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#### Differential behavior of distinct motoneuron pools that innervate the triceps surae

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e

#### 21 Abstract

22

23 It has been shown that when humans lean in various directions, the central nervous system 24 (CNS) recruits different motoneuron pools for task completion; common units that are active 25 during different leaning directions, and unique units that are active in only one leaning direction. 26 We used high-density surface electromyography (HD-sEMG) to examine if motor unit (MU) 27 firing behaviour was dependent on leaning direction, muscle (medial and lateral gastrocnemius; 28 soleus), limits of stability, or whether a MU is considered common or unique. Fourteen healthy 29 participants stood on a force platform and maintained their center of pressure in five different 30 leaning directions. HD-sEMG recordings were decomposed into MU action potentials and the 31 average firing rate (AFR), coefficient of variation (CoV<sub>ISI</sub>) and firing intermittency were 32 calculated on the MU spike trains. During the 30-90° leaning directions both unique units and 33 common units had higher firing rates (F = 31.31, p < 0.0001). However, the unique units 34 achieved higher firing rates compared to the common units (mean estimate difference = 3.48 Hz, 35 p < 0.0001). The CoV<sub>ISI</sub> increased across directions for the unique units but not for the common 36 units (F = 23.65. p < 0.0001). Finally, intermittent activation of MUs was dependent on the 37 leaning direction (F = 11.15, p < 0.0001), with less intermittent activity occurring during 38 diagonal and forward-leaning directions. These results provide evidence that the CNS can 39 preferentially control separate motoneuron pools within the ankle plantarflexors during voluntary 40 leaning tasks for the maintenance of standing balance. 41

#### 42 New & Noteworthy

43 In this study, we demonstrate that the different sub-populations of motor units within the three

44 muscles comprising the ankle plantarflexors behave differently during multi-directional leaning.

2

45	Our results suggest that the central nervous system has the capability to control distinct sub-
46	populations of motor units to meet the force requirements necessary for leaning. This may allow
47	for a precise, efficient, and flexible control strategy for the maintenance of standing balance.
48	
49	Keywords
50	High-density surface electromyography, Motor units, Firing rates, Plantarflexors, Postural
51	control
52	
53	Introduction
54	Control of the ankle plantarflexor muscles is fundamental for motor tasks such as quiet
55	standing and leaning; these tasks are accomplished largely by the medial and lateral
56	gastrocnemius (MG and LG) as well as the soleus (SOL) muscles (1-3). The motor unit (MU)
57	behaviour of these muscles during various non-weight bearing contractions have been well
58	documented (4-10). However, motor units may behave differently during postural control than
59	during non-weight bearing contractions (11).
60	Evidence is mounting for differential MU behaviour during standing compared to that
61	observed from non-weight bearing contractions. During quiet standing, the common modulation
62	of MU discharge rates in the SOL is higher compared to voluntary isometric contractions (12)
63	reflecting a stronger common drive during standing than seated contractions. Further, there is
64	greater inhibition of MUs of the MG muscle in response to sural nerve stimulation when
65	individuals are standing compared to lying supine (13). The relative contribution of recruitment
66	or rate coding to ankle plantarflexor force production depends on the posture. That is, when
67	responding to perturbations of increasing intensity during standing, MU recruitment in the

