Loss of Tissue Regenerative Capacity in Aging - The Tendon

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Abstract
Intrinsic to the process of aging is the loss of the ability to regenerate of different organs and tissues, becoming more susceptible to aggression and impaired function. Tendon aging is a complicated process following, however, similar mechanisms of other tissues - proposed model of the "Hallmarks of Aging", by López-Otín, et al [1]. Tendon structure and cellular composition is also unique, which makes scientific research in this field both a very specific and select task. Mesenchymal Stem Cells (MSCs), more specifically, Tendon stem/progenitor cells (TSPCs) may be of key importance.

Tendon disease is one of the most common entities worldwide, and is currently increasing with the augment of the human life-expectancy. This review proposes to make a short summary of the state of the art in both the aging tendon and its available treatment target potentials.

Introduction
Aging is characterized by a progressive loss of physiological integrity, leading to impaired function and increased vulnerability to death [1]. This inherited life characteristic heterogenous form organs to tissues, where tendons as highly organized connective tissues comprise bottom-up assemblies of collagen molecules [2].

Tendon cells include tenocytes are differentiated cells committed with synthesis of scleraxis (Scx), a basic loop-helix-loop transcription factor; tenomodulin + (Tnmd), a transmembrane glycoprotein, as well as with the tendon extracellular matrix (ECM) component collagen type I (Col-1) together with collagen type III, proteoglycans and glycosaminoglycans. Tendon stem/progenitor cells (TSPCs) fulfill the universal criteria of mesenchymal stem cells (MSCs): Clonogenicity, multipotency and self-renewal according with stimuli and respective cellular adaptation.

Tendon aging is a complicated process caused by multifaceted pathways and aging plays a critical role in the occurrence and severity of tendon injury [3].

Due to an increasingly aged population, the occurrence of tendon-related injuries is also increasing. No significant advances have been made in the treatment of such conditions in recent years.

The purpose of this review is to discuss the current evidence on tendon aging, its treatment, and how this can eventually create new targets for therapy.
Material and Methods

An analytical review conducted through PubMed database relied on original articles using “tendon cell aging” and “tendon cell regeneration” as main objectives. The search included only English written articles, without year of publication restrictions. The articles selected included those that reflected studies/reviews of aging mechanisms in a specific tissue: The tendon, evolving historical summing up knowledge, conductivity to cellular and matrix crosslinking, in order to allow medical understanding of the disease and its potential therapeutic targets.

Loss of Tissue Regenerative Capacity in Aging

Models to explain the process of aging have been proposed throughout years. In 2013 Carlos López-Otin, et al. proposed “The Hallmarks of Aging” enumerating nine hallmarks that represent common denominators of aging: Genomic instability, telomere attrition, epigenetic alterations, loss of proteostasis, deregulated nutrient-sensing, mitochondrial dysfunction, cellular senescence, stem cell exhaustion, and altered intercellular communication [1].

Deficient proliferation of stem/progenitor cells seemed to be detrimental the long-term resistance, while excessive proliferation might be deleterious by accelerating the exhaustion of the stem cell niches [1].

TSPCs represent a minor percentage of the overall tendon cell composition. These cells present features such as self-renewal, clonogenicity and multidifferentiation to be distinguished through the presence of stem cell markers [3].

Up-regulation of P53, P16, P14 and P21 genes related to cell cycle arrest and activation of the P53 and P21 pathways have been observed in aged TSPCs, which are thought to function as accelerators of the cellular aging process [4]. The imbalance in between suppressor genes seems to conduct to cell exhaustion in the tendon tissue.

Epigenetic changes involving alterations in DNA methylation patterns, post-translational modification of histones, and chromatin remodelling [1] support a series of significant cellular epigenetic alterations through age, viewed as age-related markers in TSPCs. These results are consistent with those obtained from other types of stem cells [3] and can be used in future studies.

Tendon Stem/Progenitor Cells

Pluripotent stem cells are supposed to differentiate into various tissue types under different genetic stimuli/conditions and serve as an internal repair system, according with embryonic layer of origin [3].

Compared with bone marrow stromal cells, TSPCs express higher levels of Oct 4, known to positively modulate mesodermal lineage differentiation and have greater ability to proliferate and get clonogenicity. Thus, TSPCs potentially represent a more appropriate cell source for the regeneration of musculoskeletal tissue, particularly tendon tissue, which has limited repair and healing abilities with traditional tenocytes [3].

