

Vastus lateralis motor unit firing rate is higher in women with patellofemoral pain

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1 Title: **Vastus lateralis motor unit firing rate is higher in females with patellofemoral**
2 **pain**

3 Running head: Motor unit firing rate in patellofemoral pain

4

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14

15 **ABSTRACT:**

16 Objective: To compare neural drive, determined from motor unit firing rate, to the vastus
17 medialis and lateralis in females with and without patellofemoral pain.

18 Design: Cross-sectional study.

19 Setting: University research laboratory.

20 Participants: Females (N=56) 19-35 years old, 36 with patellofemoral pain and 20
21 controls.

22 Interventions: Not applicable.

23 Main Outcome Measure(s): Participants sustained an isometric knee extension
24 contraction at 10% of their maximal voluntary effort for 70s. Motor units (N=414) were
25 identified using high-density surface electromyography. Average firing rate was
26 calculated between 5 and 35s after recruitment for each motor unit. Initial firing rate was
27 the inverse of the first three motor unit inter-spike intervals.

28 Results: In control participants, vastus medialis motor units discharged at higher rates
29 than vastus lateralis ($p=0.001$). This was not observed in females with patellofemoral
30 pain ($p=0.78$) due to a higher discharge rate of vastus lateralis compared to control
31 participants ($p=0.002$). No between-group differences were observed for vastus
32 medialis ($p=0.93$). Similar results were obtained for the initial motor unit firing rate.

33 Conclusions: These findings suggest that females with patellofemoral pain have a
34 higher neural drive to vastus lateralis but not vastus medialis, which may be a
35 contributor of the altered patellar kinematics observed in some studies. The different
36 neural drive may be an adaptation to patellofemoral pain, possibly to compensate for
37 decreased quadriceps force production, or a precursor of patellofemoral pain.

38 **KEYWORDS:** Patellofemoral pain; Motor unit; EMG; quadriceps; neural drive

39 **ABBREVIATIONS LIST:** PFP: Patellofemoral Pain; EMG: Electromyographic; VM:

40 Vastus Medialis; VL: Vastus Lateralis; MU: Motor Unit; ES: Effect Size; CI: Confidence

41 Interval.

42 **INTRODUCTION:**

43 Patellofemoral pain (PFP) is a musculoskeletal disorder characterized by anterior
44 knee pain during activities such as stair climbing, squatting, and sitting for long time
45 periods ¹. As lower knee extension strength is associated with PFP ² and has been
46 identified as a risk factor for PFP ³, altered neuromuscular function of the knee extensor
47 muscles is considered to play a role in the development and maintenance of PFP ⁴.
48 More specifically, as the medial and lateral components of the quadriceps apply
49 different medio-lateral forces at the patella ⁵⁻⁸, their unbalanced activation may alter
50 pressure distribution across the patellofemoral joint ⁷ as well as patellar kinematics ^{8,9}.

51 A widely investigated indicator of coordination between vasti muscle activation in
52 PFP is the relative timing of activation of vastus medialis (VM) and lateralis (VL)
53 muscles during movement ¹⁰⁻¹². Although commonly used, a systematic review
54 identified only a trend for delayed activation of the VM relative to VL and this was largely
55 accounted for by large and unexplained variability across studies ¹³. In addition,
56 although timing measures are easy to obtain and provide valuable information, they only
57 permit the identification of temporal differences in muscle activation. The force exerted
58 by a muscle is known to depend on the number and discharge rate of its active motor
59 units ^{14,15}. For this reason, altered neural drive to VM and VL may be relevant for PFP.
60 Previous studies have investigated surface electromyographic (EMG) amplitude to
61 compare vasti muscle activation between participants with and without PFP ¹⁶⁻¹⁸.
62 However, this provides only a crude indicator of the neural drive to a muscle, as surface
63 EMG amplitude is influenced by factors such as: adipose tissue thickness,
64 normalization, crosstalk, motor unit action potential cancellation, and others ¹⁹, which

65 may differ between groups. In isometric tasks, the influence of these factors can be
66 limited by estimating the neural drive from motor unit (MU) activity. Although motor unit
67 activity has traditionally been assessed using intramuscular recordings, recent
68 technological advances enable estimation of neural drive using non-invasive high-
69 density surface electromyography ²⁰.