68 gastrocnemius appears to be the main mechanism of force gradation (14), whereas in a sitting 69 position both rate coding and recruitment are used to modulate force gradation in the 70 gastrocnemius (15). Taken together, it appears that the MUs of the plantarflexors behave 71 differently during postural control than during non-weight bearing contractions. 72 For a MU to be recruited, synaptic current must reach the soma of a motoneuron for it to 73 discharge and initiate an action potential (16). When the distribution of synaptic current occurs 74 uniformly across the motoneuron pool, the first MUs to discharge are those with the small size 75 and high input resistance (17). Unequal distribution of synaptic current however may influence 76 MU recruitment. In previous work, we have shown that the ankle plantarflexor muscles have 77 unequal, regionally specific activation in response to directionally induced perturbations (18). 78 Further, we observed the spatial recruitment of MUs within the ankle plantarflexor muscles 79 during a multidirectional leaning task (19). Our findings indicate that the CNS possesses the 80 ability to recruit two distinct subpopulations of MUs. During the different directional 81 movements, *common* MUs are recruited from the MG and SOL, putatively to meet the baseline 82 torque requirements. As the directionality of the force requirements for body stabilization 83 changes, unique MUs in separate locations were recruited in the plantarflexor muscles. We 84 defined common units as MUs that were matched between different leaning directions identified 85 by waveforms that were indistinguishable from each other (19, 20). Unique units were MUs that 86 were not matched between leaning directions and were distinguishable from each other (19, 20). The CNS may rely on the uneven spatial distribution of synaptic inputs to control recruitment; 87 88 however, whether the MU firing behaviour is influenced by this nonuniform recruitment pattern 89 is unknown.

90	An interesting consideration of MU firing behaviour in the plantarflexors is the
91	intermittent firing patterns observed during standing. Evidence from ultrasound and surface
92	EMG has shown intermittent activity of the MG muscle while subjects stand at ease (3, 21, 22).
93	Observations from intramuscular EMG have determined that postural activation of the MG MUs
94	occurs intermittently, specifically at different body sway positions (23). Taken together, there
95	appears to be intermittent activation of MUs in the MG that is associated with the position and
96	velocity of the standing body. What is still unknown is whether this intermittent firing behaviour
97	continues as task difficulty increases from quiet stance to more demanding postural tasks.
98	The purpose of this study was to investigate the MU firing behaviours of the MG, LG,
99	and SOL during a multi-directional leaning task. It was hypothesized that: a) MU firing rate
100	would be higher in leaning directions that required higher force production; b) based on our
101	previous finding of common and unique units across leaning directions (19), MU firing
102	behaviour would be dependent on whether the MU was common or unique; and c) intermittency
103	would be more evident in the gastrocnemius, compared to the SOL, and would decrease in
104	response to forward-leaning directions.

105

#### 106 Methods

#### 107 Participants and Experimental Protocol

Fourteen healthy adults participated in this study (7 females, 7 males; mean  $\pm$  SD: body mass 70  $\pm$  15.3 kg; height 168  $\pm$  9.3cm; age 25  $\pm$  2.5 years). Participants were excluded if they had any health conditions that negatively impacted balance (e.g., musculoskeletal disorders, neuropathies, dizziness), or previous injury to their lower legs within the past 6 months. The experimental protocol was approved by the institutional Ethics Committee (Research Ethics 113 Board Number: 110471) and conformed to the latest amendment of the Declaration of Helsinki.

114 Participants provided written, informed consent before participating in the study.

115 Experimental data used in this study were part of a larger study examining the spatial 116 location of MU recruitment (19). Briefly, standing in their natural bipedal stance on a 117 piezoelectric force platform (9286AA Kistler, Zurich Switzerland), participants were instructed 118 to lean in different directions. To identify the limit of stability (LoS), participants were instructed 119 to lean about their ankles anteriorly, posteriorly, and laterally to both sides as far as they could 120 without taking a step. This 4-way LoS test has been shown to be more reliable in testing LoS 121 compared to other tests (24). After completing the 4-way LoS test, participants performed a quiet 122 stance trial for 30s. Then, while provided with visual feedback of their center of pressure (CoP), 123 participants were asked to lean in 5 different directions. Circular targets for each direction, 0°, 124 30°, 60°, 90° and 120°, counter-clockwise from the mediolateral axis crossing the average CoP 125 position were computed during the 30s of quiet stance. Each target was displayed one at a time 126 on a screen in front of the participant (Figure 1A, B). The targets appeared over an ellipse with 127 semiaxes corresponding to 30% LoS (or 40% LoS) in the anterior-posterior and medio-lateral 128 directions (Figure 1B). The resulting ellipses representing 60% LoS and 80% LoS, respectively, 129 imposed progressively greater contraction intensities, given the CoP distance from the ankle 130 scales with the ankle torque (25). Participants leaned toward each of the targets, maintained their 131 CoP position in the target for 35s, and then returned to their quiet stance position. After leaning 132 in a specific direction, a rest period of 15s was provided. An example of the CoP trace can be 133 found in Figure 1C. During the leaning task, participants were instructed to keep their legs and 134 body straight and lean like a pole. Participants moved in a smooth manner until their CoP 135 position was in the target for a total duration of 45s (5s towards the target, 35s in the target, and

