Continuous cyclic mechanical tension increases ank expression in endplate chondrocytes through the TGF-β1 and p38 pathway

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Abstract

The normal ANK protein has a strong influence on anti-calcification. It is known that TGF-β1 is also able to induce extracellular inorganic pyrophosphate (ePPi) elaboration via the TGF-β1-induced ank gene expression and the mitogen-activated protein kinase (MAPK) signaling acts as a downstream effector of TGF-β1. We hypothesized that the expression of the ank gene is regulated by mechanics through TGF-β1-p38 pathway. In this study, we investigated the mechanism of short-time mechanical tension-induced ank gene expression. We found that the continuous cyclic mechanical tension (CCMT) increased the ank gene expression in the endplate chondrocytes, and there was an increase in the TGF-β1 expression after CCMT stimulation. The ank gene expression significantly increased when treated by TGF-β1 in a dose-dependent manner and decreased when treated by SB431542 (ALK inhibitor) in a dose-dependent manner. Our study results indicate that CCMT-induced ank gene expressions may be regulated by TGF-β1 and p38 MAPK pathway.

Introduction

Intervertebral disc degeneration is the most common causes of low back pain. It manifests as loss of the disc space and loss of signal intensity on Magnetic Resonance Imaging (MRI). The endplate cartilage is a layer of hyaline cartilage, approximately 0.6-mm thick, lying between the vertebral body and the intervertebral disc. Endplate cartilage calcification plays an important role in the process of disc degeneration. The intervertebral disc is the largest avascular tissue in the body. One of the main pathways for nutrients to reach the avascular nucleus pulposus is by diffusion from the blood supply of the vertebral body through this endplate cartilage. Endplate calcification could impede the passage of nutrients from the blood to the intervertebral disc leading to alterations in the mechanical material properties of the disc, which can result in failure to maintain the nucleus pulposus.

The normal ANK protein has a strong function on anti-calcification. ANK is a transporter able to export intracellular inorganic pyrophosphate (iPPi) from cells and is known to be upregulated in osteoarthritis. Abnormal extracellular inorganic pyrophosphate (ePPi) metabolism has been implicated in abnormal calcification, decreased concentrations leading to basic calcium phosphate deposition in the articular tissues. Inorganic pyrophosphate (Pi) is also a substrate for alkaline phosphatase which generates the inorganic phosphate (Pi) needed for mineralization. In ank−/−mice, loss of ANK activity results in diminished ePPi levels and consequent basic calcium phosphate (BCP) calcification in joints.

Mechanical cervical traction is an intervention that is often recommended for the treatment of patients with neck pain. However, intermittent cyclic mechanical tension (ICMT) could induce the calcification of endplate chondrocytes and long-term mechanical stimulation decreased the ank gene expression as evident from our previous study. In this study, we investigated the mechanism of short-time mechanical tension-induced ank gene expression. First we focused on the ank gene expression by applying continuous cyclic mechanical tension (CCMT). This was different from the previous mechanical loading used in our earlier study, the CCMT applied continuous stimulation throughout the study period, using short-term mechanical stimulation in vitro. Transforming growth factor-beta-1 (TGF-β1) plays a significant role in regulating crystal deposition in endplate cartilage. TGF-β1 is a potent regulator of cell proliferation and a modulator of interactions of cell with their extracellular matrix (ECM). TGF-β1 is also able to induce ePPi elaboration via TGF-β1-induced ank gene expression. Besides, our study on calcification of endplate chondrocytes induced by ICMT indirectly demonstrated that TGF-β1 is a key factor to regulate the calcification process. The p38-MAPK pathway is important in mechanotransduction and signaling of ank expression. However, there is no evidence that CCMT regulates the expression of ank through TGF-β1-p38 pathway. So in the endplate chondrocytes, was it possible that CCMT could regulate the expression of ank through the TGF-β1 and p38 pathway. The role of TGF-β1 in regulating expression of ank was examined by measuring the expression upon pretreatment with TGF-β1 and SB431542, a selective activin receptor-like kinase (ALK) receptor inhibitor. The role of p38 in regulating expression of ank upon CCMT stimulation was investigated by studying the expression of phosphorylated p38 with pre-treatment of SB203580, a specific inhibitor of p38 kinase, and prior to CCMT stimulation.

