Biologics for tendon repair*

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Abstract

Tendon injuries are common and present a clinical challenge to orthopedic surgery mainly because these injuries often respond poorly to treatment and require prolonged rehabilitation. Therapeutic options used to repair ruptured tendons have consisted of suture, autografts, allografts, and synthetic prostheses. To date, none of these alternatives has provided a successful long-term solution, and often the restored tendons do not recover their complete strength and functionality. Unfortunately, our understanding of tendon biology lags far behind that of other musculoskeletal tissues, thus impeding the development of new treatment options for tendon conditions. Hence, in this review, after introducing the clinical significance of tendon diseases and the present understanding of tendon biology, we describe and critically assess the current strategies for enhancing tendon repair by biological means. These consist mainly of applying growth factors, stem cells, natural biomaterials and genes, alone or in combination, to the site of tendon damage. A deeper understanding of how tendon tissue and cells operate, combined with practical applications of modern molecular and cellular tools could provide the long awaited breakthrough in designing effective tendon-specific therapeutics and overall improvement of tendon disease management.

Keywords

Tendon; Tendon repair; Growth Factors; Cell-based therapy; Mesenchymal stem cells; Embryonic stem cells; Tendon-derived cells; Natural biomaterials; Gene therapy

1. Introduction

Tendons are unique forms of connective tissue that connect and transmit forces from muscle to bone [1]. They are able to store elastic energy and withstand the high tensile forces upon which locomotion is entirely dependent [2]. This review article is designed:

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(1) to provide background information on the clinical relevance of tendons and to remind the reader of the lengthy and incomplete nature of the native tendon repair process. This motivates the urgent need for improving the outcome of tendon repair; biologics offer attractive possibilities in this regard;

(2) to introduce the basic tissue and cellular organization of tendon and its major tendon-specific molecules (Sections 1.1–1.3);

(3) to summarize the results of studies based on the four main approaches - growth factors (Section 2.1), stem cells (2.2), natural biomaterials (2.3) and gene therapy (2.4);

(4) to discuss critically unresolved issues.

We have focused on in vivo studies of the repair of tendon injury, and only in some cases included in vitro examples to strengthen certain points.

1.1. Tendon clinical relevance

Primary disorders of tendons (tendinopathies), due to overuse or age-related degeneration, are widely distributed clinical problems in society, possibly resulting in acute or chronic tendon injuries. Hospital evidence and statistical data suggest that certain tendons are more prone to pathology than others; these are the rotator cuff, Achilles, tibialis posterior and patellar tendons, whose pathologies are often based on a degenerative process. In addition, the extensor and flexor tendons of the hand and fingers are frequently subjected to direct lacerations at all ages. Although there are no accurate figures specifically relating to tendon disorders, studies from primary care show that 16% of the general population suffer from rotator cuff-related shoulder pain [3] and this rises to 21% when the statistics shift to elderly hospital and community populations [3,4]. These numbers further increase in the sports community; for example, Kannus reported that 30 to 50% of all sporting injuries involve tendons [5].

Although there are a number of studies discussing this issue, there is still a need to clarify the classification and terminology of the different tendon pathologies. This situation is mainly due to the clinical problem that tendon biopsies are generally difficult to obtain and that this material is usually collected at the end-stage of the condition or after tendon rupture. In general, the major conditions affecting tendons are tendinitis and tendinosis; the first assumed to be accompanied by inflammation and pain, whereas the second can be caused by tendinous degeneration [6]. It is believed that these conditions are rarely spontaneous [7] and are not caused by single factors. Rather, they are the end result of a variety of pathological processes [8,9] which can ultimately lead to the main clinical problem: loss of tissue integrity with full or partial rupture of the tendon.

Many factors are likely to be involved in the onset and progression of tendinopathies. Intrinsic factors include age, gender, anatomical variants, body weight, and systemic disease. Extrinsic factors include sporting activities, physical loading, occupation, and environmental conditions such as walking surfaces or footwear [8,9]. In addition, it has been reported that genetic polymorphisms affecting collagen fiber formation [10] or even blood group [11] are associated with tendon injuries and tendinopathy.
Hence, tendinopathies represent major medical problems associated with physical activity and age-related degeneration. Unfortunately, due to hypocellularity and hypovascularity, the natural healing ability of tendons is extremely low and inefficient [12]. Nevertheless, healthy tendon tissue has the potential to heal by itself as long as the ruptured parts have contact and the well vascularized peritendinous tissue, the so-called paratenon, is intact [13]. There is continuing debate on whether to treat acute Achilles tendon ruptures operatively or conservatively [14]. Both options have their advantages. In the case of rotator cuff pathologies, the choice of therapy very much depends on the patient's age, degree of tendon degeneration and extent of laceration [15]. A ruptured patellar tendon needs to be treated operatively to restore the extensor apparatus of the knee [16]. The treatment of tibial tendon insufficiencies is stage-dependent [17].

The main surgical repair techniques aim to re-establish tendon alignment by suturing the ruptured ends together, which requires a non-degenerate tendon with healing potential. The reconstructions are limited by the tendon's biology. Sometimes an allograft is used to bridge certain defects, while use of allografts has increased in recent years [18,19]. When autografts are used, a certain donor side morbidity must not be neglected. And in both cases, ingrowth of the bridging graft is necessary, requiring good tissue conditions without degeneration. It is estimated that $30 billion are spent on musculoskeletal injuries in the United States each year and tendon/ligament injuries represent approximately 45% of these cases [20]. In addition, surgical repairs are often unsuccessful in which case the majority of these injuries become essentially chronic conditions that are prone to recur [21].

In summary, tendon disorders are common, debilitating conditions affecting both the working population and recreational athletes. Their etiology remains controversial, particularly in understanding which factors are primary and which are secondary to the disorder. Moreover, these conditions not only have an impact on peoples' quality of life, but also represent an enormous economic burden on the worldwide healthcare system. Therefore, it is of great importance to identify key molecular and cellular processes involved in the progression of tendinopathies and subsequent ruptures in order to develop effective therapeutic strategies for treating them.

1.2. Tendon molecular composition and cell niche

The extracellular matrix (ECM) of tendons is composed of collagen and a smaller fraction of elastin embedded in a hydrated proteoglycan matrix. The principal role of the collagen fibers is to resist to tension, whereas proteoglycans are primarily responsible for the viscoelastic properties of the tendon. The smallest structural unit is the collagen fibril. Each fibril is built from soluble tropocollagen molecules forming cross-links to create insoluble collagen molecules which then aggregate progressively into microfibrils, fibrils and finally into fibers. Bundles of fibers are bound together by thin layers of loose connective tissues known as the epi- and endotenon, which allow the fiber groups to glide on each other in an almost frictionless manner; they also carry blood vessels, nerves and lymphatics to the deeper portion of the tendon [2]. The smooth gliding of tendons as they move is aided by the lubricating molecule, lubricin [22]. Altogether this complex, three-dimensional, internal
ultrastructure endows the tendon with high tensile force and resilience, while preventing
damage and separation of the fibers under mechanical stress [8] (Fig. 1).

Collagen type I is the most abundant molecule in the ECM, accounting for almost 60% of
the dry mass of the tissue and approximately 95% of the total collagen [8]. Type III is the
next most abundant collagen [23]. Normally, collagen type III is restricted to the tendon
sheets; however, it is found abundantly in pathological tendons and it is also the first
collagen to be produced in high quantity during tendon healing [8]. Other collagens in
tendon include types V, VI, XII, XIV and XV.

Besides collagen fibers, the tendon ECM is composed of many other components including
elastic fibers, the ground substance and inorganic components. In general, elastic fibers
ensure tissue flexibility and extensibility, permitting long-range deformability and passive
recoil without energy input [24]. Furthermore, they are thought to be involved in the
recovery of the crimp pattern of the collagen fibers after tendon stretching [25]. The ground
substance comprises hyaluronan, proteoglycans (decorin, biglycan, fibromodulin, lumican),
structural glycoproteins and a wide variety of other molecules. The highly viscous and
hydrophilic nature of the ground substance provides spacing and further support of the
collagen fibers. Water makes up 60 to 80% of the total weight of the ground substance,
whereas proteoglycans account only for 1–2% [23].

Mature tendons are normally characterized by low cellular density (Fig. 1). Approximately
90–95% of the cellular content of tendon comprises tendon-specific cell types described in
the literature as tenoblasts and tenocytes, the latter being terminally differentiated [26].
Other cell types include the synovial cells of the tendon sheaths, chondrocytes at the
pressure and insertion sites, and vascular cells. Tendon cells are able to synthesize all
components of the tendon ECM with a peak activity during growth and a gradual decrease
during aging [26]. It is thought that the low metabolic rates with anaerobic energy
production typical of mature tendon cells can reduce the risk of ischemia and necrosis,
especially during the extended periods of tensional stresses to which tendons are usually
subjected. On the other hand, this feature is a disadvantage for tendon recovery and healing.
Conversion of tenoblasts to tenocytes might occur in response to various stimuli such as
exercise and trauma in which higher rates of proliferation and matrix remodeling are needed
[26].

In 2007, Bi et al., identified within human hamstring tendons a novel cell population of
resident tendon stem/progenitor cells (TSPC) [27]. It was shown that the TSPC exhibit
classical adult mesenchymal stem cell (MSC) criteria such as presence of specific surface
antigens, self-renewal, clonogenicity and three-lineage differentiation (adipogenic,
osteogenic and chondrogenic), but also that they express tendon-related genes such as
scleraxis and tenomodulin, and are able to form tendon and enthesis-like tissues when
implanted in vivo. The existence of TSPC was further confirmed in subsequent studies with
human, equine, rabbit, rat and mouse tendons [28–35]. Whether adult TSPCs represent a
residual population of the embryonal tendon progenitors remains still unclear. Furthermore,
there is a need for studies demonstrating the exact roles and location of TSPC during tendon
maintenance and healing as well as their exact relationship to tenoblasts and tenocytes. At
present, the direct comparison of TSPCs to tenoblasts and tenocytes is impeded by the lack of molecular markers allowing their precise identification and hence the isolation of pure subsets of cell populations along the tendon differentiation cascade. Despite the incomplete understanding of TSPC nature and function, they represent a potential cell source for treating injured tendons. How the tendon research field foresees using TSPC in tendon repair will be discussed later in the review.

