Effect of synovial fluid insulin rate on mesenchymal stem cells differentiation to chondrocyte

Abstract

The demand for effective treatment strategies of cartilage cause to autologous MSC used for the repair of cartilage defects in patient. Tissue engineering has developed strategy for the repair and regeneration of a variety of tissues. In this study surveyed effect of some factors on MSCs differentiation to chondrocyte such as present of PCL scaffold and amount of synovial fluid. Total cellular RNA was extracted and was used for analysis by quantitative RT-PCR. Cell culture for differentiation assay performed in 6 well pellets by DMEM high glucose medium with supplement of 10% FBS and synovial fluid 5% and 10% of medium add to pellets for 21 days. The scaffold surface topography was captured by SEM electron microscope. MTT assay results in this study showed that the cells proliferation increased from 1st day to 2st days on electro spun Nano-fibrous scaffold. Cellular attachment and proliferation of MSCs on PCL substrate, and control group in the first day to 2st days. MTT assay and Real-time PCR analysis and the expression of chondrogenic specific genes at pellet in comparison with control group demonstrated cell survival ability and proliferation. It is suggested that this method can be used for induction of chondrogenesis.

Keywords: mesenchymal stem cells, insulin, synovial fluid, differentiation, chondrogenic

Introduction

In recent years, effective Clinical treatment of cartilage diseases has become more promising, especially with an increase in the number of patients with skeletal system defects. Hence, new therapeutic approaches have been developed to treat patients. HMSCs or Human Mesenchymal stem cells is found in bone marrow and other mesenchymal tissues, such as cartilage and bone. Recently autologous hMSC has been used for the purposes of healing and repairing patients with cartilage defects.

Tissue engineering combines different biological fields. Such as material sciences, engineering, and surgery to supply new functional tissues by living cells, signaling molecules, and biometrics. In recent decades, these policies have been greatly developed, and several groups of researchers focus on these therapeutic approach for repair and reconstruction of a different tissues.

Articular cartilage is one of the attractive targets for tissue engineering strategies, because it has been well proven that harm to articular cartilage, without direct access to a specified resource of repairing cells, is not self-healing. Some research efforts have led to the isolation of progenitor cells and understanding chondrogenic differentiation mechanisms that exist. Protocols for the Distinction and mitotic extension of adult human bone marrow-derived hMSC and the special, culture conditions extend for their chondrogenic differentiation.

Synovial fluid (SF) lubricates articular joint surfaces and nourishes the cells of articular cartilage and act as a shock absorber also contains many hormone such as growth factors like TGFβ-1. Hyaluronic acid (HA) based polymers is a principal component of SF, that it is secreted by synoviocytes, are protease inhibitors and glycoproteins. Poly (ε-caprolactone) (PCL) is one of the convenient synthetic biomaterials, because it has many comprehensive Specification like: flexibility, biocompatibility, mechanical strength and biodegradability. It also provides a suitable bio-substrate at the microenvironment level for seeding of the cell, but PCL has hydrophobic nature and this property cause to low cell loading in the initiatory step of the cell culture and its cause to less cellular adhesion, proliferation and migration.

Material and method

In this experimental study, the hMSCs obtained from the Cell Bank of Pasteur institute of Iran. 3-[4,5-dimethylthiazol-2-yl]-2, 5 diphenyl tetrazolium bromide MTT, (Sigma) FBS (Fetal Bovine Serum, Gibco), Pen/Strep (Gibco, 100U/ml), DMEM (Dulbecco’s Modified Eagle’s Medium, Gibco), Trypsin/EDTA 0.025% (Gibco,) were purchased. Synovium was obtained from the knee of arthritic patients undergoing arthroplasty from Loghman Hakim Hospital of Tehran.

Cell viability of hMSCs measured at 1and 2, days by MTT assay. The absorbance values of Formosan solutions were measured by ELIZA method at 570nm. The scaffold surface topography was captured by SEM electron microscope. Cell culture for differentiation assay performed in 6 well pellets by DMEM high glucose medium with supplement of 10% FBS by adding synovial fluid 5% and 10% of medium that changed twice a week at the condition of 95% air, 5% CO₂ at 37°C for 21days.

Total RNA extraction carried out by using the guanidine thiocyanate extraction method described by Chomczynski and Sacchi. These RNA preparations were used for analysis by RT-PCR. Relative quantification of mRNA expression of hyaluronan enzymes (has-2) and collagen type II was performed. These related Primer gene showed in below Table 1. cDNA was synthesized by using a high-
quality cDNA reverse transcription kit. The PCR reaction performed at 94°C 5min for denaturation, and 30 cycles at 94°C 45s, 60°C 45s, and 72°C 45sand final extension at 72°C for 5min. 1.5% agarose gel electrophoresis used for PCR products (3µl) analyzed containing safe stain, and the resulting bands were visualized by UV transillumination.