Aging and Alterations in Epigenetic and Underlying Mechanisms

Aging and cell morphology

In vitro aged TSPCs exhibit cell star-like flattened shape, while youngTSPCs exhibit spindle-shaped morphology [4]. In addition, aged TSPCs are larger in size, have more podia, spread further, and exhibit more robust actin stress fibers and a higher actin content that distorts the balance of the actin cytoskeleton organization [4]. An increase in the size is often associated with cell senescence [5]. Additionally, aged TSPCs display a large, flat and heterogeneous morphology, while younger cells exhibit an uniform elongated morphology [6].

Aging and cell differentiation

The capacity of TSPCs to differentiate into tenocytes is reduced with ageing [7] consistent with the observation that capacity of differentiation of the TSPCs is markedly diminished during the aging process. Moreover, aged TSPCs are not sensitive to transforming growth factor-β3, a sublineage of the TGF-β superfamily that regulates cell growth and differentiation also engaged in matrix renewal [8].

Although, TSPCs have been reported to display a decrease in self-renewal and clonogenic capacities, multipotency has been maintained in vitro [4].

Multidifferentiation capacities of TSPCs are maintained during aging process without a conclusive determination of the trends in their variations, but most studies conclude that aging impairs the tenogenic differentiation capacity of TSPCs [3].

Aging and cell proliferation

According to Docheva, et al. (2005) [9], Tnmd is an established gene marker for the mature tendon/ligament lineage in vertebrates. Its protein has a highly conserved cleavable C-terminal cysteine-rich domain that has been identified as an important regulator for TSPCs proliferation and senescence as well as for tendon maturation. Tnmd loss-of-function in mice leads to a phenotype with distinct signs of premature aging on tissue and stem/progenitor cell levels [10].

Aged TSPCs showed a proliferation deficit after 120 days of culture and had an early plateau phase, while young TSPCs did not exhibit the plateau [4]. It has also been observed this decrease in TSPCs proliferation with increasing age, consistent with similar results observed in TSPCs from other aged vertebrate animals [5].

Another age-dependent important factor, is the ca-
pacity for these cells to form colonies (clonogenicity). Globally, clonogenic deficits in TSPCs are based in reduced colony number and lowering colony-forming unit efficiency verified in aging.

**Aging and cell viscoelasticity**

A variety of tissues express Tnmd mRNA, however Tnmd cleavage and secretion is exclusive to tissues undergoing tension, suggestive of a possible mechano-regulation mechanism [10].

Stiffening of human tendons was observed in aging or in diabetic individuals due to increased pathological cross-linking of collagen fibrils [11].

Denser, well-structured actin cytoskeleton in aged TSPC, seems to correlate with augmented cell stiffness [12]. Also a dense cytoskeletal organization might result in a larger cell size and anomalous cell shape conducting also to increase stiffness and viscosity [6].

**Mechanisms involved in the aging process**

Despite unclear molecular mechanisms involved, aging is known as a risk factor for tendon disorders. When TSPCs premature enter into senescence exhibit proliferation, differentiation and differentiation deficits. Many pathways and theories have been proposed up to date [3].

Forkhead box P1 (FOXP1) gene belongs to subfamily P of the fork head box (FOX) transcription factor family and plays important roles in various biological processes, including cell cycle progression, proliferation, and differentiation and senescence [13]. During TSPCs aging, FOXP1 was significantly reduced in both mRNA and protein levels; age-related dysfunction, such as self-renewal, migration and differentiation, appear to be a hallmark of TSPCs senescence [13].

There are also different mechanisms of the aged tendon to adapt to stressors with aging. The effect of these biological changes on the ability of resident tenocytes to respond to various stimuli is still relatively unexplored and present many avenues for future research suggesting differential pathways in males and females, supporting matrix turnover in males (increased matrix degeneration, apoptosis) but tissue preservation in females (decreased inflammation, apoptosis, matrix degeneration) [14].

**Rejuvenation of aged tendon (Stem/Progenitor Cells)**

Numerous factors are involved in altering tendon homeostasis. Macroscopic factors associated with an uncomfortable exercise intensity and microscopic factors associated with an impaired estrogen balance, deteriorated ECM conditions and inappropriate drug use, alter the features of TSPCs, particularly during aging [3].

