Microfluidics-based optimization of neuroleukin-mediated regulation of articular chondrocyte proliferation

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Abstract. Due to the low proliferative and migratory capacities of chondrocytes, cartilage repair remains a challenging clinical problem. Current therapeutic strategies for cartilage repair result in unsatisfactory outcomes. Autologous chondrocyte implantation (ACI) is a cell based therapy that relies on the in vitro expansion of healthy chondrocytes from the patient, during which proliferation-promoting factors are frequently used. Neuroleukin (NLK) is a multifunctional protein that possesses growth factor functions, and its expression has been associated with cartilage development and bone regeneration, however its direct role in chondrocyte proliferation remains to be fully elucidated. In the current study, the role of NLK in chondrocyte proliferation in vitro in addition to its potential to act as an exogenous factor during ACI was investigated. Furthermore, the concentration of NLK for in vitro chondrocyte culture was optimized using a microfluidic device. An NLK concentration of 12.85 ng/ml was observed to provide optimal conditions for the promotion of chondrocyte proliferation. Additionally, NLK stimulation resulted in an increase in type II collagen synthesis by chondrocytes, which is a cartilaginous secretion marker and associated with the phenotype of chondrocytes. Together these data suggest that NLK is able to promote cell proliferation and type II collagen synthesis during in vitro chondrocyte propagation, and thus may serve as an exogenous factor for ACI.

Introduction

Due to the lack of vascularization in articular cartilage and the low proliferative and migratory capacities of chondrocytes, cartilage remains a challenging tissue to repair (1-4). Current clinical treatment strategies involve mosaicplasty, micro-fracture, periosteum or perichondrium transplantation and fresh osteochondral allograft implantation (5). These strategies appear promising, however the long term outcomes are unsatisfactory (3,6,7). Tissue engineering has provided alternative possibilities for hyaline cartilage repair using cell based therapy, utilizing chondrocytes or adult stem cells combined with synthetic substrates and bioactive factors to prepare for the functional replacement of hyaline cartilage (8-10). Bioactive factors have been widely utilized in cell based therapy to promote cell proliferation and the production of the extracellular matrix, however current therapeutic options are far from optimal and the anticipated outcomes are rarely achieved (11). Overall, it remains to be fully understood how improvements in chondrocyte amplification may be achieved with the maintenance of the chondrocytic phenotype. The aim is to generate active and phenotypically stabilized tissue engineered cartilage in order to treat cartilage lesions.

Neuroleukin (NLK), a neurotrophic factor of spinal and sensory neurons, is additionally a growth factor in mouse salivary glands, and promotes the survival of peripheral and central neurons in culture. NLK is additionally known as autocrine motility factor (AMF) and phosphoglucose isomerase (PGI) (12,13), where the peptide sequences of NLK, AMF and PGI indicate that they are the same protein, with the differing names assigned based on previous functional analyses. NLK is secreted by tumor cells and functions as a cytokine to promote migration, invasion and proliferation of tumor cells (14-16). This paracrine function of NLK is further supported by studies on the migration of vascular endothelial cells which propagate and form new capillaries during tumor angiogenesis, in addition to the proliferation and migration of fibroblasts treated with NLK (17,18). Of note, Zhi et al (19) reported differential expression of NLK in osseous tissue and during differentiation, with elevated levels in superficial articular chondrocytes, proliferating chondrocytes, fibroblasts and osteoblasts within the fracture callus, however, NLF was absent in terminally differentiated hypertrophic chondrocytes or osteocytes. These studies suggest that NLK is involved in cartilage development and bone regeneration. In the current study, the effect of NLK on the proliferation of isolated rat articular chondrocytes was investigated in vitro and the working concentrations of NLK were optimized using a microfluidic device.

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Materials and methods

Antibodies and reagents. The following antibodies were used in immunofluorescent analyses: Mouse anti-collagen type II polyclonal antibody (1:4,000; cat. no. BA0533; Wuhan Boster Biological Technology, Ltd., Wuhan, China), Alexa Fluor® 488 goat anti-mouse secondary antibody (1:1,000; cat. no. A11001; Invitrogen Life Technologies, Carlsbad, USA). Human recombinant NLK/PGI full length protein was obtained from Abcam (Cambridge, UK; cat. no. ab87625). Trypsin was purchased from Gibco Life Technologies (Carlsbad, USA), collagenase type II from Sigma-Aldrich (St. Louis, MO, USA), and the Cell Cycle Detection kit was from Nanjing KeyGen Biotech Co., Ltd. (Nanjing, China; cat. no. KGA511).

