FULL LENGTH ARTICLE

Chondrogenic priming of human fetal synovium-derived stem cells in an adult stem cell matrix microenvironment

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Abstract Cartilage defects are a challenge to treat clinically due to the avascular nature of cartilage. Low immunogenicity and extensive proliferation and multidifferentiation potential make fetal stem cells a promising source for regenerative medicine. In this study, we aimed to determine whether fetal synovium-derived stem cells (FSDSCs) exhibited replicative senescence and whether expansion on decellularized extracellular matrix (dECM) deposited by adult SDSCs (AECM) promoted FSDSCs’ chondrogenic potential. FSDSCs from passage 2 and 9 were compared for chondrogenic potential, using Alcian blue staining for sulfated glycosaminoglycans (GAGs), biochemical analysis for DNA and GAG amounts, and real-time PCR for chondrogenic genes including ACAN and COL2A1. Passage 3 FSDSCs were expanded for one passage on plastic flasks (PL), AECM, or dECM deposited by fetal SDSCs (FECM). During expansion, cell proliferation was evaluated using flow cytometry for proliferation index, stem cell surface markers, and resistance to hydrogen peroxide. During chondrogenic induction, expanded FSDSCs were evaluated for tri-lineage differentiation capacity. We found that cell expansion enhanced FSDSCs’ chondrogenic potential at least up to passage 9. Expansion on dECMs promoted FSDSCs’ proliferative and survival capacity and adipogenic differentiation but not osteogenic capacity. AECM-primed FSDSCs exhibited an enhanced chondrogenic potential.

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Introduction

Articular cartilage possesses limited regenerative abilities when insulted with injury or degenerative disease such as osteoarthritis. Autologous chondrocyte implantation (ACI) is an effective procedure in treating articular cartilage defects; however, there are inherent drawbacks of autologous chondrocytes including limited availability and donor site morbidity which challenge the efficacy of this approach. Human adult stem cells isolated from various tissues have been proposed as promising sources for supplementing autologous chondrocytes. Among them, the synovium-derived stem cell (SDSC) has been characterized as a tissue-specific stem cell for chondrogenesis. Though having attracted extensive interest and achieved great success in the past few years, unfortunately, the miniscule quantities of adult stem cells and their dramatic loss of self-renewal abilities and accompanying cellular senescence during ex vivo expansion have forced researchers to look for additional alternatives.

Fetal stem cells have been discovered in prenatal tissues, such as umbilical cord blood or vein, amniotic fluid, placenta, and Wharton’s jelly. Normally discarded as medical waste, these perinatal tissue-derived fetal stem cells seem useful clinically for autologous transplantation for genetic disorders, and for banking in later stages of life. Fetal stem cells have been applied in clinical transplantation for genetic stability after more than 100 population doublings and postnatal sources. Recent studies also showed that human fetal MSCs from aborted fetuses possess superior proliferation capacity, better intrinsic homing and engraftment, more robust differentiation potential, and lower immunogenicity, as compared to MSCs from perinatal and postnatal sources. Recent studies also showed that human fetal MSCs maintained their karyotypic and epigenetic stability after more than 100 population doublings in vitro, making them even more stable than human ESCs.

In our earlier report, late stage (passage 9) fetal SDSCs (FSDSCs) were characterized; the chondrogenic potential of FSDSCs was compared after expansion on either conventional plastic flasks (PL) or decellularized extracellular matrix (dECM) deposited by FSDSCs. Our recent work investigated the replicative senescence of adult SDSCs (ASDSCs) resulting from ex vivo expansion on PL and the functional rescue of the chondrogenic potential of ASDSCs via expansion on dECM deposited by ASDSCs (AECM) and FSDSCs (FECM), particularly by FSDSCs. However, we did not know whether FSDSCs had replicative senescence and whether AECM had advantages over FECM and PL in priming human FSDSCs in their inherent property—chondrogenic potential. In this study, we hypothesized that, different from ASDSCs, FSDSCs did not exhibit replicative senescence and chondrogenic potential of FSDSCs could be boosted by expansion on AECM.

