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**Healing of subcutaneous tendons: Influence of the mechanical environment at suture line on healing process**

Massoud EIE. Tendon healing: Influence of mechanical environment

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**Abstract**

Tendon ruptures remain significant musculoskeletal injury. Despite advances in surgical techniques and procedures, traditional repair techniques maintain a high incidence of rerupture or tendon elongation. Mechanical loading and biochemical signalling both control tissue healing. This has led some researchers to consider using of a technique based on tension regulation at suture line for obtaining good healing. However it is unknown how they interact, and to what extent mechanics controls biochemistry. This review will open the way for understanding of the interplay between mechanical loading and process of tendon healing.

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**Key words:** Tendon; Healing; Mechanical loading; Mechanotransduction; Rupture; Repair

**Core tip:** Ruptured tendons heal poorly compared to skin, muscles and bones. Immobilization during repair has been shown to be detrimental for the healing process. Mechanical loading of the tendon callus gives rise to intracellular signalling, increase gene expression and protein synthesis. However early loading reported clinical complications. Surgical technique that based on control of the mechanical environment at the suture line provided satisfactory results. Therefore, understanding of the interplay between loading and healing process seems necessary. This review focuses on the biological processes that regulate tendon repair; Timing of mechanical loading during healing process. How do tendon cells sense mechanical forces?

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**INTRODUCTION**

This article is limited to the subcutaneous tendons, *i.e.*, tendons that do not glide through synovial sheaths, responsible for their nutrition. Describes thefunction, structure and mechanobiology of tendons, reviews the phases of tendon healing and reviewsinfluence of the mechanical environment at site of repair on the healing process.

**TENDON STRUCTURE**

Healthy tendons are brilliant white in colour and have a fibroelastic texture.Tendons demonstrate marked variation in form; generally, extensor tendons are more flattened than flexor tendons, which tend to be round or oval [1,2]. The dry mass of human tendons is approximately 30% of the totaltendon mass, with water accounting for the remaining 70% [1,3]. LeMoine *et al* [3] reported that men had a significantly greater amount of tendon dry mass than women. Many authors have explained this dissimilarity on the fact that oestrogen directly alters collagen kinetics, and inherently higher oestrogen levels in women may therefore chronically depress collagen production, as tendon cells have oestrogen receptors [4,5]. Thus, blunted collagen production via oestrogen could explain the lower amount of dry mass in female tendons, as collagen comprises 90% of dry mass [3,4].

 Tenoblasts and tenocytes constitute about 90%to 95% of the cellular elements of tendons. They lie between the collagen fibres along the long axis of the tendon [6].The remaining 5% to 10% of the cellular elements consists of chondrocytes at the insertion sites, synovial cells, andvascular cells, including capillary endothelial cells and smoothmuscle cells of arterioles [1]. Tenocytes are mature tendon cells. They are active in energy generation; synthesize collagen and all components of the extracellularmatrix network [7].

***Extracellular matrix***

Extracellular matrix (ECM) of tendons largely consist of collagens and proteoglycans and are dominated by the type I collagen as it accounts for 65% to 80% of the dry mass of tendons. However, other collagens (*e.g.*, type II, III, V, VI, IX, XI) are also present [2,8]. Collagen molecules consist of polypeptide chains(tropocollagen). Solubletropocollagen molecules form cross-links to create insolublecollagen molecules*.* Three such chains combine together to form a densely packed, helical tropocollagen molecule(triple-helix polypeptide chain). In turn, five tropocollagens constitute a microfibril, and microfibrils aggregate together to form fibrils. Fibrils are then grouped into fibres, fibres into fibre bundles (primary bundles) and fibre bundles into fascicles(secondarybundles); tertiary bundles; and the tendon itself.A collagen fibre is the smallest tendon unit that can be testedmechanically and is visible under light microscopy [1,2]. Some of the larger collections of fascicles are visible in gross dissections [2].