Chondrocyte isolation and cell culture. All experimental procedures were approved by the Institutional Committee of Animal Use and Protection (Dalian Medical University, Dalian, China). A total of 12 Sprague-Dawley rats were used in the present study, which were provided by the Experimental Animal Center of the Dalian Medical University. The rats were anesthetized with 3.6% chloral hydrate (1 ml/100 g; Kermel, Tianjin, China) by intraperitoneal injection, prior to sacrifice by cervical vertebra dislocation. The surrounding muscles of the knee joints were removed to expose the articular cartilage, which was collected using curved tissue scissors. Cartilage was separated from the femoral heads and femoral condyles of 1 month-old male Sprague-Dawley rats and cut into 1.5 mm³ sections using a surgical blade in sterile phosphate-buffered saline (PBS; Beijing Solarbio Science & Technology Co., Ltd., Beijing, China). Primary chondrocytes were isolated by digestion with 0.2% type II collagenase for 4 h at 37°C in an agitating water bath and resuspended by cervical vertebra dislocation. The microfluidic device was designed by techniques involving SU-8 photolithography and polydimethylsiloxane (PDMS) soft lithography. The transparency mask was designed by AutoCAD 2014 (Autodesk, Inc., San Rafael, CA, USA) and was used in 1:1 contact photolithography with an SU-8 photoresist consisting of patterned photoresist on a silicon wafer. Positive replicas with embossed channels were fabricated by molding PDMS against the master. The PDMS replica and a clean glass substrate were irreversibly sealed using oxygen plasma (2 Torr, 100 W) for 1 min.

Microfluidic chip operation. The inlets of the device were connected to a syringe pump (model LSP02-1B; Baoding Longer Precision Pump Co., Ltd., Baoding, China) to drive fluid flow. The outlet of the device was connected to a culture medium reservoir containing DMEM/F12. Culture media with and without NLK were simultaneously infused into the microfluidic device from the two inlets, with 2% serum containing culture medium into the left and 2% serum medium with 30 ng/ml NLK into the right. The concentration gradient of NLK was established within 30 sec. The flow speed was controlled at 0.1 ml/min. The device was stored in an incubator at 37°C with 5% CO₂ during the culture period.

CGG performance validation. By using this chip, solutions from inlets were repeatedly split at the nodes, combined with neighboring streams in a laminar fashion, and mixed by diffusion in serpentine channels. As a result, the solutions were continuously diluted and a series of concentrations was produced in the outlets of the CGG. In the current study, solution A (concentration 0) and solution B (concentration X) were injected into the CGG via the 2 inlets simultaneously, and the concentrations in the 8 outlets were as follows: 0, 1/7X, 2/7X, 3/7X, 4/7X, 5/7X, 6/7X and X. Fluorescein isothiocyanate dextran (FITC-dextran) with a molecular weight of 20,000 Da (Sigma-Aldrich) was used as a probe for the CGG performance validation according to a previous study (18). The fluorescence intensity of FITC-dextran at the 8 outlets of the CGG were imaged by confocal laser scanning microscopy (model TCSSP5; Leica Microsystems GmbH, Wetzlar, Germany) and quantified using Image-Pro Plus 6.0 software (Media Cybernetics, Inc., Rockville, MD, USA). The intensities were compared with the theoretical values achieved by the equation, and correlation factors were calculated. All the experiments were repeated three times.

Cell cycle analysis. Cell cycle phase distribution was determined by flow cytometry using propidium iodide (PI; Nanjing KeyGen Biotech Co., Ltd.) staining. For cell synchronization, cells were starved for 12 h prior to NLK treatment. Following 24 h culture in 2% FBS medium and 24/48 h NLK treatment, cells were harvested, fixed in 70% ice-cold ethanol for 12 h at 4°C, stained in 50 µg/ml PI supplemented with 1 mg/ml RNase (Nanjing KeyGen Biotech Co., Ltd.) and 0.1% Triton X-100 (Sigma-Aldrich), and analyzed by flow cytometry (Accuri C6; BD Biosciences, Franklin Lakes, NJ, USA). The histogram was used to present the percentage of cells in the G₁, S and G₂/M phases. Cell cycle distribution was analyzed by FlowJo 10.0.6 software (FlowJo, LLC., Ashland, OR, USA).