Materials and Methods

Chondrocytes isolation and culture

Primary chondrocytes were isolated from...
the lumbar spine endplate cartilage of Sprague-Dawley rats (160-180g). Cartilage of the L1-L5 endplates was carefully removed from the vertebrae and minced into small pieces (<0.03mm³). The isolation method and culture conditions of chondrocytes were used in Musumeci’s and our papers.13,14 The second passage cells were used for experiments.15 The study was carried out in strict accordance with the recommendation of the Guide for the Care and Use of Medical Laboratory Animals (Ministry of Health, P.R. China). This study protocol was approved by the Medical Laboratory Animals Care and Use Committee of Anhui Province and the Ethics Committee of Yijishan Hospital of Wannan Medical College and in accordance with the guideline for the Chinese ethical conduct in care and use of animals.

Application of cyclic strain

Endplate chondrocytes were plated at the density of 1 ¥ 10⁵ cells/cm² in 2 mL of medium on six-well flexible silicone rubber BioFlex™ plates coated with collagen type I (Flexcell International Corporation, Hillsborough, NC, USA).16 The cells were cultured for 48 h to allow them to attach and reach 80-90% confluency, at which time the growth medium was replaced, and then mechanical strain was applied. A continuous cyclic mechanical strain at 0.5 Hz sinusoidal curve at 3%, 6%, or 9% elongation was applied using an FX-4000T™ Flexercell® Tension Plus™ unit (Flexcell International Corporation) according to the method elaborated in our previous study.9 The cultures were incubated in a humidified atmosphere at 37°C and 5% CO₂ while stretching. Cells were harvested immediately after CCMT stimulation was applied. NC group means no mechanical strain was applied.

Live/Dead cell viability assay

A LIVE/DEAD viability/cytotoxicity kit (Invitrogen, Carlsbad, CA, USA) was used after the application of CCMT and removal of supernatant to confirm that the NC group and the CCMT group endplate chondrocytes remained viable and adherent. The kit was used following manufacturer specifications.

Real time PCR

Total RNA was isolated using Tirol reagent (Invitrogen) according to the manufacturer’s instructions. After reverse transcription reaction, real time PCR (RT-PCR) was performed by a Roche Light Cycler 480 system using SYBR®Premix Ex Taq™ (Takara, Dalian, China) according to the manufacturer’s instructions. The sequences of genes are shown in Table 1. All RT-PCR data were normalized to the GAPDH gene for quantitative comparisons.

ELISA

The NC and CCMT groups’ chondrocytes supernatant were collected. ELISA (Bender Med Systems, Vienna, Austria) for TGF-β1 was performed using standard protocols.

CCMT-induced ank expression leads to ePPi generation

Rat endplate chondrocytes were transfected with siRNA (ank) 24 h before CCMT stimulation, and then stimulated for 20 min of 3% CCMT. Total RNA was extracted from rat endplate chondrocytes and real-time PCR analysis was performed.

Figure 1. CCMT increased the ank gene expression in rat endplate chondrocytes. Chondrocytes were treated with CCMT at 3%, 6%, and 9% elongation for 10 min (A), 20 min (B) or 40 min (C) and the mRNA expression of ank was then measured. The protein expressions of ank were also assessed (D). The expression phase of ANK mRNA and protein after 3% CCMT. Statistically significant differences were indicated in (E). GAPDH expression was used as an internal control. GAPDH protein level acted as a loading control for Western blotting. All experiments were repeated at least three times. The columns represent the mean ± SE. *P<0.05, **P<0.01 vs nonloading.