 The principal role of the collagen fibres is to resist tension, although they still allow for a certain degree of compliance (*i.e.*, reversible longitudinal deformation). Such apparently conflicting demands are probably resolved because of the collagen is arranged in hierarchical levels of increasing complexity, beginningwith tropocollagen [1,2]. At various levels of tendon organization, including the whole tendon, fascicles and fibrils, a helical architecture (often with superimposed ‘crimp’, *i.e.*, a zigzag undulation of collagen fibrils) occurs in certain tendons. This helical organization of tendon components makes them comparable to man-made ropes and the presence of crimp contributes to their inherent flexibility [9-11]. Roukis *et al*[10] have suggested that the twisting that characterizes the tendon of tibialis posterior reduces the need for longitudinal slippage between fascicles during triplanar movements of the foot. The angle of torsion of the inner fibrils in a helical tendon fascicle may be less oblique than that of the outer fibrils and this may give the tendon regionally distinct compliance [10]. The fibres of the Achilles tendon twist through its descent, thus elastic recoil within the tendon are possible [12]. This point will be detailed under subheading elastic recoil of the tendon.

 The ground substance of the extracellular matrix network surroundingthe collagen and the tenocytes is composed of proteoglycans,glycosaminoglycans, glycoproteins, and several other small molecules [1]. Proteoglycans make up less than 1% of the dry weight of most tensile tendons [8]. Proteoglycansare strongly hydrophilic, enabling rapid diffusion of water-solublemolecules and the migration of cells [1]. Additionally, the proteoglycans have the function of providing a viscous environment, allowing the collagen fibrils, fibres or fascicles to slide relative to each other as well as to stretch and dissipate the force of sudden loads [8,13].

**TENDON ENVELOPE**

The epitenon, a fine loose connective-tissue sheath containingthe vascular, lymphatic, and nerve supply to the tendon*,* surrounds the tendon as a whole and forms the gross structure of the tendon [14,15]. The epitenon extends deeply as the endotenon, which is a thin reticularnetwork of connective tissue investing each tendon fibre. The endotenon protects tendon vasculature and allows fascicles to slide over one another in particularly malleable parts of the tendon [2,14]. Due to the epitenon is directly continuous with the endotenon, the points of continuity help to bind it firmly to the surface of the tendon [14]. The epitenon is further enclosed by paratenon, a loose areolar connectivetissueseparated from the epitenon by a thin layer of fluid to allow tendon movement with reduced friction [2,14,15]. The paratenon may be quite vascular and is a source of the blood supply to the tendon itself [8].

***Myotendinous junction***

The myotendinous junction (MTJ)is the interface between muscle and tendon; it tailored for transmitting the mechanical force generated by a muscle contraction to the extracellular matrix of the muscle and onto the tendon [16-18]. The characteristic morphology of the MTJ is folding of the sarcolemma into finger-like projections at the interface between muscle and tendon at sites of myocyte termination [16,17]. The projections increase the area of muscle-tendon contact to more thane 10 folds over the cross-section of the muscle fibre. Thus, the local stress (force per unit area) is reduced. Additionally the longitudinal arrangement of the projections ensures that the stresses experienced by the MTJ are shear stresses [19,20]. Mechanical loading of the MTJ activates cell-signalling pathways that instruct the cells located at the interface to secrete and deposit proteins to form a specialized extracellular matrix at the MTJ. However, lack of the expression of these proteins has been shown to lead to structural damage of the interface during contraction [18].

***Osteotendinous junctions***

The osteotendinous junctions (OTJ) are sites of stress concentration at the region where tendons attach to bone [21]. These regions are characterized by the presence of a unique transitional tissue called “entheses” at the interface, which can effectively transfer the stress from tendon to bone and vice versa through its gradual change in structure, composition and mechanical behaviour [21,22]. There are two types of entheses based on the how the collagen fibres attach to bone [14,21,22]. Direct insertions (also called the fibrocartilaginous entheses), such as the insertion of Achilles tendon and patellar tendon, is composed of four zones in order of gradual transition: tendon, uncalcified fibrocartilage, calcified fibrocartilage and bone [21,22]. The continuous change in tissue composition from tendon to bone is presumed to aid in the efficient transfer of load between the two materials [22]. Indirect insertions (also called fibrous entheses), such the insertion of the deltoid tendon into the humerus, has no fibrocartilage interface. The tendon passes obliquely along the bone surface and inserts at an acute angle either directly to the bone or indirectly to it *via* the periosteum [21,22]. The main factors affecting the type of insertion seem to be strain, site, length and angle of insertion. When a tendon runs parallel to the bone, the insertion is more likely to be indirect, while when the tendon enters the bone quite perpendicularly, the insertion is direct [22].