1.3. Tendon healing

The lack of detailed molecular and histopathological studies on tendons has hampered our understanding of the mechanisms underlying tendon healing. Some evidence has been obtained from animal models of experimentally induced tendon damage [36,37]. In general, the healing process of an injured or compromised tendon passes throughout three main phases containing distinctive cellular and molecular cascades (Figs. 2 and 3). These phases can overlap and their duration is dependent on the location and severity of the disease [38–40]. The initial inflammatory stage begins with the formation of a hematoma shortly after injury. Inflammatory cells such as neutrophils, monocytes and macrophages are attracted to the injury site by pro-inflammatory cytokines [38]. Secreted angiogenic factors initiate the formation of a vascular network, which is responsible for the survival of the newly forming fibrous tissue at the injury site. Despite being profuse and haphazard, the initial vascular response is essential, since it has been shown that diminution of blood supply impairs healing [41]. Next, components of the ECM, pre-dominantly collagen type III, are synthesized by recruited fibroblasts. After a few days, the proliferation stage takes place accompanied by the synthesis of abundant ECM components, including proteoglycans and collagens (mostly collagen type III), which are arranged in a random manner. Further features of this stage are increased cellularity and the absorption of large amounts of water. The remodeling stage includes two sub-stages; it begins 6–8 weeks after injury and takes around 1–2 years depending on the age and condition of the patient. The first sub-stage, consolidation, is characterized by a decrease in cellularity and matrix production, as the tissue becomes more fibrous through the replacement of collagen type III by collagen type I. Collagen fibers then start to organize along the longitudinal axis of the tendon, thereby restoring tendon stiffness and tensile strength. After approximately 10 weeks, the maturation stage starts, which includes an increase in collagen fibril crosslinking and the formation of more mature tendonous tissue.

The tendon healing process is complexly orchestrated by a variety of secreted molecules [42]. Initially, certain inflammatory cytokines, such as interleukin (IL)-6 and IL-1β, are produced by the invading inflammatory cells. Later, tissue repair is facilitated by a number of growth factors, which are released by cells located at the injury site. bFGF (basic fibroblast growth factor), BMPs (bone morphogenetic proteins)-12, -13, and -14 also known as GDFs (growth and differentiation factors) -5, -6 and -7 respectively, TGFβ (transforming growth factor beta), IGF-1 (insulin-like growth factor-1), PDGF (platelet-derived growth factor) and VEGF (vascular endothelial growth factor) are involved in different phases of the healing process with diverse molecular effects (Fig. 3). During the repair process, tendon cells are activated and both synthesize and degrade ECM components, thereby participating in the slow, continuous process of tendon remodeling [39,43,44].
Two cellular mechanisms of tendon healing, known as extrinsic and intrinsic healing, have been suggested [41,45]. It is now believed that these two mechanisms normally act cooperatively. The hypothesis is that first fibroblasts and inflammatory cells from the tendon periphery, blood vessels and circulation are attracted to the injured site contributing to cell infiltration and the formation of adhesions. Thereafter, intrinsic cells from the endotenon are activated as they migrate and proliferate at the injury site, reorganizing the ECM and giving support to the internal vascular networking [38,46]. The origin of the reparative cells remains in debate. In 2007, an enlightening study from Kajikawa et al., used a model of tendon injury applied to two different chimeric rats, one expressing green fluorescent protein (GFP) in circulating mesenchymal cells, and the other in the patellar tendon. The data were consistent with the biphasic pattern of tendon healing. This comprises an initial invasion of circulating MSCs followed by the activation of local cells, which participate in the proliferative phase and carry out the long remodeling phase [45].

In most patients, especially aged individuals, the healed tendon usually does not regain the mechanical properties of the uninjured tissue. The reduced strength of the repaired tissue compared to the native tendon results from reduced integration of collagen fibers with a higher ratio of collagen type III to collagen type I. As a consequence, the tendon thickens and stiffens to overcome the lower unit mechanical strength; thus the tendon quality and its functional activity are inferior to that of healthy tendon.

2. Current biological strategies to augment tendon repair

Experimental approaches for enhancing tendon repair consist mainly of applying growth factors, singly or in combination, stem cells in native or genetically modified form, and biomaterials, alone or cell-loaded, at the site of tendon damage. In the last decade, the number of studies investigating the functionality of the above strategies has progressively increased (Fig. 4). This section of the review will focus primarily on in vivo studies. It will critically discuss progress and the remaining open questions for future research to address in order to improve the therapy of damaged tendons. Two of the most difficult tasks that researchers are facing during regeneration of tendinous tissues are: first, to achieve the regeneration of a highly specialized and three-dimensional organized matrix, whose formation implies not only biological but also mechanical constraints; and second, when using stem or progenitor cells, to prevent inappropriate plasticity of the exogenous cells, or the transdifferentiation of the local tenocytes into undesirable lineages leading to, for example, in situ adipose, cartilaginous or bone tissue formation.

2.1. Growth factors

Tendon injury stimulates the production of a variety of growth factors at multiple stages in the healing process [40,42] leading to increased cellularity and tissue volume [47]. Increased expression of growth factors is particularly prominent in the early phases of healing [48,49]. The following growth factors are important in tendon healing: bFGF, BMP-12, -13, -14, CTGF (connective tissue growth factor), IGF-1, PDGF, TGFβ, and VEGF [49–52]. In the following section these factors are briefly introduced before describing in vitro and in vivo experiments investigating the role of the factors in tendon healing (Table 1). No human
study investigating recombinant growth factors in tendon healing has been published in the literature.

2.1.1. bFGF—Chang et al., found upregulated bFGF mRNA in mature tenocytes and in fibroblasts and inflammatory cells surrounding the healing site in the tendon sheath [53]. Being elevated early in the healing process [48,49], bFGF is well positioned to promote the early events in tendon healing [54].

2.1.2. BMP—BMP-12, -13, and -14, also known as GDF-7, -6, and -5 respectively, stimulate mitogenesis, and are established tenogenic factors with the potential of driving differentiation of MSC in vitro [55] and in vivo [56]. BMPs are elevated early in the tendon healing process, gradually decreasing thereafter [48,49]. BMP-2 plays a role at the enthesis, the anatomical junction of tendon and ligament to bone. New bone formation can be induced by BMP-2 within a tendon with comparable characteristics to the enthesis. However, in intratendinous healing this bone formation is clearly undesirable [57–59].

2.1.3. CTGF—In contrast to the previously described factors, CTGF exhibits a sustained increase in gene expression persisting over 21 days during healing of chicken flexor tendons [50]. In the rat supraspinatus injury model of Würgler-Hauri et al., CTGF was moderately expressed in both the insertion and midsubstance area throughout all time points [49].

2.1.4. IGF-I—IGF-1 induces tenocyte migration and increases synthesis of the ECM, including collagen [60]. Elevated IGF-1 mRNA and protein expression levels were found in healing rabbit ligaments 3 weeks after injury and in healing equine tendons after 4 to 8 weeks [61,62]. IGF-1 seems to be particularly important during the formation and remodeling stages of healing.

2.1.5. PDGF—Increased PDGF-levels have been found in healing tendons [63]. Elevated expression of the PDGF receptor β was found by Chan et al., to persist for over 6 months after tendon injury, potentially indicating the important role of PDGF during the entire tendon repair period [64].

2.1.6. TGFβ—Besides tendon cell migration and mitogenesis, TGFβ especially stimulates production of the ECM, including increases in the production of collagen types I and III by all the 3 isoforms TGFβ1, TGFβ2, and TGFβ3 [65]. High levels of expression and activity of TGFβ are found throughout the course of tendon-healing [66,67]. Resident tenocytes and infiltrating cells from the surrounding tendon sheath show increased expression of TGFβ1 mRNA [68]. Correspondingly, TGFβ1/3 receptor (CD 105; endoglin) expression was also found to be upregulated at the repair site [69]. Juneja et al., found a biphasic pattern of TGFβ expression corresponding to an early peak of TGFβ1 and a late peak of TGFβ3 expression during healing [70]. Heisterbach et al., also found early and late peaks of TGFβ1 expression [48]. However, there are also data indicating that TGFβ1 provokes increased fibrotic scar formation resulting in tendon adhesions [71,72]. In a rabbit model adhesions were reduced using an anti-TGFβ1 antibody, but were not further influenced by the addition of an antibody against the isoform TGFβ2 [66]. Possibly an imbalance between the TGFβ1-induced ECM-formation and tendon remodeling is responsible for the formation of
adhesions [73,74]. Thus, defining the appropriate doses and combinations of isoforms could be essential for the successful application of TGFβ in tendon healing.

2.1.7. VEGF—Angiogenesis is important in both tendon degeneration, in cases of impaired blood supply, and in regeneration, for which the best possible capillary permeability is desirable [41]. VEGF promotes angiogenesis in tendon healing [75], and its activity rises after the inflammatory phase, especially during the proliferative and remodeling phases. In a canine model of tendon transection, VEGF mRNA peaked 10 days after surgery [76].