Immunofluorescence. Cells were seeded (0.8x10⁵ cells/well) on sterile glass coverslips (0.8x0.8 cm) and cultured in 6-well plates until 70% confluent. Cells were washed twice with PBS, fixed with 4% formaldehyde (Beijing Solarbio Science & Technology Co., Ltd.) for 15 min and permeabilized with 0.1% Triton X-100 for 15 min, prior to blocking with 5% bovine serum albumin (Beijing Solarbio Science & Technology Co., Ltd.) in PBS for 1 h at room temperature. Cells were incubated with primary antibodies overnight at 4°C. Subsequently, cells were incubated with fluorescent secondary antibodies at room temperature for 1 h prior to the addition of DAPI. The nuclei were stained and visualized using a confocal microscope (Nikon A1R Si, Tokyo, Japan). The inlets of the device were connected to a syringe pump (model LSP02-1B; Baoding Longer Precision Pump Co., Ltd., Baoding, China) to drive fluid flow. The outlet of the device was connected to a culture medium reservoir containing DMEM/F12. Culture media with and without NLK were simultaneously infused into the microfluidic device from the two inlets, with 2% serum containing culture medium into the left and 2% serum medium with 30 ng/ml NLK into the right. The concentration gradient of NLK was established within 30 sec. The flow speed was controlled at 0.1 ml/min. The device was stored in an incubator at 37°C with 5% CO₂ during the culture period.
staining with 4',6-diamidino-2-phenylindole and imaging using a fluorescence microscope (model BX53; Olympus Corporation, Tokyo, Japan).

**Observation of chondrocyte morphology.** After six days of culture in a microfluidic device (designed and fabricated by Dalian University of Technology, Dalian, China) with various concentrations of NLK (0, 4.28, 8.57, 12.85, 17.15, 21.42, 25.71 and 30 ng/ml), total chondrocyte numbers were counted from several fields and morphological changes of the chondrocytes were observed using a phase contrast microscope (Olympus cx31; Olympus Corporation). Micrographs were captured from the cell culture chambers of the eight groups.

**Results**

**Microfluidic device design and gradient generation.** Dilution networks and micro-scale cell culture chambers were incorporated into a single microfluidic device, which is capable of producing multiple gradient concentrations for on-chip cell culture assays. A series of solutions with different concentrations was formed at the outlets of the CGG (0, 4.28, 8.57, 12.85, 17.15, 21.42, 25.71 and 30 ng/ml; Fig. 1A). The fluorescent probe FITC-dextran was used as an indicator for estimating the gradient produced by the CGG. From the two inlets, FITC-dextran was simultaneously infused into the CGG by a syringe pump and the resulting gradients (Fig. 1C) were imaged and quantified. A total of eight FITC-dextran gradient profiles were produced from gradient generators spanning the two inlet concentrations (Fig. 1B). The fluorescence intensities of FITC-dextran in the junctions between the CGG and the cell culture chambers were observed using a phase contrast microscope (Olympus cx31; Olympus Corporation). Micrographs were captured from the cell culture chambers of the eight groups.

**Microfluidic analyses of chondrocyte proliferation with NLK.** To evaluate the effect of NLK on chondrocyte proliferation, isolated articular chondrocytes from Sprague-Dawley rats were applied to the above-mentioned microfluidic chip with an NLK concentration gradient. Following 6 days of culture, the morphology of chondrocytes was imaged using phase contrast microscopy (Fig. 2), and a dose-dependent proliferative profile was observed following treatment with NLK between 0 and 12.85 ng/ml (Fig. 3A). Culture medium with 2% serum was used as the control. Among the eight concentrations (0, 4.28, 8.57, 12.85, 17.15, 21.42, 25.71 and 30 ng/ml), the proliferation rate plateaued at 12.85 ng/ml. The 12.85 ng/ml NLK group exhibited a 3.96-fold increase in chondrocyte proliferation.
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proliferation (Fig. 3A), with a statistically significant difference compared with the control group (P<0.01). However, there was no statistically significant difference between the 12.85 ng/ml NLK group and the higher concentrations (P>0.05). The growth curve of the chondrocytes following 12.85 ng/ml NLK stimulation is presented in Fig. 3B.

