. Student’s t-test was also used to evaluate the statistical significance. P<0.05 were considered statistically significant.

RESULTS AND DISCUSSION

ACR-induced increase in ROS: After ACR treatment (0.5 mM) for 72 h, the DCF of ROS in the BM-MSCs markedly rose compared to that of the control group (Fig. 1). It illustrated that the ROS production increased. The differences between ACR-treated and control cells were statistically significant (P<0.01).

Under normal conditions, the system of ROS generation and scavenging is maintained in a dynamic equilibrium in vivo. Previous studies showed that oxidative stress is involved in chromosomal aberrations, gene mutations, tumour formation and cancer development. ROS provides a mechanism for significant indirect genotoxicity (Vinoth et al. 2015). The above results indicated that ACR treatment increased the production of ROS in cells, creating an oxidative-stressed state. Under these conditions, ROS generation may occur at a relatively higher rate than ROS scavenging. Imai et al. 2008 showed that viral infection caused oxidative stress. In addition, previous studies have suggested that lipopolysaccharide (LPS) and dextran sodium sulphate (DSS) increase ROS production and regulate the subsequent activation of the NF-κB pathway in colonic epithelial cells (Bhattacharyya et al. 2008, 2009). A similar mechanism may exist among these xenobiotics to increase the ROS generation. Increasing ROS generation may decrease the secretion of Hsp27 and promote the activation of the NF-κB pathway and apoptosis in epithelial cells (Bhattacharyya et al. 2008). Therefore, we further examined the influence of ACR on Hsp27 and NF-κB activity in BM-MSCs.

![Fig.1. Increased reactive oxygen species (ROS) production following ACR exposure in BM-MSCs. BM-MSCs in culture was treated with ACR (0.5mM) for 72 h and the DCF of ROS was measured. Differences between the control and treated cells are statistically significant (P<0.05). Control group was BM-MSCs cultured in normal medium.]
sequence of pp65 [pcDNA3.1(+)-pp65] and the luciferase reporter gene vectors were cotransfected into the ACR-treated and control BMMSCs. ACR remarkably increased (P<0.01) the luciferase activity (Fig. 3) of the reporter gene vector (pNF-κB-Luc).

The results showed that the NF-κB pathway was activated after BMMSCs were treated with ACR. Previous researchers have implied that NFκB transcription factors play a key role in the survival and proliferation of B-cell tumours, including multiple myeloma (MM) cells (Markovina et al. 2010, Demchenko et al. 2010). In MM, the activation of NF-κB in BMMSCs may uniquely promote NF-κB activity in MM cells but not normal cells. Djouad et al. (2003) reported that the subcutaneous injection of B16 melanoma cells induces tumour growth in allogeneic recipients only with the co-injection of MSCs (Abdi et al. 2008). Whether ACR-induced NF-κB activation in BMMSCs participates in the process of carcinogenesis requires further investigation.

ACR-induced increase in IL-8 secretion from BMMSCs:

IL-8 was assayed using ELISA assay kit in control and ACR-treated BMMSCs. After 0.5mM ACR treated for 72h, the level of IL-8 secretion from BMMSCs was decreased when cells were treated with ACR (0.5mM) for 72 h. ACR increased IL-8 secretion obviously compared with control (P<0.05). (b) Expression of IL-8 in ACR(0.5mM) treated BMMSCs and control assessed by real-time PCR. Error bars indicate the SE (n = 3) of relative mRNA expression levels of IL-8 to GAPDH, determined by real-time quantitative PCR. Compare with control group, gene expression of IL-8 was increased significantly (P < 0.05). The experiments were repeated three times.

Cell cycle and apoptosis: The cell cycle was examined using a cell cycle stainsolution by flow cytometry. As shown in Fig. 5, no obvious differences were observed between ACR-treated and control BMMSCs. The apoptosis
was measured using annexin V and the dead cell apoptosis kit after ACR treatment for 72 h. The fluctuation of apoptotic rates was not observed after ACR treatment (Fig. 6).

The anti-oxidation stress activities of Hsp27 play a remarkably role in anti-apoptosis and may promote the proliferation of BM-MSCs. Conversely, cells that were treated with high concentrations of ACR might decreased Hsp27 expression, which promoted ROS generation and activation of the NF-κB pathway to induce apoptosis. However, the apoptosis-promoting function of Hsp27 was generally restricted because the protective effect of HSP was predominant. This microregulated balance is the key to cell death or survival and functions as the transition centre between cell necrosis and apoptosis.

In conclusion, ACR is gaining more notoriety as a common compound in our daily life. The current report studied the immune reaction of ACR-treated BM-MSCs. We hypothesised that BM-MSCs stimulated with 0.5 mM ACR displayed increase in the secretion of ROS, phospho-Hsp27 and IL-8 and the activation of NF-κB as well as no obvious changes in apoptosis and proliferation. Further investigation is required to detect the influence of high concentrations of ACR on BM-MSCs. The conclusions of the current study may help elucidate the mechanism of ACR on immune responses.

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