 The majority of tendons attach not only to bone, but also to adjacent fascia. This is a basic strategy for dissipating stress concentration at entheses and thus reducing the risk of failure or local wear and tear. One of the classic examples of subcutaneous tendons that have both bony and fibrous attachments is the quadriceps tendon. This not only attaches to the superior pole of the patella, but also sends a sheet of fibres anterior to the patella that become continuous with the patellar tendon [23].

**TENDON NUTRITION**

Tendons are still vascularized, and the presence of vessels is important for the normal functioning of tendon cells and the ability of tendons to repair [2]. Tendons receive their blood supply from three sources: The peritendinous tissues (the extrinsic source) that have a richer blood supply than do the tendons themselves [24]. In the tendon itself, the vessels run longitudinally, parallel to the fascicles and within the endotenon, anastomoses between parallel vessels are common [25]. Intrinsic sources include vessels that enter tendons at their myotendinous junctions and at entheses [2]. Nevertheless, the direct role of the blood vessels in tendon nutrition has been called into question. Edwards, has reported that tendons may be cut and transplanted with impunity [25]. Recently, many investigators havepointed out that diffusion from surrounding tissues may play a significant role in metabolic exchange in intact tendons [26-28].

**TENDON INNERVATION**

Tendon innervation originates from cutaneous, muscular, andperitendinous nerve trunks [1]. The majority of nerve fibres are located within the paratenon and not the tendon itself [29]. Paratenon nerves form rich plexuses that send a few branches penetrating the epitenon. These branches are described to cross the myotendinous junction and to continue into the endotenon septa [30]. Deep in the tendon tissue proper, where innervation is reported to be relatively scarce, the nerves follow the blood vessels running along the axis of the tendon [2,30]**.** Four types of nerve endings have been identified: free nerve endings, Ruffini corpuscles, Pacinian corpuscles and Golgi tendon organs [30]. Vessel-associated fibres are autonomic nerves that immunolabel for neuropeptide Y and noradrenaline (vasoconstrictive factors) and for vasoactive intestinal peptide (VIP) a vasodilator factor. It has been suggested that the nerve fibres regulate blood flow within the tendon**.** Further, free nerve fibres containing substance P and calcitonin gene-related peptide (CGRP) might be involved in collecting sensory information (including pain) and relaying this to the central nervous system [29]. Zaffagnini *et al*[31] have reported the presence of Ruffini and Pacinian corpuscles within the pes anserinus tendons, particularly at their tibial attachment sites. Benjamin et al. confirm that Pacinian corpuscles can be found on the surface of subcutaneous entheses [32].

**Biomechanical properties of the tendon**

Tendons transmit force from muscle to bone and act as a bufferby absorbing external forces to limit muscle damage [1]. We will discuss response of the tendon to mechanical stimuli at fibrillar and cellular levels*.* At rest, a tendon has a wavy configuration, a result of crimping of the collagen fibrils. The stress–strain curve of tendons usually exhibits three distinct regions [33], which can be correlated to deformations at different structural levels (Figure 1). In the first region that usually called the toe region, a very small stress is sufficient to strain (elongate) the tendon up to 2% of its length and the straightening of the macroscopic crimp in the collagen fibrils [34]. In the second region of the curve, at higher strains, the stiffness of the tendon increases [13,35]. If the strain placed on the tendon remains at less than 4%, the tendon behaves as mechanical spring and returns to its original length and crimps when un-loaded [35]. The most probable processes are thought to be the ability of the fascicles to slide independently against each other. This allows them to transmit tension despite the changing angles of a joint as it moves and allows tendons to change shape as their muscles contract [2,36]. Sliding within fascicles occurs between fibrils and this may account for up to 50% of the longitudinal deformation (*i.e.*, strain) of a tendon [37]. Sliding of fibrils or fascicles relative to each other occurs within the proteoglycan-rich matrix surrounding them [13]. Presence of the endotenon between fascicles and/or fibre bundles facilitates the sliding movement [2, 14, 38]. Lubricin, a molecule often associated with joint lubrication, is also present between the fascicles of certain tendons [39].

