Tensiomyographical responsiveness to peripheral fatigue in quadriceps femoris (#44194)

First revision

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Tensiomyographical responsiveness to peripheral fatigue in quadriceps femoris

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Background: Fatigue influences athletic performance and can also increase the risk of injury in sports and most of the methods to evaluate it require an additional voluntary effort. Tensiomyography (TMG), which uses electrical stimulation and a displacement sensor to evaluate muscle contraction properties of one or more muscle bellies, has emerged as a technique that can assess the presence of peripheral and central fatigue without requiring additional voluntary efforts. However, the evaluation of the TMG's ability to detect fatigue is limited, both at the level of muscle bellies and statistical methods. Thus, the aim of the present study was twofold: (i) to examine and compare the tensiomyographical responsiveness to quadriceps femoris (QF) fatigue by multiple statistical methods and (ii) to analyze sex differences in the variation produced by fatigue in TMG parameters. **Methods:** Thirty-nine recreational athletes participated (19 males/20 females; aged 22 \pm 2 years). TMG parameters of QF bellies and maximal voluntary isometric contraction (MVIC) were measured before and after a fatigue protocol. TMG parameters used were maximum radial deformation (Dm), contraction time between 10-90% of the Dm (Tc), contraction velocity between 10-90% (Vc) and of the first 10% (V10) of the Dm. Internal responsiveness of TMG to fatigue was analyzed by paired t-test and standardized response mean (SRM). External responsiveness was examined by correlations, regression models, and receiver operating characteristic (ROC) curves.

Results: All TMG parameters, except for Tc of rectus femoris and vastus medialis, showed large internal responsiveness. In adjusted regression models by sex, only Dm and V10 of rectus femoris were statistically associated (p < 0.05) with b coefficients of 0.40 and 0.43, respectively. r2 explained the 22% of the total variance. In addition, these parameters could discriminate between QF with and without fatigue. **Conclusion:** Since the QF is the main strength contributor during multiple physical activities, clinicians and trainers will be

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able to discriminate the presence of fatigue and the magnitude of changes in the QF strength by TMG evaluation.



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15 16

Abstract

- 17 **Background:** Fatigue influences athletic performance and can also increase the risk of injury in
- 18 sports and most of the methods to evaluate it require an additional voluntary effort.
- 19 Tensiomyography (TMG), which uses electrical stimulation and a displacement sensor to
- 20 evaluate muscle contraction properties of one or more muscle bellies, has emerged as a technique
- 21 that can assess the presence of peripheral and central fatigue without requiring additional
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- 27 participated (19 males/20 females; aged 22 ± 2 years). TMG parameters of QF bellies and
- 28 maximal voluntary isometric contraction (MVIC) were measured before and after a fatigue
- 29 protocol. TMG parameters used were maximum radial deformation (Dm), contraction time
- 30 between 10-90% of the Dm (Tc), contraction velocity between 10-90% (Vc) and of the first 10%
- 31 (V10) of the Dm. Internal responsiveness of TMG to fatigue was analyzed by paired t-test and
- 32 standardized response mean (SRM). External responsiveness was examined by correlations,
- 33 regression models, and receiver operating characteristic (ROC) curves.
- 34 Results: All TMG parameters, except for Tc of rectus femoris and vastus medialis, showed large
- 35 internal responsiveness. In adjusted regression models by sex, only Dm and V10 of rectus
- 36 femoris were statistically associated (p < 0.05) with b coefficients of 0.40 and 0.43, respectively.
- 37 r2 explained the 22% of the total variance. In addition, these parameters could discriminate
- 38 between QF with and without fatigue. Conclusion: Since the QF is the main strength contributor



39 during multiple physical activities, clinicians and trainers will be able to discriminate the

40 presence of fatigue and the magnitude of changes in the QF strength by TMG evaluation.

