Induction of chondrogenesis of human placenta-derived mesenchymal stem cells via heparin-grafted human fibroblast derived matrix

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Abstract

Background: Formation of mature and functional articular cartilage is still challenging in cartilage tissue engineering. This study investigates the potential of using heparin-grafted decellularized extracellular matrix (ECM) as a novel growth factor delivery platform towards human placenta-derived mesenchymal stem cells (hPMSCs) chondrogenic differentiation. Human fibroblast-derived extracellular matrix (hFDM) is naturally obtained from in vitro-cultured human lung fibroblasts via a mild decellularization process. hFDM was then conjugated with heparin via N-(3-Dimethylaminopropyl)-N′-ethylcarbodiimide hydrochloride (EDC) chemistry and subject to transforming growth factor (TGF-β1) immobilization. Once heparin grafted-hFDM (hFDM-hep) and hPMSCs were co-embedded into collagen gel, they were examined for in vitro and in vivo chondrogenesis of hPMSCs for 4 weeks.

Results: We identified heparin moieties on hFDM via toluidine blue O assay and Fourier transform infrared spectroscopy, respectively. We found out that collagen spheroids containing hFDM-hep and TGF-β1 exhibited a sustained release of growth factor for 28 days in vitro. Chondrogenesis of hPMSCs in vitro was supported by accumulated glycosaminoglycan (GAG) content and upregulated chondrogenic specific markers (collagen II, aggrecan, Sox9). Meanwhile, PKH26 - labeled hPMSCs incorporated collagen with either hFDM or hFDM-hep was pre-conditioned in a chondrogenic media for 3 days and subcutaneously implanted in the back of nude mice for 4 weeks. The implanted collagen spheroids containing both hPMSCs and hFDM-hep retained more viable hPMSCs and showed higher level of chondrogenic differentiation, based on immunostaining of collagen type II over collagen alone or Col/hFDM group. In addition, histological examination showed more positive signals of GAG via Safranin-O staining.

Conclusion: TGF-β1-immobilized hFDM-hep can provide an appropriate microenvironmnet for chondrogenic differentiation of hPMSCs in 3D collagen spheroid.

Keywords: Chondrogenesis, Human fibroblast-derived extracellular matrix (hFDM), Human placenta-derived mesenchymal stem cells (hPMSCs), Transforming growth factor (TGF-β1)
repair capacity so that complete recovery of articular cartilage is a major hurdle [4].

Tissue engineering has emerged as a thriving new field in medical sciences. It is an interdisciplinary field in which the principles of engineering and life science are integrated to generate biological substitute for improvement, maintenance, or restoration of organ functions [5, 6]. Formation of mature and functional articular cartilage is still challenging in cartilage tissue engineering [7, 8]. Tissue engineering generally requires three dimensional (3D) scaffold for cells and bioactive molecules. A typical extracellular microenvironment, for instance, ECM is very important in cell adhesion, migration and differentiation [9]. During tissue development, ECM is dynamically remodeled in accordance with cellular function [10]. In this sense, ECM based tissue engineering strategies are used widely for the regeneration of heart, trachea, muscle, tendon with matrices derived from bladder [11]. As a useful technique in preparing ECM, decellularization of tissues or cells is supposed to remove cells and cellular debris while retaining the bioactive cues that reside in the ECM [12]. The applications of ECM have been a great interest in tissue engineering such as, ECM incorporated scaffold, hybrid ECM, and grafted ECM [13–15]. Numerous studies demonstrate that mesenchymal stem cells (MSCs) contain multipotent cells and they can differentiate into bone, fat, cartilage, muscle, and neurons when induced by appropriate biological cues [16, 17]. MSCs are isolated from bone marrow, peripheral blood, placenta, amniotic fluid, umbilical cord blood, adipose tissue, and may perhaps be isolated from other sources in which they are resident components [18, 19]. Among these sources, placenta is an attractive source to isolate MSCs with multi-lineage differentiation capacity [20]. Because the placenta tissues are discarded after birth, they can be effectively utilized for research as well as clinical application with much less ethical concern [21].

In this work, the objective is to investigate the chondrogenic potential of human fibroblast-derived extracellular matrix (hFDM) and its derivative, heparin-grafted hFDM (hFDM-hep) as a transforming growth factor (TGF)-β1 delivery carrier in the induction of chondrogenesis of human placenta-derived mesenchymal stem cells (hPMSCs). To do this, hFDM was conjugated with heparin via EDC chemistry and subject to TGF-β1 immobilization. Once hPMSCs and hFDM or hFDM-hep were mixed with collagen gel, they were examined for in vitro and in vivo chondrogenesis of hPMSCs for 4 weeks. We hypothesize that TGF-β1 tethered hFDM-hep may provide an appropriate microenvironment for chondrogenic differentiation of hPMSCs.

