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THE ROLE OF DETRAINING IN TENDON MECHANOBIOLOGY

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The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest

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Keywords

Tendon, tenocyte, detraining, sudden detraining, Systematic Literature Review

Abstract

Word count: 306

Introduction: Several conditions such as training, aging, estrogen deficiency and drugs could affect the biological and anatomophysiological characteristics of the tendon. Additionally, recent preclinical and clinical studies examined the effect of detraining on tendon, showing alterations in its structure and morphology and in tenocyte mechanobiology. However, there is a paucity of data examining the impact that cessation of training may have on tendon. In practice, we do not fully understand how tendons respond to a period of training followed by sudden detraining. Therefore, within this review, we summarize the studies where tendon detraining was examined.

Materials and methods: A descriptive systematic literature review was conducted by searching three databases (PubMed, Scopus and Web of Knowledge) on tendon detraining. Original articles in English from 2000 to 2015 were included. In addition, the search was extended to the reference lists of the selected articles. A public reference manager ("www.mendeley.com") was used to delete duplicate articles.

Results: An initial literature search yielded 134 references (www.pubmed.org: 53; www.scopus.com: 11;

www.webofknowledge.com: 70). 15 publications were extracted based on the title for further analysis by two independent reviewers. Abstracts and whole articles were then reviewed to detect if they met inclusion criteria.

Conclusions: The revised literature comprised 4 clinical studies and an in vitro and three in vivo reports. Overall, the results showed that tendon structure and properties after detraining are compromised, with an alteration in the tissue structural organization and mechanical properties. Clinical studies usually showed a lesser extent of tendon alterations, probably because preclinical studies permit an in-depth evaluation of tendon modifications, which is hard to perform in human subjects. In conclusion, after a period of sudden detraining (e.g. after an injury), physical activity should be restarted with caution, following an appropriate rehabilitation program. However, further research should be performed to fully understand the effect of sudden detraining on tendons.

Ethics statement

(Authors are required to state the ethical considerations of their study in the manuscript including for cases where the study was exempt from ethical approval procedures.)

Did the study presented in the manuscript involve human or animal subjects: No

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30 ABTRACT

Introduction: Several conditions such as training, aging, estrogen deficiency and drugs could affect the biological and anatomo-physiological characteristics of the tendon. Additionally, recent preclinical and clinical studies examined the effect of detraining on tendon, showing alterations in its structure and morphology and in tenocyte mechanobiology. However, few data evaluated the importance that cessation of training might have on tendon. Basically, we do not fully understand how tendons react to a phase of training followed by sudden detraining. Therefore, within this review, we summarize the studies where tendon detraining was examined.

38 Materials and methods: A descriptive systematic literature review was carried out by searching 39 three databases (PubMed, Scopus and Web of Knowledge) on tendon detraining. Original articles in 40 English from 2000 to 2015 were included. In addition, the search was extended to the reference lists 41 of the selected articles. A public reference manager ("www.mendeley.com") was adopted to remove 42 duplicate articles.

Results: An initial literature search yielded 34 134 references (www.pubmed.org: 17 53;
www.scopus.com: 8 11; www.webofknowledge.com: 9 70). 11 15 publications were extracted
based on the title for further analysis by two independent reviewers. Abstracts and complete articles
were after that reviewed to evaluate if they met inclusion criteria.

47 Conclusions: The revised literature comprised 4 clinical studies and an *in vitro* and two three *in* vivo reports. Overall, the results showed that tendon structure and properties after detraining are 48 compromised, with an alteration in the tissue structural organization and mechanical properties. 49 Clinical studies usually showed a lesser extent of tendon alterations, probably because preclinical 50 studies permit an in-depth evaluation of tendon modifications, which is hard to perform in human 51 subjects. In conclusion, after a period of sudden detraining (e.g. after an injury), physical activity 52 should be taken with caution, following a targeted rehabilitation program. However, further 53 research should be performed to fully understand the effect of sudden detraining on tendons. 54