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Introduction

- 43 Fatigue is defined as a decline in muscular performance which produces a reduction in strength
- and power generation (Ditroilo et al., 2011). It can be further explained by factors related to the
- 45 central nervous system as changes at the spinal level (Gandevia, 2001) or by peripheral factors
- associated to the muscle, such as failure of transmission at the neuromuscular junction (Allen,
- 47 Lamb & Westerblad, 2008). Its manifestation can vary in subjects with different training
- 48 backgrounds (Garrandes et al., 2007), type of muscle contraction performed (Kay et al., 2000), or
- 49 even between sex (Albert et al., 2006; Martin & Rattey, 2007; Ansdell et al., 2017).
- 50 Since fatigue influences athletic performance (Thorlund et al., 2008; Ditroilo et al., 2011) and
- 51 can also increase the risk of injury in sports (Zebis et al., 2011; Liederbach et al., 2014), its
- 52 study has been of interest. Multiple methods have been used to induce fatigue, both central
- fatigue in several muscle groups or peripheral fatigue in a specific muscle (García-Manso et al.,
- 54 2011; Hunter et al., 2012; Macgregor et al., 2016; Wiewelhove et al., 2017, 2018). Thus, fatigue
- has been evaluated after short term (Macgregor et al., 2016; Abelairas-Gómez et al., 2018) and
- long duration efforts, such as several days of intense training sessions (Wiewelhove et al., 2017),
- and also after isolated long sessions (2-12h approximately) (Lepers et al., 2002; García-Manso et
- 58 al., 2011; Wiewelhove et al., 2018).
- 59 The most used fatigue evaluation methods have been based on changes in maximal voluntary
- 60 isometric contractions (MVICs) (Lepers et al., 2002; Zebis et al., 2011), muscle activation
- 61 (Garrandes et al., 2007; Thorlund et al., 2008), kinematics and kinetics measurements
- 62 (Liederbach et al., 2014; Tam et al., 2017), biochemical markers (Gorostiaga et al., 2012), or
- 63 muscular contractile properties (García-Manso et al., 2011; de Paula Simola et al., 2016). In a
- 64 situation of fatigue, most of these methods would require an additional voluntary effort. Their
- application therefore would not be practical or safe facing the possible presence of central
- 66 inhibition (Graven-Nielsen et al., 2002), or the possibility of increase any extant muscular
- 67 damage (Macgregor et al., 2016).
- 68 Tensiomyography (TMG), which uses electrical stimulation and a displacement sensor to
- 69 evaluate muscle contraction properties of one or more muscle bellies (Valencic & Knez, 1997),
- 70 has emerged as a technique that can assess the presence of peripheral and central fatigue without
- 71 requiring additional voluntary efforts (García-Manso et al., 2011; de Paula Simola et al., 2016).
- 72 Peripheral fatigue has been evaluated by TMG for specific muscle group from both lower and
- value of the result of the res
- al., 2016). In contrast, central fatigue has been evaluated only in the lower limb, being
- 75 quadriceps femoris (QF) the most studied muscle group (García-Manso et al., 2011; de Paula
- 76 Simola et al., 2015, 2016; Giovanelli et al., 2016; Raeder et al., 2016; Wiewelhove et al., 2017).
- 77 Responsiveness is defined as the ability of a tool to detect important clinical changes over time
- 78 (Guyatt et al., 1989). Since this characteristic is essential to assess fatigue by TMG, it has been



79 analyzed by multiple studies (García-Manso et al., 2011; Hunter et al., 2012; de Paula Simola et al., 2015, 2016; Giovanelli et al., 2016; Macgregor et al., 2016; Raeder et al., 2016; Wiewelhove 80 et al., 2017; Abelairas-Gómez et al., 2018). Most of these studies evaluated one muscle belly and 81 they used one or two statistical methods of either internal responsiveness (e.g. paired t-test and 82 83 effect size) or external responsiveness (correlation with reference measure or regression models) Internal responsiveness is the ability of a measure to change over a set period and external 84 responsiveness reflects the extent to which changes in a measure over a specified time frame 85 related to corresponding changes in an external reference measure of health status (Husted et al., 86 2000). Overall, TMG of those evaluated muscle bellies has shown to be internally and externally 87 responsive in assessing central fatigue (García-Manso et al., 2011; de Paula Simola et al., 2015, 88 2016; Giovanelli et al., 2016; Raeder et al., 2016; Wiewelhove et al., 2017), and internally 89 responsive to peripheral fatigue (Hunter et al., 2012; García-Manso et al., 2012; Macgregor et al., 90 91 2016; Abelairas-Gómez et al., 2018). However, to the best of our knowledge, the external 92 responsiveness of TMG has not been yet assessed for peripheral fatigue, and therefore comparisons between internal and external responsiveness has not been established. 93 Furthermore, to our knowledge, TMG responsiveness has not been simultaneously evaluated in 94 multiple bellies, neither analyzed by by multiple statistical indicators of responsiveness. At the 95 same time, understanding the mechanisms behind the changes in TMG parameters caused by 96 fatigue in both sexes, is also an area of research that needs further development. 97 Therefore, the primary objective of our study was to examine and compare the responsiveness of 98 TMG parameters to QF peripheral fatigue of three muscle bellies [rectus femoris (RF), vastus 99 lateralis (VL), and vastus medialis (VM)] by multiple statistical methods. A secondary objective 100 101 was to examine whether there are differences between sex in the variation produced by fatigue in TMG parameters. Our hypotheses were: QF bellies have different responsiveness to peripheral 102 fatigue; and the changes of TMG parameters are similar between males and females. 103

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Materials & Methods

Study design

A single group pretest-posttest design was used, which involved repeated TMG and MVIC measures of the dominant lower limb QF before and after a fatigue protocol within the same session. Participants were physiotherapy students recruited by email using the University of Valencia Intranet. This study was conducted from April to July 2018. All measurements were carried out between 10 a.m. and 2 p.m in the clinical research laboratory of the Department of Physiotherapy (University of Valencia) at an ambient temperature 21–22 °C. An experienced examiner in the measurement techniques evaluated the participants. He was a physiotherapist who had used TMG and hand dynamometers both in research and in clinical practice for several years. Before participation, participants were informed of the study procedures and their possible associated risks. All of them provided written informed consent. This study was completed



following the principles outlined in the Declaration of Helsinki and it was approved by the Ethics Committee of the University of Valencia (Spain) (H1523633864087).

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Participants

- Thirty-nine recreational athletes were evaluated. All participants performed exercise 3 times per week and practiced activities such as running, swimming, cycling, or central strength training.
- The specific inclusion criteria were: (a) aged between 18 and 30 years, (b) not surgically
- operated on the lower limb, (c) without pain in the lower limb in the 2 months before data
- 126 collection, and (d) performing physical exercise a minimum of 3 days per week. The exclusion
- criteria were: (a) practicing a specific sport as an amateur or professional, (b) contraindication to
- the use of electrodes due to injury or allergy to the adhesive, and (c) nontolerance to electrical

129 stimulation.