Methods

Cell culture

Human placenta MSCs were obtained with informed consent and approval of the institutional review board of the School of Medicine, Sungkyunkwan University. The culture medium was Dulbecco’s modified Eagle’s medium-low glucose (DMEM; Gibco, USA) with 10% fetal bovine serum (FBS; Gibco) and 100 U/mL penicillin and 100 μg/mL streptomycin (P/S). For cell passage, the culture dish were washed with phosphate-buffered saline (PBS; Gibco) and incubated with TrypLE™ Select (Invitrogen, 12604-013) for 5 min at 37 °C in incubator. Dissociated cell suspensions were removed, than pelleted by centrifugation (1000 rpm, 5 min). Once the supernatant was discarded, cells were resuspended in culture media.

Preparation of human fibroblast-derived matrix (hFDM)

WI-38 human lung fibroblasts (ATCC, CCL-75) were cultured at the cell density of 2 × 10⁴ cells/cm² on the tissue culture dish (100 mm diameter) for 7 days in the DMEM supplemented with 10% FBS and 1% P/S. Once confluent, cell-loaded culture dish was washed twice with PBS, incubated briefly in a detergent solution containing 0.15% Triton X-100 (AMRESCO, Inc., Dallas, USA) and 10 mM NH₄OH (Sigma; St. Louis, MO, USA) at 37 °C, and then treated with 50 U/mL DNase I and 50 μg/mL RNase A (Invitrogen) for 1 h. After the decellularization process, ECMs were collected into centrifuge tubes and stored at 4 °C for future usage.

Heparin grafting onto hFDM

hFDM was washed with PBS and saturated with 0.05 M 2-(Nmorpholino)ethanesulfonic acid hydrate (MES) buffer (pH = 5.5) (M2933, Sigma). 0.25% (w/v) heparin working solution is prepared by adding heparin sodium (Acros, 41121-0010) to a freshly prepared solution of 0.05 M N-(3-dimethylaminopropyl)-N’-ethylcarbodiimide hydrochloride (EDC; E7750, Sigma) and 0.06 M N-hydroxysuccinimide (NHS, 130672, Sigma) in MES solution; the EDC/NHS/MES solution was vigorously mixed and left for 10 min before the use. Eventually 2 mL of heparin working solution was added to hFDM samples in a 6-well plate. They were incubated at room temperature overnight while in rotating state.

Characterization of hFDM-hep

The hFDM, hFDM-hep, and heparin sodium powder were analyzed using an attenuated total reflection (ATR)-4100 FTIR spectrometer (JASCO, Tokyo, Japan). The absorption spectra ranges between 650 and 2000 cm⁻¹. The baseline was automatically corrected using a background scan obtained in the absence of sample. Heparin conjugated hFDM (hFDM-hep) was also observed via toluidine blue O staining. 0.005%
toluidine blue O (Sigma, T3260) solution was prepared in 0.01 N hydrochloric acid with 0.2% (w/v) sodium chloride (NaCl). The hFDM-hep was reacted with 2 mL of 0.2% NaCl and 0.5 mL of toluidine blue O solution for 1 h under shaking condition. The appearance of purple color indicates the presence of heparin on the hFDM. For quantitative analysis of heparin, we examined toluidine blue O-reacted solutions at 630 nm via Multiskan microplate reader (Thermo Scientific, Rockford, IL).

Preparation of TGF-β1 immobilized hFDM and release test
Once hFDM-hep was ready in 6-well plates, 100 ng of TGF-β1 in PBS (1 mL) was added and incubated for 4 h at room temperature under a gentle shaking to incorporate TGF-β1 onto hFDM-hep. To monitor TGF-β1 (Peprotech) level tethered on hFDM, hFDM-hep was visualized using anti-human TGF-β1 antibody (Peprotech, 500 M-66) and Alexa Fluor 488 goat anti-mouse IgG (Invitrogen, A11017) and observed using fluorescent microscope (Olympus, CKX-41). For TGF-β1 release study in vitro, collagen (control) with TGF-β1 (100 ng/mL) and collagen spheroids were visualized by incubation with rhodamine phalloidin (Invitrogen, A11017) and observed using fluorescent microscope.

Evaluation of hPMSCs viability and cell morphology
hPMSCs (P9) were cultured on tissue culture plate (TCP), hFDM, and hFDM-hep substrates, respectively. After 24 h of culture, we used LIVE/DEAD® Viability/Cytotoxicity Kit (Invitrogen, 03224) to evaluate cell viability. Live or dead cells were visualized in green and red fluorescence, respectively using fluorescent microscope.