At strain levels between 4% and 8%, the tendon becomes progressively easier to extend but its length still return to its original value. However, the wave pattern does not reappear [34]. On the other hand, recent work has suggested that strain values of 6% and even up to 8% may be physiological. Within the physiological range, particularly towards the higher range, microscopic degeneration within the tendon may start to occur, especially with repeated and/or prolonged stressing [40]. Beyond 8% to 10% strain,macroscopic failure occurs from intrafibril damage by molecular slippage [40-42]. The probable process has previously been investigated using synchrotron radiation diffraction. Initially collagen fibril elongation occurs as a result of molecular elongation. When the stress increases the stretching of the collagen triple helices and the cross-links between the helices, a considerable gliding of neighbouring molecules occur [43,44].

***Response of tendon cells to mechanical load***

There is now considerable evidence to suggest that tendons and tendon cells can respond to altered mechanical load. In man, collagen synthesis in the patellar tendon increases by nearly 100% as a result of just a single bout of acute exercise, and the effect is still evident 3 days later [45]. At a cellular level, there seems to be no difference in the response of tenocytes to mechanical load between cells that have been extracted from different tendons, *e.g.*, those associated with antagonistic muscles [46]. However, in a given tendon, different stress patterns provoke different cellular reactions depending on the amount and duration of the tensional stress applied. Cell proliferation, for example, is stimulated by short periods of repetitive tension, but inhibited by more extended periods [47]. The response seems to depend on gap junctional communication between neighbouring cells, for when gap junctions are blocked, the cells no longer increase collagen synthesis in response to stretching forces applied *in vitro*. The modulation of ECM synthesis involves two types of gap junctions: those characterized by the presence of connexin 32 and those containing connexin 43. The former junctions stimulate and the latter inhibit collagen synthesis [48]. In addition to its effects on collagen synthesis, the repetitive stretching of tenocytes *in vitro* up regulates proinflammatory cytokine production and the gene expression of mediators such as Cox-2, PGE2 and MMP-1 [49,50]. Smaller levels of repetitive tensile stress reduce the production of proinflammatory agents. Thus, repetitive small magnitude stretching seems to be anti-inflammatory, whereas large magnitude stretching is pro-inflammatory. If the findings also prove to be applicable *in vivo*, then it follows that moderate exercise may be beneficial for reducing tendon inflammation [50]. It is interesting to note that tenocytes themselves may produce IL-1β, especially if they are located next to a site where the tendon is injured. Expression is highest 1 d after injury but can persist for several days [51]. The significance of IL-1β production in an injured tendon is that it can induce the expression of a wide range of pro-inflammatory agents such as Cox2, MMP1, MMP3, MMP13, ADAMTS-4 and IL-6. It also triggers the further expression of IL-1β mRNA [52] and this is presumably a mechanism for rapidly raising its local concentration. It should be noted, however, that in addition to such actions, IL-1β reduces the elastic modulus of tenocytes by disrupting actin filaments [53]. The authors suggest that this acts as a protective mechanism against mechanical overuse of tendon cells during healing. How do tendon cells sense mechanical forces? This question will be answered in details in this review.