Materials and methods

dECM preparation

Human FSDSCs were obtained from ScienCell™ Research Laboratories (Carlsbad, CA) and ASDSCs were obtained from Asterand (North America Laboratories, Detroit, MI). Both cell types were used to prepare dECMs, termed FECM and AECM, respectively, as described previously. Briefly, PL was precoated with 0.2% gelatin (Sigma–Aldrich, St. Louis, MO) at 37 °C for 1 h and seeded with passage 3 SDSCs at 6000 cells per cm². After cells reached 90% confluence, 250 μM of L-ascorbic acid phosphate (Wako Chemicals USA Inc., Richmond, VA) was added for 10 days. The deposited matrix was incubated with 0.5% Triton X-100 containing 20 mM ammonium hydroxide at 37 °C for 5 min to remove the cells; they were stored at 4 °C in phosphate-buffered saline (PBS) containing 100 U/mL penicillin, 100 μg/mL streptomycin, and 0.25 μg/mL fungizone until use.

Cell expansion and morphology

PL expanded passage 3 FSDSCs (PL3) were plated at 3000 cells per cm² on FECM, AECM, or PL for one passage with growth medium containing alpha-minimum essential medium (αMEM), 10% fetal bovine serum (FBS), 100 U/mL penicillin, 100 μg/mL streptomycin, and 0.25 μg/mL fungizone. Expanded FSDSCs were termed FE4, AE4, and PL4. Cell number was counted in 175 cm² flasks (n = 6) using a hemocytometer. To observe cell morphology, FE4, AE4, and PL4 were fixed in 4% paraformaldehyde; the cell membrane was labeled with Vybrant® Dil Cell-labeling solution (Life Technologies, Grand Island, NY) and mounted with Prolong® Gold antifade reagent with 4',6-diamidino-2-phenylindole (DAPI) (Life Technologies). Both FECM and AECM were immunolabeled using monoclonal antibody for type I collagen (Sigma–Aldrich) conjugated with fluorescein isothiocyanate (FITC) and visualized with a Nikon TE2000-S Eclipse inverted microscope (Melville, NY).

Evaluation of cell proliferation and apoptosis and resistance to oxidative stress

Before cell expansion, PL3 cells were labeled with CellVue® Claret (Sigma–Aldrich) at 2 × 10⁻⁶ M for 5 min according to the manufacturer’s protocol. After 8 days, expanded cells were collected and measured using a BD FACS Calibur™ flow cytometer (dual laser) (BD Biosciences, San Jose, CA). Twenty thousand events of each sample were collected using CellQuest Pro software (BD Biosciences) and cell proliferation index was analyzed by ModFit LT™ version 3.1 (Verity Software House, Topsham, ME).

Apoptosis of expanded cells was detected using Annexin V-FITC Apoptosis Detection Kit (BioVision Inc., Milpitas, CA). Briefly, 2 × 10⁵ detached cells from each group (n = 3) were labeled with FITC conjugated annexin V and propidium iodide for 15 min. Samples were measured using FACS Calibur (BD Biosciences) and analyzed using the FCS Express software package (De Novo Software, Los Angeles, CA).
Expanded cells were incubated with 1 mM hydrogen peroxide (H₂O₂) at 37°C for 1 h. To measure intracellular reactive oxygen species (ROS), cells were incubated with 1 μM 2′,7′-dichlorodihydrofluorescein diacetate (H₂DCFDA) (Life Technologies) for 15 min. The plates were read on a FLUOSTar OPTIMA (BMG Labtech Inc., Cary, NC) with an excitation wavelength of 485 nm and emission of 530 nm. Samples were assayed in triplicate.