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56 Key words: tendon, tenocyte, detraining, sudden detraining, systematic literature review

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62 INTRODUCTION

Tendons are a specialized tissues that join muscle to bone and are composed by extracellular 63 collagen fibers arranged in regular arrays (Aslan, 2008). This mechanosensitive tissue shows 64 detailed mechanical properties that allow it to adapt and respond to loading transmitted by muscles 65 66 (Fang, 2015). This load transfer provide the principal mechanical stimulus for tendon cells (Kondratko-Mittnacht 2015). These tensile loads are diverted to tendon cells through different 67 matrix compartments and components. At cellular level, by various transmembrane structures and 68 pathways, they are transduced from the exterior to intracellular biochemical responses (Maeda, 69 2015; Kondratko-Mittnacht 2015). 70

While physiologic loads are required to maintain tendon homeostasis, (Galloway, 2013) unusual 71 72 loading could direct to tendon injury, either through an acute traumatic injury or chronic, 73 degenerative process (i.e., tendinopathy) resulting from an increase of microdamages and an altered cell/matrix response (Arnoczky, 2007; Magnusson, 2010). Histopathologicaly, tendinopathy is a 74 unsuccessful healing response, represented by altered tenocytes proliferation, disruption and 75 76 impaired organization of collagen fibers, increase in non collagenous matrix and neovascularization (Maffulli, 2011). In the chronic stage of tendinopathy, inflammation is 77 78 absent or minimal, nevertheless it could play a role only in the initiation, but not in the 79 propagation and progression, of the disease process (Maffulli, 2010). Even if tendinopathies also comprise conditions of damage to the tendon without symptoms, these pathologies 80 frequently occur with pain in the injured tendon, which is accentuated or appears during 81 palpation of the affected area or during active and passive movements involving the tendon 82 83 (Franceschi, 2014). Tendon injury may not only lead in the lack of mobility or irregular joint 84 kinematics, but could also result in damages to tissues adjacent to the joint. Muscle atrophy subsequent to tendon rupture is a frequent complication found by physicians and orthopedic 85 86 surgeons. This condition proves significantly weaker musculature resulting in unfavorable functional consequences, with a consequent reduction in muscle force generation (Sandri, 87 88 2008; Zhang, 2013). Despite previous studies showed complete histological and biochemical 89 characteristics of tendons rupture and some of these have been included into the clinical 90 scenario, little is known concerning the mechanical response of muscles to tendon injury (Sandri, 2008; Zhang, 2013; Jamali, 2000; Charvet, 2012). However, recently Zhang et al. 91 92 demonstrated that tendon rupture has a supplementary influence on muscle biomechanics in comparison to disuse (Zhang, 2013). 93

94 Due to their poor healing ability, tendon injuries represent an increasing problem in orthopedics as 95 physicians are faced with a growing demand in sports and recreation and in the aging population

(Kaux, 2011). Thus, primary disorders of tendons are a widely distributed clinical problem in 96 97 society and hospital evidence and statistical data suggest that some tendons are more susceptible to pathology than others; these are the rotator cuff, Achilles tibialis posterior and patellar tendons. 98 Although there are no specific figures in relation to tendon disease, several studies show that 16% 99 of the population is affected from tendon pain (Urwin, 1998) and this rises to 21% when the 100 statistics shift to elderly hospitals and community populations (Urwin, 1998, Chard, 1991). These 101 numbers supplementary enhance in the sports community, in fact it was reported that 30 to 50% of 102 all sporting injuries involve tendons (Kannus, 1997). Ordinarily, the major conditions affecting 103 104 tendons are tendinitis and tendinosis; the first assumed to be accompanied by inflammation and 105 pain, whereas the second can be caused by tendinous degeneration (Maffulli, 1998). It is assumed 106 that these conditions are seldom spontaneous (Gibson, 1998) and are not caused by single factors. Rather, they are the end result of a variety of pathological processes (Riley, 2004; Rees, 2006) 107 108 which can ultimately lead to the main clinical problem: loss of tissue integrity with full or partial tendon rupture. 109