2.1.8. Effects of different growth factors on tendon healing—Based on the presence and influence of growth factors on tendon healing a number of studies has been published with the aim of understanding the influence of growth factors on tendon biology in vitro and on tendon healing in vivo (Table 1). For in vivo studies, the growth factors can be applied by local injection, percutaneously or operatively, or by implanting scaffolds or even suture material [77–79] containing growth factors.

Growth factors are rapidly cleared following local injection, but their persistence may be prolonged using scaffolds or coated suture material. There have been few investigations of growth factor release by coated suture material and scaffolds in tendons, but there have been several studies investigating the local application of growth factors. Local injection of TGFβ into the healing site of patellar tendons in rats significantly increased the load to failure [80]. Comparable results were found in flexor tendons of rabbit treated with VEGF, as long as the plantaris tendon was preserved. In this study expression of TGFβ was significantly elevated early in the healing course. It remains unclear whether the positive effect was caused by the VEGF therapy itself, the increased TGFβ expression provoked by VEGF, or both [81].

Interestingly native cells from different areas of the tendon tend to react differently when treated with TGFβ. Type I collagen expression is down-regulated and type III expression up-regulated in endotenon cells compared to cells from the epitenon or the tendon sheath [82]. Possibly the up-regulation of collagen type III and the down-regulation of collagen type I by cells in the endotenon marks the beginning of tendon healing induced by TGFβ [81]. As well as differential expression of collagens by epi- and endotenon cells, increased mRNA expression for VEGF was found at the healing site of flexor tendons but not at the epitenon [83,84]. Increased cell proliferation and collagen production was also provoked by PDGF and bFGF. The effect was amplified by a combination of both factors. In this study VEGF and BMP-2 did not have the same positive effect [85]. Treatment with PDGF increased total DNA, collagen crosslinking and hyaluronic acid content resulting in improved functional movement, but did not improve the tensile properties of the healed tendon [86]. Combined treatment with bFGF and PDGF increases fibronectin deposition as part of the provisional matrix and angiogenesis/revascularization in canine flexor tenocytes [87]. Moreover PDGF stimulates the synthesis of proteoglycan, collagen, non-collagenous protein and DNA [88]; bFGF was shown to accelerate intratendinous healing in patellar tendons [89]. Also IGF-1 stimulates matrix synthesis and cell proliferation of tenocytes in vitro [90,91].

The positive effects of growth factors are not limited to intratendinous lesions; PDGF increased failure load of the medial femurotibial ligament [92] and several growth factors...
had a positive effect on the tendon–bone healing site of the rotator cuff [93,94]. Combinations of growth factors seem more potent than individual growth factors delivered singly. For example, in an in vitro model using rabbit flexor digitorum tenocytes, the highest cell proliferation rates were achieved by combining bFGF, IGF-1, and PDGF. The combination was effective at lower doses than when the growth factors were delivered singly [95].

2.1.9. Autologous sources of growth factors—As noted above, it seems that the interaction of several different growth factors is important in tendon healing. This could explain the observed effectiveness of concentrates of autologous growth factors, such as those contained within platelet rich plasma (PRP) or autologous conditioned serum (ACS) [96–99]. In contrast to purified growth factors, PRP has already been used clinically in patients with either tendinopathy or tendon injury. Gosens et al., found significant pain reduction in 36 patients with patellar tendinopathy after PRP injection [100]. In a prospective randomized controlled study of 27 patients after ACL-reconstruction, PRP had a positive effect on donor site healing in the harvested patellar tendon leading to pain reduction and smaller defect size in MRI controls after 6 months [101]. Moreover, the patellar tendon graft itself seemed to remodel faster during ACL-reconstruction after additional application of PRP [102]. In a rat Achilles tendon model percutaneous administration of PRP 6 h after transection and resection of 3 mm tendon resulted in an increased tendon callus and strength by about 30% after 1 week, which persisted for as long as 3 weeks after injection [96]. However, after surgery to repair acute Achilles tendon rupture in patients Schepull et al., did not find improvement in healing after additional administration of PRP to the healing site [103]. In chronic Achilles tendinopathy de Vos et al. failed to find differences in pain or activity level between patients treated with PRP or saline [104].

A later systematic review de Vos et al. found no evidence of efficacy using PRP to treat chronic lateral epicondylar tendinopathy [105]. Along these lines, Hall et al. stated in their review on PRP entitled “platelet rich placebo?” that the only reasonable use for PRP was for therapy of refractory cases of lateral epicondylar tendinopathy, but not for other tendinopathies or tendon repair [106]. However, because PRP is a variable, poorly characterized cocktail of growth factors and other substances it is difficult to draw strong conclusions.

Several different devices are approved by the FDA (U.S. Food and Drug Administration) in the United States for generating PRP, resulting in different compositions of growth factors and even cells (leucocytes and erythrocytes). Moreover, PRP contains components other than growth factors, including interleukins, chemokines, proteinases, inhibitors of proteinases, adhesion molecules, sphingolipids, thromboxanes, purine nucleotides, serotonin, calcium, and many other mediators. PRP is considered to have anti-inflammatory properties, but some components, such as IL-1, -6, and -8, are pyrogens [107,108]. Thus the precise combinations and concentrations of the different factors within PRP are important determinants of the properties of this autologous blood product. This could explain the lack of activity described by Schepull et al. [103] and de Vos et al. [104].
randomized controlled trials using PRP formulations of standard, reproducible composition are needed to determine whether PRP is useful in treatment of tendon disorders [109].

2.2. Stem cells

Cell-based tissue engineering is one of the most attractive and widely explored approaches for musculoskeletal regeneration. This strategy relies on reparative cells, alone or in combination with biocompatible scaffolds, which are delivered intra-operatively to the site of tissue damage. Selecting the appropriate cell type is one of the most important factors to be considered in such applications.

With regards to tendon engineering, several cell types, including MSCs from different tissue sources (bone marrow (BM), adipose tissue (AD), embryonic stem cells (ESCs), induced pluripotent stem (iPS) cells and TSPCs) are suggested as suitable targets (reviewed in [110–115]).

2.2.1. BM-derived MSCs—MSCs for tendon tissue engineering can be easily obtained from a BM aspirate. Although they represent only 0.001–0.01% of the total cell population, they can be expanded to higher numbers in vitro [116]. When appropriately stimulated, BM-MSCs can differentiate into various mesenchymal cell types, including osteoblasts, chondrocytes and adipocytes [117]. Attempts to commit BM-MSCs to the tenogenic lineage have been based on treatment with growth factors such as GDF-5 (BMP-14) and GDF-7 (BMP-12) [118,119], or upon genetic transduction with BMP-2 and active SMAD8, BMP-12, BMP-13 or scleraxis cDNA [120–122].

Overall, these attempts have been moderately successful; although the treated BM-MSCs adopted a tendon-like cell phenotype in vitro, it is still unclear whether the phenotype remains stable when the cells are implanted into a tendon lesion. One very attractive, potential feature of BM-MSCs is the possibility that they are hypoimmunogenic, therefore allogeneic transplantation may not require immunosuppression; furthermore they can exert immunomodulatory effects on various blood cell types resulting in anti-inflammatory impact during tissue repair [123]. It has been also suggested that these cells exercise in vivo potent trophic and stimulatory functions on local progenitors, thus contributing to tissue regeneration in this alternative manner, rather then differentiating on site into tissue-specific cell types [124]. Possible difficulties when using BM-MSCs for tissue repair include painful BM harvesting procedures, lengthy periods for cell expansion, uncontrollable differentiation in vivo into undesirable cell lineages and reduced qualities with donor age [123].

In comparison to other tissue sources, BM-MSCs are the best studied and characterized, and therefore the most frequently evaluated cell type for the repair of tendon tissue [125]. The majority of the in vivo models consist of partial or complete surgical transection or collagenase-induced lesion of horse, rabbit or rat tendons. The tendon types that are typically investigated include Achilles, patellar and digital flexor tendons. A summary of relevant in vivo studies, based on BM-MSC therapy of tendon injury, and their outcomes is given in Table 2. Taken together, these studies demonstrated improved histological and biomechanical properties of the tendon, indicating an increased rate of tendon healing and maturation. However, in many of the models ectopic bone formation was described and
when biomechanically tested, the regained tendon strength was approximately 20–60% that of an uninjured tendon. In addition, only few studies have examined tendon healing after 6 weeks, thus the long-term effects of therapy on tendon strength, functional quality and performance or re-occurrence of the injury are unknown.

So far only few clinical trials have been conducted with BM-MSCs for therapy of tendons. Mazzocca et al. [126] isolated BM-MSCs from 11 patients during arthroscopic rotator cuff surgery. After cell expansion and treatment with insulin, the authors showed that the BM-MSCs gain features similar to those of tendon cells. In this study, however, the isolated cells were investigated in vitro and no implantation in the injured tendons was performed. Non-fractioned iliac-derived BM mononuclear cells have been injected into tendinous lesions in 14 patients with complete rotator cuff tear. After 12 months, the patients were evaluated with the UCLA (University of California, Los Angeles) score and MRI, both showing improved tendon healing and integrity. Only one patient had deterioration of tendon strength and pain after 1 year [127]. Despite the very preliminary nature of the above studies, the results suggested that BM-derived cells can be isolated, stimulated towards the phenotype of tendon cells and introduced into tendon defects. However, the tendon field is in great need of carefully designed, pre-clinical studies using large animal models aiming to: (1) monitor the fate of the implanted stem cells using different labeling techniques; (2) examine cell dose-dependent effects; (3) evaluate tendon properties after longer periods of times; and (4) standardize protocols and procedures, thus allowing direct comparison between different studies. Subsequent to this research, multicentre clinical trials can be initiated to validate the true potential and optimal mode of application of stem cells for the repair of human tendons. This approach is facilitated by the fact that BM-MSCs are already approved for human use in graft versus host disease, and are in a large number of human clinical trials for other indications. They are also used in veterinary medicine to treat several disorders, including teninopathies.