Glycosaminoglycan (GAG) assay
The GAG contents after induction of chondrogenic differentiation were measured by quantifying the amount of sulfated GAG using 1,9-dimethylmethylene (DMB) blue dye binding assay. DNA content of each sample was evaluated by using Quant-it Picogreen dsDNA Assay kit (Invitrogen, Molecular Probes). The GAG content of each sample was normalized to that of DNA content.

Quantitative real time polymerase chain reaction (Q-PCR)
Q-PCR was carried out to determine the gene expression level of chondrogenic markers. Total RNA was isolated from the samples (n = 3, each group) using TRIzol RNA Isolation Reagents (Invitrogen). The synthesis of first-strand cDNA was obtained from a solution of RNA extracts, primers and reverse transcription (RT) reaction mixture. The reaction product (1 μL) was then subject to the polymerase chain reaction (PCR) using Maxime PCR PreMix (Intron). PCR was performed via Applied Biosystems 7300 Real-Time PCR system using Taqman primers and probes. The relative gene expression was
calculated using ΔΔC_t method, where each sample was internally normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH).

Chondrogenic markers tested in this study are SRY-box containing gene 9 (SOX 9), type II collagen (Col II), aggrecan, and type I collagen (Col I). Target genes and their primers are as follows: SOX 9: AAAGGCAAGCA AAGGAGATG (forward) and TGGTGTTCTGAGG CACAG (reverse); Col II: AAGGCTCCAGAAGCAT CACC (forward) and ATCCCTAGGGCAGTGATCG (reverse); Aggrecan: TCTGTAACCCAGGCTCAAC (forward) and TGGAGTACCTGGTGCTCTC (reverse); Col I: CTGGATGCCATCAAAGTCTT (forward) and AATC CATCGGTCTAGTGTCT (reverse); and control: GAPDH: GGGCTCTCCAGAACATCATC (forward) and TTCTA GACGGCAGGTCAGGT (reverse). The raw data were first normalized to GAPDH, and then normalized to collagen samples (0 day). The results were reported as a fold change relative to that of collagen (0 day).

Subcutaneous implantation of collagen spheroids
For animal study, three different groups of collagen spheroids (Col, Col/hFDM, Col/hFDM-hep) (n = 4, each group) were prepared separately. To make collagen spheroids, hPMSCs suspension were mixed with collagen (3 mg/mL), 50 ng/ml TGF-β1 and hFDM or hFDM-hep together in an ice-bath. Collagen droplets (20 μL) were then pipetted into a 6-well culture dish and cultivated at 37 °C. Six nude mice were anaesthetized and an incision was made at the dorsum to create a subcutaneous pocket. Each sample was transplanted and the skin incisions were closed using 4.0 non-absorbable silk sutures (AILEE). Before transplantation, the hPMSCs were pre-labelled via PKH 26 cell tracking dye and the collagen spheroids were pre-conditioned in the chondrogenic media for 3 days. After 4 weeks of post-implantation, the animals were sacrificed by cervical dislocation and the skin tissues at the implantation sites were harvested. All the animal experiments were approved via the Institutional Animal Care and Use Committee of Korea Institute of Science and Technology (IACUC, 2017-016).

Histology and immunofluorescence
For histological analysis, all the collagen spheroids were fixed in 4% paraformaldehyde solution for 3 h. The samples were dehydrated with serial concentrations of ethanol (50 to 100%), washed with Histo-Clear II (National Diagnostics, Atlanta), and embedded into paraffin blocks. These samples were sectioned in 5 μm thickness, then deparaffinized with Histo-Clear II and ethanol on slide glasses. For Safranin-O staining, the samples were stained with weigert’s iron hematoxylin solution for nuclei staining for 10 min and rinsed, then stained with Fast Green (Sigma, F7528) for 5 min for cytoplasm staining, quickly washed in acetic acid. Finally the slide glasses were stained with Safrarin-O solution (Sigma, S8884) for 5 min, dehydrate and cleared with xylene.

For von Kossa staining, the slides were rinsed with distilled water and incubated with silver nitrate solution (1%, w/v) in a clear glass jar under ultraviolet light (60 watt) for 1 h. After washed with distilled water, unreacted silver nitrate was removed with sodium thiosulfate solution (5%, w/v) for 5 min and the samples were counterstained with nuclear fast red (Sigma, N3020) for 5 min. For dehydration, samples were washed and went through graded ethanol and cleared in xylene. Those samples were observed using an optical microscope (Zeiss).