Suppression of proteoglycan and collagen synthesis in cultured tenocytes can be induced by glucocorticoids [54,55]. These are among the substances commonly used by clinicians to suppress inflammation in patients with tendon injuries. Glucocorticoids can also suppress tenocyte proliferation and progenitor cell recruitment [56]. If such effects also occur *in vivo*, then this may explain why the integrity of the tendon as a whole may be affected by corticosteroid treatment. In contrast to corticosteroids, nitric oxide generally benefits tendon healing and enhances collagen synthesis [57]. Nitric oxide synthetases are normally expressed at low levels and are up regulated by mechanical stimuli [58,59]. The absence of nitric oxide from tendons during wound healing is associated with prolonged inflammation [60].

**ELASTIC RECOIL OF TENDONS**

Tendons can recoil elastically when a stretching force is removed. The elastic recoil property seems to be structurally related to crimp and/or knots within fibrils in regions where fibrils are twisted or bent. When a tendon is physiologically stretched *in vivo*, the crimp numbers within it may decrease by nearly 50% [61].

The ability of tendons to stretch and recoil enables them to save energy in running by allowing the limb to have shorter muscle fascicles or slower muscle fibres that can generate force more economically [12]. The fibres of the Achilles tendon spiral through 90 degrees during its descent, such that the fibres that lie medially in the proximal portion become posterior distally. In this way, elongation and elastic recoil within the tendon are possible, and stored energy can be released during the appropriate phase of locomotion. In addition, this stored energy allows the generation of higher shortening velocities and greater instantaneous muscle power than could be achieved by contraction of the triceps surae alone [12,15] . The stiffness of tendons varies with sex, age and physical activity. The Achilles tendon of women can recoil elastically more than that of men [62]. Stiffness is greater in young men than it is in young boys however; it decreases with training in adults [63,64]. The greater compliance of tendons in young boys may be reducing the risk of sport injuries [63]. The elasticity of a fatigued tendon tends to be greater as evidenced by its ability to lengthen further with the same load [65].

Studies in goats have shown that the muscle rather than the tendon that provides the extra length within the muscle–tendon unit necessary for limb lengthening by distraction osteotomy. While the muscle may elongate by almost 10% of its initial length, the tendon only does so by 3%–4% [66]. Elastic recoil of tendon stumps will elongate the spontaneously healed tendon. This emphasizes necessity for tension relieve at site of repair in the early phase of tendon healing.

**TENDON RUPTURE**

Tendon rupture occurs spontaneously or following direct trauma such as severance of a tendon by sharp objects or caught between bones and traumatizing agent. A spontaneous rupture may be defined as a rupture that occurs during movements and activities that should not - and usually do not - damage the involved musculotendinous units[67]. Although many investigators reported that spontaneous rupture of a tendon is preceded by degenerative changes, however there is little agreement with regard to its aetiology. Degenerative changes of the tendon have been linked to genetic abnormalities of the collagen tissues [68], chronic diseases or metabolic disorder [69] and neurological conditions [70]. Fluoroquinolones and locally or systemically administered corticosteroids have been implicated in the aetiology of the tendon rupture [71 -74]. In the literature, there are four basic types of tendon degeneration: hypoxic, mucoid degeneration, lipomatosis, and calcification of the tendon. Extensive tendolipomatosis by itself may lead to rupture of the tendon without degenerative changes in the collagen tissues [75]. For reasons that are not clear, most reported cases of tendolipomatosis have been in the quadriceps or patellar tendon. The described histopathological changes predispose the tendon to rupture through decrease of its tensile strength[67]. Strength of the tendons and resistance to tensile forces are related to the angles of tendon crimps in providing a resistance to sudden elongation and to the diameter of the collagen fibres [61,76]. Järvinen *et al*[77] investigated the crimp angle and the diameter of the collagen fibres in spontaneously ruptured tendons and compared them to healthy tendons. They concluded that the collagen fibres in ruptured tendons are substantially thinner than in normal tendons. The crimp angle of the collagen fibres is also significantly decreased in ruptured tendons [77].

**HEALING OF TENDON RUPTURE**

After tendon rupture, the body restores tendon continuity through a cascade of events can be divided into three overlapping phases: the tissue inflammation, cell proliferation, and remodelling phases [78-80].