Table 1. Sequences of primers used in the real time PCR.

| Genes   | Forward primer | Reverse primer | Accession number | Product length(bp) |
|---------|----------------|----------------|------------------|--------------------|
| TGF-β1  | CATCCATGACATGCAACGGACCCTT | ACAGAAGTTGCGATGTAACCTT | NM_021578.2 | 220 |
| Ank     | CAAGAGAGACAGGGCCAAAG | AAGGCAGCGAGATACAGGAA | NM_053714.1 | 177 |
| GAPDH   | CTCAACTACATGCTCATACTGTTCCA | CTTCGCCATTTCTAGCCTTGACT | NM_017008.3 | 81 |
Western blotting analysis and protein kinase inhibitor assay

Cells were lysed on ice for 30 min in lysis buffer containing 50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1% Nonidet P-40, and 0.1% SDS supplemented with protease inhibitors (10 µg/mL leupeptin, 10 µg/mL pepstatin A, and 10 µg/mL aprotinin). For western blotting analysis, 15 µg of sample was resolved on 12% SDS-PAGE and electro-transferred onto nitrocellulose membranes (Whatman, Piscataway, NJ, USA). The primary antibodies used were anti-ERK1/2 (rabbit monoclonal anti-p44/42 MAPK, Cell Signaling Technology, Inc., Danvers, MA, USA); anti-p38 (rabbit monoclonal anti-p38 MAPK, Cell Signaling Technology); anti-SAPK/JNK (rabbit monoclonal anti-SAPK/JNK, Cell Signaling Technology); anti-Phospho-p38 [rabbit monoclonal anti-phospho-p38 MAPK (Thr180/Tyr182), Cell Signaling Technology]; anti-phospho-ERK1/2 [rabbit monoclonal anti-phospho p44/42 MAPK (Thr202/Tyr204), Cell Signaling Technology]; anti-phospho-SAPK/JNK [rabbit monoclonal anti-phospho-SAPK/JNK (Thr183/Tyr185), Cell Signaling Technology]; anti-SMAD2/3 (Smad2/3 Antibody, Cell Signaling Technology) at a dilution of 1:1000; anti-P-SMAD2 (Phospho-Smad2 (Ser245/250/255) Antibody, Cell Signaling Technology) at a dilution of 1:1000; anti-TGF-β Receptor II [TGF-β Receptor II (K105) Antibody, Cell Signaling Technology] at a dilution of 1:1000; anti-smad7 (ab90085, Abcam, Cambridge, MA, USA) at a dilution of 1:1000; anti-ank (ab58698, Abcam) at a dilution of 1.25:1000. For normalization of protein loading, GAPDH (Cell Signaling Technology) antibody was used at 1:5000 dilutions. Infrared labeled secondary antibodies goat antibodies anti-rabbit IRDye 800 (Li-Cor Biosciences, Lincoln, NE, USA) was added to bind to the primary antibody. The bound complex was detected using the Odyssey Infrared Imaging System (Li-Cor Biosciences). The images were analyzed using the Odyssey Application Software, version1.2 (Li-Cor Biosciences) to obtain the integrated intensities. For the kinase assays, endplate chondrocytes were pretreated with 5-50 µM of the specific inhibitor of p38 kinase (SB203580, InvivoGene, San Diego, CA, USA) and a selective ALK receptor inhibitor (SB431542, Sigma, St. Louis, MO, USA) for 30 min and subsequently exposed to CCMT (0.5Hz, sin3%) for 20 min. The expression levels of ank were analyzed by real time PCR and Western blot.

Silencing experiments with small interfering RNA

The siRNA sequences (designed by Shanghai GenePharma Co.Ltd) and were ank sense 5’-CUGGCCAACACGAACAAC-3’ and antisense 5’-UGUUGUUCGUGUUGGCCAG-3’ were used at final concentrations of 50 nM. Ank siRNA transfection after CTS stimulation 20 min (0.5Hz, sin3%). Briefly, siRNA and lipofectamin2000 were diluted separately in serum free medium, and then diluted lipofectamin was added to siRNA. After 20 min incubation at room temperature, cells were washed with PBS.
and incubated for 4 h with siRNA-lipofectamin mix. Then, after disposal of the mix, a medium containing 10% FBS was added to that culture dish. Two days later, Ank mRNA expression was used as a negative control to check for the specificity of siRNA effects.