70 The aim of this study was to compare neural drive to the vastus medialis and
71 lateralis in females with and without PFP during a submaximal, isometric task. On the
72 basis of theories that propose a role for unbalanced activation of the vasti muscles, we
73 hypothesized that MU firing rate of VM would be lower in participants with PFP relative
74 to asymptomatic controls, or VL would be higher, or both.

75

76 **METHODS:**

77 Thirty-six females with PFP and 20 asymptomatic females (control participants)
78 were recruited for the study from the community and from local physiotherapy clinics. To
79 be included in the PFP group, participants had to be: female, 19-35 years old, with
80 retro- or peri-patellar knee pain of intensity equal or greater than 3/10 for at least 1
81 month aggravated by any of the following activities: sitting for long time periods, stair
82 ambulation, squatting, running, kneeling or jumping. They also needed to report pain or
83 discomfort to at least one of the following tests: patellar palpation, patellar compression,
84 resisted knee extension with knee close to full extension, or isometric knee extension
85 while applying pressure proximally to the patella. These criteria were similar to those
86 used in other studies ^{10,17,21}; the screening was performed by a physiotherapist with
87 more than 2 years of clinical experience in musculoskeletal assessment. Asymptomatic

88 controls must not have had any knee pain in the last 12 months. Participants were
89 excluded from either group if they had chronic neuromuscular disorders affecting the
90 legs or previously had lower-limb surgery. All participants provided written informed
91 consent before the start of the experimental session. The study was approved by the
92 institution's Clinical Research Ethics Board.

93 Body mass and height were measured, and age, time of onset of pain and
94 average pain intensity in the previous week (11-point Numerical Rating Scale) were
95 obtained by self-report for each participant. Physical activity (General Physical Activity
96 Questionnaire, GPAQ ²²) and functional limitation (Anterior Knee Pain scale ²³) were
97 evaluated using validated questionnaires. The test leg was the most painful knee. For
98 control participants, the leg was determined randomly before the testing session.

99 The protocol consisted of recording high-density surface EMG signals from both
100 vasti during a submaximal, isometric task. The electrode grids were placed according to
101 anatomical references as described previously ²⁴. The medial and lateral edges of VM
102 and VL were identified using ultrasound imaging^a and were marked on the skin. VM and
103 VL innervation zones were located using a linear electrode array (16 silver bar
104 electrodes, 10-mm interelectrode distance^b) and marked on the skin. Two electrode
105 grids (semidisposable adhesive matrix^b) were placed on the skin so that the innervation
106 zone was aligned between the second and third column, and all the electrodes were
107 placed on the muscle of interest. Each grid comprised 64 electrodes arranged in 5
108 columns and 13 rows with an electrode missing in one of the corners, 8 mm inter-
109 electrode distance and was held in place using bi-adhesive foam. In the VM, for
110 instance, the longer dimension of the electrode grid (approximately 10 cm) spanned the

111 distal-medial to the proximal-lateral region of the muscle (fig. 1); for this reason, the grid
112 placement provided EMG signals representative of different regions within each vastus
113 muscle. Reference electrodes (2x3.5 cm; conductive hydrogel^e) were placed on the
114 patella and both sides of the knee.

115 Isometric knee extension torque was measured using an isokinetic
116 dynamometer^d. Participants were secured to the chair; the hip and knee angles were 85
117 and 45 degrees of flexion, respectively. Resistance was applied approximately 2 cm
118 proximal to the medial malleolus. Participants performed 3 maximal voluntary
119 contractions (MVC) of knee extension with verbal encouragement, with a rest period of
120 at least 60s between trials. Contraction intensity was increased to maximum over
121 approximately 1-2 s and was maintained for at least 3 s before relaxation. The peak of
122 the torque profile was extracted from each trial. The highest torque of the three values
123 was considered the maximal knee extension strength, and normalized to body mass.
124 The submaximal task consisted of a single 70 s knee extension at 10% MVC.
125 Participants were provided with real-time feedback of their knee extension torque and
126 target.