***Inflammatory phase***

This phase starts immediately post injury and persist for about 24 h[78]. In this phase, injured blood vessels that are in the tendon envelope cause the formation of a hematoma, which activates the release of various chemotactic factors such as vasodilators and proinflammatory molecules [78,79]. The chemotactic factors attract inflammatory cells (*e.g.*, neutrophils, monocytes, and macrophages) that migrate to the wound site and clean the site of necrotic materials by phagocytosis. Tendon fibroblasts recruited to the site begin to synthesize various components of the ECM [81]. Moreover, during this phase the angiogenic factors initiate the formation of a vascular network [82]. These processes include an increase in DNA and in ECM, which establishes continuity and partial stability at the site of injury [78].

***Proliferative phase***

In this phase that lasts a few weeks; tendon fibroblasts synthesize collagen and other ECM components and deposit them to the wound site [78]. These components are initially arranged randomly within the ECM, which at this time is composed largely of type III collagen [83]. An extensive blood vessel network is present, and the wound has a scar-like appearance [84]. During this phase, the repair tissue is highly cellular and contains relatively large amounts of water and an abundance of ECM components [78,79].

***Remodelling phase***

This phase that begins by nearly 6th week after injury is characterized by decreased cellularity, reduced matrix synthesis, decrease in type III collagen, and an increase in type I collagen synthesis [78,79]. Type I collagen fibres are organized longitudinally along the tendon axis and are responsible for the mechanical strength of the regenerate tissue [85]. During the later remodelling phase, covalent bonding between collagen fibres increases consequently tendon stiffness and tensile strength increased. In addition, both the metabolism of tenocytes and tendon vascularity decline [78,79].

**ROLE OF FIBROBLAST IN TENDON HEALING**

It is known that the fibroblasts during healing generate and exert force on the ECM. This force is referred to as fibroblast contraction, which is essential for wound closure [86]. However excessive cell contraction, may lead to tissue scarring. On the other hand, inhibiting fibroblast contraction results in impaired wound healing [87]. Therefore, an optimal level of fibroblast contraction is desirable to facilitate wound closure while minimizing scar tissue formation.

Cell contraction involves the actin cytoskeleton [88]. The interaction between actin and myosin generates cell contraction [89], and the contractile forces transmit through actin filaments to integrins and to the ECM [90].

In the literature, most studies concerned with cell contraction focus on skin fibroblasts [78]. Using a cell force monitor, contractile forces of tendon and skin fibroblasts were measured over time. It was found that tendon and skin fibroblasts exhibited different patterns of contraction, where tendon fibroblasts produced a lower maximum contraction force than skin fibroblasts [91]. In healing tissues, myofibroblasts are thought to play a major role in tissue contraction. These cells have phenotypic characteristics of both fibroblasts and smooth muscle cells, including the formation of stress fibres parallel with the long axis of the cell [92] Mechanical loading also influences myofibroblast differentiation. Increased tension on granulation tissue in rats increases the formation of stress fibres and the expression levels of a-SMA and ED-A fibronectin [93], which are two protein markers of myofibroblasts [78].

**HOW DO TENDON CELLS SENSE MECHANICAL FORCES?**

As described above tendon cells respond to mechanical forces by altering gene expression, protein synthesis, and cell phenotype. These early adaptive responses may proceed, initiate long-term tendon structure modifications, and thus lead to changes in the tendon’s mechanical properties. After that we in need to understand how do tendon cells sense mechanical forces and convert them into cascades of cellular and molecular events that eventually lead to changes in tendon structure? This question comprises definition of mechanotransduction, nevertheless necessitates review of mechanotransduction mechanisms. Cellular components implicated in the transduction of mechanical forces are extracellular matrix, cytoskeleton, integrins, G proteins, receptor tyrosine kinases (RTKs), mitogen-activated protein kinases (MAPKs), and stretching-activated ion channels.

*ECM*

 ECM is composed of cell-produced proteins and polysaccharides [94,95]. It defines tissue shape, structure, and act as the substrate for cell adhesion, growth, and differentiation [96]. Mechanical loading increases ECM protein production by promoting release of growth factors which mediate collagen secretion [97,98] .