Many intrinsic and extrinsic factors such as aging, gender, anatomical variants, obesity, systemic 110 diseases, estrogen deficiency, drugs, sporting activities, physical loading, occupation, and 111 112 environmental conditions could affect the biological and anatomo-physiological characteristics of the tendon (Sandberg, 2015; Frizziero, 2014; Galdiero, 2014; Oliva, 2014a, 2014b; Snedeker, 113 2014; Abate 2014; Hast, 2014; Boivin, 2014; Berardi, 2014; Franchi, 2013; Frizziero, 2013; 114 Malliaras, 2013; Moerch, 2013; Torricelli, 2013; Frey, 2007; Holmes, 2006; Torricelli, 2006; 115 Nakama, 2005;). Thus, over the past decade, tendon and tenocyte adaptations in relation to 116 immobilization, training, aging and medications have been the center of an growing number of 117 studies (Maffulli, 2003; Sharma, 2005; Stanley, 2008; Torricelli, 2006; Torricelli, 2013). 118

While proper mechanical loads at physiological levels are typically helpful to tendons in terms of 119 enhancing its mechanical properties, recent preclinical and clinical studies examining the effect of 120 detraining on tendon, showed alterations in its structure and morphology and in tenocyte 121 mechanobiology. However, there is a paucity of data that evaluated the impact that detraining may 122 123 have on tendon. Thus, it has not yet been understood how tendons behave to a period of training followed by cessation of training. Nevertheless, to guide rehabilitation and/or athletic programs it is 124 125 necessary to elucidate tendon adaptation after sudden detraining. Therefore, within this descriptive systematic literature review, we summarize the studies where tendon detraining was examined. 126

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128 MATERIALS AND METHODS

Descriptive literature review

According to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) 130 a systematic search was carried out for this descriptive literature review (see Figure 1 for details) in 131 three databases (www.pubmed.org, www.webofknowledge.com, www.scopus.com). The keywords 132 were "tendon detraining", "tendon detraining associated damage", "tendon sudden detraining", 133 "tendon disuse", "tendon discontinuous training associated damage", "tendinopathies and 134 discontinuous training", "tendinopathies and detraining", "tenocytes detraining associated 135 damage", "tenocytes discontinuous training associated damage". We sought to identify studies 136 in which tendon detraining was examined. Publications from 2005 to 2015 (original articles in 137 138 English) were included. The reference lists from the articles included in this review were analyzed to recognize additional studies that were not found by the initial search. A public reference manager 139 140 ("www.mendeley.com") was used to delete duplicate articles.

141

142 **RESULTS**

An initial literature search yielded 34 134 references. Seventeen Fifty-three articles were identified 143 144 using www.pubmed.org, 9 70 articles using www.webofknowledge.com and 8 11 articles were found in www.scopus.com.The resulting references were submitted to a public reference manager 145 (Mendeley 1.13.8, "www.mendeley.com") to delete duplicate articles. Of the 20 76 remaining 146 articles, 11 publications were extracted based on the title for further analysis. Abstracts and 147 whole articles were then reviewed to ascertain whether the publication met the inclusion criteria and 148 seven 8 articles (3 4 preclinical studies, 1 in vitro and 2 3 in vivo, and 4 clinical studies) were 149 considered appropriate for the review (Figure 1). From the reference lists of the included articles, 150 no- supplementary publications were identified. We did not perform meta-analyses of the selected 151 studies, but quoted the results in a descriptive fashion. 152

- 153
- **Figure 1:**Literature search strategy and criteria.