127 High-density surface EMG signals were collected as monopolar recordings (128-
128 channel EMG-USB^b). Signals were amplified 500-1000 times, filtered (band-pass 10-
129 750 Hz) and digitized at 2048 Hz using a 12-bit A/D converter. Knee extension torque
130 was acquired simultaneously using the same amplifier. Butterworth filters (4th order; 10-
131 400 Hz for the EMG signals; low-pass 10 Hz for the torque) were applied to the signals
132 before processing.

133 Motor unit discharges were identified separately for VM and VL using a
134 previously described method²⁵ reliable between sessions²⁷ and valid when compared to
135 a gold-standard, intramuscular electromyography²⁶. An example of motor unit
136 identification from surface EMG signals can be observed in figure 1. Motor unit firing
137 patterns were reviewed visually and firing rates >30 Hz or <3 Hz were manually
138 excluded²⁷. Similar to a previous study²⁸, the MU template was created by averaging
139 epochs of 40ms around each MU discharge. The peak-to-peak amplitude was
140 calculated for each of the 13x5 channels to identify the location of each MU, i.e.: where
141 it was represented with highest amplitude. Motor unit recruitment was identified as the
142 first of four consecutive discharges <500 ms apart. The initial MU firing rate was
143 calculated as the inverse of those first three MU inter-spike intervals. The neural drive
144 was quantified two ways: as the initial firing rate at recruitment, and the average firing
145 rate, calculated as the average firing rate between 5s and 35s after motor unit
146 recruitment. Additional parameters used to describe the population of MUs identified
147 were: MU recruitment threshold, calculated as the torque value coincident with the first
148 motor unit discharge (see above); MU location, calculated as the proximal-distal
149 coordinate of the channel with largest peak value (along the longest dimension of the
150 electrode grid, fig.2).

151 Statistical analyses were performed using SPSS v. 22^e. After logarithmic
152 transformation of average firing rate and initial firing rate, the assumptions of normally
153 distributed data (Shapiro-Wilk's test) and equal variances across groups (Levene's test)
154 were met. Demographic variables and knee extension strength were compared between
155 groups using unpaired T-tests. Differences in MU firing rates between *Groups* (PFP,

156 control) and *Muscles* (VM, VL) were tested using a two-way ANCOVA, separately for
157 average firing rate and initial firing rate. To account for the effect of the MU recruitment
158 threshold on average firing rate and initial firing rate, recruitment threshold torque was
159 included in the model as a covariate. Effect sizes (ES) were calculated using Cohen's *d*,
160 separately for each comparison. Two-way ANOVA was used to determine whether MU
161 recruitment threshold torque or MU location differed between *Groups* or *Muscles*. Post-
162 hoc tests were corrected for multiple comparisons using Bonferroni corrections.
163 Statistical significance was set at $p < 0.05$.

164

165 **RESULTS:**

166 When compared to controls, participants with PFP were of similar age, height,
167 weight and physical activity but had higher BMI and lower knee extension strength
168 (Table 1). Twenty-six participants with PFP reported bilateral symptoms. After visual
169 inspection, a total of 414 MUs were identified and included in the analyses. The number
170 of identifiable MUs included for each participant ranged from 2-10 (mean
171 MU/participant=4.8; total N=96) for the VM and 1-12 (mean 4.3; N=86) for the VL of
172 controls, and 0-8 (mean 3.0; N=104) for the VM and 0-8 (mean 3.6; N=128) for the VL
173 of participants with PFP. No MUs were identified from the VM of one participant with
174 PFP and from the VL of 3 participants with PFP.

