Enhanced Effect of Tendon Stem/Progenitor Cells Combined With Tendon-Derived Decellularized Extracellular Matrix on Tendon Regeneration

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Abstract
Decellularized extracellular matrices have been clinically used for tendon regeneration. However, only a few systematic studies have compared tendon stem/progenitor cells to mesenchymal stromal cells on the tendon-derived decellularized matrix. In the present study, we prepared extracellular matrix derived from porcine tendons and seeded with tendon stem/progenitor cells, embryonic stem cell-derived mesenchymal stromal cells or without stem cells. Then we implanted the mixture (composed of stem cells and scaffold) into the defect of a rat Achilles tendon. Next, 4 weeks post-surgery the regenerated tendon tissue was collected. Histological staining, immunohistochemistry, determination of collagen content, transmission electron microscopy, and biomechanical testing were performed to evaluate the tendon structure and biomechanical properties. Our study collectively demonstrated that decellularized extracellular matrix derived from porcine tendons significantly promoted the regeneration of injured tendons when combined with tendon stem/progenitor cells or embryonic stem cell-mesenchymal stromal cells. Compared to embryonic stem cell-mesenchymal stromal cells, tendon stem/progenitor cells combined with decellularized matrix showed more improvement in the structural and biomechanical properties of regenerated tendons in vivo. These findings suggest a promising strategy for functional tendon tissue regeneration and further studies are warranted to develop a functional tendon tissue regeneration utilizing tendon stem/progenitor cells integrated with a tendon-derived decellularized matrix.

Keywords
Tendon stem/progenitor cells, mesenchymal stem cells, decellularized matrix, tendon tissue engineering, rat, in vivo

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### Introduction

Tendons are connective tissues that transmit force from muscle to bone. They are injured frequently due to rigorous activities and sports. It is well known that tendons naturally heal with the formation of fibrotic scars rather than regeneration tissue owing to their vascular deficiency and lack of highly differentiated tenocytes, with limited proliferation capability and metabolic rate. Therefore, the natural tendon healing process always leads to poorer tendon quality and mechanical properties. A variety of treatment modalities have been employed, such as xenografts, autografts, allografts, tendon-to-bone fixation, and tendon prostheses. However, these options are accompanied by limited donor resource, poor integration graft and high cost of medication.

Tissue engineering provides an alternative with a high potential for treating injured tissue. Cells (stem cells), scaffolds, and bioactive molecules are three fundamental elements; their internal coordination and external interplay with the surrounding tissue of the implantation site induce the native, regenerative healing of injured tissues and organs. Scaffolds play a crucial role during the tissue-regeneration process; the various scaffolds include artificial synthetic and naturally derived materials. Recently, native organs have become a popular alternative as they contain extracellular matrix in which the composition, organization, mechanical properties, and/or geometry support the viability and phenotype of tissue-specific cells. Considering that many native organs are not suitable for transplantation, particularly for xenograft, decellularized organs and scaffolds show enormous potential to work as instructive matrices for tissue-engineering applications. For tendons, many such scaffolds have been approved by the US Food and Drug Administration.

Currently, stem cells with the capability of multi-lineage commitment and self-renewal are an ideal cell source for tissue engineering. For tendons, tendon stem/progenitor cells (TSPCs) and mesenchymal stem cells (MSCs) are the most popular. Since their discovery in rats in 2007, TSPCs have been isolated from various mammals. As a local stem cell, mounting evidence shows that TSPCs have the potential for tendon tissue engineering. TSPCs could regulate inflammation in tendon healing via JNK and STAT3 signaling, and serve as a potential target for the prevention or treatment of tendinopathy via inhibiting the P16 tenogenics of the microRNA signaling pathway. The shortcomings of TSPCs are the lack of sources and the difficulty in obtaining them. In contrast, MSCs are relatively easy to acquire and can be isolated from a variety of tissues. They can commit toward the expected cell lineages in situ, or secrete trophic molecules, for instance, chemotactic molecules and growth factors, which can recruit extra reparative cells into the lesion site.

A previous study showed that TSPCs possessed more potential for tenogenic expression compared to MSCs in vitro, however, there has not yet been any systematic study that compares TSPCs to MSCs on the tendon-derived decellularized matrix. In the present study, we hypothesized that the TSPCs as the resident stem cells of tendons and integrated with a tendon-derived decellularized matrix will express high tenogenics, produce more tendon-like tissue, and improve structural and biomechanical properties of regenerated tendons much better than MSCs or without seed cells in vivo.