136 5s back to their natural bipedal stance). To familiarize themselves with the procedure,

137 participants first conducted an unrecorded practice trial. If the participant did not maintain the

138 instructed body position or was unable to keep their CoP steady in the target for more than 25s,

139 the trial was discarded and repeated. The trial was retained and analyzed once the correct leaning

140 posture was maintained.

141

142 High-Density Surface EMG Electrode Placement

143 Placement of the High-Density surface Electromyography (HD-sEMG) was guided using 144 an ultrasound imaging system (LogicScan 64 LT-1T; Telemed, Vilnus, Lithuania). With 145 participants lying prone and the ankle in the neutral position, the medial and lateral edges of the 146 gastrocnemius and SOL, the insertion of the MG and LG on the Achilles tendon, the fascial 147 space between the MG and LG, and the distal edges between the two muscles were identified 148 and marked on the skin. The skin was abraded and cleaned before grid placement. Semi-149 disposable grids with 32 electrodes at an interelectrode distance of 10mm (LISiN, Torino, Italy) 150 were affixed to each of the MG and LG (one grid on each muscle). The grids were placed on the 151 proximal belly of the muscle, 10mm above the distal edges of the superficial aponeurosis of the 152 MG and LG to avoid EMG detection from the distal region of gastrocnemius where propagation 153 of action potentials is observed along the muscle fibres (18, 19, 26). A 64-electrode matrix, 154 comprised of two adjacent 32-electrode semi-disposable grids with an interelectrode distance of 155 10mm (LISiN, Torino, Italy), was placed over the SOL muscle. The 64-electrode grid was 156 centered on the Achilles tendon and placed 10mm below the distal insertion of the MG (Figure 157 1D). The grids were held in place with bio-adhesive foam, and conductive paste ensured optimal

- 158 contact between the skin and the electrodes. Three reference electrodes were placed on the
- 159 medial malleolus (SOL), the patella (MG), and the fibular head (LG).
- 160
- 161 HD-sEMG and Forceplate Recordings

162 Signals were detected in a monopolar configuration using a modular, wireless HD-sEMG 163 amplifier (27). This device is composed of four modules (one for each gastrocnemius head, and 164 two for SOL), each detecting, conditioning, and transmitting 32 EMG signals (amplification: 165 192V/V; sampling frequency: 2048 Hz; 16-bit A/D Converter). A Wi-Fi link from the EMG 166 acquisition modules to a personal computer enabled the detected signals to be transmitted, 167 visualized, and stored in real-time. 168 CoP coordinates in the sagittal and frontal planes were computed from the ground 169 reaction forces, sampled at 2048 Hz using a 16-bit A/D Converter (±2.5 V input voltage range). 170 A pulse generator (LISiN, Torino, Italy) sent a signal at 3Hz (0V – 3.3 V amplitude) to both 171 EMG amplifier and force platform to synchronize the systems. Signals were aligned on the first, 172 rising edge of the trigger signal. 173 174 **Data Analysis** 175 Motor unit analysis

176 All EMG data analyses were performed using MATLAB R2020b (The MathWorks, Inc.,

177 Natick, MA, USA). Single-differential EMGs between pairs of adjacent electrodes along each

- 178 column of the matrix were computed from the monopolar signals and visually inspected for
- 179 quality. Channels presenting contact problems or power line interference were linearly
- 180 interpolated with the data collected by the surrounding 8 channels. After verifying the quality of