ECM transmits mechanical loads, store and dissipate loading-induced elastic energy. Moreover, mechanical deformations in the ECM can transmit to the actin cytoskeleton and cause the remodelling of the actin cytoskeleton [99,100], which is known to control cell shape, affect cell motility, and mediate various cellular functions including DNA and protein syntheses [101].

***Cytoskeleton***

Cytoskeleton is composed of microfilaments, microtubules, and intermediate filaments [101,102]. The cytoskeleton responds to extracellular forces, participates in transmembrane signalling, and provides a network for translocating signalling molecules. Mechanical forces applied to the cell surface transmit directly to the cytoskeleton and cause changes in its structure [103]. Consequently, these changes due to applied mechanical forces can initiate transduction cascades within the cell through the activation of integrins and the stimulation of G protein receptors, RTKs, and MAPKs [78].

***Integrins***

Integrins are transmembrane protein composed of a and b subunits [104]. It mediates mechanotransduction between the extracellular matrix and the cell through ‘‘outside in’’ and ‘‘inside-out’’ fashions [104,105]. Mechanical forces stimulate the conformational activation of integrins in cells and increase cell binding to the extracellular matrix [106].

***G proteins***

G proteins are of membrane proteins that are involved in mechanotransduction. Mechanical forces may simultaneously activate G proteins and integrins [107].

***RTKs and MAPKs***

RTKs and MAPKs are a class of cell membrane proteins that are phosphorylated when subjected to cyclic stretching or shear stress. They can travel into the nucleus and alter gene expression [107,108]. It has been shown that cyclic stretching activates MAPKs in patellar tendon fibroblasts [109].

***Stretching-activated ion channel***

The activation of these channels permit calcium (Ca2+) and other ions (*e.g.*, sodium and potassium) to influx followed by membrane depolarization [110].

Mechanical stretching induced Ca2+ signal transmission involves the actin microfilament system because an actin polymerization inhibitor was found to abolish Ca2+ responses induced by mechanical stimulation [111]. This means that calcium is an important mediator in cellular mechanotransduction.

**TIMING OF MECHANICAL LOADING**

It has been agreed that injured tendons require mechanical loading for optimal healing, but early loading is not without risks of adverse effects. Loading might create excessive damage to the repair tissue leading to failure of the healing process or mostly plastic deformation (elongation) of the callus with subsequent tendon lengthening [112]. Lengthening during tendon healing is one potential clinical adversity. It is therefore important to understand the interplay between loading and healing process.

The inflammatory phase of tendon repair seems to prepare the way for the formation of a fibrous callus. If early inflammation is inhibited the fibrous callus can lose a third of its strength, due to inferior material properties (lower stress at failure). During the inflammatory phase, there is little mechanical strength and mostly plastic deformation [112]. About effect of loading during inflammatory phase,Eliasson*et al*[113] noticed suppression of genes related to inflammation and extracellular matrix components. Moreover, they observed the loaded tendons by third dayalso had a lower expression of collagens III and I than unloadedones, emphasizing, again, the value of mechanicalprotection in the early phase [113].Once some elasticity has been obtained in the early fibrous callus, however, deformed tissues will resume their pre-load shape, and cyclic loading will lead to biological signals [114]. It is remarkable that the investigators have suggested the period between 4th and 6th week postoperative for institution of mechanical loading at the tendineous interface as well at the tendon-bone junction [114-117]. Eliasson*et al*[113] reported that later during healing, loading was related to a higher expression of extracellular matrix-related and tendon-specific genes, perhaps suggesting that the tissue was to some extent undergoing transformation from scar to tendon regenerate.

