Carbenoxolone inhibits mechanical stress-induced osteogenic differentiation of mesenchymal stem cells by regulating p38 MAPK phosphorylation

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Abstract. The aim of the present study was to explore the effects of pannexin1 (Px1) protein channels on osteogenic differentiation of mesenchymal stem cells (MSCs) under mechanical stress stimulation. MSCs were isolated from Sprague Dawley rats (3 weeks old, weighing 100-120 g) and cultured in vitro. A safe concentration of carbenoxolone was determined (CBX, an inhibitor of Px1 channels; 100 µM) on MSCs using the Cell Counting Kit-8 (CCK8) method. MSCs were divided into 6 groups: Control, stress (4,000 µ strain), and stress following 3, 6, 12, and 24 h pretreatment with CBX. Stress groups were stimulated with mechanical stress for 15 min. Alkaline phosphatase (ALP) activity, type I collagen expression, intracellular calcium ion (Ca²⁺) concentration, Px1 expression, p38 mitogen-activated protein kinase (MAPK) and extracellular signal-regulated phosphorylation were determined. ALP activity was increased in the stress group, and this was prevented by pretreatment with CBX. Similarly, stress-induced increases in type I collagen expression, Ca²⁺ concentration, Px1 expression, and p38 MAPK phosphorylation decreased in the presence of CBX. ERK phosphorylation was decreased by stress, however was not affected by CBX treatment. Altogether, the results suggest that mechanical stress promoted the osteogenic differentiation of MSCs, and this promotion was inhibited by pretreatment with CBX, possibly through regulating the phosphorylation of p38 MAPK.

Introduction

Mesenchymal stem cells (MSCs), derived from the mesoderm, have a strong self-replicative ability and multi-differentiation potential. MSCs are used in tissue engineering, cytokines alternative therapy, gene therapy, and other aspects of organ transplantation (1), and as biomaterials in the repair of bone, cartilage, tendons, and glia (2). MSCs differentiate into osteoblasts and then into bone cells, accounting for approximately 95% of bone. Bone cells can act as mechanical stress receptors, and regulate the activities and functions of both osteoblasts and osteoclasts in the form of endocrine cells (3).

A previous study has shown that mechanical stress stimulation promotes osteogenesis (4). Mechanical stress stimuli can cause cell morphology changes, cell projection increases, formation of intercellular gap junctions (GJs), and cytoskeletal structural changes (5). GJs are composed of two adjacent hemichannels which anchor to each other, forming a hydrophilic channel (6) for intercellular communication via small molecules, such as inorganic ions (calcium and potassium), amino acids and glucose (7,8). GJs are dynamic structures, and a variety of factors are involved in regulating their opening and closing, such as intracellular pH, calcium ion (Ca²⁺) concentration, and membrane potential (9). GJs have functions in cell metabolism and differentiation, transmission of nerve impulses and conduction of information, coordination of consistency between cellular activity, material transport, and electrical excitation conduction (10).

There are three GJ protein families: i) innexin, mainly expressed in invertebrates; ii) connexin, mainly expressed in vertebrates; and iii) a class of genes in vertebrates which are similar to invertebrate innexin, and named pannexin (Px) by
Panchina et al. (11). There are three Px subtypes: Px1, Px2, and Px3. Px is widely expressed in human tissues and organs, such as osteoblasts, chondroblasts, spleen cells, skin tissues, the kidney, and the central nervous system (12). Px forms homodimers in the cell membrane, which can be induced to open by hypoxia, low permeability, mechanical stress, cell depolarization, and Ca\(^{2+}\) concentration increases to mediate ATP release and intercellular Ca\(^{2+}\) wave transmission, regulating blood flow or the immune response (13,14). Px1 is a newly discovered mechanical stimuli-sensitive channel protein; however, whether Px1 participates in MSC osteogenic differentiation and signal transduction after mechanical stimulation has been rarely reported. This study aimed to explore the relationship between Px1 channels and MSC differentiation under mechanical stress stimulation, and examine the specific molecular mechanism.