155 **Preclinical studies**

This revised literature comprised 3 4 preclinical studies, an *in vitro* and two three *in vivo* reports, 156 respectively on tenocytes from patellar tendon (Salamanna, 2015) and on patellar (Frizziero, 2011 157 and 2015) and gastrocnemius (Foutz, 2007) tendon of detrained rats animals (Salamanna, 2015; 158 Frizziero, 2011 and 2015). Concerning the *in vitro* study patellar tendon tenocytes from rats 159 subjected to training and to sudden detraining were examined. Rats were trained for 10 weeks on a 160 treadmill (speed of about 25 m/min, corresponding to ~65-70% VO₂max) and successively caged 161 without exercise for further 4 weeks. Tenocytes from patellar tendon were cultured to evaluate 162 morphology, viability, proliferation and metabolic activity. It was found that detraining in the short-163

term alters tenocyte synthetic and metabolic activity (C-terminal-propeptide of type I collagen, 164 collagen III, fibronectin, aggrecan, tenascin-c, interleukin-1ß, matrix-metalloproteinase-1 and-3). 165 These results indicated that tenocytes do not merely have a passive role but play an important 166 function during detraining (Salamanna, 2015). Similarly results were found by the same authors 167 also when the patellar tendons of detrained rats were studied by histology and histomorphometry 168 (Frizziero, 2011 and 2015). In fact, the studies showed alteration in tendon morphology and also in 169 its enthesis due to discontinuation of training. These alteration involved proteoglycan content, 170 collagen fiber organization with an increase of collagen III and a decrease of collagen I, which 171 172 means less resistance to stress, and a related increased risk of rupture. Differently from the above 173 mentioned studies, Foutz et al. investigated the mechanical adaptability responses due to 174 disuse on the biomechanical properties of the gastrocnemius tendon of chicks (Foutz, 2007). Chicks were trained for 3 weeks on a treadmill (speed of 0.22 m/s, for 5 min) and successively 175 176 immobilized in a whole body suspension system for further 2 weeks. It was found that structural strength and toughness of the gastrocnemius tendon were reduced by 10 and 30%, 177 178 respectively, whereas the material strength, material toughness, and material stiffness of the tendon increased by approximately 75, 65, and 70%, respectively. These results showed that 179 180 the chicken gastrocnemius tendon reacts to mechanical disuse as foretold by the mechanobiology process (Foutz, 2007). 181

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183 Clinical studies

184 The PubMed, Web of Knowledge and Scopus search strategy identified 4 clinical papers that examined the impact that detraining may have on tendons. Several studies showed that tendon 185 characteristics influence the performances during stretch-shortening cycle exercises (Bojesen-186 Moller, 2005; Kubo, 2007; Stafilidis, 2007); thus, information on the time course of changes in 187 tendon characteristics during training and detraining is critical for the progress of performances in 188 the athletic field. To evaluate the time course of modifications in mechanical and morphological 189 properties of tendon during detraining, Kubo et al (Kubo, 2010) examined these variables in eight 190 volunteered men that executed unilateral knee extension exercise in a seated position. Subjects were 191 192 trained 4 times per weeks for 3 months and detrained for the following three months. Results of this study showed that tendon stiffness was significantly increased after 3 months of training, while the 193 194 maximal elongation was unaltered. Conversely, during the detraining period, tendon showed greater values of maximal elongation compared to the post-training, and tendon stiffness decreased to the 195 pre-training levels after 2 months of detraining (Kubo, 2010). With a similar methodology, the same 196 authors in 2012 focused more specifically on the alterations found in the human Achilles tendon 197

during training and detraining (Kubo, 2012). In addition, they measured the blood volume and 198 oxygen saturation of tendon, and evaluated the serum concentrations of markers of collagen type I 199 synthesis. Results were similar to the previous study ones: the elongation values did not change 200 after training but increased significantly during detraining; tendon stiffness increased only after 201 202 three months of training and rapidly decreased during detraining. Thus, Authors showed that during detraining, the sudden decrease in tendon stiffness might be linked to modifications in the structure 203 of collagen fibers within the tendon. In addition, no significant alterations in blood supply or 204 collagen synthesis were observed (excluding an increase in procollagen peptides after 2 months of 205 206 training) (Kubo, 2012).