### Materials and Methods

This study was approved by the Zhejiang University Institutional Animal Care and Use Committee, and we strictly followed the current laws for animal experiments. The schematic drawing shown in Fig. 1 was used to illustrate the process of this study.

#### Preparation of Porcine Tendon Decellularized Matrix

The decellularization protocol we used is similar to our previous study. Firstly, fresh porcine tendons were cut into 10 mm pieces, washed in 50 ml of 0.1 M phosphate buffered saline (PBS) for 15 minutes, then soaked in liquid nitrogen for 2 minutes and immersed in a 37°C solution of 500 ml 0.9% saline for 10 minutes. This procedure was repeated five times. Before being placed in the 24-well plates, the tendon pieces were cut into 80 μm-thick sections with a cryostat microtome (Microm HM550, Waltham, MA, USA). They were then washed three times with ml of 0.1 M PBS, for 30 min each time. Next, 100 μl of 200 unit ml-1 DNase (Roche, Basel, Switzerland) was placed into each well, then they were incubated at 4°C for 14 hours followed by washing for 30 min with 1 ml of 0.1 M PBS at 4°C. Before the experiments, hematoxylin and eosin (H&E) staining confirmed the resident cells were completely removed.

#### Stem Cell Isolation and Culture

The human TSPCs used in this study were acquired according to our previous study. Briefly, the cells were planted in 10 cm dishes and cultured at 37%, 5% CO2 with complete medium consisting of Dulbecco’s modified Eagle’s medium, 10% fetal bovine serum, and 1% penicillin-streptomycin (all from Gibco, Carlsbad, CA, USA). The non-adherent cells were removed with PBS and the medium exchanged every 2 to 3 days. Cells at passage 3 were used for experiments. Then 3 days before surgery, the cells at a concentration of 1×10^5 cells per ml were seeded on the scaffold. Before being used for the experiments, the multi-lineage differentiation potential and clonogenicity of these stem cells were confirmed in this study.
In Situ Rat Achilles Tendon Repair Model

The hind limbs of 18 skeletally mature female Sprague Dawley (SD) rats (Zhejiang University Laboratories, Hangzhou, China) weighing between 200 g and 220 g were used for this experiment. All the rats were treated with cyclophosphamide (150 mg/kg) 24 h before the operation. After general anesthesia, a full tear wound was created and the Achilles tendon was removed to create a defect of 6 mm in length. The decellularized extracellular matrix (ECM) scaffolds (10 × 10 mm, thickness 80 μm), seeded with TSPCs (ECM + TSPCs group, N = 12, 5 × 10^5 cells per scaffold) or MSCs (ECM + MSCs group, N = 12, 5 × 10^5 cells per scaffold) or without seeding cells (ECM group, N = 12) were rolled into the gap wound, then sutured to the remaining Achilles tendon using a non-resorbable suture (6-0 nylon). We next irrigated the wound and closed the skin. The animals were allowed free cage activity after surgery. Then 4 weeks post-surgery, specimens were collected successively.

Histological Staining and Immunohistochemistry

The harvested, regenerated tendon tissue was promptly immersed in 10% (v/v) neutral buffered formalin (Xinghan Ltd, Zhengzhou, China) and dehydrated through an alcohol gradient, then embedded in paraffin blocks. For standard histological evaluation, 7 μm sections were stained with H&E (N = 3 per group). To examine the general appearance of the collagen fibers, Masson’s Trichrome staining (N = 3 per group) was also performed according to standard procedures. Polarizing microscopy (N = 3 per group) was used to assess mature collagen fibrils. The general histological score (fiber structure, fiber arrangement, rounding of nuclei, inflammation, vascularity, cell population) of the H&E staining result was calculated in this study and the method was according to Shen et al. For immunohistochemical analysis (N = 3 per group), mouse anti-rat monoclonal antibody against collagen I (1:200 dilution; Abcam, Cambridge, UK) was used to assess the expression of collagen I in repaired Achilles tendon.

Determination of Collagen Content

The amount of collagen in the repaired tendon was quantified using a collagen assay kit (Jiancheng Ltd, Nanjing, China). We digested the lyophilized tendons (N = 3 per group) with a hydrolysis regent at 95% for 20 min, and serially diluted acid-soluble collagen type I (provided by the kit) to generate the standard curve according to the manual. The concentration was obtained through absorbance at 550 nm via a microplate reader (Molecular Devices, San Jose, CA, USA).