Materials and methods

Isolation and culture of rat MSCs. The Ethics Committee and Animal Management Committee of The First Affiliated Hospital of Dalian Medical University (LCKY2013-18) approved all experiments. MSCs were isolated from the femurs and tibias of Sprague Dawley (SD) rats (SPF grade, 3 weeks old, weighing 100-120 g). Briefly, rats were euthanized by ether anesthesia with 1.5% pentobarbital sodium (375 mg/kg) followed by cervical dislocation. Muscle tissue and cartilage were removed, and bone marrow suspensions were obtained from marrow cavities using an injector. Subsequently, the bone marrow suspensions were dispersed and centrifuged at 200 x g for 5 min. After removing the supernatant, the pellet was suspended in Dulbecco's modified Eagle's medium (DMEM; HyClone; GE Healthcare Life Sciences, Logan, UT, USA) with 100 U/ml penicillin and 100 U/ml streptomycin (HyClone; GE Healthcare Life Sciences) in a 10-cm dish, and cultured for 48 h. Cell passaging was performed and the third generation was used in the following experiments.

Experimental protocols. First, a safe concentration of CBX (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) was determined using the CCK8 method according to the instructions provided with the CCK-8 cell proliferation and cytotoxicity assay kit (cat. no. CA1210; Beijing Solarbio Science & Technology Co., Ltd., Beijing, China). This concentration (100 \(\mu\)M) was used for subsequent experiments. MSCs were divided into 6 groups: A control group that was neither stressed nor treated with CBX; a stress group, which was only treated with stress; and four groups that were stressed after CBX treatment for 3, 6, 12, and 24 h. Stress loading was performed using a four-point bending device, as previously described (15). The stress loading strength was 4,000 \(\mu\)N strain with a frequency of 0.5 Hz for 15 min. The treated MSCs were cultured in DMEM with 10% fetal bovine serum (Gibco, Carlsbad, CA, USA) in a 37°C incubator (SANYO, Osaka, Japan).

Detection of alkaline phosphatase (ALP) activity and intracellular ATP content. MSCs were sonicated with an ultrasonic cell disruptor (SENIXIN, Shanghai, China) and centrifuged at 13,800 x g for 10 min. The optical density (OD) of the cells was read with a UV spectrophotometer (USA Thermo Electron Corporation, San Jose, CA, USA) at 520 nm. ALP activity was calculated using an Alkaline Phosphatase Assay kit (cat. no. P0321; Beyotime Institute of Biotechnology, Jiangsu, China), according to the manufacturer's instructions. Intracellular ATP content was determined utilizing an ATP bioluminescence kit (cat. no. GN202-01; YPH-Bio, Beijing, China), according to the manufacturer's instructions.

Immunofluorescence observation of type I collagen expression. The expression of type I collagen in MSCs was observed by immunofluorescence. MSCs were fixed in 4% paraformaldehyde at 25°C for 15 min, and then 0.1% Triton X-100 was added for 15 min. After blocking with 5% bovine serum albumin (Beijing Solarbio Science & Technology Co., Ltd.) for 1 h, MSCs were incubated with a rabbit-anti-mouse polyclonal type I collagen antibody (1:500; cat. no. AB765P; Merck KGaA, Darmstadt, Germany) for 2 h and then incubated with DyLight 488 AffiniPure goat-anti-rabbit IgG (H+L) (1:500; EarthOx, San Francisco, CA, USA) for 1 h. After washing thrice with phosphate-buffered saline (PBS), theMSCs were mounted with the anti-quenching sealing agent Fluoromount-G (Southern Biotech, Birmingham, AL, USA) and observed under a fluorescence microscope (Nikon, Tokyo, Japan).

Flu-3AM Ca\(^{2+}\) fluorescence probe to detect the intracellular Ca\(^{2+}\) concentration. Intracellular Ca\(^{2+}\) concentrations were detected using the Flu-3AM Ca\(^{2+}\) fluorescence probe. Briefly, the treated cells were incubated with 3 \(\mu\)M Flu-3AM reagent (Beyotime Institute of Biotechnology, Jiangsu, China, cat. no. SI056) for 15-60 min at 37°C. After washing thrice with PBS, the cells were reincubated once to ensure the complete conversion of Flu-3AM to Flu-3, and observed using a fluorescence microscope (Nikon, Tokyo, Japan).