2.2.2. AD-MSCs—Subcutaneous adipose tissue is a source of stem cells that are very similar morphologically and molecularly to BM-MSCs. The AD-MSCs are also multipotent; however they are less efficient at osteogenic and chondrogenic differentiation, but excel in adipogenesis compared to BM-MSCs [128,129]. When treated with IGF-1 and TGFβ or GDF-5, AD-MSCs upregulate the expression of tendon-related gene markers, such as scleraxis and tenomodulin [55,130]. Kryger et al. [131] compared AD-MSCs to tendon-derived cells and BM-MSCs, and found comparable scaffold adherence and proliferation potential, hence suggesting AD-MSCs as alternative cell type for tendon tissue repair. When injected into horse superficial digital flexor tendons with collagenase-induced tendinitis, AD-MSCs improved collagen fiber organization and overall tendon structure [132]. Increased yield loads and energy absorption were described in a rabbit injury model of deep digital flexor tendons treated with AD-MSCs [133]. In sum, AD-MSCs are widely available, multipotent cells that are simple to obtain without high morbidity, and represent an attractive source of cells for tendon tissue engineering. Nevertheless, there has been little exploration of using AD-MSCs for tendon therapy [134]. A major difficulty is to restrict AD-MSC differentiation within the tendon defect site to the tendon cell lineage, avoiding their indigenous preference towards forming adipocytes. Nevertheless, the risk of heterotopic
ossification should be less than when using BM-MSCs [134]. Like BM-MSCs, AD-MSCs are already in clinical trials for other indications.

2.2.3. ESCs and iPS cells—ESCs constitute the inner cell mass of blastocysts and are able to produce all different cell lineages from the three germ layers [135]. Unlike MSCs, they can be passaged indefinitely. Therefore in various tissue and organ regenerative models, including cardiovascular, neuronal and pancreatic repair, ESCs have considerable advantages [135]. Despite some clear benefits of ESCs over adult stem cells, their application raises social and moral issues regarding disassembly of embryonal tissues [135]. Other obstacles that have to be resolved when using ESCs include teratoma formation and spontaneous differentiation. Therefore, in the recent years iPS cells, generated by genetic reprogramming of adult lineage committed cells, have been suggested as alternatives to ESCs [136]. These cell types have overlapping characteristics and yet iPS cells can overcome many current ethical concerns in ESC-based therapy.

So far ESCs and iPS cells have not been extensively studied for tendon repair due to the lack of protocols to differentiate these cells into the tendon lineage. Non-differentiated ESCs have been injected in horse tendon lesions 1 week after collagenase-induced tendonitis, and histology and ultrasound analyses demonstrated improved lesion size [137]. Guest et al., [138] directly compared the survival of ESCs to MSCs after implantation into surgically created tendon defects in the horse. Interestingly, ESCs were detectable up to 90 days post operatively, while less than five percent of MSCs survived. However, in these studies it remains unclear if ESCs successfully differentiated into tendon progenitors. Chen et al. [139] were the first to propose stepwise differentiation of human ESCs into tenocytes via an MSC intermediate step. Cell sheets of ESC-derived MSCs were engineered into tendon-like layers under static mechanical load in vitro and used to repair a window defect in the patellar tendon. The implanted cells were detectable at least 4 weeks after surgery, and the ESC-MSC-treated tendons were larger than the controls and contained continuous collagen fibers and cells resembling tenocytes. Importantly, because of the stepwise differentiation procedure, the risk of teratoma formation is greatly reduced and, indeed, was not observed in vivo.

A study by Xu et al. [140], was the first to report a positive effect of human iPS cell-derived neural crest stem cells combined with a fibrin gel on the healing of rat patellar tendon window defects. Histological and mechanical analyses demonstrated improved matrix synthesis and superior mechanical properties of defects treated with iPS cells. Interestingly, the authors also found that the transplanted cells produced fetal tendon-related matrix proteins, stem cell recruitment factors, and tenogenic differentiation factors, and accelerated the host endogenous repair process. The above results suggest that ESCs and, presumably iPS cells, can be applied safely in tendon regeneration after controlling their differentiation pathway.

2.2.4. Tendon-derived cells—Although knowledge of the differentiated cells resident in the tendon tissue has increased, still little is known about their precursors. Furthermore, the field is still lacking clear terminology on the different subsets of tendon cells. This is mostly due to difficulties to purify, expand, maintain and compare populations of pure stem cells,
progenitors, tenoblasts and tenocytes. For this reason we have given this section a unifying
title.

Stem/progenitor cells of mesenchymal origin are of great interest in understanding tendon
development and the healing processes. As mentioned previously, Bi et al., [27]
demonstrated that human and mouse tendons harbor an unique cell population that has both
stem cell but also tendon-specific characteristics. For example, these tendon-derived cells
expressed high levels of scleraxis, cartilage oligomeric matrix protein (COMP), tenascin-C
and tenomodulin, all tendon-related factors. Because the cells of this population showed
heterogeneity in their stem cell properties, the authors named them tendon stem/progenitor
cells (TSPC). When compared to BM-derived MSCs, TSPC were closely related, but not
different in terms of molecular marker profile and in vivo behavior. When the cells were
applied in vivo, TSPC formed tendon- and enthesis-like structures, whereas BM-MSC
formed bone- and BM-like structures. However, in this study, the cells were not used in a
clinically related, tendon defect model. In addition, this study showed that TSPC reside
within a unique niche, where the two extracellular proteoglycans biglycan and fibromodulin
control their functions by modulating BMP signaling.

The double knockout of these two proteoglycans is characterized by higher tendon
cellularity together with decreased collagen fibril thickness. TSPC isolated from these mice
had augmented clonogenicity and cell proliferation, but reduced collagen type I and
scleraxis expression. Lastly, Bi et al. [27] were the first to show that there is a link between
distorted TSPC functions and tendon pathology, since TSPCs within the biglycan/
fibromodulin-deficient tendon niche were far more responsive to BMP signaling, leading to
TSPC favoring the osteogenic lineage. In turn, this resulted in so-called in-tendon
ossification. Thus, the above data suggest that the molecular environment provided by the
niche is essential for the correct maintenance and differentiation of the stem/progenitor cells
during tendon development and repair.

Studies by Tempfer et al., [28] and Kohler et al., [29] also demonstrated the existence of a
TSPC population within human supraspinatus and Achilles tendons, respectively. Several
articles have suggested that tendon-derived stem cells (TDSC) can be isolated, expanded and
eventually used in regenerative strategies (reviewed in [141,142]). Purification and
expansion of a cell population containing only TDSCs is still difficult, because we lack
molecular markers discriminating the discrete steps of tendon cell lineage differentiation
from primitive stem cells via progenitors to mature tenocytes, as well as the incomplete
differentiation of the primary cells. Because of this, we have used in the text the term TSPC.
In order to unite and validate the existing data, the tendon field urgently requires: (1) to
standardize the protocols for TSPC enrichment; (2) to develop appropriate methods to
separate stem cells from progenitors; (3) to establish efficient methods for achieving
terminal tenogenic differentiation in vitro which will permit validation of TSPC properties;
and (4) to determine if TSPC differentiation in vitro reflects their differentiation capacity in
vivo.

The discovery of TSPCs had a major impact in the field, since TSPCs might be involved in
tendon tissue homeostasis and repair; alternatively, they can be used for practical purposes
in tissue engineering strategies for injured tendons. Still, there remains the need to clarify whether embryonic tendon progenitors and TSPCs are identical cell populations as well as to generate solid data concerning TSPC location and function in vivo. Tempfer et al., [28] have shown that cells expressing simultaneously tendon and pericyte-associated marker genes are localized to the perivascular space of tendon tissue, hence suggesting that this niche might be the source of local stem/progenitor cells. Still, tendons are poorly vascularized, hence the contribution of perivascular cells to the regulation of tendon cell fate and functions might be less pronounced than in tissues with high blood supply. Interestingly, Mienaltowski et al., [36] reported the existence of two different stem/progenitor populations within the peritenon and tendon proper of mouse Achilles tendons. More studies are required to reconstitute carefully the regional cell composition of tendons and the interconnections between different cell types. Improving our knowledge on the above questions can provide novel, fundamental understanding not only of the development of tendon tissues, but also of their sustainability and repair.

In terms of practical application, there are several challenging issues to solve prior the use of tendon-derived cells for tendon repair. Allogeneic cells may lead to an immune reaction, whereas autologous tendon-derived cells will avoid immune complications, but may lead to a comorbid state in the patient. Furthermore, during in vitro expansion, tendon-derived cells may undergo phenotypic drift. Yao et al., [143] have shown in human tendon cells that the ratio between collagen III and I increases with progressive passaging, while decorin expression significantly decreases. Schulze-Tanzil et al., [144] suggested that the differentiated state could be preserved if the cells are grown in three-dimensional pellets. A combination of ultrastructural, biochemical and molecular analysis indeed demonstrated that the phenotypic identity of the tendon-derived cells was retained when the cells were cultured in this fashion [144]. Therefore, it will be important to establish in vitro culture systems, based on three dimensional cultivation or by using mechanical strain, which can support unaltered TSPC identity over longer periods of time.