**Effect of evaluation of culture substrates on FSDSC chondrogenic potential**

Plastic expanded FSDSCs (3 × 10^5) from passage 2 and passage 9 were centrifuged at 500 g for 5 min in a 15-mL polypropylene tube to form a pellet. After overnight incubation (day 0), the pellets were cultured in a serum-free chondrogenic medium consisting of high-glucose Dulbecco’s modified Eagle’s medium (DMEM), 40 μg/mL proline, 100 mM dexamethasone, 100 U/mL penicillin, 100 μg/mL streptomycin, 0.1 mM ascorbic acid-2-phosphate, and 1× ITS™ Premix (BD Biosciences) with the supplementation of 10 ng/mL transforming growth factor β3 (TGF-β3, PeproTech Inc., Rocky Hill, NJ) in a 5% O₂ incubator as long as 21 days. The pellets from day 21 were assessed using AB staining for sulfated GAGs and IHC labeling for type II collagen, TaqMan® real-time PCR for ACAN and COL2A1, biochemical analysis for DNA and GAG amounts, and biomechanical testing for pellet stiffness and Young’s Modulus (YMOD).

**Effect of evaluation of culture substrates on FSDSC adipogenic and osteogenic potentials**

For adipogenesis, expanded cells on varied substrates were seeded at 10,000 cells/cm² and cultured for 21 days in adipogenic medium (growth medium supplemented with 1 mM dexamethasone, 0.5 mM isobutyl-1-methyloxanthine, 200 mM indomethacin, and 10 mM insulin). The cultures (n = 3) were fixed in 4% paraformaldehyde and stained with a 0.6% (w/v) Oil Red O (ORO) solution (60% isopropanol, 40% water) for 15 min. Intracellular lipid-filled droplet-bound staining was photographed under a Nikon TE300 phase-contrast microscope (Nikon, Tokyo, Japan). Adipogenic marker genes [lipoprotein lipase (LPL) and peroxisome proliferator-activated receptor gamma (PPARγ)] were quantified using TaqMan™ real-time PCR.

For osteogenesis, expanded cells (n = 3) were seeded at 8000 cells/cm² and cultured for 21 days in osteogenic medium (growth medium supplemented with 0.1 mM dexamethasone, 10 mM β-glycerol phosphate, 50 mM ascorbate-2-phosphate, and 0.01 mM 1,25-dihydroxyvitamin D₃) were collected for alkaline phosphatase (ALP) activity assay with a reagent kit (Sigma–Aldrich) by measuring the formation of p-nitrophenol (p-NP) from p-nitrophenyl phosphate following the manufacturer’s instructions. p-NP was quantified based on the spectrophotometric absorbance at 405 nm and enzymatic activity was expressed as millimoles of p-NP/min/mg DNA. For evaluation of calcium deposition, induced cells (n = 3) were fixed with 70% ice-cold ethanol for 1 h and then incubated in 40 mM Alizarin Red S (ARS) at pH 4.2 for 20 min with agitation. After rinsing, matrix mineral-bound staining was photographed. Quantification of staining was performed by incubating cells with a solution of 10% acetic acid and 20% methanol for 15 min to extract the calcium-chelated ARS stain. Samples were analyzed for absorption at 450 nm, which was normalized to total DNA amount for standardization.

**Histology and immunolabeling**

Representative pellets (n = 2) were fixed in 4% paraformaldehyde at 4 °C overnight, followed by dehydrating in a gradient ethanol series, clearing with xylene, and embedding in paraffin blocks. 5-μm sections were stained with AB staining (counterstained with fast red) for sulfated GAGs. For immunohistochemistry, the sections were immunolabeled with a primary antibody against type II collagen [II-I] and type II collagen [II-II] by using a secondary antibody of biotinylated horse anti-mouse IgG (Vector, Burlingame, CA). Immunoreactivity was detected using Vectastain ABC reagent (Vector) with 3,3′-diaminobenzidine as a substrate.