Based on most recent development in regenerative medicine, Dale, et al. [15] induced human embryonic stem cells to differentiate into tendon-like cells in the presence of exogenous bone morphogenetic protein (BMP) 12 and BMP 13 and directed parthenogenetic stem cells to differentiate into tenocytes, similar to embryonal development.

**Discussion and Conclusions**

In human musculoskeletal injuries, specifically in tendon injuries, there is no optimal treatment strategy till this day. TSPCs can be the beginning of a new era in the treatment of these kind of injuries - either by being stimulated or by being transplanted.

Evidence suggests that tendon resident cells, including tendon MSCs (including TSPCs) may be the main orchestrators directing tendon-regenerative processes. Such biological response upon injury may be further boosted by the administration of non-tendon MSCs.

An important role for increased Rho has been reported, associating coiled-coil forming protein kinase (ROCK) activity in accelerating the ageing progress of aged TSPC, changing back to a morphology similar to young TSPCs upon treatment with Y-27632, a common ROCK inhibitor [4]. These changes which are observed in tendon with increasing age, may inherit a high density of intrafibrillar covalent cross-links and this could regulate tensile fracture resistance mechanics [16], making it more rigid and less elastic - propitious to rupture. Treating aged TSPC with ROCK-inhibitor, might reverse these age-related changes and rejuvenating effect on cell morphology and stiffness may be acquired. We assume that cellular stiffness is a suitable marker for cell aging and ROCK a potential target for therapeutic applications of cell rejuvenation [12] - also a target for further research in the future.

As important as it is to understand the mechanisms of aging in the tendon, it is also paramount to understand how does the aged tendon respond to different stressors.

Moderate exercise ameliorates the depletion of the TSPC pool by upregulating the expression of cell proliferation and stem cell markers coupled with decreased lipid deposition, proteoglycan accumulation and calcification formation, and it is beneficial for delaying the undesirable effects of age [17].

Recently, the discovery of induced pluripotent stem cells (iPSCs), particularly cells isolated from mature adult, inspired researchers to develop potential therapies to cure clinical diseases and ponder the eternal topic of regaining our youth. Thus, iPSCs provided inspiration to reverse the stem cell fate by modulating the factors that influence cell growth [3]. Mechanical stretching improved the tenogenic differentiation of both these iPSCs and pre-MSCs [18].

When tenocytes are exposed to a certain amount
NSAIDs, the differentiation of MSCs to tenocytic lineage gets impaired and drawn toward adipocytic lineage [19]. Different concentrations of ascorbic acid have been used in tenocyte cultures to enhance collagen synthesis - ascorbic acid is a well-characterized antioxidant that can promote collagen biosynthesis and prevent free radical formation [19]. Because of its acidity, the exact concentration of ascorbic acid to use is still unknown.

Inflammation does not seem to affect the proliferation rate of the isolated TSPCs and the tenogenic marker gene expression [20]. Supporting the theory: that the long head of the biceps tendon with and without tendinitis may be a novel source of tendon derived stem cells, which might facilitate treatment of degeneration and induction of regeneration in shoulder surgery, like J Schmalzl, et al. [20] proposed.

Pioglitazone can be used to enhance the therapeutic effects of mesenchimal stem cells for tendon repair, as shown by the increased proliferation, migration, remolding gene expression, and ECM secretion of tenocytes [21].

Various types of biophysical stimulations, such as ultrasonication and shock wave, were experimentally confirmed to be beneficial for promoting the differentiation and proliferation of MSCs [22].

As an effective, non-invasive, cheap, and safe ultrasonication, low-intensity pulsed ultrasound (LIPUS) was shown to accelerate bone-tendon healing, the mechanism of which might be related to its influence on the local mechanical and biologic environment and thus its ability to induce lineage-specific differentiation of the MSCs around the healing interface [22].

To this date, tendon aging can be correlated with TSPC aging. Limited studies, have been done to explore all theelaborate cell repair mechanisms in the tendon.

As life expectancy augments, so do tendon injuries. No inovative treatments/strategies have appeared in recent years. It is of utmost importance that we understand effectively how does in fact the tendon ages and adjust accordingly: Alternative guided treatment strategies that can be offered to clinicians around the globe.

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