Western blot analysis. Total proteins were extracted in cell lysis buffer [1% Triton X-100, 100 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 0.2 mM NaVO\(_4\), 10 mM NaF, Protease Inhibitor (Sigma-Aldrich; Merck KGaA), 0.5% NP-40]. Centrifugation at 13,800 x g at 4°C for 10 min, the supernatant was collected. The protein concentration was estimated by Bradford assay utilizing a Bio-Rad Protein Assay Kit II (cat. no. 5000002; Bio-Rad Laboratories, Inc., Hercules, CA, USA). Further, 50 \(\mu\)g of total protein was separated on 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Subsequently, the proteins were transferred onto polyvinylidene difluoride membranes and blocked with 5% skim milk for 1 h at 25°C. Rabbit Px1 polyclonal antibody (1:1,000; cat. no. ab214313; Abcam, Cambridge, MA, USA), rabbit p38 phosphorylation polyclonal antibody (1:1,000; cat. no. ab47363; Abcam), rabbit ERK phosphorylation polyclonal antibody (1:1,000; cat. no. ab214362; Abcam), rabbit anti-mouse polyclonal type I collagen antibody (1:1,000), and β-actin mouse monoclonal antibody (1:5,000; cat. no. AF0003; Beyotime Institute of Biotechnology) were used as primary antibodies, and were incubated overnight with the membranes at 4°C. Subsequently, membranes were incubated with goat anti-rabbit IgG (1:5,000; cat. no. A0208; Beyotime Institute of Biotechnology) or goat anti-rat horseradish peroxidase-conjugated secondary antibodies (1:5,000;
cat. no. A0192; Beyotime Institute of Biotechnology) for 2 h at 25˚C. Finally, proteins were detected using the ECL Plus Western Blotting Detection System (GE Healthcare, Little Chalfont, UK) and visualized using the Bio-Rad gel imaging system (Bio-Rad Laboratories, Inc.). Bands were analyzed using Labwork 4.6 software (UVP Products, Upland, CA, USA).

Statistical analysis. Data are expressed as the mean ± standard deviation (SD). Differences within groups were analyzed utilizing a one-way repeated measures analysis of variance. P<0.05 was considered to indicate a statistically significant difference. SPSS 16.0 software (SPSS, Inc., Chicago, IL, USA) was used for all statistical analysis.

Results

MSCs morphology. Fig. 1 shows the morphology of the MSCs extracted and cultured in whole bone marrow culture. Primary MSCs were initially rounded (Fig. 1A) and then gradually attached and became fusiform (Fig. 1B and C). After the cells were passaged thrice and the confluence reached 80% (Fig. 1D), they were used in experiments.

MSCs ALP activity after treatment with CBX. The stress group showed a significantly higher level of ALP than the control group (P<0.05, Fig. 2). Cells pretreated with CBX showed significantly lower levels of ALP than the stress group (P<0.05, Fig. 2); however, no time-dependent trends were observed.

Expression of type I collagen. Stress stimulation significantly promoted the expression of type I collagen (P<0.05, Fig. 3). When pretreated with 100 µM CBX, type I collagen expression was significantly lower than in the stress group (P<0.05). These results suggest that CBX inhibits the expression of type I collagen.

Detection of intracellular Ca²⁺. Intracellular Ca²⁺ fluorescence intensity significantly increased in the stress group compared with the control group (P<0.05, Fig. 4), while the fluorescence intensity dramatically decreased with CBX treatment compared with the stress group (P<0.05, Fig. 4). Of these, the group treated with 100 µM CBX for 24 h showed the lowest Ca²⁺ level. These results suggest that intracellular Ca²⁺ exchange is blocked by CBX.

ATP release from MSCs. The release of ATP significantly increased in the stress group compared with the control group (P<0.01, Fig. 5). However, after 6-24 h treatment with CBX, ATP release was significantly lower than in the stress group (P<0.05, Fig. 5). These results demonstrate that ATP release can be reduced by CBX.

Pxl expression and p38 and ERK phosphorylation. As shown in Fig. 6, the expression of Pxl and the phosphorylation of p38 and ERK protein kinases were estimated by western blot analysis. There was a dramatically increased expression of Pxl in the stress group (P<0.01, Fig. 6A). However, stress combined with 6-24 h pretreatment with CBX showed significantly lower Pxl expression than with stress alone (P<0.05, Fig. 6A). These results further confirm that Pxl expression is inhibited by CBX. The stress and stress combined with 3 h CBX treatment groups showed higher p38 phosphorylation (P<0.01, Fig. 6B), while longer CBX treatment (6-24 h) resulted in significantly lower p38 phosphorylation (P<0.05, Fig. 6B). These results indicate that CBX inhibits the phosphorylation of p38. ERK phosphorylation was significantly reduced in the stress group compared with the control group (P<0.01, Fig. 6C). Stress combined with 3 h CBX treatment resulted in dramatically higher ERK phosphorylation compared with the stress group (P<0.01, Fig. 6C), but significantly reduced phosphorylation with longer CBX treatment (6-24 h; P<0.05, Fig. 6C). This suggests that there is no direct link between Pxl and ERK signaling.