**Biochemical analysis for DNA and GAG amount**

Representative pellets (n = 4) were digested at 60°C for 4 h with 125 μg/mL papain in PBE buffer (100 mM...
phosphate, 10 mM ethylenediaminetetraacetic acid, pH 6.5) containing 10 mM cysteine. To quantify cell density, the amount of DNA in the papain digestion was measured using the QuantIT™ PicoGreen® dsDNA assay kit (Life Technologies) with a CytoFluor® Series 4000 (Applied Biosystems, Foster City, CA). GAG was measured using dimethylmethylene blue dye and a Spectronic® BioMate™ 3 Spectrophotometer (Thermo Fisher Scientific, Waltham, MA) with bovine chondroitin sulfate (Sigma–Aldrich) as a standard.

**TaqMan® real-time PCR**

Total RNA was extracted from pellets (n = 4) using an RNase-free pestle in TRIzol® (Life Technologies). Two µg of mRNA were used for reverse transcriptase with the High-Capacity cDNA Archive Kit (Applied Biosystems) at 37 °C for 120 min. Chondrogenic marker genes [ACAN (Assay ID A1QJAP5) and COL2A1 (Assay ID Hs00156568_m1)] and adipogenic marker genes [LPL (Assay ID Hs00173425_m1) and PPARG (Assay ID Hs00115513_m1)] were customized by Applied Biosystems as part of their Custom TaqMan® Gene Expression Assays. Eukaryotic 18S rRNA (Assay ID HS99999901_s1 ABI) was carried out as the endogenous control gene. Real-time PCR was performed with the iCycler iQ™ Multi Color RT-PCR Detection kit and calculated by computer software (Perkin–Elmer, Wellesley, MA). Relative transcript levels were calculated as $\Delta \Delta Ct = \Delta E - \Delta C$, $\Delta E = Ct_{\text{exp}} - Ct_{18S}$, and $\Delta C = Ct_{\text{ctrl}} - Ct_{18S}$.

**Biomechanical testing**

Representative pellets (n = 6) were placed into a reservoir filled with PBS and loaded onto the custom miniature stepper motor driven compression device. This device used a 10 g load cell and a miniature differential variable reluctance transducer (DVRT) for displacement readout. A small preload of 0.0001 N was applied followed by a 10% strain at 1 s duration. From the linear portion of the load displacement curve, the Stiffness and YMOD were calculated for the spherical shaped pellet according to Rodriguez and colleagues.19

**Statistical analysis**

A Mann–Whitney U test and a linear model with contrast analysis were used for pairwise comparison. All statistical analyses were performed with SPSS 13.0 statistical software (SPSS Inc., Chicago, IL). p values less than 0.05 were considered statistically significant.

**Results**

**Cell expansion enhanced FSDSCs’ chondrogenic potential**

To determine whether cell passaging on plastic flasks would affect expanded cells’ chondrogenic potential, human FSDSCs from passage 2 and passage 9 were evaluated after

**Fig. 1** Later passage FSDSCs exhibited an enhanced chondrogenic potential. Passage 2 (P2) and P9 FSDSCs were chondrogenically induced in a pellet culture system for 14 days. The effect of passaging on FSDSC chondrogenic capacity was evaluated using (A) histology, including pellet size (mm) and Alcian blue staining for sulfated GAGs, (B) biochemical analysis for DNA and GAG contents of pellets with DNA ratio indicating cell viability and a ratio of GAG to DNA indicating chondrogenic index, and (C) real-time PCR for chondrogenic marker gene expression (ACAN and COL2A1). Data are shown as average ± standard deviation (SD) for n = 4. *p < 0.05 indicated a statistically significant difference.
14-day chondrogenic induction. The pellets from passage 9 cells were bigger in size with a shiny surface; AB staining showed comparable intensity to those from passage 2 cells (Fig. 1A). Biochemical analysis data showed that, despite a higher DNA ratio (by day 0) in 14-day pellets than those from passage 9 cells, the pellets from passage 2 cells exhibited a lower amount of GAG per pellet and a lower ratio of GAG to DNA at both day 7 and day 14 (Fig. 1B). Real-time PCR data showed that, compared to a continuing increase in passage 9 cells, both ACAN and COL2A1 increased at day 7 but decreased at day 14 in passage 2 cells (Fig. 1C).