Recently McMahon et al (McMahon, 2013) evaluated the patella tendon properties during 207 208 detraining (1 months), after a 3-months period of training with different strains. The patella moment arm, the perpendicular distance between the tibiofemoral contact point and the mid-portion of the 209 210 tendon, was estimated using dual-energy x-ray absorptiometry (DEXA) scan images. Tendon elongation and stiffness were measured by ultrasonic analyses and tendon forces were calculated as 211 212 the ratio between the measured torque and the patella moment arm. Furthermore, they evaluated the circulating transforming growth factor (TGF)-B1 levels as it is associated to exercise-induced 213 214 response to mechanical loading of muscle and tendon. The authors found no significant alterations in patella tendon dimensions or circulating TGF-\beta1 levels following training or detraining. 215 However, the training groups with the muscle-tendon complex at a lengthened position or over a 216 wide range of motion better maintained adaptations compared to the training in a shortened position 217 subsequent to detraining, with a pattern of slower loss of progress at the early phase of detraining in 218 219 all training groups.

Finally Kannas and colleagues (Kannas, 2014) analyzed the effect of 4 weeks of detraining on the mechanical properties of medial gastrocnemius aponeurosis into two groups that performed plyometric training on incline and plane ground. They evaluated the aponeurosis strain of medial gastrocnemius and found that it decreased after detraining; the ankle muscle tendon complex properties withdrew to the pre-training values with lower performances. These findings suggested that after four weeks of detraining, ankle muscle tendon complex properties withdraw to the pretraining values with lower performance (Kannas, 2014).

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228 **DISCUSSION**

The tendon is a connective tissue responsible for the transmission of force from the muscular tissue to the bones, promoting body movement. It is not a static tissue, preferentially it adapts itself in compliance to the level, direction and frequency of the load that is applied to it with a process ofremodeling possibly executed by tenocytes.

It was shown that appropriate mechanical loads are useful to tendons by improving their anabolic 233 processes and it is undertaken or prescribed for different reasons such as sports performance, 234 general health, functional maintenance, recovery (e.g., following injury, illness/diseased states) and 235 also to compensate the effects of ageing. However, extreme mechanical loads are harmful to 236 tendons by bringing catabolic processes such as matrix degradation. Immobilization or disuse of 237 tendons also leads catabolic effects on it. Differently there are few data that examined the impact 238 239 that detraining may have on tendons. Thus, the present descriptive systematic literature review tried 240 to summarize the effects of discontinuing physical activity on tenocyte metabolism and/or in tendon 241 morphology in order to elucidate the mechanism behind these changes.

All examined studies, both preclinical and clinical, observed that discontinuing activity negatively 242 243 influence tendon structure and morphology, albeit with differences in the training and/or detraining protocols, in the types of tendons, in subjects involved, in the study design or in the experimental 244 245 setting involved. The results of all these studies suggested that after a period of sudden detraining 246 (such as after an injury) physical activity should be restarted with caution and with appropriate 247 rehabilitation programs because cessation of activity causes modifications in tenocytes and tendons metabolism, morphology, i.e. in collagen type I and III synthesis, collagen organization, cellularity, 248 vascularity, proteoglycan content, tear density, mechanical properties. 249

Notwithstanding the alterations highlighted in the reviewed articles after tendon detraining, some limitations of the examined studies should be also considered. In fact, this systematic review has as its main focus not only to bring together major works involving major changes in morphological and structural properties of tendons during detraining, but also to examine the methodological process on which the articles were based to assess the trustworthiness of the results found.