To date, only few studies have used tendon-derived cells in tendon repair models. Cao et al., [145], used autologous tenocytes to bridge tendon partial defects in hens and observed 14 weeks postoperatively that the engineered tendons displayed histologically a structure very similar to that of normal tendons. Similar conclusions were reached by Chen et al., [146] after introducing autologous patellar tenocytes cultured on porcine bioscaffolds into massive rabbit rotator cuff defects and analyzing tissue regeneration at 4 and 8 weeks. The authors concluded that implantation of tenocyte-loaded scaffolds results in superior tendon healing when compared to the control group receiving only scaffold. However, autograft tendon controls were still better then the engineered tenocyte constructs, suggesting that further optimization of the technology is required. Stoll et al., [147] investigated the healing of a partial Achilles tendon in the rabbit after filling defects with Achilles tenocytes loaded onto polyglycolic acid (PGA)/fibrin scaffolds. The authors generated novel scoring systems for macroscopic, histological and elastic fiber assessment of the progress of tissue repair. Although no clear advantage of the tenocyte-scaffold group was detected at 6 and 12 weeks post operation, this study provides a useful multifaceted scoring system for characterization and cross-study comparison of tendon healing models.
Tenocyte-based regeneration of full size Achilles tendon defects in rats has been compared to that based on BM-MSCs [148]. For bridging the defect ends, PGA and collagen type I scaffolds seeded with one or the other cell type were used and fixed with a frame suture. After 16 weeks, DNA from the implanted cells was detected in the regenerated tissue, consistent with their long-term survival. Despite evidence of central ossification in all study groups, biomechanical tests revealed that samples loaded with tenocytes had significantly better failure strength/cross-section ratios compared to defects receiving BM-MSCs or inseeded scaffolds. Ni et al., [149], were the first to describe histologically, biomechanically and ultrasonographically that TSPCs can improve the repair of a rat patellar tendon window defect. No ectopic bone formation was detected up to 4 weeks post-injury. How implanted tendon-derived cells influence local cells at the site of tendon damage is unclear and needs further clarification. More studies are also required to understand the fate and long-term effects of tendon-derived cells during tendon repair, and to compare the benefits of using these cells rather than MSCs from different sources. The main advantage of tendon-derived cells is that, being native in origin, when re-implanted into tendon defects, they will better accommodate to familiar environment and are likely to survive longer and differentiate more easily into terminal tenocytes.

Taken together, this area holds much promise for tendon therapy after clarifying many open questions. It will be of great importance to investigate further the identity, genetic marker profile, localization and in vivo functions of TSPCs as well as to carry out well-designed pre-clinical experiments to determine the role of TSPCs during the progression of tendon diseases and subsequent repair processes.

2.2.5. Unresolved issues in cell-based therapy for tendons—The involvement of each of the above cell types has to address a number of important challenges, such as determining: the amount of cells needed; whether combination with growth factors or genetic modification is helpful and, if so, which growth factors or genes to use; which scaffolds, if any, to use; the optimum time point of delivery. To define the ideal cell number, dose-dependent in vivo studies will be important. Most likely the optimal number of cells will vary relative to the size of the tendon defect, the type of cell type used and the particular tendon in need of repair. Therefore, it is necessary in forthcoming research to define and understand the exact mechanisms underlying the in vivo performance of different cell types. Crucial questions include: do the implanted cells provide trophic support and stimulate local progenitors or do they commit on-site into the tenogenic lineage? A useful approach to address such questions will be to develop or employ strategies to label and track the implanted cells during the different healing stages. In addition to determining the optimal ratio of implanted cells to defect dimensions, the field has to resolve how to obtain and multiply in a short time frame the required amount of stem cells, without harming their innate ability to differentiate or causing uncontrolled differentiation. Co-delivery of stem cells with growth factors or using genetically altered stem cells can enhance their qualities and navigate them quicker and more effectively into the preferred differentiation cascade. However, such manipulations could have poorly controlled side effects on local stem and progenitor cell populations.
The seeding of the stem cells onto different matrices can improve their maintenance and amplification at the site of tendon injury. Furthermore, the topography and mechanical properties of the carriers can be designed in a way to direct stem cell differentiation towards tendon progenitors or even mature tenocytes. The precise timing of stem cell delivery can be determined by the cell type and carrier combination. For example, when MSCs are to be used, implantation can take place already at the inflammatory stage since this cell type might exert a beneficial effect by participating in immunomodulation, reducing inflammation and stimulating local stem cells and tenoblasts. When tendon-derived cells are to be used, it might be more appropriate to introduce them at the later, proliferative stage when they can speed the endogenous healing process.

With regards to the timing of the complete procedure the following factors should be considered: (1) hospitalization time of the patient; (2) time to obtain, enrich and multiply the cells; (3) if the cells are modified or stimulated, the time necessary to carry out such procedures; (4) if the cells are applied in combination with biomaterials, the time for scaffold loading and cell adhesion. Ideal protocols should contain only a few short steps. For example, one way to speed up the preparation will be to purify the reparative cells from the primary cell milieu with cell-specific surface antigens. Finally, stem cells of autologous, allogenic or xenogenic nature should be compared. Each of these variants holds advantages and disadvantages. In autologous applications, it is important to determine donor site morbidity and the fitness of the stem cells since tissue aging or pathology can distort stem cell functions. Recent investigations focusing on BM-MSC demonstrated apparent age-related changes such as a reduced proliferation and clonogenicity as well as altered differentiation potential [150,151]. Baxter et al., [152] observed rapid telomere shortening and earlier entry into growth arrest and senescence; Kasper et al., additionally found reduced antioxidant defense, altered cytoskeleton organization and lower migratory capacity of aged BM-MSCs [153]. We have recently reported that human TSPCs from aged and degenerated tendons have significantly reduced proliferation capacity and premature entry into senescence, decelerated motion and delayed wound closure due to dysfunctional actin dynamics [29]. Hence, similar to other tissues, aging and disease exhaust the local stem cell pool in terms of size and functional fitness. Therefore, when such autologous cells are candidates for use in tendon repair in aged individuals, various possibilities to correct their endogenous deficits or to pre-activate them ex vivo or in situ via growth factor stimulation or gene therapy have to be carefully considered [154]. The development of suitable allograft donor cells would obviate many of these problems. However, the use of allogenic cells can also be associated with difficulties such as obtaining sufficient donor material, donor background diseases, prolonged storage of the cells and a possible deterioration of cell quality during storage. One major advantage of xenograft cells is that they can be available in large numbers and ready for use at any moment. However, their use is limited while possible zoonotic diseases and xenograft reactions have to be considered and addressed.

2.3. Biomaterials

Currently, tendon repair involves the use of autologous or allogeneic tendon transfer, which can restore tendon function in the affected area. However, both options have drawbacks, the first related to donor site morbidity and the second to risk of immune rejection. In addition,
rarely does the transferred tendon material match the tensile properties of the repaired tissue. Therefore, a number of biomaterials have been explored as alternatives to tendon transfer for tendon tissue engineering (reviewed in [155,156]). Some of these materials have been borrowed from the neighboring fields of cartilage and bone tissue engineering, and some have been specifically designed to resemble as close as possible the structural and biomechanical features of native tendon tissue. The ideal scaffold should cover a number of requirements such as: (1) to be biocompatible; (2) to support cell attachment and growth; (3) to have high surface area; (4) to promote tenogenic differentiation pathway; (5) not to induce host inflammatory responses; (6) when not biodegradable, to mimic native tendon architecture and mechanical properties. Furthermore, the scaffold should be easily reproducible, scalable, have good storage properties and, ideally, able to be customized.

Natural biomaterials include: collagen; silk; fibrin; hyaluronic acid; elastin; alginate; chitosan; porcine small intestine submucosa (SIS); human, porcine or bovine dermis; and decellularized tendon xenografts [155–157]. Most biomaterial studies have investigated how MSCs or tendon-derived cells respond to these materials in terms of cell adhesion, cell proliferation and survival over time, gene expression and differentiation [155–157]. Some of the studies have taken a step further into in vivo testing of the materials, alone or in combination with cells, and have examined host tissue reactions or tendon healing process (refer also to [115,157]). Some examples of studies on collagen-based scaffolds and xenografts will be discussed here.

### 2.3.1. Collagen-based materials

Collagen gels and composites, most frequently loaded with BM-MSCs, have been used for repair of different tendon gap models, as indicated in Table 1. In the articles of Young et al., [158] and Awad et al., [159] experimental groups treated with cell/gel implants achieved higher strength compared to suture-only controls. Interestingly, in the second study no additional benefit of increasing cell density in the collagen type I gel was found [159]. Another study showed that reducing cell to collagen ratio by 20-fold actually improved cell viability, lowered the degree of ectopic bone formation and enhanced the biomechanical properties of patellar tendon 12 weeks post-operatively [160]. It was suggested that material implants should exhibit physical properties similar to normal tendon tissue, but should be degradable. This would allow support and protection of the introduced cells in the early phases of the healing, but also replacement of the scaffold over time during de novo production of tendon matrix [160].

As mentioned earlier, critical design criteria for the ideal tendon graft requires the material to exhibit the mechanical properties of normal tendon, to facilitate functional integration and also to promote native tendon regeneration. Nanotechnology-based approaches allow development of various biomimetic scaffolds such as nanofibers and nanocomposites. Specifically, aligned nanofibers from collagen type I hold advantages because of their potential to mimic the matrix architecture of native tendon and, in turn, to regulate cellular responses. In vitro studies with cell-loaded aligned collagen I [161,162] convincingly showed that the aligned scaffold topography can induce a cell morphology similar to that of tenocytes, achieve matrix alignment and promote the upregulation of tendon-related genes such as scleraxis and collagen type XIV. Furthermore, the in vivo investigation by Kishor et
al., [161] reported that braided, aligned collagen type I fibers introduced in longitudinally incised rabbit patellar tendons undergo limited degradation and associate with a low-grade granulomatous inflammation. Additionally, quantitative histology revealed that the cross-sectional areas of tendons treated with the aligned scaffolds were larger and stiffer than controls. In sum, the above studies suggested that aligned nanofibers are superior to randomly oriented biomaterials, because they are biocompatible and, moreover, can stimulate the implanted cells to differentiate towards the favorable tenogenic lineage. Thus they have the potential to be used as carriers for tendon tissue engineering applications. One critical limitation of these scaffolds is scale-up to dimensions relevant for the repair of human tendon.