dECM expansion promoted FSDSCs' proliferative and survival capacity

To evaluate the proliferative effect exerted by AECM and FECM expansion, human FSDSCs were seeded for one passage on three different substrates: PL, AECM, or FECM. Cells grown on PL exhibited random organization and spindle morphology (red color); in contrast, cells expanded on dECMs were more elongated and well organized by lining up with matrix fibers in which type I collagen was positively detected (initial color was green and merged color was yellow) (Fig. 2A). Six-day-expansion on FECM yielded the highest cell number followed by AECM with PL having the lowest (Fig. 2B). This finding was supported by proliferation index (PI) data, in which AECM expansion yielded a 1.8-fold increase and FECM expansion yielded a 2.5-fold increase compared with PL expansion (Fig. 2C).

Our flow cytometry data showed that expanded FSDSCs from all groups exhibited a similar percentage of apoptotic cells (Fig. 2D). To evaluate cell viability after expansion on these substrates, 1 mM H2O2 was applied to measure ROS scavenging capacities in expanded FSDSCs. Both FEM and AECM expanded cells produced about half the ROS compared to PL upon stimulus (Fig. 2E), indicating dECM expanded FSDSCs had higher viability in a harsh environment.

To evaluate the effect of dECM expansion on MSC surface markers of FSDSCs, our flow cytometry data showed that in both AECM and FECM expanded FSDSCs, CD29 and CD90 were down-regulated in median fluorescence intensity (median) with CD105 down-regulated in both median and percentage. Interestingly, SSEA4 was up-regulated in both

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**Fig. 2** dECM expanded FSDSCs exhibited an enhanced proliferation and resistance to oxidative stress, especially for cell expansion on FECM. (A) FSDSCs were expanded on three substrates (PL, AECM, and FECM) for one passage; the cell membrane was labeled with Vybrant Dil Cell labeling solution (red) and the nuclei were stained using DAPI (blue), the dECM fibers were immuno-labeled for type I collagen (green). (B) Cell number was counted in 175 cm² flasks (n = 6) using a hemocytometer. (C) Flow cytometry was used to measure the proliferation index of expanded ASDSCs. (D) Flow cytometry was used to measure percentage of apoptotic cells after expansion on the three substrates. (E) Generation of ROS following hydrogen peroxide incubation was evaluated. Relative fluorescence emitted by fluorescent dye 2',7'-dichlorofluorescein diacetate was measured and is presented as means ± standard deviation (SD) of the mean for n = 3. *p < 0.05 indicated a statistically significant difference.
percentage and median while integrin β5 was up-regulated only in median (Fig. 3).

AECM primed FSDSCs exhibited an enhanced chondrogenic potential

To determine the priming effects of dECM expansion on FSDSCs’ chondrogenic potential, expanded cells were chondrogenically induced for up to 21 days. At the early stage, 10-day pellets from PL3, PL4, and FE4 were similar in size while those from AE4 were slightly bigger; the pellets from dECM expansion were intensely stained with Alcian blue for sulfated GAGs and immunolabeled for type II collagen compared to those from PL expansion (Fig. 4A). Chondrogenic marker gene (ACAN and COL2A1) expression further revealed that dECM expansion up-regulated chondrogenic genes with AE4 being the highest, followed by FE4 (Fig. 4B). Consistent with the above cell expansion data in Fig. 1C, PL4 cells exhibited a higher level of ACAN and COL2A1 than PL3 cells after 10-day chondrogenic induction (Fig. 4B).