175 An interaction effect between *Group* and *Muscle* was observed in the MU
176 average firing rate analysis ($p < 0.05$; fig.3). A higher average firing rate was observed in
177 the PFP group compared to controls for VL (8.8 ± 1.7 Hz vs. 8.2 ± 1.6 Hz, $p = 0.002$, ES:
178 0.34, 95% CI: [0.03 0.13]) but not for VM (8.9 ± 2.0 Hz vs. 8.8 ± 1.6 Hz, $p = 0.93$, ES: 0.07,

179 95% CI: [-0.05 0.06]). VM had a higher average firing rate than VL in controls (8.8±1.6
180 Hz vs. 8.2±1.67 Hz, $p=0.001$, ES: 0.33, 95% CI: [0.04 0.15]), but no difference between
181 the two vasti was observed in females with PFP (8.9±2.0 Hz vs. 8.8±1.7 Hz, $p=0.78$,
182 ES: 0.04, 95% CI: [-0.04 0.05]). Similarly, an interaction effect between *Group* and
183 *Muscle* was observed in the MU initial firing rate analysis ($p=0.001$; fig.3). A higher initial
184 firing rate was observed in the PFP group for VL (7.4±2.1 Hz vs. 6.4±1.7 Hz, $p<0.001$,
185 ES: 0.49, 95% CI: [0.07 0.21]) but not for VM (7.1±2.0 Hz vs. 7.1±1.7 Hz, $p=0.55$, ES:
186 0.03, 95% CI: [-0.09 0.05]). VM had a higher initial firing rate than VL in controls
187 (7.1±1.7 Hz vs. 6.4±1.7 Hz, $p=0.002$, ES: 0.40, 95% CI: [0.05 0.20]), but no difference
188 between the two vasti was observed in PFP (7.1±1.9 Hz vs. 7.4±2.1 Hz, $p=0.17$, ES:
189 0.17, 95% CI: [-0.11 0.02]). Neither *Muscle* nor *Group* influenced the recruitment
190 threshold torque ($p>0.2$) or MU position ($p>0.15$, fig.3).

191

192 **DISCUSSION:**

193 This study found differences in MU firing rate across individual heads of the
194 quadriceps between females with and without PFP. In females without PFP, VM motor
195 units discharged at higher rates than VL. This difference was not observed in those with
196 PFP and was explained by a higher VL firing rate. We suggest that the greater neural
197 drive to the VL may contribute to altered patellofemoral kinematics, which is proposed to
198 be relevant for PFP.

199 The evidence of higher neural drive to the VL in females with PFP implies a role
200 of vasti muscle activation in the adaptation to, or in the development of, PFP. Our
201 findings are strengthened by the fact that differences in neural drive cannot be attributed

202 to the location of the motor unit within the muscle or its recruitment threshold, as neither
203 differed between groups or muscles. Previous studies identified altered timing and
204 amplitude of surface EMG in PFP ^{10,12,16,18} and with experimental knee pain, ²⁹⁻³¹. Our
205 findings appear to concur with studies that reported earlier activation for VL rather than
206 delayed activation of VM in reflex contractions ³² and when participants with PFP were
207 asked to rise onto their toes ³³. Overall, this study further expanded this research,
208 showing that the distribution of neural drive between VM and VL, measured as motor
209 unit firing rate, differs between females with and without PFP.

210 Changes in muscle activation with pain and in musculoskeletal disorders are
211 thought to be a purposeful adaptation to avoid pain in the short-term by altering joint
212 kinematics ³⁴. Previous studies on cadavers identified altered patellar kinematics ⁷ and
213 pressure distribution within the patellofemoral joint ⁸ when the relative load of VM and
214 VL was manipulated. In vivo studies showed that anesthetic block of the VM results in
215 altered patellar kinematics ⁹, and studies using EMG timing and amplitude identified
216 associations between VM/VL activation and patellar tilt ^{12,16}. Considering the results of
217 these studies, a greater neural drive to the VL may result in larger force produced by the
218 lateral component of the quadriceps. However, caution must be used when inferring
219 forces from muscle activation because muscle force depends on both neural activation
220 and peripheral factors at the muscle level ³⁵. As individuals with PFP appear to have
221 smaller cross-sectional areas of the quadriceps muscles as a whole (systematic review
222 by Giles and colleagues ³⁶), the neural drive is likely an important contributor to the
223 relative amount of force produced by VM and VL. However, other factors such as

224 between-group differences in fiber type composition and structural parameters of the
225 quadriceps should also be considered could also play a role.