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Procedures

Before starting the session, height was measured using a 1-millimeter sensitivity flexible tape measure, while body mass and body mass index (BMI) were assessed using a standardized body composition analyzer (Tanita BC 418 MA, Tanita Corp, Tokyo, Japan). Next, TMG parameters were measured and then, participants performed a warm-up, which consisted of 10 minutes cycling at comfortable speed (80 revolutions per minute) with low resistance and the performance of three submaximal isometric contractions of isometric knee extension (Martins et al., 2017). Following this, the MVIC test was performed. After the fatigue protocol, the order of the tests was reversed, and the strength test was performed first to reduce the time between MVIC and TMG tests in acute fatigue. A schematic representation of the experimental

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[Please insert Figure 1 about here]

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Tensiomyography measurements

procedures is reported in the Figure 1.

First, participants were placed supine and resting on the stretcher. The knee was placed at 120° of flexion (considering full extension at 180°), fixing such position with a triangular foam cushion (García-García et al., 2016; Martín-San Agustín et al., 2018). The area where the TMG sensor and electrodes were placed was shaved and cleaned with gauze and alcohol. The position of the sensor for each QF belly was determined using the anatomical criteria described in the literature (Dahmane et al., 2005; Tous-Fajardo et al., 2010; Rey, Lago-Peñas & Lago-Ballesteros, 2012). This position was marked with a permanent marker so that it would remain throughout the evaluation. The sensor was finally placed on this point perpendicularly to the thigh and the electrodes were placed at 5 cm distance from it, forming an imaginary straight line along the belly (Figure 2).

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[Please insert Figure 2 about here]

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160	The contractile properties of each belly were evaluated during an maximal elicited contractions
161	with the TMG electro stimulator (TMG-100 System). Starting from 20 mA with 1ms pulses,
162	each stimulation was increased by 10mA until achieving the maximum radial deformation (Dm)
163	of the muscular belly. A time of 10s was left between stimuli to minimize fatigue or potentiation
164	effects (Krizaj, Simunic & Zagar, 2008). Before data acquisition, a pilot test was done to verify
165	the functioning of the TMG. For each belly, spatial and temporal parameters were measured:
166	Dm, contraction time between 10 and 90% of the Dm (Tc), contraction velocity between 10 and
167	90% of the Dm (Vc), and contraction velocity of the first 10% of the Dm (V10). TMG has
168	proven to be a method with a high relative [ICC for Dm (0.91-0.99), Tc (0.70-0.98), and VC >
169	0.95] and absolute (low coefficient of variations for Dm, Tc, and VC) reliability (Martín-
170	Rodríguez et al., 2017; Lohr et al., 2018).

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Maximal voluntary isometric contraction test

- 174 MVIC of the QF was measured by a MicroFET2 handheld dynamometer (Hoggan Health
- 175 Technologies Inc., Salt Lake City, UT). Participants were seated in an isokinetic dynamometer
- 176 (Prima Plus, Easytech, Italy) with their torso and hips tied so they were stable, and with a 90° hip
- 177 flexion. MVIC was evaluated in 90° knee flexion, considering 0° the complete extension (Figure
- 178 3). MicroFET2 was fixed with a rigid belt perpendicular to the ankle 5 cm above the malleoli,
- 179 with a pad between the tibia and the dynamometer to minimize the discomfort caused by the

180 contact (Hansen et al., 2015).

- 181 After the warm-up, participants completed three MVIC for 5s, with a 60-second rest after each
- 182 repetition. Through verbal stimuli, participants were instructed to exert and maintain the
- 183 maximum effort during the session. MicroFET2 has proven to be a valid method to measure the
- MVIC of the QF with an excellent inter-examiner reliability (ICC: 0.93, 95% CI 0.83; 0.97) and
- a minimal detectable change (MDC) of 14.1 N*m (95% CI, 9.23; 22.01) (Hansen et al., 2015).

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[Please insert Figure 3 about here]

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Fatigue protocol test

- After performing the baseline measurements, participants were requested to implement a
- protocol based on a 60s fatiguing isometric contraction at 70% MVC (Melchiorri & Rainoldi, 2011). The experimental setup was the same as the one adopted during the MVIC test. The
- handheld dynamometer, previously set at 70% MVIC, was used to display the feedback
- 194 (Melchiorri & Rainoldi, 2011). It was considered that the fatigue was achieved when the
- 195 reduction of the MVIC was higher than the upper limit of the MDC reported in a previous study
- 196 (22.01 N*m) (Hansen et al., 2015).