**Radiometric assay for extracellular inorganic pyrophosphate**

The ePPi levels were measured using the differential adsorption of UDP-(6-3H) glucose (GE Healthcare), and its reaction product 6-phospho-(6-3H) gluconate on activated charcoal, as previously described. The standard concentrations, ranging from 10 to 400 pmol of PPi, were included in each assay. After adsorption of the reaction mixture on charcoal, and centrifugation at 14,000 rpm for 10 min, 100 μL of the supernatant was removed carefully and counted for radioactivity in 5 mL of BioSafe II (Research Products International Corp, Mt. Prospect, IL, USA). Results were expressed as picomole of ePPi per microgram of total cell proteins.

**Transfection of SMAD7 expression plasmid**

Cells were exposed to each biochemical agent for 1 h before 3% CCMT stimulation. SMAD7/pcDNA3.1(+) expression plasmid, which contained rat SMAD7 coding sequence was transiently transfected into end-plate chondrocytes by lipofectamine2000 (Invitrogen). After 24 h post-transfection, cells were exposed to mechanical loading.

**Statistical analysis**

As to the data presented for Figures 1-5, statistical comparisons were made by performing Student’s t-test to check differences between non-loading (NC) and CCMT groups. Statistical comparisons were made by performing ANOVA with SPSS (SPSS, Inc., Chicago, IL, USA version 10.0), followed by a Student-Newman-Keuls post-test in Figures 1 and 3-5. A P value <0.05 was considered as statistically significant.

**Results**

The CCMT increased the ank gene expression in rat endplate chondrocytes. To confirm that CCMT did not cause cell death of endplate chondrocytes, we investigated the viability of endplate chondrocytes by Live/Dead assay after exposure to CCMT. Our results showed that the NC and CCMT group endplate chondrocytes remained adherent, with no change in viability following the application of CCMT (Figure 6). We applied three levels of elongation of CCMT to test whether different levels had the same effect on the ank gene expression of endplate chondrocytes. RT-PCR showed the up-regulation mRNA level expression of ank in 3%, 6%, and 9% groups after loading for 10 min (Figure 1A), 20 min (Figure 1B) and 40 min (Figure 1C). As the 3% elongation group showed the largest increase in the expression of ank mRNA, it was used in the subsequent experiments. Western blot (Figure 1D) showed the up-regulation of ANK protein expression in 20 min and 40 min groups. At last, we analyzed the expression phase of ANK mRNA and protein after 3% CCMT. As shown in Figure 1E, after 3% CCMT treatment, ANK mRNA expression increased at 10 min, which peaked at 20 min, and then decreased at 40 min. Besides, 3% CCMT treatment upregulated ANK protein at 20 min, and maintained at a high level until 40 min. Because both ANK mRNA and protein increased to the same level, so we chose the 3% elongation and 20 min group in the subsequent experiments.
CCMT-induced ank expression leads to ePPi generation

The effect of ANK on ePPi levels was assayed in the rat end plate chondrocytes transfected with ank siRNA after CCMT stimulation. Data showed that ANK siRNA could effectively knockdown ANK expression (Figure 2A) and down-regulate CCMT-induced ePPi expression (Figure 2B).

CCMT increased TGF-β1 expression

To examine if TGF-β1 was involved in the regulation of ank upon CCMT stimulation, we examined the expression of TGF-β1. Our results showed that there was a significant increase in the TGF-β1 expression of both mRNA (Figure 3A) and protein (Figure 3B) levels in the CCMT group compared to those in NC group.