Additionally the collagen spheroids were cryosectioned for immunofluorescence for collagen type II. The specimens were washed two times with PBS, blocked for 45 min with 3% BSA. Once they were incubated overnight with a mouse monoclonal anti-Col II (sc-7763; Santa cruz) (1:50) at 4 °C, the samples were washed three times with PBS, incubated for 1 h with Alexa Fluor 488 goat anti-mouse IgG (1:200) at room temperature, and then rinsed with PBS. DAPI staining was also conducted for nucleic detection. The fluorescent signals of PKH26 and Col II were visualized using confocal microscope (Olympus FV1000).

Statistical analysis
Statistical analysis was performed using the unpaired student t-test. All data represented the mean values and standard errors. Statistical significance was marked as * (p < 0.05), ** (p < 0.01), or *** (p < 0.001).

Results
Preparation and characterization of matrix-bound heparin
After the decellularization process was done, we found out that cells completely lost their original morphologies after detergent and enzyme treatments (Fig. 1a). In addition, hematoxylin staining of decellularized ECM was also confirmed via nanofiberous context without the presence of cells (Additional file 1: Figure S1). Upon the application of EDC chemistry, the purple color on hFDM-hep was apparent as indicated by toluidine blue staining, the slides were rinsed with distilled water and incubated with silver nitrate solution (1%, w/v) in a clear glass jar under ultraviolet light (60 watt) for 1 h. After washed with distilled water, unreacted silver nitrate was removed with sodium thiosulfate solution (5%, w/v) for 5 min and the samples were counterstained with nuclear fast red (Sigma, N3020) for 5 min. For dehydration, samples were washed and went through graded ethanol and cleared in xylene. Those samples were observed using an optical microscope (Zeiss).

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colors) on hFDM-hep substrate were more pronounced than on hFDM (Fig. 1e).

Cell viability, proliferation and morphology
Our results showed that TCP, hFDM and hFDM-hep substrates had similar level of hPMSCs viability, showing little cytotoxicity of each substrate as indicated via live/dead staining after 24 h of cell culture (Fig. 2a). For cell proliferation measurement in 72 h, we found that proliferation rate of hPMSCs was comparable to each other (Fig. 2b). In addition, we observed cell morphologies at 3 and 24 h, respectively (Fig. 2c). However, little difference was found at both time points.

MSCs encapsulated collagen spheroids and release test of TGF-β1 in vitro
For hPMSCs encapsulation in 3D environment, collagen spheroids with MSCs and hFDM-hep were prepared (Fig. 3a). They were intended to examine cells viability and their distribution in the spheroids. For this purpose, we labeled hPMSCs with live cell tracking dye (PKH-26, red) and the positive signals of PKH-26 in the spheroids were detectable at different time points (day 0, 1, and 2 day) (Fig. 3b–d). Meanwhile, when the release of TGF-β1 (100 ng) encapsulated in collagen (control), and collagen spheroids containing hFDM or hFDM-hep was examined, respectively individual release profile showed quite different patterns for up to 28 days, especially with a moderate amount of early release for hFDM-hep (Fig. 3e). It suggests that heparin is effective in tethering TGF-β1 and thus in controlling the release rate of TGF-β1.

Chondrogenic differentiation of hPMSCs in vitro
Based on the results of GAG contents measurement, Col alone seems to be not very effective as the GAG concentration stayed low at 2 and 4 weeks (Fig. 4a). Col/hFDM was more effective than collagen group but little GAG increment was found at 4 week. It is notable however that Col/hFDM-hep was the most effective and it showed the highest level of GAG concentration at 4 week. Current results suggested that Col/hFDM-hep may hold an active TGF-β1 and thus lead to advanced chondrogenesis of hPMSCs.

Gene expression level of selected chondrogenic markers shows that at 2 week the expression levels were similar to each other but significant changes were found at 4 week (Fig. 4b). The results unveil that hFDM
presented the highest expression of Sox9 and Col/hFDM-hep revealed higher upregulation of aggregcan and collagen type II than the other groups. Collectively, these results suggested that hFDM-hep exerts a very positive effect on the in vitro chondrogenic differentiation of hPMSCs.

**Chondrogenic differentiation of hPMSCs in vivo**

Chondrogenic differentiation of the specimens implanted in vivo was further examined histologically upon retrieved from the implantation sites. According to Safranin-O staining, we found that sulfated GAGs were highly accumulated in Col/hFDM-hep compared to Col and Col/hFDM (Fig. 5a). No significant difference was noticed between Col and Col/hFDM group until 4 weeks. To investigate the chance of osteogenesis differentiation, von Kossa staining was performed. The data showed that there was no indication of calcified mineral deposits (no black staining) (Fig. 5b). When the question about whether sulfated GAGs can be derived from hFDM was examined, we saw that sulfated GAGs were barely detectable from hFDM itself via Safranin-o staining (data not shown).