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Statistical analysis



199 Baseline data were summarized as means and standard deviations (SD) for continuous variables and as absolute and relative frequencies for categorical variables. Variables were checked for 200 normality with the Kolmogorov-Smirnov test and homogeneity of variances with Levene's test. 201 A summary was also provided for participants with and without fatigued OF. 202 203 Paired t-tests were used to compare changes in the TMG parameters and MVIC within each sex group. These changes were also compared between sex groups by using non-paired t-tests. 204 Internal responsiveness was determined by the paired t-test and supplemented with an effect size 205 statistic, as recommended by Husted et al. (2000) [30]. To reduce the probability of getting false 206 positives, we increased the acceptance level from 0.05 to 0.01 for paired t-test because multiple 207 comparisons were made on the same data set. Of the current effect size statistics we used the 208 standardized response mean (SRM), which provides an estimate of the magnitude of change that 209 is not influenced by sample size (Navarro-Pujalte et al., 2018). It was calculated as 210 211 (MeanFollowup MeanBaseline)/Standard deviationFollowup-Baseline and the 95% confidence 212 intervals were calculated using the bootstrapping estimation method. Values of 0.20, 0.50, and 0.80 or higher have been proposed in the literature (Husted et al., 2000) to represent small, 213 moderate, and large responsiveness, respectively. Besides, we calculated the percentage of 214 participants that exceeded MDC. This statistic examines the extent to which change score 215 216 exceeds the amount of variability accounted by measurement error (Pardasaney et al., 2012), which is calculated as $SEMx1.96x\sqrt{2}$, where SEM is the standard error of measurement. 217 External responsiveness was determined by correlations, regression models, and receiver 218 operating characteristic (ROC) curves (Husted et al., 2000). The external criterion for assessing 219 the external responsiveness of the TMG tool was the magnitude of change in the MVIC. 220 221 We assumed that: (i) changes in the external criterion (MVIC) in participants with fatigue would be associated with changes in the TMG parameters; (ii) participants without fatigue would have 222 the smallest change in the TMG parameters (and therefore change in these TMG parameters can 223 be useful to classify participants' QF as fatigued or not fatigued). To test the first hypothesis, 224 225 correlations and simple and multiple linear regression models were used. In the regression models the explanatory variable was the change of each TMG parameter while the response 226 variable was the change in MVIC between before and after protocols. Each model was controlled 227 by sex, and comparisons were carried out between the presence or absence of this control. 228 229 Goodness-of-fit of the model was assessed by r2. To test the second hypothesis, we calculated the area under the ROC curve (AUC), which represents the probability that the measure of 230 correctly classifying participants has (Husted et al., 2000). An AUC > 0.70 was used as a generic 231 benchmark to consider acceptable its discriminant ability (Menaspà, Sassi & Impellizzeri, 2010). 232 233 For sample size calculation, we selected the multiple regression as the main statistic of responsiveness because it allowed us to examine change relationships controlling by a covariate 234 relevant in our study (sex). Regarding this statistic, we used the usual rule of thumb that 15 235 participants per predictor are needed for a reliable equation in multiple regression models 236 (Tabachnick & Fidell, 2007). We recruited a minimum of 30 participants assuming a maximum 237 238 of 2 explanatory variables (TMG parameter and sex). Statistical significance was set at p < 0.05.



239 240	All analyses were performed using the Statistical Package for the Social Sciences software program (SPSS version 24.0; IBM SPSS, Chicago, IL, USA).
241	
242	Results
243	Participants' characteristics
244	Baseline characteristics of participants are listed in Table 1. A total of 35 (89.7%) participants
245	achieved QF fatigue after the application of the fatigue protocol. They were 19 of 20 females
246	(95%) and 16 of 19 males (84.2%). Participants with and without fatigue showed no significant
247	differences (p >0.05) in any of their baseline characteristics.
248	
249	[Please insert Table 1 about here]
250	
251	Changes associated with the fatigue protocol
252	Participants with peripheral fatigue (n=35) had a significant decrease (31.5%) on their MVIC
253	after the fatigue protocol (from 203.3 N*m to 138.9 N*m). Table 2 shows that both sex groups
254	had a similar pattern of change: males reduced 30.8% and females 32.1%. Table 2 also shows
255	patterns of change by sex groups for TMG parameters of the RF, VL, and VM. All these
256	parameters, except for the Tc of the RF and VM, had significant differences within but not
257	between sex groups.
258	Figure 4 shows changes in TMG parameters for all participants with peripheral fatigue. All
259	parameters, except for Tc, showed a significant difference ($p < 0.001$) for the three bellies of the
260	QF. Dm's decrease ranged from 18.22% to 21.65%; Vc decreased from 15.62 to 22.20%, and
261	V10 decreased from 14.80% to 23.77%.
262 263	[Please insert Table 2 about here]
264	[1 lease misert Table 2 about here]
265	[Please insert Figure 4 about here]
266	[rease insert Figure 1 about here]
267	Internal and external responsiveness
268	Internal and external TMG responsiveness to fatigue of QF bellies is shown in Table 3. Internal
269	responsiveness statistics suggest that all TMG parameters, except for Tc of RF and VM, showed
270	large internal responsiveness (SRM> 0.8) among participants with QF fatigue. Dm and V10 in
271	RF were the parameters in which most of the participants exceeded the MCD (91.3% and 97.1%,
272	respectively). Only Dm, Vc, and V10 of the RF showed to be linearly associated with changes in
273	the MVIC. After controlling by sex, adjusted models typically provided b coefficients and r2
274	with small variations regarding their respective unadjusted model (range 0.01 to 0.05).
275	Consequently, Dm and V10 of RF were still statistically associated with b coefficients of 0.40
276	and 0.43, respectively. Moreover, the models of these parameters explained the 22% of the total
277	variance.