181	all EMG signals, monopolar signals were filtered (Butterworth, 2 <sup>nd</sup> order, 20-350 Hz) and
182	decomposed with the blind source separation method (28) implemented in the DEMUSE tool
183	software (v. 4.9; The University of Maribor, Slovenia). Signals for each leaning direction and
184	LoS condition were decomposed separately, over the central 20s of the 35s period when CoP was
185	maintained at the target location. This decomposition procedure can identify MU discharges over
186	a wide range of contraction intensities and has been extensively validated using experimental and
187	simulated signals (29, 30). After decomposition of the EMG signals, the firing instants of
188	identified MUs were used to trigger and average single differential EMGs over 30ms epochs,
189	providing the surface representation of single MU action potentials (31). The results of the
190	decomposition were inspected visually, according to Power et al. (32), and noisy waveforms
191	were discarded. Examples of retained MUs are shown in Figure 2.
192	For each retained MU, the average firing rate (AFR) was calculated. Interspike intervals
193	(ISIs) < 25ms were discarded for subsequent analysis because these ISIs are likely due to
194	decomposition errors (30) and would have a large impact on the AFR. The coefficient of
195	variation of the ISI ( $CoV_{ISI}$ ) was calculated as the standard deviation of the ISI divided by the
196	mean ISI and is reported as a percentage.
197	To determine whether the MUs were activated intermittently, an intermittency index was
198	established. It has been shown previously that very frequently, the postural MUs in the MG
199	muscle discharge below the physiological minimal tonic rate (<4 Hz) during quiet stance (23).
200	These occurrences of discharge rates lower than 4 Hz are considered as representing instances of
201	intermittent recruitment of MU (i.e., units were successively de-recruited and re-recruited). To
202	standardize the amount of intermittent activity for a single MU, we calculated the degree of
203	intermittency:

204 
$$Degree \ of \ Intermittency = \frac{Number \ of \ intermittent \ occurrences}{Time \ MU \ was \ active \ (s)}$$

205 where the "*Number of intermittent occurrences*" is the number of cases in which the MU 206 firing rate is below 4 Hz, and the "*Time MU was active*" is the total time that the MU was 207 actively recruited, from the instance of the first firing to the last firing of the corresponding MU. 208 This procedure standardized the intermittency of MU activity and allowed comparisons between 209 different MUs by evaluating the number of intermittent occurrences per second.

210

#### 211 Motor unit tracking

212 A MU tracking analysis was conducted separately for the 60% and 80% LoS conditions. 213 MU action potentials that were common between the different leaning directions were identified 214 by focusing on both the shape and amplitude of the waveform of MU action potentials. For each 215 retained MU, the average rectified value (ARV) was calculated across 30ms epochs centered on 216 individual action potentials. Each retained MU was paired to all other MUs identified during 217 different leaning directions within the same muscle and LoS condition for each participant (e.g., 218 a MU identified in the 0° leaning direction in the 60% LoS condition within the MG is compared 219 to all other units identified in the MG at the  $30^\circ$ ,  $60^\circ$ ,  $90^\circ$ , and  $120^\circ$  leaning directions of the 60%220 LoS condition within a single participant). The action potentials for each pair of MUs were 221 aligned in time by maximizing their cross-correlation function. The mean square difference was 222 calculated between the two sets of time-aligned action potentials, averaged across channels, and 223 then normalized with respect to the mean ARV of the two sets of action potentials. This 224 normalization procedure is utilized to reduce the bias of waveform amplitude between two MU 225 pairs (33). Finally, pairs of MU action potentials with a mean square difference smaller than 10% 226 were considered common (19, 20). An expert operator visually confirmed all MUs that were

identified as common by the algorithm. This provided us with a group of MUs that will be
referred to as *common* units that were present in two leaning conditions or more. When a MU
had a mean square difference greater than 10% with all MUs it was paired with, the MU was
considered to be a uniquely recruited unit not found in other leaning directions. These MUs will
be called *unique* units from this point. An example of a pair of retained *common* MUs is
provided in Figure 3.