**CLINICAL APPLICATIONS FOR MECHANICAL LOADING OF THE REPAIRED TENDONS**

Because of the great distraction forces that arise from the muscles, the long-lasting devices have been used for reinforcement of the repaired tendon [118-120]. However, prolonged reinforcement adversely influences the healing progress. Current knowledge indicated the appropriate time during the healing process for loading to start, but need to find a suitable method for initiation of the mechanical loading during the healing process. In our practice, we use an absorbable reinforcement device for tension regulation at the suture line. The utility of absorbable devices for reinforcement has been mentioned in several studies [121-123]. In two published studies [124,125], we used a reinforcement device made of Vicryl (polyglactin 910) suture, which has initial tensile strength equal or superior to nonabsorbable sutures [126,127]. Thereby, it serves initially as “a suture line tension-relieving suture”. By the fourth postoperative week, the suture loses about 75% of its tensile strength [128]; fortunately, this coincides with the remodelling phase.

We used the technique for repair of fresh rupture of the patellar and Achilles tendon in two groups of patients (Figures 2, 3). Our philosophy based on protection of the tendon callus in the early healing phase. The spontaneous loss of tensile strength of the suture let the callus exposed to the muscle tone, which continually changes. Continuous change of the muscle tone, theoretically, equals in its effect the cyclic mechanical loading. The group was treated for patellar tendon rupture resumed their pre-injury activities at an average of 6.1 mo, knee motion reported no extension lag or flexion deficit and radiologically no patella alta, patella baja or degenerative changes in the patellofemoral joints were noted [124]. The group was treated for Achilles tendon rupture returned to pre-injury daily activates by fourth month and reported no tendon lengthening or reruptures [125]. Moreover, doubtless we can attribute the noticed preservation of thigh and calf girths to our technique. The lack of tension on the immobilized musculotendinous unit is a major factor in the development of atrophy in the muscles [129,130]. This is due to the muscle spindle relaxes and afferent impulses to type-I fibres ceases. The human soleus muscle contain high portion of type-I muscle fibres, as they are responsible for postural tone and are continually activated while the person is standing [131]. Our technique is designed to protect the suture line, and places the muscle fibres under tension as long as the Vicryl suture preserves its tensile strength (Figures 2, 3). By end of the fourth week, patients were allowed active joint motion and weight bearing as tolerated.

In addition to alteration of the morphological character, lack of tension in immobilized muscle as well alters its physiological properties that appear as liability to rerupture. In a study of transected sheep Achilles tendons that had spontaneously healed, the rupture force was only 56.7% of normal at twelve months [132]. One possible reason for this is the absence of mechanical loading during the period of immobilization [1]. Immobilization reduces the water and proteoglycan content of tendons and increases the number of reducible collagen cross links [133,134]. Collagen fascicles from stress-shielded rabbit patellar tendons displayed lower tensile strength and strain at failure than did control samples [135]. Additionally, as detailed above lack of tension in the musculotendinous unit leads to structural damage of MTJ [18].

**CONCLUSION**

Perception of tendon biology and the biological processes that regulate tendon repair have progressed to great extent, however many challenges need to be addressed to bring about a successful treatment strategy. Simplicity of surgical technique that based on control of the mechanical loading at the suture line may reduce the requirement for the demanding tissue engineering particularly in simple circumstance. Moreover, this technique may open the way for earlier plaster removal and institution of more vigorous rehabilitation programmes; thereby, the morbidity period can be reduced.

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**Figure 1 Stress–strain relationship for progressive loading of a tendon showing three distinct regions (toe, linear and partial failure) prior to complete rupture approximate stress forces (MPa) and strain values (% strain) are shown.** Reprinted from [40] with permission from the Oxford University Press.

**Figure 2** **Drawing illustrates repair of the patellar tendon and making of the “suture line tension regulating suture”.** A: The modified Kessler suture in the proximal portion of an avulsed patellar tendon; B: The modified Kessler suture and the reinforcement device before tying the threads into a knot; C: The final appearance after tying the threads into knots. Reprinted from [124] with permission from theSpringer.

**Figure 3 Drawing illustrates repair of the ruptured Achilles tendon.** A: Thread of the reinforcement suture when it passes through the osseous tunnel in the calcaneus and by interweaving through the proximal stump of the Achilles tendon; B: Tendon stumps that were held together by Kessler’s suture; C: Final appearance of the reinforcement suture and the repaired Achilles tendon. Reprinted from [125] with permission from theElsevier.