226 Higher discharge rates of VL and similar discharge rate for VM suggest that
227 neural drive to the quadriceps as a whole is higher in PFP. This is in contrast with the
228 clinical belief that the quadriceps muscle is inhibited in PFP, and it could inform the
229 mechanisms that should be targeted in future intervention studies. Functionally, the
230 greater neural drive to VL (or potentially vastus intermedius or rectus femoris – not
231 measured in this study) could be an attempt to compensate for a decreased overall
232 force production capability of the knee extensors, observed as smaller quadriceps
233 cross-sectional area ³⁶. Due to its architecture, VL mainly produces a force vector
234 towards knee extension ⁵ and may be more efficient than VM to generate forces due to
235 its greater physiological cross-section area ³⁷. However, as the VL also applies a
236 laterally-directed force vector on the patella ⁵⁻⁷, a selective increase of neural drive
237 could potentially result in increased lateral forces applied to the patella. In line with this,
238 some studies reported increased lateral patellar spin and translation ^{38,39} and higher
239 joint reaction forces in the lateral patellofemoral compartment in PFP ⁴⁰. The potential
240 association between altered neural drive to the quadriceps and altered force production
241 capabilities at the muscle could be observed in the cross-over effects of interventions
242 targeting the two dysfunctions ⁴². Future studies should investigate the association
243 between force production capability of the quadriceps and neural drive to the VL in PFP.

244 The notion that unbalanced vasti activation may be due to greater neural drive to
245 VL rather than VM inhibition may potentially have clinical implications. Traditionally,
246 putative imbalanced activation of VM and VL in PFP is treated by enhancing the

247 activation of the medial component using techniques such as taping⁴³ and therapeutic
248 exercise intended to preferentially target the distal region of the VM⁴². Less frequently,
249 interventions such as taping^{44,45} and botulinum injections⁴⁶ are aimed at reducing the
250 activation or force produced by the VL. This study suggests that reducing neural drive to
251 the VL as opposed to increasing neural drive to the VM may result in muscle activation
252 patterns similar to individuals without PFP. Techniques that reduced/inhibit VL activation
253 have a potential role in rehabilitation. This might be achieved in clinical practice using a
254 variety of techniques, for instance, with taping techniques^{44,45}. Given the assumed link
255 between muscle activity and resultant joint kinematics, our findings support the notion
256 that reducing drive to VL may have positive clinical outcomes. Future longitudinal
257 studies should also evaluate if reducing the neural drive to the VL improves
258 patellofemoral kinematics and kinetics in PFP, resulting in less degeneration of the
259 lateral patellar facet⁴¹. The findings of this study may also be relevant for prevention. If
260 differences in neural drive were present before the development of PFP, screening and
261 early treatment may reduce the incidence of PFP; this however should be carefully
262 evaluated in prospective studies. Overall, this study suggests that neural drive may be
263 an important variable of interest in PFP, and further research into its clinical and
264 biomechanical implications is warranted.

265

266 **LIMITATIONS:**

267 Due to the cross-sectional design of our study, whether the greater drive to the
268 VL is an adaptation to, or precursor of, PFP cannot be determined. More research is
269 needed to understand what drives the greater neural drive to the VL and if this motor

270 control alteration can be observed in tasks other than isometric contractions. The
271 association between altered neural drive and differences in force production capabilities
272 are not examined in the current study and should be assessed to make informed
273 inferences on force production. It should be noted that changes in MU firing rate provide
274 an accurate, but only partial, representation of changes in the neural drive. Changes in
275 motor unit recruitment strategies, such as the number and population of active motor
276 units, have been described with experimental knee pain ⁴⁸ and are not accounted for by
277 changes in firing rates. In addition, only females with PFP were tested in this study to
278 limit the confounding effect of sex-differences in anatomy, muscle strength and
279 neuromuscular strategies. For this reason, these findings are only generalizable to
280 females with PFP; future studies should investigate whether similar findings are
281 observed in males with PFP.