255 In relation to the results obtained in the *in vitro* study examined in this review (Salamanna, 2015), 256 that showed a decrease of tendon mitochondrial area, rough endoplasmic reticulum area, C-terminal propeptide of type I collagen, fibronectin, aggrecan and tenascin-c synthesis and presence of 257 258 inflammatory cytokine production, we have to consider that tenocytes from animals subjected to sudden detraining were studied. In addition, results were obtained in in vitro cultured cells, which 259 260 were not any longer structured into tissues, but in monolayer and static conditions. Thus, it is 261 probable that the performance of explanted tendon cells is not equal to the performance of tendon cells in their native matrix environment in vivo (Fu, 2008; Leigh, 2008). However, these results 262 indicated that the tendon does not operate as a inert connector between muscles and bone, but 263 264 dynamically responds to mechanical loading.

The two three preclinical studies examined in this review employed a rat or chicken animal model 265 266 that may not be fully representative of human conditions but the invasive analyses conducted in these studies permitted a depth investigation for the advancement of knowledge of many aspects on 267 tendon response to detraining (Frizziero, 2011 and 2015; Foutz, 2007)). Moreover, looking at the 268 literature, rat and rodents are the most used animals when mechanical load with treadmill running is 269 used (Warden, 2009; Lui, 2011). In fact, the results of these in vivo studies demonstrated that the 270 adopted running protocol did not induce tendinopathy or other pathologic changes in hindlimbs. 271 Another methodological process that must be considered is that in these studies all morphometric 272 273 parameters were measured by 2D image analysis, while other investigation methods, such as micro-274 MRI, may allow a more in-depth understanding of tendon structure. However, as for the reviewed 275 in vitro paper, these in vivo results provide interesting data for both sports medicine practitioners and orthopedic surgeons, wishing to prevent the pathological or degenerative modification that 276 277 affect these structures.

Great variability was noted in the four clinical studies (Kubo, 2010 and 2012;McMahon, 2013; 278 279 Kannas, 2014) that analyzed the effects of detraining. In fact, these studies involved different tendons (Achilles, gastrocnemius, patellar), different types of exercise (isometric knee extention, 280 resistance training, plyometric training on incline and plane ground), different training and 281 detraining periods (3 and 4 months) and different types of analyses (Dual Energy X-Ray 282 Absorptiometry, ultrasonography, electromyography). Furthermore, it is important to point out that 283 284 the different effects of detraining on tendons depends not only on the above mentioned variables, but also on the patient intrinsic characteristics, that are affected by age, gender, drug assumption, 285 286 the presence of systemic or genetic or endocrine diseases (i.e. obesity, diabetes, Cushing syndrome, hypercholesterolemia, osteoporosis). In fact, recently it was shown that proliferation and synthetic 287 activity of tenocytes are negatively affected by aging and estrogen deficiency (Torricelli, 2013). In 288 addition, clinical studies did not permit a depth understanding of the alteration in tendon 289 290 metabolism and morphology (i.e. expression of type I collagen, fibronectin, aggrecan and tenascin-291 c synthesis and/or presence of inflammatory cytokine, cellularity, vascularization, fibers 292 arrangements ect). However, despite these limitations these clinical studies indicate that tendons 293 may be susceptible to detraining. These findings could have a direct relevance to functional 294 rehabilitation practices showing that after a period of sudden detraining, physical activity should be restarted with caution. 295

Despite the fact that the examined studies showed a potential negative effect of detraining on tenocytes and tendons, there is a paucity of preclinical and clinical studies that examined the importance that cessation of training may have on tendon. These results should be confirmed by

other preclinical and clinical research in order to completely comprehend the effect of detraining on 299 tendons. In particular, several aspects should be further studied and refined in order to improve our 300 understanding on the role of detraining in tenocytes and tendon mechanobiology: 1) standardization 301 of the training and detraining protocols in both preclinical and clinical research; 2) development of 302 303 systems that reproduce tendon detraining in culture with high reliability to native tendon; 3) comprehend how tenocytes respond to detraining and how they mechano-regulate their response; 4) 304 305 evaluate the presence of altered tendon structure and/or morphology due to detraining in its various stages; 5) evaluation of the role of other tissues (bone, muscle, nerve, vascularity, etc.) on tendon 306 mechanobiology during detraining. Finally an integrated, collaborative multi-disciplinary multiscale 307 approach is likely to yield the greatest advances in this field. 308