2.3.2. Xenografts—A possible way to overcome the difficulty of generating stable scaffolds in large sizes is to use xenograft tissues which have matching and customized proportions similar to those of human tendon defects. FDA-approved porcine SIS devices (Restore and CuffPatch) have been used in a number of laboratory studies of rotator cuff and Achilles tendon injury models performed in dog [163–165], rabbit [166] and rat [167]. Although the properties of healthy tendon were not fully restored, the studies reported positive histological and mechanical outcomes in comparison to non-treated defects. Furthermore, upon analyzing SIS degradation patterns, it was found that SIS is subjected to rapid degradation in the first 4 weeks after surgery, which suggests that it can serve as a temporary scaffold for quick cellular infiltration [165]. Following these encouraging results, multiple clinical studies were conducted with patients undergoing rotator cuff or Achilles tendon surgery (reviewed in [155]). Earlier investigation suggested successful tendon reconstruction with SIS devices in 11 out of 12 patients up to 2 years after the surgery [168]. However, subsequent investigations found that SIS-treated groups had no augmented properties and that SIS incorporation did not improve the rate of tendon healing [169–171]. The reasons for this discrepancy are not entirely clear and the major side effect reported in the above studies was a non-infectious effusion.

Decellularized tendon, of allograft or xenograft origin, is another tissue with promise for tendon repair. This application consists of harvesting tendon pieces from cadavers or animals which, after decellularization and slicing, are re-seeded with BM-MSCs, and finally packaged together into a single scaffold. Interestingly, when cultivated on such matrices, BM-MSCs exhibit a phenotype resembling tendon cells, suggesting yet again that the appropriate nano-topography and stiffness can enforce lineage differentiation [172–174]. It is logical to conclude that the best choice for tendon repair is tendon or ligament ECM; however, there are several unresolved difficulties with the use of decellularized tendon scaffolds, such as the poor cell repopulation of the deeper tendon layers and the observation that the decellularization procedure reduces the mechanical properties of the grafts.

Scaffolds derived from human cadaver (GraftJacket), bovine (TissueMend and Bio-Blanket) and porcine dermis (Permacol) have a rich collagenous matrix, retain native dermal ECM architecture and vascular channels, and have been approved by FDA for the reinforcement of soft tissues. More than a few studies have shown their efficacy in rotator cuff and Achilles tendon repair. For example, Adams et al., [175] studied the use of human cadaver dermis in a canine infraspinatus injury model and found robust tendonous tissue formation at
the site of scaffold implantation after 6 months. Bond et al., [176] reported improved shoulder mobility and decreased pain after implantation of human dermis in patients with rotator cuff tears. Positive outcomes after applying human dermis in the repair of large rotator cuff defects were also documented by Rotini et al., [177] and Snyder et al., [178]. The healing effect of human dermis materials on Achilles tendon ruptures was shown in a series of clinical studies published by Lee et al., [179] and Lee et al., [180]. Early return to physical activity, improved foot strength, reduced chronic pain and no re-ruptures were observed up to 30 months after surgery. Interestingly, Valentin et al., [181] examined histologically the host tissue response to porcine SIS and human, bovine and porcine dermis devices in a rodent model. The authors found that each biomaterial leads to a distinct form of tissue remodeling in terms of cellularity, vessels, inflammation and matrix organization. Interestingly, xenograft dermis scaffolds degraded slowly and were associated with higher inflammatory score and accumulation of denser and less organized fibrous tissue. Therefore, future controlled comparison studies are necessary to clearly define the advantages and limitation of each biomaterial.

In conclusion, natural biomaterials show significant promise for enhancing tendon repair especially in combination with reparative cells. At present, the tendon field still lacks classical biologics or tendon-specific drugs to aid repair. Therefore biomaterials, especially scaffolds mimicking native tendon architecture, represent a smart alternative option to solve one of the major problems of the field, namely to drive the reparative cells into the appropriate tendon cell lineage. However, the available data do not permit definitive conclusions, and often the use of identical materials, for example dermis scaffolds, produces variable results between different studies [175,181]. Thus, more investigations are required to improve and standardize the properties of biomaterials and to evaluate their role in the clinical practice of tendon repair.

2.4. Gene therapy

2.4.1. Concepts—It has been recognized for a long time that gene therapy has the potential to promote the repair and regeneration of damaged tissues, including tendons [182–185]. In this context, gene transfer is not used to compensate for a defined genetic defect, but instead to serve as a biological delivery system for the encoded gene products.

Gene transfer holds many advantages over traditional methods for delivering biologicals to sites of injury. In particular, it allows the local, focal production of gene products within and around the lesion in a sustained and potentially regulated fashion. Moreover, proteins synthesized locally as a result of gene transfer are likely to have undergone authentic post-translational processing leading to greater biological activity and a reduced risk of triggering neutralizing immune reactions. Gene transfer is particularly useful for delivering gene products with an intracellular site of action, including non-coding RNA molecules, signaling molecules and transcription factors; SMAD8 and scleraxis are relevant examples of the last two [120,122].

2.4.2. Vectors—Vectors are used to transfer genes (usually cDNAs) to target cells. Because viruses are naturally able to transfer with high efficiency their genes to the cells
they infect, they have been widely used as vectors. For this purpose, the viral genome is manipulated to remove sequences required for replication and virulence, while retaining those needed for infectivity. (The ability to replicate is retained in certain cancer gene therapy applications.) Therapeutic genes can be spliced into the genetic space generated by these manipulations to produce a viral vector that, in principle, can infect a target cell and deliver its genetic payload to the nucleus without replicating or causing adverse events.

Recombinant viruses so far studied experimentally for gene delivery to tendons and ligaments include adenovirus, lentivirus, retrovirus, adeno-associated virus (AAV). The main properties of these four vectors are compared in Table 3, bearing in mind that the many modifications made progressively to these vectors make simple generalizations increasingly difficult. Gene transfer with a viral vector is known as transduction.

Because clinical grade viral vectors are expensive and complicated, there is continuing interest in non-viral vectors for gene delivery. These raise less safety issues, are usually simpler to manufacture, have less restrictions in carrying capacity, often lower immunogenicity and should make quicker progress through the regulatory process for human use. Non-viral vectors can be as simple as naked, plasmid DNA. Often, the efficiency of gene transfer is improved by combining the DNA with a polymeric carrier or by using a physical stimulus such as electroporation. Non-viral gene transfer is known as transfection.

The properties of viral and non-viral vectors used in regenerative orthopedics have been reviewed in several recent publications (refer to [186–188]).

2.4.3. Strategies—Regardless of the vector, there are two general gene delivery strategies, in vivo and ex vivo. For in vivo delivery, the vector is introduced directly into the body by injection or other form of direct application. Because the cellularity of tendon is low, in vivo administration in this way should not lead to high levels of transgene expression. Nevertheless, there exist several examples of its successful application in animal models of tendon healing (Table 4).

An alternative in vivo application strategy uses a scaffold impregnated with vector; this is known as a gene-activated matrix (GAM). This concept has been applied to tendons by associating adenovirus vectors with a gelatin sponge [189] and by using allograft tendon as a scaffold for AAV in a process known as “allograft revitalization” [190].

During ex vivo delivery, cells are genetically modified outside the body and then injected or otherwise implanted at the appropriate site. Ex vivo delivery combines gene therapy with cell therapy and is increasingly popular when progenitor cells, such as MSCs, are used.

Although the methods of in vivo gene delivery are simpler than ex vivo delivery, the latter is presumed to be safer because viruses are not introduced directly into the body. Because the recombinant viruses used for this type of gene therapy cannot replicate, the cells that carry them do not shed infectious particles. It can, however, be argued that the cells used in ex vivo gene delivery may have been cultured in media containing xenogeneic components,
thereby introducing an element of risk, although the same would be true of ex vivo cell therapy in general.

Also, as noted, ex vivo gene delivery offers the possibility to combine the power of cell therapy with that of gene therapy. However, clinical application of such an approach is constrained by the present need to use autologous cells, which makes the process expensive and cumbersome. Development of suitable allogeneic cell lines for this purpose would greatly expedite the process.

To expedite ex vivo delivery, there is interest in developing technologies where suitable tissues that harbor accessible progenitor cells are harvested, genetically modified and reimplemented during one surgery [191,192]. Using a technique that was first developed for bone healing [193] genetically modified muscle grafts have been employed for tendon healing in animal models [194,195].

Although regulated transgene expression has not yet been explored in the context of tendon gene therapy, the availability of inducible promoters allows consideration of this approach. This reflects the likelihood that optimal healing may require the level of transgene expression to vary during the healing process. Also, such promoters allow the theoretical possibility of expressing one or more genes at different times from a polycistronic vector.

2.4.4. Progress—Early experiments confirmed the ability of various viral [196–199] and non-viral vectors [200–203] to deliver marker genes to ligaments and tendons by in vivo and ex vivo means. This work has been comprehensively reviewed by Hildebrand et al., [204]. Once marker gene delivery was achieved, it became possible to investigate the results of transferring genes with therapeutic potential.

As summarized in Table 4, most published studies using animal models of tendon repair have taken the approach of delivering a growth factor, especially one expected to promote the differentiation of progenitor cells into tenocytes. Promising results have been reported with BMP-12/GDF-7 [194,205] and BMP-14/GDF-5 [190,206,207], but not BMP-13/ GDF-6 [208], even though all three of these induce tenogenesis in other systems [56,209] and BMP-13 gene transfer to MSCs induces ligamentogenesis in vitro [121]. It is possible that mechanical factors account for this discrepancy [210]. Transfer of scleraxis has been shown to promote the differentiation of MSCs into tenocytes in vitro [122] and, when used ex vivo with MSCs, to enhance healing of the rotator cuff in a rat model [211]. Similar results were reported using a combination of BMP-2 and SMAD8 cDNAs to promote tenogenesis [120].