Meanwhile as the in vivo specimens were subject to immunofluorescence of collagen type II, the expression of Col II was much stronger with Col/hFDM-hep than...
Fig. 4 GAG contents and chondrogenic markers expression. 

a The GAG concentrations normalized to DNA content of each sample; b Chondrogenic gene expression of positive markers (aggrecan, Sox9, collagen type2) and negative marker (collagen type1) after 2 and 4 weeks of culture in chondrogenic medium. (*p < 0.05, **p < 0.01, ***p < 0.001)

Fig. 5 Histological analysis of subcutaneously implanted collagen spheroids at 4 weeks. 

a Safranin-O staining and b von Kossa staining. The boxed area appears in higher magnification. Scale bar is 500 and 200 μm, respectively
Col and Col/hFDM, along with strong PKH-26 positive signals (Fig. 6). These data suggested that there were more chance of viable hPMSCs at 4 weeks in Col/hFDM-hep and that hFDM-hep can provide a suitable microenvironment for chondrogenesis of hPMSCs.

**Discussion**

In this study, we investigate induction of hPMSCs for chondrogenesis in 3D collagen spheroid with the support of hFDM-hep and TGF-β1. In our early studies, cell-derived ECM offered favorable microenvironment for MSCs differentiation into osteogenesis and chondrogenesis [22, 23]. The development of ideal vehicles for growth factors delivery towards MSCs differentiation has been of great interests. Heparin is well known for its highly sulfated glycosaminoglycan which has a binding affinity to various growth factors such as fibroblast growth factor, vascular endothelial growth factor (VEGF), and TGF-β1 [24]. Previously we reported the use hFDM-hep proved advantageous in delivering an angiogenic growth factor (VEGF) and prompting much better angiogenic activity [25]. In this work, we have tested TGF-β1 immobilized hFDM-hep. TGF-β1 promotes chondrocyte differentiation at early stage by modulating proliferation and increasing alkaline phosphatase activity and proteoglycan synthesis [26, 27]. TGF-β1 on hFDM-hep can be released in a sustained manner for up to 28 days, suggesting that the interaction between heparin and TGF-β1 is secure and effective. The present data demonstrate that current release system of TGF-β1 could be useful in inducing the chondrogenic expression. In addition to TGF-β1 effect on chondrogenesis, the role of heparin itself should be addressed. There have been some reports that incorporation of heparin in the hydrogels can enhance chondrocyte phenotype or re-differentiation of dedifferentiated chondrocytes [28, 29]. The mechanism is reasoned that chondrogenic activity of heparin might be associated with the intrinsic nature of heparin that would secure endogenous growth factors (i.e., TGF-β) secreted from the cells, which can promote the chondrogenesis as a result.

**Conclusions**

This study successfully fabricated heparin-grafted hFDM and TGF-β1 tethering onto it. hFDM-hep proves nontoxic and supports the growth of hPMSCs. Encapsulated in collagen spheroids with hPMSCs, hFDM-hep showed an improved chondrogenic potential in vitro and in vivo. This may be due to prolonged bioactivity of TGF-β1, where it was immobilized onto hFDM-hep. Taken together, TGF-β1-immobilized hFDM-hep can provide a beneficial microenvironment for chondrogenesis of MSCs and thus find further applications in cartilage tissue engineering.

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**Fig. 6** Analysis of chondrogenesis of hPMSCs from subcutaneously implanted collagen spheroids at 4 week. PKH-26 labeled cells (red) and immunofluorescence of Col II (green), and merged images with DAPI staining (blue). Scale bar is 50 μm.
Additional file

Additional file 1: Figure S1. Optical image of decellularized hFDM after hematoxylin staining. It shows a nanofibrous ECM structure without the presence of cells. (DOCX 2570 kb)

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Availability of data and materials

All data generated or analyzed in this study are included in this published article.

Authors’ contributions

KP developed the concept and designed experiments. YKN performed the whole process of experiments. ADSDC, PD and KP extensively contributed on the manuscript preparation. All authors read and approved the final manuscript.

Ethics approval

All the animal experiments were approved via the Institutional Animal Care and Use Committee of Korea Institute of Science and Technology (IACUC, 2017-016).

Consent for publication

Not applicable.

Competing interests

The authors that they have no competing interests.

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