**NLK increases the proportion of proliferative state chondrocytes.** To further characterize the pro-proliferative effect of NLK on chondrocytes, flow cytometry was conducted to assess cell cycle distribution. Chondrocytes were starved with serum-free medium for 12 h to synchronize the cells, followed by treatment with 2% FBS for 24 h and 12.85 ng/ml NLK with 2% FBS for 24 and 48 h. Cells were collected and analyzed post treatment. As presented in Fig. 4A, an accumulation of NLK-treated cells in the S phase of the cell cycle was observed compared with the control cells at 24 and 48 h. In addition, the G2/M phase distribution was increased following NLK treatment. Quantification indicated that the percentage of sub G1 phase cells was reduced from 82 to 61%, and the proportion of cells in S and G2/M phases was significantly greater in NLK-treated cells (P<0.01; Fig. 4B). These data further indicate that NLK promotes proliferation of articular chondrocytes in vitro.

**NLK stimulates type II collagen synthesis in chondrocytes.** As demonstrated above, NLK is able to serve as a bioactive molecule for chondrocytes. Therefore, it was investigated whether NLK is involved in phenotype maintenance of chondrocytes. To determine the response of chondrocytes to...
Figure 4. NLK increases the proportion of proliferative chondrocytes. Chondrocytes were treated with NLK for 24/48 h, 24 h 2% fetal bovine serum treatment was used as the control. Cells were stained with propidium iodide, prior to flow cytometric analysis. (A) Representative histograms from three independent experiments. Cell cycle distribution was analyzed by FlowJo software. (B) Percentage of cells in G0/G1, S, and G2/M. NLK treatment led to a reduction of cells in the G0/G1 phase, however an increase of cells in the S and G2/M phases. Quantification from 3 independent experiments. Values presented are the mean ± standard deviation. *P<0.05 and **P<0.01 vs. control. NLK, neuroleukin.

Figure 5. NLK stimulates type II collagen synthesis in chondrocytes. (A) P0, P1 and P2 chondrocytes were cultured with or without NLK (12.85 and 25 ng/ml) and type II collagen expression was detected with immunofluorescence. (B) The percentage of type II collagen positive chondrocytes was calculated. NLK stimulates collagen II synthesis in P1 and P2 cells compared with the control. Quantification from 3 independent experiments. Values presented are the mean ± standard deviation. **P<0.01 vs. control. Scale bar = 100 µm. NLK, neuroleukin; P, passage.
NLK on type II collagen synthesis during in vitro culture. Immunofluorescence analysis was conducted. Chondrocytes were examined from passage 0 (P0; 24 h following isolation) to passage 2 (P2; 6-7 days following isolation) by staining with the type II collagen antibodies. As presented in Fig. 5A, following 12 h serum-starvation, P0, P1 and P2 chondrocytes were cultured with or without NLK (12.85 and 25 ng/ml), and cells with positive type II collagen stains were counted. A reduction in the fluorescent signal of collagen II was observed during prolonged culture in vitro due to the dedifferentiation of primary chondrocytes, characterized by a change from the expression and secretion of Collagen I instead of Collagen II. A significant increase in collagen II positive cells was observed following NLK treatment in P1 (22-41%) and P2 (5-17%) groups compared with the control (without NLK). However, no significant difference was observed in P0 or between NLK treatment groups (Fig. 5B). A reduction in the fluorescent signal was observed during prolonged culture in vitro due to dedifferentiation of primary chondrocytes.

Discussion

Clinical management regarding cartilage repair may be classified into three main categories: Marrow stimulation-based, osteochondral transplantation, and cell-based repair techniques (5). ACI is a cell-based tissue regeneration technique, representing a novel biological approach for the treatment of cartilage defects (21-23). However, due to dedifferentiation of autologous chondrocytes during expansion in vitro and following implantation, ACI may subsequently result in scar tissue akin to fibrocartilage (22,24). Therefore, studies have attempted to combine ACI with biomaterials of natural or synthetic origin as scaffolds, together with the use of bioactive factors providing proliferative stimuli, in order to reconstruct a microenvironment in vitro which allows the cells to grow as in their native tissue (25,26). Bioactive factors function as key components in cell-based therapy. Growth factors that exist in or are secreted by bone marrow mesenchymal stem cells (BMSCs) and chondrocytes such as bone morphogenic protein, vascular endothelial growth factor and transforming growth factor-β do not only upregulate chondrogenic markers, however, additionally are associated with ossification (27-29). This may lead to the formation of fibrocartilage with inferior mechanical properties and limited durability (29,30).