Discussion

This study indicates that Pxl closely participates in the process of osteogenic differentiation under mechanical stress. Mechanical stress stimulation increased ALP activity, type I collagen expression, and ATP and Ca²⁺ release, promoting osteogenic
differentiation, and these effects were prevented by the addition of the Px1 channel inhibitor CBX. In addition, a positive relationship between Px1 and p38 MAPK signaling was noted.

ALP, also known as mineralization-associated protein, is critical for mineralization during bone formation (16). Akbari and colleagues demonstrated that ALP can serve as a sign of osteogenic differentiation (17). Ca\textsuperscript{2+} deposition is regarded as another marker of osteogenic differentiation (18). A rapid increase in intracellular osteoblast Ca\textsuperscript{2+} concentration occurs in response to external forces (19). Osteoblast cytoskeletal reorganization and gene expression changes are associated with increased stimulation, and are mainly dependent on internal inositol triphosphate-mediated release (20). This increase in Ca\textsuperscript{2+} results from intracellular Ca\textsuperscript{2+} release and extracellular Ca\textsuperscript{2+} flow. In this study, under the stimulation of mechanical stress, ALP activity and cellular Ca\textsuperscript{2+} concentration were significantly increased in MSCs, indicating...
ATP is an important energy molecule in cells under physiological conditions, and an important cell-cell signaling molecule. Under conditions of hypoxia, low permeability, cell deformation, and other stimuli such as depolarization, ATP release increases. There are two explanations for this: ATP, like other neurotransmitters, or in conjunction with other neurotransmitters, could be released via vesicles; ATP release could also be mediated through some kind of a channel. Lu et al suggested that connexin-containing hemichannels are involved in ATP release (21). The Px1 channel has been reported to mediate ATP release from taste cells, airway epithelial cells, red blood cells, and other cell types (22-24). Under physiological conditions, certain stimuli, such as mechanical stress and hypotonic solutions, can induce normal cells to release ATP (25). Previous study indicated that Px hemichannels may mediate the release of both ATP and Ca\(^{2+}\) waves (26).

Recent studies have reported the presence of functional GJs between osteoblasts (27,28). The process of phosphorylation and dephosphorylation of GJ proteins acts as a switch for GJ function. In vitro cultured bone cells under stress display increased phosphorylation levels, indicating increased connections between osteoblasts. In addition, GJs participate in the regulation of osteoblast function and signaling. Px hemichannels can be induced to open by hypoxia, low permeability, mechanical stress, cell depolarization, increased intracellular Ca\(^{2+}\) concentration, and other factors (26).

MAPK signaling pathways are important components of intracellular signal transduction and are involved in a variety of cellular processes, and p38 MAPK in particular plays an important role in the process of differentiation into bone cells (29). The p38 MAPK signal transduction pathway controls three stages in the early differentiation of bone cells, including proliferation, extracellular matrix maturation, and matrix mineralization. In the matrix maturation stage, ALP gene expression levels peak as an early differentiation marker. After the onset of osteoblast differentiation, type I collagen secretion peaks, further promoting bone cell differentiation. The p38 MAPK pathway transduces cellular stress signals to promote osteoblast maturation, including the regulation of ALP expression during osteoblast differentiation (29). This study found that mechanical stress could increase the phosphorylation of p38, but this was decreased after pretreatment with a Px1 inhibitor. Thus we speculate that p38 signaling is important for MSC osteoblast differentiation. In contrast, no changes in ERK phosphorylation were observed after Px1 inhibition, indicating that there was no direct relationship between Px1 channels and ERK signaling.

Altogether, our results indicate that Px1 channels might play a crucial role in the process of osteogenic differentiation. Notably, this may involve signal transduction between Px1 channels and the p38 MAPK phosphorylation.

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