All the experiments were performed in triplicate and data analyzed by a Dunnet one way (ANOVA) by using software SPSS version 16.0 and the data reported as the mean value with the standard deviation of ± 0.05.15

**Table 1** Primer sequence for has-2, collagen type II and beta-actin gene

| Gene name          | Sequence                        |
|--------------------|---------------------------------|
| Has-2              | 5-CAGACAGGCTGAGGAGCAGCTTTAT-3   |
|                    | 5-GGATACATAGAAACCTCTCAACATGC-3 |
| Collagen type II   | 5-CCGCCGCTTCACCTACAGC-3         |
|                    | 5-TTTGTATTCAATCCTGCTTGC-3      |
| Beta-Actin         | 5-GGCACCCAGCACAATGAAG-3         |
|                    | 5-CCGATCCACACGGAGTTACCTTG-3    |

**Result**

Chemical and biochemical component of synovial fluid had measured by different method such as enzymatic, colorimetric and Eliza method that show in Table 2. These results show that synovial fluid as a supplementary medium contains requiring component for cell proliferation and differentiation.

**Table 2** Synovial fluid chemical and biochemical component

| Test             | Amount       |
|------------------|--------------|
| Glucose          | 41mg/dl      |
| Total protein    | 4.1mg/dl     |
| Albumin          | 1.7mg/dl     |
| Insulin          | 100µIu/ml    |
| Hyaluronic acid  | 2-4mg/ml     |

HMSCs cultured for 21 days on PCL scaffold at present and absent of CAP and different amount of synovial fluid carried out. MTT test for cell viability and mRNA expression for differentiation were assessed. MTT assay results in this study showed that the cells proliferation increased from 1day to 2days on electro spun nanofibrous scaffold. MTT test for different conditions of cell culture showed at Figure 1.

The image of the electron microscope from the PCL nanofibrous scaffold surface and the cellular connection to the scaffold surface is shown in Figure 2. The scaffold causes the cells seed better when they are cultured.

Cell proliferation and cellular attachment of hMSCs on PCL at the present of synovial fluid 10%, Higher rate than to the synovial fluid 5%, and. These results confirmed that increase in amount of synovial fluid has effect on cell proliferation, viability and differentiation due increase of insulin rate. For hMSCs, the cell proliferation gently increased on PCL nanofibrous from 1 to 2days.

RT-PCR analyses: An expression analysis for the known chondrogenic markers has-2 and collagen type II genes using RT-PCR was performed for hMSCs differentiation. This method showed that has-2 and collagen type II were induced during the course of hMSCs differentiation into chondrogenic lineage. In present of PCL substrate and adding 5% and 10% synovial fluid increased collagen type II expression at days 14 and 21, when compared with the cells cultured in the control groups. The expression of Collagen type II gene in primary days was low but increase piecemeal that show in Figure 3. Has-2 gene expression is lately and expressed at days 14 and 21 that show in Figure 4.
Discussion

Chondrogenesis can be made in several different ways that each of them has limitations. One of the complexities of chondrogenesis is the study of a simple laboratory system. HMSCs can be differentiate into multiple connective tissues, such as cartilage, ligament, muscle, bone, fat, and tendon. Several scientists have presented reports of cartilage differentiation of hMSCs in a pellet micro mass culture containing bioactive substances such as TGF–β, BMP–6 and dexamethasone. In this study We showed that human synovial fluid of the human’s knee has a potentially effective potential for chondrogenesis differentiation of hMSCs on PCL. The PCL scaffold as a surface for cell support provides conditions for Adhesion, growth, proliferation and differentiation of cells.

SF can acts as a sample for finding perfect combination of differentiation factors and deserves more research on its constituent parts and its potential for chondrogenesis.

RT-PCR analysis result for collagen 2 and Has-2 gene expressions at 14, 21days in 5% and 10% of medium synovial fluid show that collagen 2 gene expressed increased from 14 to 21days and Has-2 gene expressed at 14 and 21days. And 10% of synovial fluid has more effect on differentiation to chondrocyte at the present of PCL. HAS-2 is the major expressed isoform in chondrocytes and cartilage; there are appreciable levels of HAS-1 in contrast to reports by Nishida et al.

The results suggest the fact that SF Includes such factors as insulin and hyaluronic acid which enhance and expansion of cartilage of MSCs and progenitor cells. SF also includes the following subtypes of fibroblast Growth Factors (FGF), TGFβ, and Bone Morphogenic Proteins (BMP), which are capable of creating a reciprocal and mutual differentiation. Although, stimulation by the SF is not enough to detect HAS-2 or type II collagen or some aggrecan on protein level, But identifiable at mRNA level. SF has a great potential to persuade chondrogenic differentiation. Approximately in synovial fluid hyaluronic acid concentration is 2–4 mg/ml. Prolactin hormone is one of the effective factors in the human synovium that causes proliferation and chondrogenic differentiation of hMSCs and becoming cartilage.

SF also consist insulin in high concentration and TGF–β1 and also contains glucose and protein that cause to cell proliferation and differentiations. High concentration of insulin cause to glucose inters to the cells and initiate anabolic pathways and cells inter to proliferation cycle and differentiations state.

Conclusion

In this study, our obtained results from MTT assay, and the expression of chondrogenic specific genes in comparison with control groups demonstrated cell survival ability, proliferation and cell differentiation by synovial fluid on PCL. It is suggested that this method can be used for induction of chondrogenesis.

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None.

Conflict of interest

Author declares no conflict of interest.

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