282

283 **CONCLUSIONS:**

284 Motor unit firing rate of the vastus lateralis, but not medialis, during low-force,
285 isometric contractions differs between females with and without PFP. Neuromuscular
286 control of individual quadriceps heads could be considered a possible target for future
287 interventions aimed to prevention and management of PFP.

288

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290 REMOVED FOR BLINDED REVIEW

291

292

293 **SUPPLIERS:**

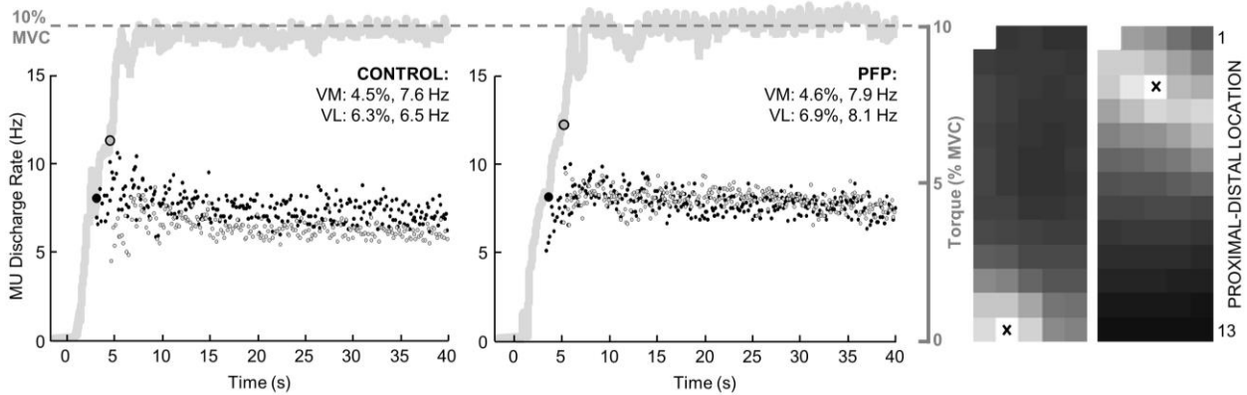
- 294 a. LogicScan 64 LT-1T; Telemed, Vilnius, Lithuania
- 295 b. OTBIOelettronica, Torino, Italy
- 296 c. Kendall, Covidien, Mansfield, MA, USA
- 297 d. System 4 Pro; Biodex Medical Systems, Shirley, NY, USA
- 298 e. IBM Inc., Armonk, NY, USA
- 299