Transmission Electron Microscopy

Achilles tendon specimens (N = 3 per group) were fixed via standard procedures for transmission electron microscopy (TEM; Tecnai G2 F20 S-TWIN, FEI, Hillsboro, Oregon, USA) to assess the diameter and alignment of collagen fibril. Methods and processes are the same as previously
described. To acquire an accurate representation of the fibril diameter distribution, we measured more than 500 collagen fibrils for each specimen.

**Mechanical Testing**

Mechanical testing (N = 5 per group) was carried out through an Instron tension/compression system with Fast-Track software (Model 5543, Instron, Canton, MA). Measurements of the tendon’s cross-sectional area were performed via two Vernier calipers at 5 mm proximal to the conjunction of the bone and tendon. The bone end of the tendon was secured by a specially designed restraining jig and the tendon end was pinched with a clamp. The Achilles tendon-calcaneous complex (ACC) was then rigidly fixed to custom-made clamps. After applying a preload of 0.1 N, each ACC underwent pre-conditioning by cyclic elongation of between 0 and 0.5 mm for 20 cycles at 5 mm per min. This was followed by a load-to-failure test at an elongation rate of 5 mm per min. The load-elongation behavior of the ACCs and failure modes were recorded. The structural properties of the ACC were represented by stiffness (N/mm), ultimate load (N), energy absorbed at failure (mJ) and stress at failure. For each ACC, the greatest slope in the linear region of the load-elongation curve over a 0.5 mm elongation interval was used to calculate the stiffness.

**Statistical Analysis**

Statistical significance between groups was assessed by one-way analysis of variance followed by post-hoc Scheffe test using SPSS 22.0 (IBM, Amun, New York, USA). All values of P < 0.05 were accepted as statistically significant.

**Results**

**Characterization of Decellularized Porcine Tendon**

After the decellularization, most of the cells were removed and only the native collagen structure was present on H&E staining (Fig. 2A). MSCs or TSPCs were seeded to the scaffold (Fig. 2B) and kept in an incubator at 37%, 5% CO2. Then 3 days later, the combination of cells and scaffolds were rolled up, thus an in vitro engineered tendon was generated, in which the cells and scaffolds were alternately layered, as could be observed upon H&E staining (Fig. 2C). Subsequently, the engineered tendons were transplanted into a rat model of an Achilles tendon defect.

**TSPCs Promoted Tendon Regeneration Structurally**

Then 4 weeks post-surgery all rats were euthanized, and the inflammatory reaction, immunological reaction or scaffold dislocation were observed in the Achilles tendon repair site. Masson trichrome and H&E staining were employed to evaluate the reparative effect in both the repaired zone and junction zone for each group. In the repaired zone, Masson trichrome staining manifested more organized collagen fiber structures in the ECM + TSPCs group when compared with other two groups (Fig. 3A). Meanwhile, H&E staining also showed more fibroblast-like cells that closely resembled the native tenocyte in the ECM + TSPCs group (Fig. 3B). Around the collagen fibers, vascular components could be found in each group. It showed the good histocompatibility of the ECM scaffold for tissue engineering tendons. Polarized light imaging confirmed the better effect of ECM + TSPCs as there was a notable increase of mature collagen fibers in comparison to the other two (Fig. 3C). At the junction of the repaired tendons, more disarranged collagen fibers and scar tissue could be seen in control groups of both the H&E and Masson trichrome staining (Fig. 4A–B). The histological score (Fig. 4C) was lowest in the ECM + TSPCs group (6.64 ± 1.33 vs. 9.09 ± 0.92 in ECM + MSC group and 10.61 ± 0.71 in ECM group). These data collectively demonstrated the combination of decellularized porcine tendon and TSPCs promoted the regeneration of injured tendons to form a better structure.
TSPCs Produced More Collagen

We monitored the expression level of the tendon-related collagen content of repaired tendons. Immunohistochemical staining showed the highest density of collagen type I in the ECM + TSPC group (Fig. 5A). Moreover, both TSPCs and MSCs deposited more collagen type I compared to the cell-free group. The collagen content assay (Fig. 5B) showed a similar trend (that had less than 0.124 ± 0.012 mg/mg in ECM + TSPC group vs. 0.12 ± 0.006 mg/mg in the ECM + MSC group and 0.11 ± 0.003 mg/mg in the ECM group). These data suggested stem cells produced more collagen.