233

#### 234 Statistical Analysis

235 All statistical analyses were conducted in R (v.5.12.1, R Development Core Team, 2009). 236 To analyze MU behaviour, we performed a multilevel mixed model linear regression analysis 237 using the package lme4 (v. 1.1-27.1) (34). As previously discussed in detail (35, 36), linear 238 mixed models have several advantages that are particularly suitable for MU experimental 239 designs: 1) they allow the whole sample of extracted MUs to be analyzed and not just the mean 240 observations for each subject and direction. This allows better evaluation of the data variance 241 compared to conventional ANOVA statistics; 2) they account for the non-independence of 242 observations (i.e., different MUs coming from the same participant) with correlated error; 3) they 243 separately treat the effects caused by the experimental manipulation (fixed effects) and those that 244 were not (random effects), increasing the generalizability of the findings. Linear mixed 245 modelling computes and utilizes estimated marginal means of the data. Further, in a nested or 246 hierarchical data structure, averaging is performed separately in each nesting group (in our case, 247 by participant). This procedure creates estimated marginal means that are based on the statistical 248 model, not directly from the raw data, and therefore the estimated marginal means are different 249 than the means of the raw data.

#### 250 Motor Unit Behaviour

251 Diagnostic plots of the residuals from all models were inspected for violations of the 252 assumptions of normality and homoscedasticity. The AFR and CoV<sub>ISI</sub> data met the assumptions, 253 however, the degree of intermittency did not meet homoscedasticity and normality of residuals 254 assumptions. The intermittency values were transformed by a log (x + 1) transformation for the 255 data to fit the assumptions of the model. The aggregated data set were analyzed by three separate linear mixed-effects models, using the fixed effects of leaning direction (0°, 30°, 60°, 90°, 120°), 256 257 LoS (60% and 80%), muscle (MG, LG, and SOL), and MU subgroup (unique or common units). 258 Starting with the maximal random structure, including by-participant and by-item random slopes 259 and intercepts, this structure was then reduced to the optimal structure that could be supported by 260 the data following the steps of Bates (2015). Likelihood ratio tests were used to find the optimal 261 model structure; the random slope model improved the model. We considered the random 262 intercept over participants as a random factor and the random slope of the MU subgroup and 263 muscle. We could not include a random slope of leaning direction or LoS condition because of 264 singular fit violation (i.e., multiple collinearities). 265 Statistical significance of fixed effects was determined using type III Wald F tests with 266 Kenward-Rogers degrees of freedom with the ANOVA function from R's car package (v.

267 3.0.12). Post hoc pairwise comparisons (with Bonferroni's correction) were performed using the

estimated-means contrast, as employed in R's emmeans package (v. 1.7.2). Post hoc comparisons

269 were applied between each leaning direction (all-pairwise comparisons), and within the same

270 leaning direction for the LoS condition, muscle group, and MU subgroup. Confidence intervals

around the parameter estimate differences were calculated via parametric bootstrapping with

- 5000 iterations. The post hoc results are reported with the mean estimated difference (M), 95%
  confidence intervals (CI) and adjusted *p*-values.
- 274 Balance Performance

To determine whether the participants' balance performance during the leaning task differed across the leaning directions, the CoP 95% confidence ellipse area was compared using a one-way repeated measures ANOVA for 60% and 80% LoS separately. The 95% confidence ellipse area is the area of the smallest ellipse containing 95% of the data points collected during the central 20s of the 35s task (when CoP was maintained in the target). It is used to interpret sway displacement and is a measure of performance during postural tasks (37).

281

#### 282 Results

The total number of identified MUs for each muscle and LoS condition can be found in Table 1. There were no differences in the CoP 95% confidence ellipse area in any leaning directions (*p*-values ranging from 0.23 to 0.94), indicating that participants performed the leaning task appropriately and similarly among leaning directions.