300 **FIGURES:**



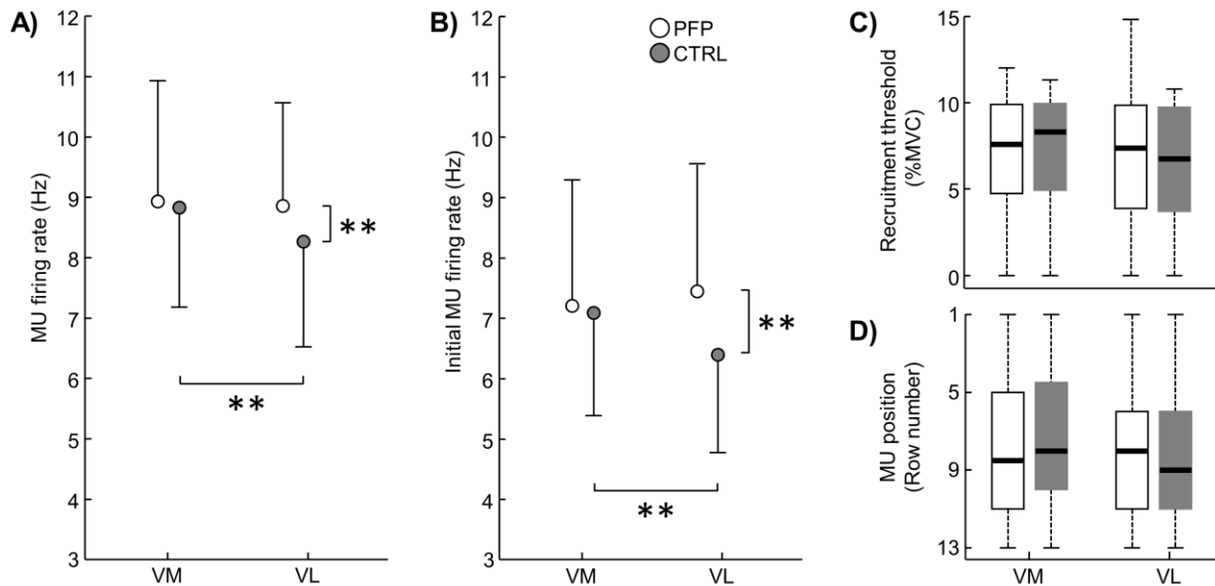
301
 302 Fig.1: Experimental set-up and example of motor unit identification. Left: Placement of
 303 the electrode grids; the dashed line depicts the location of the innervation zones across
 304 both muscles. Middle: Double differential EMG signals from 10 channels of the VM of a
 305 control participant; three of the motor units automatically identified are highlighted with
 306 grey boxes (A, B, C). Right: the triggered-average surface EMG representation of each
 307 motor unit.

308



309

310 Fig.2: Motor unit discharges and location. Left: firing rate of one motor unit from VM
 311 (black) and VL (gray) in one control participant and one with PFP. The torque signal is
 312 shown as a thick, light grey line; the recruitment threshold of the VM and VL MU is
 313 indicated as a black and a gray circle respectively. Right: examples of motor units
 314 located distally and proximally within the VL. Crosses on each surface EMG amplitude
 315 distribution identify the peak of the distribution; the Y coordinate of that channel was
 316 considered to be the proximal-distal location within the muscle.
 317



318
 319 Fig.3: A) MU firing rate while holding 10% MVC. B) Initial MU firing rate. C) MU
 320 Recruitment threshold torque. D) MU position. Statistical significance for post-hoc
 321 comparisons is indicated. ** p<0.01.

322

323 **TABLES:**

324

325 Table 1: Participant characteristics and knee extension strength. Pain intensity was
 326 subjectively rated indicating a number between 0 (no pain) and 10 (worst imaginable
 327 pain). Anterior knee pain scores of 100 indicate maximal function and no pain. KES:
 328 knee extension strength; nKES: normalized knee extension strength.

329

	CTRL	PFP	T-test
AGE, years	25.6 (4.3)	26.7 (4.1)	$p=0.38$
HEIGHT, cm	167.7 (8.5)	166.4 (7.9)	$p=0.59$
BODY MASS, kg	58.2 (8.5)	62.3 (8.9)	$p=0.10$
BMI, kg/m²	20.6 (1.7)	22.5 (2.9)	$p=0.01^*$
PHYSICAL ACTIVITY ²², METmin/week	3153 (2034)	4018 (2961)	$p=0.20$
PAIN ONSET, months (interquartile range)		12-60	
PAIN INTENSITY, out of 10	0 (0)	4.1 (1.5)	
ANTERIOR KNEE PAIN SCORE ²³, out of 100	100 (0)	74.3 (8.1)	

nKES, Nm/kg	2.3±0.4	1.9±0.5	<i>p</i><0.01*
KES, Nm	135.3±32.9	116.5±30.6	<i>p</i><0.05*

330

331

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