At the later stage, 21-day pellets were evaluated for chondrogenic differentiation and functionality. Similar to the data at day 10, the pellets from PL3, PL4, and FE4 groups were comparable in size; the pellets from dECM expansion, particularly from AE4, were bigger and intensely stained for sulfated GAGs and type II collagen (Fig. 5A). The pellets from AE4 exhibited the highest level of chondrogenic marker genes (ACAN and COL2A1) followed by those from PL expansion (ACAN and COL2A1); pellets from the FE4 group had the lowest level (ACAN and COL2A1) (Fig. 5B). Our biochemistry data showed that, after chondrogenic induction, AECM expansion yielded higher cell viability (DNA% by day 0) though it was not significantly different; AECM expansion yielded the highest GAG amount per pellet, followed by those from FECM expansion with PL expansion having the least with a comparable ratio of GAG to DNA among groups (Fig. 5C). Our functionality data suggested that dECM expansion yielded FSDSC pellets with comparable Stiffness and Young’s modulus with those from PL expansion (Fig. 5D).

dECM expanded FSDSCs exhibited an enhanced adipogenic potential rather than osteogenic capacity

To determine the rejuvenation effect of dECM expansion on FSDSC adipogenic and osteogenic potentials, expanded cells were incubated in corresponding lineage induction medium for 21 days. Our adipogenic induction data showed that expanded FSDSCs from varied substrates all stained positive for ORO (Fig. 6A); FE4 exhibited the highest levels of LPL and PPARG mRNAs followed by AE4 (LPL); PL expansion had the lowest levels (Fig. 6B). Our osteogenic induction data showed that expanded FSDSCs from varied substrates all stained positive for ARS (Fig. 6C) and ALP (Fig. 6E). Quantitative assessment for relative ARS density (Fig. 6D) and ALP activity (Fig. 6F) did not show a significant difference between groups.

Discussion

To achieve successful cartilage repair, stem cells for application in cartilage regeneration should be obtained easily, non-invasively, and in sufficient quantities and then transplanted safely and efficiently to a host. 20 Fetal stem cells are a promising source; however, there are limited studies investigating the potential of these sources in cartilage regeneration. Our group was the first to characterize FSDSCs through expression of MSC markers

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Fig. 3  Flow cytometry was used to measure both percentage and median of MSC surface markers (CD29, CD90, and CD105), SSEA4, and integrin β5 in human FSDSCs after one passage on three substrates (PL, AECM, and FECM).
CD90, CD105, CD29, and SSEA4 and multi-lineage differentiation potential. In this study, we found that FSDSCs exhibited an enhanced chondrogenic potential during cell expansion at least up to passage 9. Expansion on dECM promoted FSDSCs' ability in proliferation and resistance to oxidative stress, particularly on FECM, while expansion on AECM promoted FSDSCs' ability in chondrogenic differentiation. Interestingly, expansion on dECM also enhanced FSDSCs' ability in adipogenesis rather than osteogenesis.

Our previous reports showed that adult stem cell passaging on conventional plastic flasks could cause replicative senescence, evidenced by a down-regulation of the ratio of GAG to DNA (chondrogenic index), ACAN, and COL2A1 in porcine SDSCs from passage 3 to 5 and in human bone marrow stromal cells (BMSCs) from passage 5 to 8. To the best of our knowledge, for the first time, the findings in this study indicated that replicative senescence did not apply to fetal source SDSCs, which exhibited an enhanced chondrogenic capacity after expansion from passage 2 to 9. The discrepancy between fetal and adult SDSCs also existed in their response to dECM. In a recently published parallel study, despite a higher level of GAG amount in each pellet from FSDSCs compared to ASDSCs (38.89 ± 2.59 μg versus 24.60 ± 1.42 μg) after a 21-day chondrogenic induction, FSDSC expansion on either AECM (1.52-fold increase) or FECM (1.38-fold increase) yielded pellets with a lower level of GAG amount per pellet than those from ASDSC expansion (2.78- and 4.42-fold increase, respectively). Consistent with a previous report that fetal BMSCs expressed higher osteogenic genes and deposited a higher level of calcium compared to adult BMSCs in nude mice, this finding suggested that human FSDSCs exhibited higher chondrogenic potential than human ASDSCs despite more sensitivity of ASDSCs to chondrogenic priming from dECM expansion. Interestingly, we found the most chondrogenic capacity was achieved for ASDSCs by expansion on FECM and for FSDSCs by expansion on AECM, indicating that the shortage of “stemness” signals in adult SDSCs could be supplemented by fetal matrix while the shortage of