Other investigators have transferred cDNAs encoding growth factors that are not specifically associated with differentiation of tenocytes, but which may enhance cellularity, vascularity or the deposition of extracellular matrix. Examples include TGFβ [195], bFGF [212], VEGF [213] and PDGF-B [214,215]. In general, the results have been encouraging.

Because repair and regeneration are progressive, multi-step processes, there is also interest in delivering more that one growth factor. However, in the studies of Hou et al. [213], although ex vivo transfer of TGFβ cDNA improved healing in a rabbit Achilles injury...
model, VEGF did not and added nothing to the effectiveness of TGFβ. It is probable that different growth factors will be needed at different times during the healing process, in which case their genetic transfer will require staged delivery of vectors with different transgenes, or single delivery of a polycistronic vector with different, inducible promoters. The latter will be very difficult to navigate through the regulatory process and into human clinical use.

Inhibiting inflammation is an alternative approach to promote repair and regeneration, and successful use of IL-10 cDNA in this regard has been reported [216]. Anti-inflammatory gene products may also be of therapeutic use in tendinitis. IGF-1 gene transfer has been explored as a way of augmenting tendon structure in tendinitis [217].

Additional uses of gene transfer in treating tendinopathies are listed in Table 5.

There is interest in using osteogenic genes to enhance the incorporation of tendon into the osseous insertion site after reconstructive surgery [218–220]. Because the tendon at this site has a fibrocartilagenous zone, there is also the possibility to improve the function of the regenerate repair by promoting the formation of cartilage in this important area using chondrogenic genes [221,222].

Another application seeks to prevent the formation of fat or bone within tendon, a risk when using multipotent progenitor cells as agents of repair. In this case, the ex vivo use of tengenetic cDNAs, such as those encoding scleraxis [122,211], SMAD8 [120] or the appropriate BMP would steer the cells towards tenogenesis and prevent them from differentiating along adipogenic or osteogenic lineages. Heterotopic ossification can also occur without the addition of such cells as a result of injury. Inhibitory species of non-coding RNA that knockdown Runx2 or SMAD4 have shown promise in blocking this process [223, 224]. Knockdown of decorin has been explored as a way of inhibiting scar formation [225].

**2.4.5. Translation**—Approval of clinical protocols for orthopedic applications of gene therapy is a long, expensive and tedious process [226]. Although the literature, summarized here, holds promise of success for gene-based technologies to regenerate tendons, the data are preliminary and restricted to acute, small animal models. The optimal vector, transgene, promoter and delivery mechanism still need to be determined. Efficacy then needs to be confirmed in large animal models. Safety is a major issue for all gene therapy protocols, especially those involving non-lethal pathologies; the pharmacology and toxicology testing of gene therapeutics is complicated. The cost of a therapeutic is also a large factor in today’s economic environment, which is one reason to favor expedited approaches that do not require the ex vivo propagation of autologous cells [191,192].

**3. Concluding remarks**

Repair of tendon injuries is a lengthy process that frequently results in poor structural, mechanical and functional quality of the healed tissue. At present, the clinical options for treating tendon injuries are often unsatisfactory, especially in elderly populations. Therefore, several alternative strategies are being explored. The concept of biological therapy is
attractive, because it makes use of the body’s intrinsic potential to repair and heal its
damaged tissues [191]. When tissue regeneration is approached in this manner, the result is
expected to be natural, complete and lasting. As noted in this review, promising
experimental data support the concept of using proteins, genes and cells, often in
conjunction with biological scaffolds, as agents of the biological repair of tendons.
However, these strategies are still in the stage of pre-clinical development and optimization.
To reach their full potential and become realistic clinical options, further research focusing
on solving critically important challenges is needed (Graphical abstract).

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Abbreviations

| Abbreviation | Definition                                |
|--------------|-------------------------------------------|
| AAV          | Adeno-associated virus                    |
| ACS          | Autologous conditioned serum              |
| ACL          | Anterior cruciate ligament                |
| AD           | Adipose tissue                            |
| bFGF         | Basic fibroblast growth factor            |
| BM           | Bone marrow                               |
| BMP          | Bone morphogenetic protein                |
| COMP         | Cartilage oligomeric matrix protein       |
| CTGF         | Connective tissue growth factor           |
| ECM          | Extracellular matrix                      |
| ESC          | Embryonic stem cell                       |
| GAM          | Gene-activated matrix                     |
| GDF          | Growth and differentiation factor          |
| GFP          | Green fluorescent protein                 |
| IGF-1        | Insulin-like growth factor-1              |
| iPS          | Induced pluripotent stem                  |
| IL           | Interleukin                               |
| MSC          | Mesenchymal stem cell                     |
| MT1-MMP      | Membrane-bound matrix metalloprotniase 1  |
| PGA          | Polyglycolic acid                         |
**PDGF** Platelet-derived growth factor

**PRP** Platelet rich plasma

**Runx2** Runt-related transcription factor 2

**SIS** Small intestine submucosa

**SMAD** Small body size mothers against decapentaplegic

**Sox-9** Sex determining region Y - box 9

**TDSC** Tendon-derived stem cell

**TGFβ** Transforming growth factor beta

**TSPC** Tendon stem/progenitor cell

**VEGF** Vascular endothelial growth factor

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Fig. 1.
A schematic drawing of basic tendon structure. The collagen molecules are organized hierarchically in fibrils, fibers and fascicles. The cellular content is dominated by the tenocytes, which are terminally differentiated cells. Tendons contain stem and progenitor cell populations, whose exact location is still debated (therefore indicated with a ?). Different sheets, endotenon and epitenon (loose connective tissues), and paratenon (fatty areolar tissue) are shown as well as blood vessels and nerves.
Based on [227].
The tendon repair process in humans. The healing of ruptured tendons passes through three main phases containing distinctive cell and molecular cascades. These phases overlap and their duration depends upon the location and severity of the tendon injury. Currently, the tendon research field is actively exploring the use of growth factors, genes, stem cells and biomaterials, alone or in various combinations, for enhancing tendon healing. Mostly, the appropriate times of application are in the first two stages (indicated by white arrows), and depend on the type of growth factors, genes, stem cells or biomaterials implemented. Based on [47].
Fig. 3.
Key molecular, cellular and matrix changes occurring during the three main phases of tendon repair. Each healing stage is characterized by involvement of different growth factors, activation of certain cell types and production of essential matrix proteins, which collectively contribute to the replacement of the initial fibrous tissue with more a tendonous regenerate. Based on [45,46].
Fig. 4.
Studies on the use of biologics for tendon repair. Article counts were carried out after searching in PubMed using the following key words: tendon repair/healing in combination with growth factors, stem cells, biomaterials and gene therapy. The articles include in vivo and in vitro studies, and some articles scored in more than one category. The search results demonstrate that in the last decade the tendon research field has progressively expanded as represented by the continuous increase in the number of articles focusing on different strategies for enhancing tendon tissue healing. Such cumulative efforts may lead to the development of efficient biologics for tendon repair.
Table 1
Summary of in vitro and in vivo studies on growth factors.

| Growth factor | Tendon                  | Type of study | Model    | Reference |
|---------------|-------------------------|---------------|----------|-----------|
| bFGF          | Flexor tendon           | In vitro      | Canine   | [87]      |
|               | Flexor tendon           | In vitro      | Canine   | [85]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [95]      |
|               | Patellar tendon         | In vitro      | Rat      | [68]      |
|               | Supraspinatus tendon    | In vivo       | Rat      | [93]      |
| BMP 12        | Achilles tendon         | In vivo       | Rat      | [194]     |
| BMP 2         | Flexor tendon           | In vitro      | Canine   | [85]      |
|               | Extensor tendon         | In vivo       | Canine   | [57]      |
|               | Flexor tendon           | In vivo       | Rabbit   | [58]      |
|               | Infraspinatus tendon    | In vivo       | Rabbit   | [59]      |
| BMP 2, 7, 12  | Infraspinatus tendon    | In vivo       | Sheep    | [94]      |
| IGF           | Flexor tendon           | In vitro      | Rabbit   | [90]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [91]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [95]      |
|               | Rotator cuff            | In vitro      | Rat      | [214]     |
| PDGF          | Flexor tendon           | In vivo       | Canine   | [86]      |
|               | Flexor tendon           | In vitro      | Canine   | [87]      |
|               | Flexor tendon           | In vitro      | Canine   | [85]      |
|               | Flexor and peroneal tendon | In vitro  | Rabbit   | [88]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [95]      |
|               | Patellar tendon         | In vitro      | Rat      | [64]      |
|               | Rotator cuff            | In vitro      | Rat      | [214]     |
|               | Rotator cuff            | In vivo       | Sheep    | [78]      |
|               | Medial femur-tibial ligament | In vivo    | Rat      | [92]      |
| TGFβ          | Patellar tendon         | In vivo       | Rabbit   | [80]      |
|               | Achilles tendon         | In vivo       | Rat      | [76]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [82]      |
|               | Flexor tendon           | In vitro      | Rabbit   | [69]      |
|               | Flexor tendon           | In vivo/in vitro | Rabbit | [77]      |
|               | Achilles tendon         | In vivo       | Rat      | [195]     |
|               | Flexor tendon           | In vitro      | Rabbit   | [65]      |
|               | Flexor tendon           | In vivo       | Rat      | [71]      |
|               | Flexor tendon           | In vivo       | Murine   | [72]      |
| TGFβ1         | Flexor tendon           | In vitro      | Murine   | [73]      |
|               | Flexor tendon           | In vivo       | Murine   | [74]      |
| TGFβ1, 2, 3   | Infraspinatus tendon    | In vivo       | Sheep    | [94]      |
| VEGF          | Flexor tendon           | In vitro      | Canine   | [83]      |
|               | Flexor tendon           | In vitro      | Canine   | [84]      |
|               | Flexor tendon           | In vitro      | Canine   | [85]      |
| Growth factor | Tendon          | Type of study | Model  | Reference |
|---------------|-----------------|---------------|--------|-----------|
| ACS           | Flexor tendon   | In vitro      | Rabbit | 81        |
|               | Achilles tendon | In vivo       | Rat    | 98        |
| PRP           | Achilles tendon | In vivo/In vitro | Rat | 96        |
|               | Flexor tendon   | In vitro      | Equine | 99        |
|               | Flexor tendon   | In vivo       | Equine | 97        |
|               | Review          | In vivo       | Human  | 105       |
|               | Review          | In vivo       | Human  | 106       |
|               | Review          | In vivo       | Human  | 109       |
Table 2
Tendon repair with bone marrow-derived MSC.