NLK is a multifunctional protein with intra- and extracellular functions (13,31,32). It has been studied in neoplastic cells and their normal counterpart cells for the expression, secretion and distribution, suggesting that a wide range of normal and neoplastic cells express and secrete NLK (33-35). Despite the extensive investigation of the pro-metastatic effects in tumor cells, the role of NLK in various normal tissues remains to be fully elucidated. The expression of NLK has been reported to be associated with cerebellar and hippocampal neurons, which are implicated in memory and learning (36,37). NLK is reported to stimulate migration, propagation and new capillary formation for human umbilical vein endothelial cells (37,38). In addition, fibroblasts gain increased proliferative and migratory abilities following treatment with NLK (17). These results provide evidence that NLK may act as a secreted factor to promote cell motility and proliferation in normal tissues. In a previous study, differential mRNA display was used and RNA populations isolated from osseous tissues were compared, from which it was reported that NLK was involved in cartilage development and bone regeneration (21). A high level of NLK expression was identified in superficial articular chondrocytes in 1, 4 and 8-month-old normal mice, in addition to in proliferating chondrocytes. However, in hypertrophic chondrocytes NLK is not detected (19). Furthermore, NLK expression was augmented during fracture healing. From day 3 post fracture, strong NLK labeling was detected in osteoblasts of the newly formed trabecular bone, proliferating chondrocytes of the soft callus and fibrous periostium. In the late stage of fracture healing, immature osteocytes and proliferating chondrocytes had differentiated towards mature osteocytes and hypertrophic chondrocytes, when the expression of NLK was reduced (19). The spatial and temporal expression of NLK in cartilage and osteoblast development indicates that NLK serves a role during cartilage development.

Microfluidics, known as “lab-on-a-chip”, brings the benefits of integration, miniaturization and automation to numerous research areas (39,40). As a novel cell culture platform, microfluidics provides spatial and temporal control of exogenous stimuli via the integration of various functional units (41). Furthermore, microfluidic devices have been successfully used for the long-term culture of cell lines and primary cells (42,43). By using microfluidic chips, experimental conditions are flexible and may be optimized by changing either the type of growth factor or the concentration of the input. Additionally, microfluidics make it possible to perform parallel experiments to optimize conditions whilst using a limited number of cells. A previous study reported data generated from a microfluidic platform, which included the effects of fluid mechanics on articular chondrocytes and BMSCs and the optimization of biofactor concentrations (20,44). As NLK is additionally implicated in tumor cell physiology, its overdosage during in vitro chondrocyte expansion may pose a risk of adverse side effects. In order to maximize its positive effects on chondrocyte proliferation while avoiding detrimental side effects, the current study sought to use microfluidics to optimize the concentration of NLK in chondrocyte in vitro propagation. Using a microfluidic device, 8 concentrations of NLK were generated precisely in 30 sec, and the effect of a range of concentrations of NLK on chondrocyte proliferation was evaluated over a 6 day period. A minimal concentration of 12.85 ng/ml NLK was demonstrated to affect chondrocyte growth, leading to a ~4-fold increase. This effect of NLK was further supported by the flow cytometry results of cell cycle analysis (Fig. 4).

Redifferentiation commonly occurs during in vitro expansion of chondrocytes, during which they gradually lose their typical round shape and acquire a spindly fibroblast-like form, with concurrent reductions in cartilaginous protein synthesis such as type II collagen (29,45,46). In the current study, immunofluorescence was conducted to examine the effect of NLK on the synthesis of type II collagen by chondrocytes. Treatment with NLK leads to a significant increase in the number of positively stained cells relative to untreated cells following in vitro passaging, indicating an inhibitory effect of NLK on dedifferentiation. The optimized concentration of 12.85 ng/ml was not significantly different compared with a
saturated concentration of 25 ng/ml, confirming the data from the microfluidic device.

In the present study, NLK was reported to act as an exogenous factor to promote the proliferation of articular chondrocytes during in vitro culture. A microfluidic device was used to evaluate a range of concentrations of NLK on chondrocyte proliferation, with a concentration of 12.85 ng/ml demonstrated to be optimal. Furthermore, this concentration of NLK may additionally prevent dedifferentiation of cultured chondrocytes in vitro. Future studies will aim to provide mechanistic insights into NLK regulation of chondrocyte growth, in addition to assessing the application of NLK during ACI.

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