278 279	The AUC analysis suggests that changes of several TM G parameters (Dm in RF and VL, Tc in VL, and V10 in RF and VM) were >0.70 and could discriminate between QF with and without
280	fatigue. Also, the overlapping among their 95%CI suggests that none of these TMG parameters
281	is superior to the others to discriminate fatigue.
282	
283	[Please insert Table 3 about here]
284 285	
286	Discussion
287	To our knowledge, this is the first study to evaluate the internal and external TMG
288	responsiveness across a variety of QF muscle bellies to changes induced by peripheral fatigue.
289	We found that TMG parameters Dm and V10 of the RF showed both internal and external
290	responsiveness.
291	In our study, multiple statistical methods to evaluate the internal responsiveness (paired t-test and
292	SRM) and external responsiveness (correlations, regression models and ROC) of the TMG were
293	used, which is line with the recommendations of Husted et al. (2000). In previous studies, most
294	of these statistics have been used to evaluate only the TMG ability of change to fatigue (García-
295	Manso et al., 2011; de Paula Simola et al., 2015). Thus, to the best of our knowledge, this is the
296	first study to use several statistical methods to assess internal and external responsiveness.
297	Furthermore, since most of the previous studies assessing fatigue by TMG have only evaluated
298	isolated muscle bellies (García-Manso et al., 2011; Hunter et al., 2012; de Paula Simola et al.,
299	2015, 2016; Giovanelli et al., 2016; Macgregor et al., 2016; Raeder et al., 2016; Wiewelhove et
300	al., 2017)), our study presents novel findings in the evaluation of TMG across multiple muscle
301	bellies.".
302	Regarding the internal responsiveness, large and negative SRM of the TMG parameters were
303	found in most of the muscle bellies. Overall, our results are consistent with previous studies that
304 305	induced peripheral and central QF fatigue (i.e. selective QF fatigue or caused in the entire lower limb musculature). Therefore, the reduction of RF TMG parameters is consistent with previous
306	studies using peripheral (Carrasco et al., 2011) or central fatigue (de Paula Simola et al., 2015),
307	finding them reductions in Dm, VC, or V10 after fatigue due to cycling or strengthening. On the
308	other hand, the changes in VL and VM are also consistent with studies using central fatigue
309	caused by strengthening programs (de Paula Simola et al., 2016; Raeder et al., 2016). In addition
310	Dm results showed consistence with other studies that induced peripheral fatigue in muscles such
311	as the biceps brachii (Hunter et al., 2012; García-Manso et al., 2012) or the gastrocnemius
312	medialis (Macgregor et al., 2016). These findings could be explained by changes in the pH
313	(Hunter et al., 2009) and in different cellular molecules (e.g. Na+ or K+) (Brody et al., 1991),
314	which cause damage in the sarcolemma and the reduction of the electrical stimulus, with a
315	possible decrease in muscle displacement.
316	This study showed that Dm and V10 of RF had an acceptable external responsiveness in relation
317	to our external criterion, namely changes in the strength evidenced by MVIC. As reflected by the



- 318 regression coefficients, there was a moderate relationship between the amount of change in TMG
- 319 parameters and strength scores. This relationship is consistent with a previous study using central
- 320 fatigue (de Paula Simola et al., 2015). Furthermore, Dm and V10 were relevant according to sex,
- 321 which can be explained by the fact that our sample showed similar change magnitudes in both
- 322 TMG parameters and strength scores.
- 323 The fatigue protocol used in this study was highly effective (most of the QF showed fatigue).
- 324 Males and females had similar strength change scores (Table 2). Previous studies reported
- 325 different strength change scores between sexes when intensities between 25-50% of MVIC were
- used (Clark et al., 2005; Ansdell et al., 2017). In our study, an intensity of 70% of MVIC was
- 327 used, suggesting that as the contraction intensity increase, the sex differences in muscle fatigue
- decrease, (Hunter, 2014). Therefore, future investigations should examine whether sex
- 329 differences in strength changes are detected by sex differences in the TMG changes.
- Our present study also showed that TMG has discriminative ability to classify the participants'
- QF as having fatigue or not after the application of the protocol. Dm and V10 of the RF also
- were two of the four parameters with this discriminative ability. This finding is partially
- consistent with previous studies (Wiewelhove et al., 2017), who examined AUC of RF after
- central fatigue in elite young athletes. Nevertheless, while AUC values of V10 shown in this
- 335 study was similar to their results, AUC values of Dm was higher than previously published
- (Wiewelhove et al., 2017). Differences may be explained by the different type of fatigue (central
- 337 fatigue caused by several training sessions of high-intensity interval training vs peripheral fatigue
- by an MVIC test) or by the athletes' training background (junior tennis players vs recreational
- athletes). Other parameters with that discriminative ability were Dm and Tc of VL, and V10 of
- VM. Since this ability was not previously analyzed in these muscle bellies (VL and VM), results
- of the actual study supplements earlier findings which have only evaluated AUC for external
- responsiveness of the TMG in RF (Wiewelhove et al., 2017) and it provides evidence to expand
- 343 the application of the TMG to discriminate fatigue.
- Actual study has several limitations. First, we used a fatigue protocol based on MVIC, which
- induces peripheral fatigue. Therefore, our findings would be limited to be extrapolated to others
- fatigue situations (e.g. concentric contractions). Second, our study was conducted with
- recreational athletes (i.e. anyone participating in an aerobic or athletic activity at least three times
- per week) (Heinert et al., 2008). Since the contractile properties of the muscle are conditioned by
- 349 the type of exercise performed (Loturco et al., 2015), future research should compare our results
- with findings from athletes of different sports.
- Our study found that most of the TMG parameters showed an acceptable internal responsiveness
- of OF peripheral fatigue evidenced by a reduction of the MVIC. In contrast, only Dm and V10 of
- 353 RF showed external responsiveness. Therefore, our study illustrates that the use of only internal
- or external responsiveness may lead to incomplete conclusions (Husted et al., 2000). In this way,
- professionals should use both, as recommended by Husted (Husted et al., 2000).
- 356 This study showed that Dm and V10 of RF measured by TMG were both internally and
- externally responsive to changes between before and after a peripheral fatigue protocol. Since the