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Table 1:

Table 1:							
Experimental set-up	Type of tendon	Control group	Training protocol	Detraining protocol	Analysis	Main results	Reference
In vitro model	Rat patellar tendon tenocyte	Untrained patellar tendon tenocyte Trained patellar tendon tenocyte	10 week on a treadmill (~65–70% VO ₂ max)	Caged without exercise for 4 weeks	Transmission- electronic-microscopy, C-terminal-propeptide of type I collagen, collagen III, fibronectin, aggrecan, tenascin-c, interleukin-	Altered tenocyte synthetic and metabolic activity.	Salamanna et al 2015
					1β, matrix- metalloproteinase-1 and-3.		
<i>In vivo</i> model	Chicken gastrocnemius tendon	No control group	3 week on a treadmill (speed of 0.22 m/s, for 5 min)	Controls or immobilized for 2 weeks.	Tendon midregion cross-sectional area and biomechanical properties	Gastrocnemius tendon responds to mechanical disuse as predicted by the mechanobiology process	Foutz et al. 2007
In vivo model	Rat patellar tendon	Untrained patellar tendon Trained patellar tendon	10 week on a treadmill (~60% VO ₂ max)	Caged without exercise for 4 weeks	Collagenfiberorganizationandproteoglycan content.	Low proteoglycan content and collagen fiber organization .	Frizziero et al 2011
In vivo model	Rat patellar tendon	Untrained patellar tendon Trained patellar	10 week on a treadmill (~65–70% VO ₂ max)	Caged without exercise for 4 weeks	Structureandmorphology (modifiedMovinscore, tear	Altered structure and morphology with the highest Movin score	Frizziero et al 2015

tendon		density, collagen type I	values, the highest	
		and III).	percentage of	
			collagen III and the	
			lowest of collagen I	

312313 Table 2:

Type of	Patients	Training protocol	Detraining protocol	Analysis	Main results	Reference
tendon						
Patellar	8 (training group);	Unilateral isometric knee	Return to usual levels of	- Tendon elongation by	Greater values of tendon	Kubo et al 2010
tendon	6 (control group).	extension, 4 times/week, 3	physical activity, 3	ultrasounds;	elongation, decrease in	
		months.	months.	- Cross-sectional areaby MRI.	tendon stiffness during	
					detraining.	
Achilles	9 (training group);	Unilateral (left side)	Return to usual levels of	- Tendon elongation by	Tendon elongation	Kubo et al 2012
tendon	7 (control group).	isometric plantar flexion	physical activity, 3	ultrasounds;	increased and stiffness	
		exercise, 4 times/week, 3	months.	- Cross-sectional areaby MRI;	rapidly decreased after	
		months.		- Blood supply and oxygen	detraining.	
				saturation;		
				- Serum concentration of BAP and		
				P1P by ELISA.		
Patellar	10 (training with the	Resistance training, three	4 weeks of detraining	- Patella moment arm by DEXA;	No significant alterations	McMahon 2013
tendon	MTC at a shortened	times per week, 8 weeks.		- Tendon elongation and stiffness	in patella tendon	
	position);			by ultrasounds;	dimensions or circulating	
	11 (MTC at a lengthened			- Circulating TGF-β1 levels by	TGF-β1 levels following	
	position);			ELISA.	training or detraining in	
	11 (wide range of				any of the groups.	

	motion); 10 (control group).					
Achilles	10 (training on inclined	Plyometric training	4 weeks of detraining	Aponeurosis strain of MG	Strain was decreased	Kannas 2014
tendon	ground)				from 22.7% (± 0.05) to	
	10 (training on plain				16.3% (± 0.05) after	
	ground)				detraining period.	

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