Fig. 3. Repaired zone of engineered tendons 4 weeks after surgery. (A) Masson trichrome staining of each group. (B) Typical H&E staining of repaired zones in a section of each group. (C) Polarized light microscopy images of each group. Scale bars = 50 μm (A, B, C), 200 μm (inset of A, B, C).

Fig. 4. Junction zone (square frame of each picture) of engineered tendons 4 weeks after surgery. (A) Masson trichrome staining of each group (N: normal zone, R: repaired zone). (B) Typical H&E staining of junctional zone of each group. (C) Histological evaluation was performed for each group (ECM vs ECM + TSPC: 10.61 ± 0.71 vs 6.64 ± 1.33, P < 0.05). Scale bars = 50 μm (A, B), 200 μm (inset of A, B).
Ultrastructural Morphology

To correlate histological evidence with the ultrastructure morphology of repaired Achilles tendons (ATs), specimens were observed under TEM. The ECM\textsuperscript{+}TSPC group displayed thicker collagen fibrils (Fig. 6A) in comparison to other groups. The fibrils’ average diameter was 23.5 nm (816 fibers) in the ECM\textsuperscript{+}TSPC group, 21.7 nm (984 fibers) in the ECM\textsuperscript{+}MSC group, and 21.1 nm (522 fibers) in ECM group (Fig. 6B). The diameter distribution showed 19.85\% of the ECM\textsuperscript{+}TSPC group’s regeneration fibrils were larger than 30 nm, which was higher than that of the ECM\textsuperscript{+}MSC group (8.33\%) and the ECM group (8.05\%). Those results indicate that TSPCs are beneficial for the generation of larger fibrils.

TSPCs Promoted Biomechanical Properties of Repaired AT

To further correlate tissue structural features with their biomechanical properties, we performed mechanical testing. Most of the specimens fractured at the muscle-tendon junction or the tendon-calcaneal interface. Only one of the ECM\textsuperscript{+}MSC group specimens fractured at the middle of the AT. The ECM\textsuperscript{+}TSPC group (23.24\pm 5.54 N/mm) showed significantly more stiffness than the ECM\textsuperscript{+}MSC group (17.89\pm 4.48 N/mm) and the ECM group (14.40\pm 7.09 N/mm) (Fig. 7A). The maximum force (Fig. 7B) of the ECM\textsuperscript{+}TSPC group (59.79\pm 8.05 N) was significantly higher than ECM group (40.83\pm 14.38 N), and stress at failure (Fig. 7C) manifested a similar trend (6.91\pm 1.11 Mpa vs. 4.32\pm 2.13 Mpa, P < 0.05). The assays of energy absorbed at failure (Fig. 7D) of the ECM\textsuperscript{+}TSPC group were obviously higher than the ECM\textsuperscript{+}MSC group (131.44\pm 32.63 mJ vs. 90.10\pm 34.21 mJ, P < 0.05). The lack of statistical significance may be due to the small sample size (N = 5 per group). The ECM\textsuperscript{+}TSPC group exhibited better biomechanical properties than the other two groups.

Discussion

In this study, we used a decellularized mammalian tendon extracellular matrix scaffold with TSPCs or MSCs to repair injured tendons, and demonstrated that: (1) the tendon ECM scaffold easily acquires and promotes tendon regeneration especially when combined with TSPCs or MSCs, which can be used as a suitable bioscaffold for tendon tissue engineering; and (2) seeded TSPCs showed enhanced improvement of biomechanical properties and morphological structure in regenerated tendons compared to MSCs.

In recent years, bioscaffolds’ physical properties have been given more attention and various sources of decellularized matrix have been used in tendon tissue engineering\textsuperscript{20}. In our engineered tendons with stem cells, the cells and decellularized tendon matrix were arranged in layers and were in close contact with each other. Such an intimate contact may be a key inducer in directing the site-specific differentiation.
Fig. 6. (A) Ultrastructure of tissue-engineered tendons after 4 weeks post-surgery. (B) Histogram of the distribution of collagen fibril diameters (N > 500). Scale bars = 100 nm.