358	QF is the main strength contributor during cycling (Raasch et al., 1997) or running
359	(Montgomery, Pink & Perry, 1994), the fatigue evaluation after an effort is essential to manage
360 361	recovery of the athlete and the intensity of subsequent training sessions. Thus, clinicians and trainers should be able to direct the fatigue evaluations without making new efforts with TMG,
362	taking into consideration Dm and V10 parameters in RF to discriminate the presence of
363	peripheral fatigue and the magnitude of the strength changes and, in this way, be able to regulate
364	training loads (e.g. in the presence of peripheral fatigue, decrease intensity or activities that
365	involve the QF).
366	
367	Conclusions
368	According to the results, it can be concluded about positive responsiveness of the TMG in
369	peripheral fatigue of the QF, demonstrating that the Dm and V10 parameters of the RF present
370	acceptable responsiveness to fatigue. Therefore, by using the TMG, it is possible to determine
371 372	whether the QF shows peripheral fatigue or not, and to relate changes in the parameters with the reduction of strength. Thus, clinicians and trainers should be able to direct the fatigue evaluations
373	without making new efforts with TMG, facilitating the regulation of training loads. Finally,
374	future studies should examine the responsiveness of TMG to other types of fatigue and in other
375	sports.
376	•
377	
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Table 1(on next page)

Baseline characteristics of the participants in total and separated by fatigued condition.

Date represents mean and standard deviation unless otherwise noted. BMI: body mass index; Dm: maximal radial displacement; Tc, contraction time; Vc: contraction velocity between 10-90% of the Dm; V10: contraction velocity of the first 10% of the Dm; QF: quadriceps femoris; MVIC: maximal voluntary isometric contraction



Baseline Characteristics	Total (n=39)	Fatigued participants (n=35)	Non-fatigued participants (n=4)		
Males/females, N (%)	19 (48.7%)/20 (51.3%)	16 (45.7%)/19 (54.3%)	3 (75%)/1 (25%)		
Age (years)	22 (2)	22 (2)	21 (1)		
Physical activity (minutes)	316.5 (180.8)	314.6 (186.7)	332.5 (136.9)		
Anthropometric					
Body mass (kg)	67.37 (13.42)	66.10 (11.12)	78.55 (12.05)		
Stature (cm)	173.3 (9.50)	172.5 (9.09)	180.7 (11.24)		
BMI (kg/m2)	22.22 (2.72)	22.02 (2.71)	24 (2.53)		
QF strength					
MVIC (N*m)	207.56 (74.19)	203.31 (75.82)	244.72 (50.24)		
Tensiomyography parameters	` ,	, ,	, ,		
Rectus femoris					
Dm (mm)	10.26 (1.42)	10.32 (1.44)	9.76 (1.28)		
Tc (ms)	25.45 (4.04)	25.69 (3.95)	23.39 (4.84)		
Vc (mm/s)	327.96 (58.59)	326.62 (69.76)	339.70 (53.04)		
V10 (mm/s)	43.07 (5.32)	43.08 (5.39)	42.93 (5.33)		
Vastus lateralis					
Dm (mm)	5.74 (1.11)	5.63 (0.94)	6.64 (2.04)		
Tc (ms)	21.37 (3.02)	21.54 (3.11)	19.87 (1.35)		
Vc (mm/s)	217.78 (50.10)	211.58 (39.81)	271.95 (97.28)		
V10 (mm/s)	25.31 (5.18)	24.73 (4.21)	30.46 (9.98)		
Vastus medialis					
Dm (mm)	4.57 (0.85)	4.52 (0.64)	5.08 (2.01)		
Tc (ms)	19.60 (1.82)	19.61 (1.90)	19.48 (1.04)		
Vc (mm/s)	187.22 (33.12)	185.08 (26.57)	205.93 (73.31)		
V10 (mm/s)	23.22 (4.03)	22.97 (2.89)	25.37 (10.19)		

1



Table 2(on next page)

Differences within and between sex groups in the TMG parameters and MVIC after fatigue protocol.

SD: standard deviation; Dm: maximal radial displacement; Tc, contraction time; Vc: contraction velocity between 10-90% of the Dm; V10: contraction velocity of the first 10% of the Dm; QF: quadriceps femoris; MVIC: maximal voluntary isometric contraction.