CCMT increased ank expression via the TGF-β1 pathway

Western blot analysis showed that the expression levels of TGF-βR up-regulated after 3% CCMT stimulation (Figure 4A). Biochemical studies have shown that SMAD7 blocks signal transduction of TGF-β. To determine whether the regulation of ank by TGF-β1 upon CCMT stimulation was dependent on SMAD2/3, endplate chondrocytes were transfected with SMAD7/pCDNA3.1(+) or pCDNA3.1 vector. Western blot analyses showed that the expression levels of SMAD7 up-regulated in endplate chondrocytes transfected with SMAD7/pCDNA3.1(+) (Figure 4B). After 1 day’s transfection, cells were exposed to 3%CCMT, cells with SMAD2 expression showed lower p-SMAD2 expression than vector transfected group after 3% CCMT stimulation, but the p-P38 and Ank expression had no significant changes (Figure 4C). Endplate chondrocytes were pretreated with TGF-β1 or SB431542 for 30 min prior to CCMT and then were treated by TGF-β1 or SB431542 during CCMT stimulation. Date showed that the ank expression in mRNA (Figure 4E,G) and protein (Figure 4D,F) levels significantly increased when treated by TGF-β1 in a dose-dependent manner, and decreased when treated by SB431542 compared with the NC group.

CCMT-stimulated phosphorylation of MAPKs in endplate chondrocytes

Phosphorylation of p38 at 20 min after CCMT increased compared to the NC group (Figure 5A), phosphorylation of p44/42 and JNK did not increase after CCMT. The effects of MAPK inhibitors on proteinase expression were analyzed by RT-PCR (Figure 5C) and protein (Figure 5D). SB203580 (10 μM) down-regulated CCMT-induced ank expression; U0126 and JNK inhibitor II did not influence ank induction (data not shown); and SB431542 (10μM) down-regulated CCMT-induced p-p38 expression (Figure 5B). Above data suggest that CCMT may activate the p38 pathways in endplate chondrocytes.

Discussion

The ANK is a multipass transmembrane protein thought to serve either as an anion channel or as a regulator of such a channel. Previous studies showed that in ank−/− mice, loss of ANK activity results in diminished ePPi
levels and consequent basic calcium phosphate (BCP) calcification in joints. The autosomal recessive ank mutation caused spontaneous bony (BCP) ankylosis of peripheral and axial joints, subsequent destructive arthritis, osteophyte formation, and death owing to immobility.6,7 We also found that the change of ank expression is intimately related with endplate chondrocytes calcification in our previous study.2

Previous studies also showed that intermittent mechanical tension promotes osteogenesis of mesenchymal stem cells and osteo-progenitor cells.10,18 Our previous work showed that ICMT could induce the calcification of endplate chondrocytes and long-term mechanical stimulation decreased the ank gene expression.3 Mechanical cervical traction is an intervention that is often recommended for the treatment of patients with neck pain.19 Previous study demonstrates that short-term mechanical cervical traction is beneficial for neck pain. CCMT increases the ank expression as opposed to ICMT stimulation results. It indicates that short-term mechanical cervical traction has a strong function on anti-calcification through up-regulation of ank expression.

TGF-β1 plays an essential role in the regulation of ank gene expression. It has been reported that TGF-β1 increases ank gene expression through Ras/Raf-1/extracellular signal-regulated kinase pathways in chondrocytes and that TGF-β1 is immediately released after mechanical tension stimulation.21,22 A number of mechanisms through which mechanical forces induce proximal biological changes in chondrocytes have been proposed. A recent study found that integrins αvβ3, αvβ5, αvβ6, and αvβ8 appear to change the conformation of the latent TGF-β1 complex by transmitting cell traction forces.20 External mechanical tensile loads transduced by ECM can stimulate NIH3T3 cells through integrins, which then activates multiple downstream signaling pathways, including the TGF-β1 pathway,23 resulting in cellular changes such as growth, adhesion, migration, ECM synthesis, and proliferation.22,24