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**Fig. 4** AECM promoted expanded FSDSCs' early chondrogenic differentiation. PL expanded passage 3 FSDSCs (PL3) were grown on PL, AECM, or FECM for one passage; the expanded cells were PL4, AE4, and FE4. The above four cells were chondrogenically induced in a pellet culture system for 10 days. (A) Before histological staining, pellet size was measured with a scale bar in mm; Alcian blue (AB) was used to stain sulfated GAGs and immunohistochemistry labeling (IHC) was used to detect type II collagen (Col II). (B) Real-time PCR was used to evaluate chondrogenic marker gene expression (ACAN and COL2A1) in day 10 pellets. Data are shown as average ± standard deviation (SD) for n = 4. *p < 0.05 indicated a statistically significant difference.
in promoting cell proliferation and preventing apoptosis, which might be explained by the discrepancy in chemical composition of these two matrices. Our proteomics data showed that FECM had more fibrillin-2, tenasin C, and clusterin than AECM. Despite being largely restricted to developing fetal tissues, enhanced expression of fibrillin-2 and tenasin C has been observed in adults with fibroproliferative conditions, such as wound healing and scleroderma. AECM had more biglycan, decorin, dermatopontin, elastin, periostin, thrombospondin-1, and TGF-β1 than FECM. Decorin and biglycan were reported to reduce proliferation of pre-adipocytes, partly by induction of apoptosis. Dermatopontin has been implicated in chondrogenic differentiation. Notably, periostin and TGF-β are involved in osteochondrogenesis. Thrombospondin-1 signaling through CD47 inhibits self-renewal by regulating c-Myc and other stem cell transcription factors. The above data showed that, compared with the role of FECM during cell expansion, AECM has more matrix components favoring cell differentiation and apoptosis rather than cell proliferation.

In this study, we also found that dECM expansion enhanced FSDSC adipogenic potential rather than osteogenic capacity. This finding is in line with our parallel study, in which ASDSCs were compared for tri-lineage
differentiation capacities after expansion on FECM and AECM. A possible explanation is that SDSCs are an adult stem cell inherently favoring chondrogenic and adipogenic differentiation rather than osteogenic differentiation.4,5
dECM expansion augmented the inherent differentiation potential of a specific stem cell; for instance, dECM deposited by SDSCs favored expanded SDSCs’ chondrogenic capacity21 while dECM deposited by BMSCs benefited expanded BMSCs’ endochondral bone formation.22

In conclusion, even though there is a history of fetal stem cell transplantation since 1928 and hundreds of clinical trials using various types of fetal transplants have been performed worldwide,44 there are few studies investigating fetal stem cells from synovial tissue for chondrogenic potential. Different from adult stem cells with replicative senescence, in this study, we found that passing human fetal SDSCs could yield enhanced chondrogenic capacity. Our study also suggests that a mature microenvironment can render fetal stem cells with lineage specific differentiation. The knowledge from this study will contribute to future cartilage regeneration and engineering using fetal source SDSCs in the treatment of osteoarthritic cartilage defects.

Competing interest statement
The authors report no conflict of interest.

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