	Males				Females			
Muscle	Baseline	Estimod	Differences		Baseline	Fatigued	Differences	
			Mean (SD); p	%	Daseinie	ratigueu	Mean (SD); p	%
QF strength								
MVIC (N*m)	272.1 (51.0)	187.3 (40.1)	84.7 (37.8); <0.001	30.8	145.4 (30.7)	98.1 (24.4)	47.3 (22.3); <0.001	32.1
Rectus femoris								
Dm (mm)	9.91 (1.66)	7.46 (1.87)	2.45 (1.27); <0.001	25.2	10.67 (1.16)	8.71 (1.76)	1.95 (1.13); <0.001	18.7
Tc (ms)	24.58 (4.25)	24.52 (6.37)	0.06 (3.28); 0.941	1.1	26.62 (3.52)	27.63 (5.43)	-1.01 (4.42); 0.334	4.1
Vc (mm/s)	330.01 (78.95)	250.71 (66.81)	79.30 (48.65); <0.001	21.8	373.76 (39.15)	256.21 (51.02)	67.55 (42.26); <0.001	20.9
V10 (mm/s)	43.17 (6.55)	32.78 (7.72)	10.39 (5.35); <0.001	24.4	43.01 (4.37)	33.01 (5.13)	10.00 (4.20); <0.001	23.2
Vastus lateralis								
Dm (mm)	5.47 (1.18)	4.48 (0.76)	0.99 (1.10); 0.003	20.5	5.78 (0.70)	4.10 (1.15)	1.68 (0.90); <0.001	29.5
Tc (ms)	21.69 (3.05)	19.93 (4.31)	1.76 (2.44); 0.011	8.6	21.42 (3.24)	19.04 (1.88)	2.38 (2.15); <0.001	10.4
Vc (mm/s)	203.67 (49.77)	179.33 (66.24)	24.35 (43.77); 0.042	12.8	218.24 (28.76)	170.24 (37.41)	48.00 (43.15); <0.001	20.9
V10 (mm/s)	24.28 (5.04)	20.46 (6.78)	3.82 (4.33); 0.003	17.3	25.10 (3.45)	18.65 (4.66)	6.45 (4.55); <0.001	25.3
Vastus medialis								
Dm (mm)	4.69 (3.91)	3.91 (0.78)	0.78 (0.59); <0.001	16.3	4.37 (0.50)	3.51 (0.69)	0.86 (0.53); <0.001	19.8
Tc (ms)	20.25 (1.78)	19.96 (2.66)	0.28 (1.97); 0.573	1.4	19.07 (1.88)	18.26 (1.88)	0.81 (1.64); 0.045	3.9
Vc (mm/s)	186.06 (30.93)	159.90 (25.72)	29.16 (22.46); <0.001	14.9	184.26 (23.12)	153.76 (29.26)	30.50 (26.86); <0.001	16.2
V10 (mm/s)	23.76 (3.19)	21.09 (3.95)	2.67 (2.97); 0.003	11.2	22.31 (2.51)	18.33 (3.40)	3.98 (2.74); <0.001	17.8



Table 3(on next page)

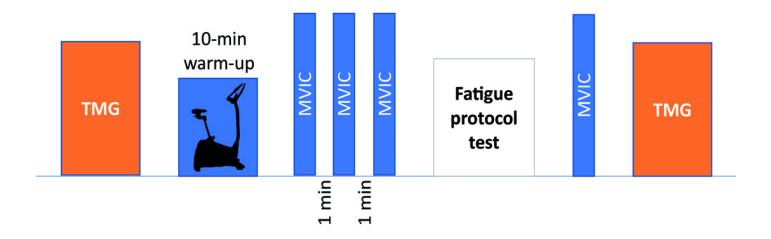
Responsiveness statistics for the TMG parameters

SRM: standardized response mean; CI: confidence interval; MCD: minimal detectable change; SE: standard error; AUC: area under curve; Dm: maximal radial displacement; Tc, contraction time; Vc: contraction velocity between 10-90% of the Dm; V10: contraction velocity of the first 10% of the Dm. † Adjusted by sex

Internal responsiveness			External responsiveness				
– Muscle	Paired t-test	SRM	%	Correlation method (Pearson's r and 95%	Linear regression method†		AUC (95% CI)
	(p)	(95% CI)	MCD	CI); <i>p</i>	b(SE); <i>p</i>	r2	
Rectus femoris							
Dm (mm)	0.001	-1.83 (-2.31; -1.47)	91.3	0.42 (0.12; 0.65); 0.004	0.40 (0.14); 0.007	0.22	0.73 (0.57; 0.86)
Tc (ms)	0.439	0.13 (-0.24; 0.39)	15.9	0.10 (-0.22; 0.40); 0.276	0.14 (0.15); 0.363	0.06	0.62 (0.45; 0.77)
Vc (mm/s)	0.001	-1.65 (-1.98; -1.30)	79.7	0.33 (0.02; 0.58); 0.020	0.26 (0.13); 0.052	0.13	0.59 (0.42; 0.74)
V10 (mm/s)	0.001	-2.20 (-2.65; -1.78)	97.1	0.45 (0.15; 0.67); 0.002	0.43 (0.15); 0.006	0.22	0.73 (0.57; 0.86)
Vastus lateralis							
Dm (mm)	0.001	-1.33 (-1.74; -0.82)	79.7	0.18 (-0.14; 0.47); 0.133	0.10 (0.12); 0.403	0.05	0.81 (0.65; 0.92)
Tc (ms)	0.001	-0.87 (-1.27; -0.41)	65.2	0.12 (-0.12; 0.48); 0.111	0.23 (0.19); 0.238	0.07	0.92 (0.79; 0.98)
Vc (mm/s)	0.001	-0.86 (-1.21; -0.46)	43.5	0.09 (-0.23; 0.39); 0.298	0.03 (0.11); 0.782	0.04	0.55 (0.39; 0.71)
V10 (mm/s)	0.001	-1.17 (-1.56; -0.71)	68.1	0.12 (-0.20; 0.42); 0.224	0.06 (0.12); 0.638	0.04	0.67 (0.50; 0.81)
Vastus medialis							
Dm (mm)	0.001	-1.46 (-1.84; -1.07)	76.8	0.12 (-0.21; 0.42); 0.116	0.09 (0.20); 0.643	0.04	0.65 (0.48; 0.79)
Tc (ms)	0.069	-0.34 (-0.72; 0.02)	42	-0.14 (-0.43; 0.18); 0.200	-0.28 (0.28); 0.331	0.06	0.52 (0.36; 0.68)
Vc (mm/s)	0.001	-1.17 (-1.50; -0.79)	68.1	0.17 (-0.15; 0.46); 0.143	0.17 (0.19); 0.364	0.06	0.68 (0.52; 0.82)
V10 (mm/s)	0.001	-1.14 (-1.47; -0.76)	71	0.26 (-0.06; 0.53); 0.054	0.25 (0.19); 0.194	0.08	0.76 (0.60; 0.88)