Some studies indicate that the dependent variable of human is probably 1000 μstrain in physiological status, and no more than 4000 μstrain in vigorous activity; 1000 μstrain is equal to 1% elongation, and 6% and 9% elongation mechanical strain is unacceptable for human.24 We choose therefore the 3% mechanical strain. Our results showed that 3% CCMT stimulation increases the expression of both ank and TGF-β1. Our data also showed that compared with an NC group, ank mRNA and protein levels significantly increase in a dose-dependent manner when cells are treated with TGF-β1 and decrease in a dose-dependent manner when cells are treated with SB431542. These results suggest that during CCMT, TGF-β1 alteration can regulate ank gene expression. Many down-stream signaling molecules are regulated by TGF-β1. Among these molecules is the MAPK family, which is known to regulate ank gene expression25 and is a central conduit through which mechanical forces are transduced into biological responses in cartilage.26 MAPK pathways are activated by the static and dynamic compression of cartilage, which simultaneously induce intratissue fluid flow, pressure gradients, cell and matrix deformation.27

In the current study, we investigated the inhibitory effects of MAPK on ank expression. CCMT phosphorylated p38 MAPK, which implies that CCMT activates the p38 pathways in endplate chondrocytes. SB203580 could attenuate the induction of both ank mRNA and protein level expression. In the downstream mechanism of TGF-β1-induced ank up-regulation, p38 might play a central role in the response to CCMT. SB431542 could decrease CCMT-induced p-p38 expression. In our experiments, we found that when the p38 inhibitor SB203580 was added, Ank protein expression was higher than that in the control group. However, when the TGFβ1 receptor inhibitor SB431542 was added, Ank protein expression was still higher than that in the control group, suggesting that the TGFβ1 and p38 pathways may have a synergetic effect on Ank expression. CCMT-induced ank expression may occur through both the TGF-β1 and p38 signaling pathways. Dynamic shear-activated p38 at later time points and static compression caused transient activation of p38, with maximal phosphorylation occurring at 10 min. ERK1/2 pathway activation was similar to p38 pathway activation, with peak phosphorylation occurring at the earliest time points (10-60 min). Static compression stimulated JNK phosphorylation at the relatively late time point of 1 h;27 p38 may be differentially sensitive to CCMT compared with ERK1/2 and JNK. Our experiments confirmed that 3% CCMT can up-regulate the expression of TGF-β1 receptors. The overexpression of Smad7 obviously inhibited Smad2/3, whereas the p-p38 and Ank expression levels were not affected. This finding indicates that p38 activation does not depend on the Smad2/3 pathway and that there may be other pathways that are responsible for these changes.

In this study, ank expression was highly up-regulated by CCMT-activated TGF-β1, which subsequently activated p38, at 20 min, but not ERK or JNK. This discrepancy might be partly explained by the different cell types, time course of mechanical stimulation and mechanical strain (tensile or compressive) used in these in vitro experiments.25-28 We found that ICMT decreases ank expression in our previous study; however, CCMT increased ank expression in this study. These contrasting effects of mechanical loads may be due in part to different temporal dynamics and MAPK activation magnitudes, which may have led to load-dependent activation of downstream transcription factors.

Study limitations

There were several limitations in the current study. First, in the article, we speculated that CCMT increased the TGF-β1 expression through integrins pathway. Unfortunately, the expression of integrins upon CCMT was not detected. Second, CCMT may regulate the ank gene expression via multiple signaling pathways. However, we only study the TGF-β1-MAPK signaling pathway changes. Third, it is still unknown whether our results with endplate chondrocytes in vitro could represent the biologic behavior of human endplate chondrocytes against mechanical stress in vivo.

In conclusion, the results of the current study demonstrate that CCMT-induced ank gene expression is regulated by the TGF-β1 and p38 MAPK pathways in endplate chondrocytes. Since improving the function of endplate can contribute to postpone the intervertebral disc degeneration process, this information may provide a new approach to treat the intervertebral disc degeneration by appropriate mechanical tension.

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