Fig. 7. Biomechanical properties of repaired tendons at 4 weeks post-surgery. The stiffness (A), maximum force (B), stress at failure (C), and energy absorbed at failure (D) of the TSPC-treated group was higher than other two groups. *P < 0.05. N = 5 per group.
of stem cells because the biophysical and biochemical properties of the decellularized matrix is relatively native, such as the native parallel arrangement of collagen fibers of the ECM leading to the right direction of engraftment stem cells and suppressing the wrong direction of commitment. Our previous study showed the tendon ECM not only enhanced tenogenic differentiation of TSPCs, but also significantly suppressed osteogenic-lineage differentiation.

In comparison to the study by Kryger et al., both studies proved tenocytes and MSCs could be used to successfully repopulate acellularized tendons for tendon repair. The present study goes deeper into the in vivo application of these cells for tissue engineering. Furthermore, the bioactive factors of tendon extracellular matrix could quickly release into the microenvironment of stem cells and promote cell proliferation. Zhang et al. recently reported tendon extracellular matrix effectively stimulated the proliferation of rabbit patellar tendon stem cells in vitro.

Plenty of evidence has documented that tissue-specific stem cells are best for tissue engineering. Numerous studies demonstrated that TSPCs promoted tissue regeneration in vivo. As the local resident stem cells, TSPCs showed higher potential in repairing injured tendons compared to MSCs. This is consistent with our study, wherein the ECM + TSPCs group possessed a better anatomical structure, synthesized more collagen, and had more powerful biomechanical properties than ECM + MSCs. The regenerated tendons were closer to normal tendon tissue structure and mechanical properties than ECM + MSCs. The regenerated tendons were closer to normal tendon tissue structure and mechanical properties, and regulated inflammation in tendon healing via JNK and STAT3 signaling. However, it is known that the tenogenic differentiation capacity of TSPCs significantly decreases with advancing age. The expression of transcript factor Fos observably decreased in early post-natal rat Achilles tendons with time. Loss of tenomodulin (Tnmd) possibly resulted in augmented senescence and reduced self-renewal of TSPCs. Only an allogeneic cell source of TSPCs can be used for tissue repair because it is hard to obtain autologous cells without causing morbidity of the donor site. Hence, the aging and the small quantity will limit the development of TSPCs in the future.

The MSCs’ specialty of directed differentiation and paracrine mechanisms has important implications for tendon repair and regeneration. Many kinds of tissue-derived MSCs used to repair injured tendons have achieved good curative effects. In this study, the ECM + MSCs group possessed better functional healing of injured tendons than ECM group. This is consistent with previous results that show the tendon tissue matrix promotes the tenogenic-lineage differentiation of MSCs. Even though their therapeutic effect is lower than TSPCs, their wide variety of sources, ease of isolation and culture, and suitability for large-scale industrial development underline their huge potential.

The selection of suitable seed cells is critical to tendon tissue engineering. Although stem cells perform similarly to some extent, there are some unique properties of tissue-specific stem cells that may influence the tissue engineering outcome. TSPCs expressed a higher level of adipogenic, osteogenic, and chondrogenic besides tenogenic genes, suggesting they possessed the capacities of adipogenesis, osteogenesis, and chondrogenesis differentiation. When an optimized condition is obtained, the complete capability of TSPCs in tendon tissue engineering is achieved, which shows its potential. Large animal models may be required, with long-term follow up and cell tracking to appreciate the functional benefits in future.

Our study collectively demonstrated that decellularized extracellular matrix derived from porcine tendon significantly promoted injured tendon regeneration when combined with TSPCs or ESC-MSCs. Compared with ESC-MSCs, TSPCs combined with decellularized matrix showed more improvement in the structural and biomechanical properties of regenerated tendons in vivo. This study developed a practical strategy for functional tendon tissue regeneration and further studies are warranted to utilize TSPCs integrated with a tendon-derived decellularized matrix for tendon regeneration.

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Author Contribution
Haixin Song and Zi Yin contributed equally to this study.

Ethical Approval
This study was approved by the Zhejiang University Institutional Animal Care and Use Committee.

Statement of Human and Animal Rights
This study was approved by the Zhejiang University Institutional Animal Care and Use Committee, and we strictly followed the current laws for animal experiments.

Statement of Informed Consent
Statement of Informed Consent is not applicable for this article.

Declaration of Conflicting Interests
The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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