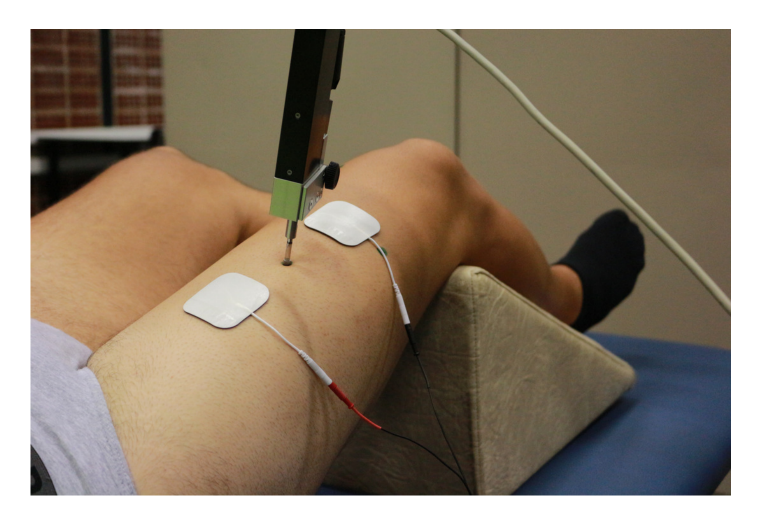
Schematic representation of experimental procedures.

TMG: tensiomyography; MVIC: maximal voluntary isometric contraction.



Tensiomyographical measurement of rectus femoris.

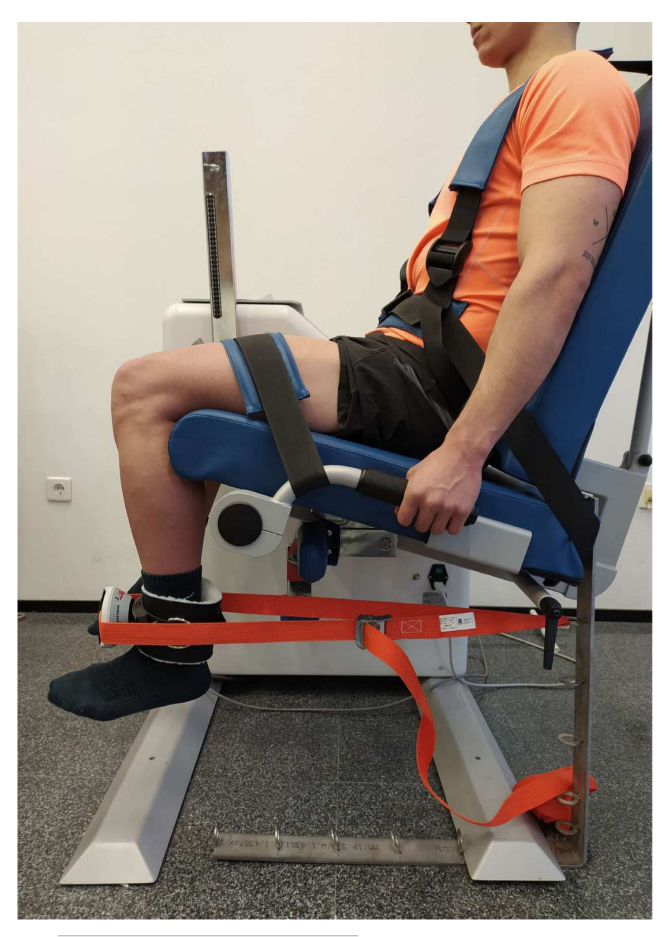
Photo credit: Rodrigo Martín-San Agustín





Maximal voluntary isometric contraction test for quadriceps femoris.

Photo credit: Rodrigo Martín-San Agustín



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Differences in TMG parameters of quadriceps bellies between pre- and post-fatigue in all participants.

(A) Differences in Dm, (B) in Tc, (C) in VC, and (D) in V10. *Significant differences set at p<0.01; Specific p-values are shown in table 3.