| Tendon type            | Model                                                                 | Conclusion                                                                                                                                    | Reference |
|------------------------|-----------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------|
| Achilles tendon        | Rat; surgical cut; MSCs cultured at hypoxic and normoxic conditions, analysis at 2 and 4 weeks | Superior biomechanical testing in tendons treated with MSC cultured in hypoxic conditions                                           | [228]     |
|                        | Rat; surgical cut and enthesis destroyed; suture and MSC injection; analysis at 15, 30 and 45 days | Improved healing and biomechanical properties; enthesis comparable to control                                                               | [229]     |
|                        | Rat; surgical cut; total BM cell or MSC injection in DMEM; analysis at 1, 2 and 3 weeks | Biomechanical properties of tendon treated with BM cells comparable to normal tendon; MSC second best                                       | [230]     |
| Rabbit; surgical cut; MSC-fibrin; follow up at 1, 3, 6 and 12 weeks | At 3 weeks improved histological and biomechanical properties with no difference at 12 weeks                                              | [231]     |
| Rabbit; surgical transection; knitted poly-lactide-co-glycolide scaffold loaded with MSCs; analysis at 2, 4, 8 and 12 weeks | Higher rate of tissue formation and remodeling was observed early on with restored function similar to native tendon | [232]     |
| Rabbit; hallucis longus tendons transferred into calcaneal bone tunnel; MSC treatment; analysis at 2, 4 and 6 weeks | Improved healing of the insertion of tendon to bone in the early stage                                                                      | [233]     |
| Rabbit; surgical cut; MSCs in collagen gel; analysis at 12 weeks | Constructs with lower cell density displayed superior biomechanical properties                                                             | [234]     |
| Rabbit; surgical transection; MSC-collagen implants; analysis at 4, 8 and 12 weeks | Improved collagen organization and increased load properties                                                                            | [158]     |
| Patellar tendon        | Rat; surgical transection; MSCs with fibrin injection; analysis at 10 and 20 days | Biomechanical properties were not significantly improved but tendons displayed more mature organization without ectopic ossification | [235]     |
| Rabbit; surgical cut; MSCs in dog decellularized tendon composites; analysis at 2 weeks | More dense collagen fibers, higher cellularity and matrix without ectopic ossification were observed                                           | [236]     |
| Rabbit; surgical cut; MSCs in a gel-sponge composite; analysis at 12 weeks | MSC survived in multilayer composite and expressed tendon phenotype                                                                      | [173]     |
| Rabbit; surgical defect; MSCs from young and aged rabbits in collagen I gels; analysis at 12 weeks | Superior cellular alignment, but maximum force and stress compared to gel only                                                             | [237]     |
| Rabbit; surgical cut; implanted MSCs; analysis at 2, 3, 5 and 8 weeks | No significant difference in biomechanical properties of tendons treated with young or aged MSC                                           | [238]     |
| Rabbit; surgical cut; MSC-collagen implants; analysis at 6, 12 and 26 weeks | MSC survived and differentiated into tendon-like spindle cells                                                                            | [239]     |
| Rabbit; surgical cut; MSCs in collagen gel; analysis at 4 weeks | Ectopic ossification developed in approx. 25% of the tendons with MSC-collagen implant; no significant differences in mechanical properties across different seeding densities | [159]     |
| Rabbit; surgical cut; MSCs in collagen gel; analysis at 4 weeks | Better biomechanical properties but no significant improvement of tendon microstructure                                                   | [240]     |
| Superficial digital flexor tendon | Horse; naturally-occurring tendon injury; MSCs with marrow supernatant; analysis at 6 months | Treated group exhibited normalization on a biochemical, morphological and compositional level                                                 | [241]     |
| Tendon type                                                                 | Model                                                                 | Conclusion                                                                                     | Reference |
|----------------------------------------------------------------------------|----------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|-----------|
| Horse; collagenase-induced tendinitis lesion; MSC injection; analysis at 8 weeks |                                                                                   | Increased tensile stiffness in MSC-treated group, but similar histological scores to controls without MSC | [217]     |
| Horse; collagenase-induced tendinitis lesion; MSC injection                |                                                                                   | Repaired tendon architecture comparable to healthy tendon                                      | [242]     |
Table 3
Salient properties of vectors used in experimental studies of tendon healing.

| Vector          | Key properties                              | Advantages                                      | Disadvantages                               | Comment                                      |
|-----------------|---------------------------------------------|------------------------------------------------|---------------------------------------------|----------------------------------------------|
| Adenovirus      | Non-integrating                             | Straightforward production                      | Inflammatory                                | Widely used in clinical trials               |
|                 | Multiple serotypes                          | Efficient                                      | Antigenic                                   | One well publicized death                    |
|                 | Double stranded DNA genome                  | Transduces non-dividing cells                   |                                              |                                              |
| Adeno-Associated| Recombinant AAV is non-integrating          | Transduces non-dividing cells                   | Difficult to produce                        | Possible to engineer AAV with double         |
| Virus           | Wild-type AAV has single stranded DNA genome| Wild-type AAV causes no known disease           | Small carrying capacity                      | stranded DNA genome                          |
|                 | Multiple serotypes                          | Non-inflammatory                               |                                              | Increasingly popular for clinical trials     |
|                 | Wild-type AAV causes no known disease       |                                              |                                              | because of safety                            |
| Retrovirus      | RNA genome                                  | Straightforward to produce                     | Transduction requires host cell division    | Usually used in ex vivo gene delivery        |
|                 | Integrating                                 | Amphotropic virus has wide host range          | Risk of insertional mutagenesis             | Has been widely used in clinical trials      |
|                 |                                            |                                              |                                              | Insertional mutagenesis has caused leukemia |
| Lentivirus      | RNA genome                                  | Transduces non-dividing cells                   | Risk of insertional mutagenesis, but        | Increasing use in clinical trials            |
|                 | Wild-type virus is integrating               | Very high levels of transgene expression        | non-integrating vectors developed           |                                              |
Table 4
Use of gene transfer to promote tenogenesis in animal models of tendon injury.

| Gene           | Vector | Delivery mode | Animal model         | Reference |
|----------------|--------|---------------|----------------------|-----------|
| BMP-14/GDF-5   | Adenovirus | In vivo      | Rat, Achilles        | [206]     |
| BMP-13/GDF-6   | Adenovirus | In vivo      | Rat, Achilles        | [207]     |
|                | AAV    | In vivo      | Mouse, flexor tendon | [190]     |
| BMP-12/GDF-7   | Adenovirus | Ex vivo/MSC  | Rat, rotator cuff    | [208]     |
|                | Adenovirus | Ex vivo/muscle | Rat, Achilles      | [205]     |
| TGFβ, VEGF     | Adenovirus | Ex vivo/MSC  | Rabbit, Achilles     | [213]     |
| TGFβ           | Adenovirus | Ex vivo/muscle | Rat, Achilles      | [195]     |
| Scleraxis      | Adenovirus | Ex vivo/MSC  | Rat, supraspinatus   | [211]     |
| SMAD8, BMP-2   | Liposome | Ex vivo/MSC  | Rat, Achilles        | [120]     |
| bFGF           | AAV    | In vivo      | Chick, flexor tendon | [212]     |
| PDGF-B         | Liposome | In vivo      | Rat, patellar tendon | [215]     |
|                | Nanoparticle | In vivo     | Rat, Achilles Tendon  | [243]     |
|                | Retrovirus | Ex vivo/tendon fibroblasts | Rat, rotator cuff | [214]     |
| Interleukin-10 | Lentivirus | In vivo     | Mouse, patellar tendon | [216]     |
Table 5
Use of gene transfer to address additional aspects of tendinopathy.

| Gene          | Vector  | Delivery mode | Species, indication                  | Reference |
|---------------|---------|---------------|--------------------------------------|-----------|
| Sox-9         | Adenovirus | In vivo       | Rabbit, bone–tendon junction          | [222]     |
| IGF-1         | Adenovirus | Ex vivo/MSC   | Horse, tendinitis                     | [217]     |
| BMP-4         | Lentivirus | In vivo       | Rat, tendon insertion site            | [218]     |
| BMP-2         | Adenovirus | In vivo       | Rabbit, tendon insertion site         | [219]     |
| MT1-MMP       | Adenovirus | Ex vivo/MSC   | Rat, tendon insertion site            | [221]     |
| siRNA-Runx2   | Adenovirus | In vivo       | Mouse, heterotopic ossification       | [223]     |
| siRNA-Runx2, SMAD4 | Adenovirus | In vivo       | Rat, heterotopic ossification         | [224]     |
| shRNA-Decorin | Lentivirus | Ex vivo/tendon cells | Rat, patellar tendon               | [225]     |