287 Average Firing Rate

Individual MU Average Firing Rates (AFRs) and associated distributions for the MG, LG and SOL during the 5 leaning directions are depicted in Figure 4. The AFR of the common units is distributed unimodally (with a rightward skew) with a peak occurring at approximately 8-12 Hz. The AFR of the unique units, on the other hand, had two peaks occurring at approximately 12-15 Hz and 22-24 Hz. However, this second peak was modest in the unique MUs of the SOL muscle. Note that, in gastrocnemius, a larger proportion of the higher AFRs occurred during the  $30^{\circ}-90^{\circ}$  leaning directions (indicated by the density of the symbols at the higher firing

295 frequencies in Figure 4).

296 The estimates of the MU AFR generated by the statistical model are depicted in Figure 5. 297 There were main effects of direction (F = 31.31, p < 0.0001), muscle (F = 29.35, p < 0.0001) and 298 MU subgroup (F = 254.53, p < 0.0001) on the AFR. There was no statistically significant effect 299 of LoS on AFR, suggesting that recruitment of MUs is utilized to meet the force requirements of 300 the higher contraction intensities. The higher number of unique units during the 80% LoS 301 condition than in the 60% LoS condition confirms this (Table 1). No interaction effects were 302 observed. 303 A detailed description of the post hoc test results (including 95% CI and *p*-values) are 304 reported in Table 2. Post hoc tests for the leaning direction showed that for all three muscles and 305 MU subgroups the AFR was higher at 30°, 60°, 90° directions compared to the 0° and 120° 306 leaning directions, 30° (0°: M = 1.65 Hz; 120°: M = 2.32 Hz), 60° (0°: M = 1.52 Hz; 120°: M = 307 2.15 Hz) and 90° (0°: M = 1.27 Hz; 120°: M = 2.09 Hz) leaning directions. The post hoc tests for 308 the muscle effect showed that during each of the leaning directions the MUs in the MG (M = 309 1.44 Hz) and LG (M = 1.39 Hz) were discharging with higher AFRs compared to the SOL. 310 Finally, the post hoc test for MU subgroup showed that unique units were discharging with 311 higher firing rates compared to the common units during all leaning directions and muscle 312 groups (M = 3.48 Hz).

313

314 *Coefficient of Variation of the Interspike Interval* 

The estimates of the CoV<sub>ISI</sub> are displayed in Figure 6. The CoV<sub>ISI</sub> values increased across leaning directions (F = 23.64, p < 0.0001), and were larger during the 80% LoS condition (F = 317 30.26, p < 0.0001). The directional effect was dependent on the MU subgroup, indicated by the 318 interaction between direction and MU subgroup (F = 23.58, p = 0.0005). Additionally, there was 319 muscle effects (F = 47.97, p < 0.001) however, these effects were expected based on previous 320 literature.

321 A detailed description of the post hoc test results (including 95% CI and *p*-values) are 322 reported in Table 3. In the unique units there was an increase in  $CoV_{ISI}$  during the 60° (M = 2%) 323 leaning direction compared to the  $0^{\circ}$  target. In addition, the CoV<sub>ISI</sub> during the  $30^{\circ}$  (M = 4%),  $60^{\circ}$ 324 (M = 5%) and 90° (M = 4%) leaning directions were higher compared to the 120° leaning 325 direction. The post hoc test for the MU subgroup interaction showed that the unique units'  $CoV_{ISI}$  increased higher from the 0°-60° (M = 1.5%) and 120°-60° (M = 4%) leaning direction 326 327 compared to the common. The post hoc test for LoS showed higher CoV<sub>ISI</sub> values during the 328 80% LoS condition at all leaning directions in all three muscles of both MU subgroups (M = 329 2%). The MG (Unique: M = 3%; Common: M = 6%) and LG (Unique: M = 4%; Common: M =4%) had higher CoV<sub>ISI</sub> than the SOL in both MU subgroups during all leaning directions. 330 331 Finally, the unique units had higher CoV<sub>ISI</sub> than the common units during all leaning directions 332 (M = 7.1%).

333

#### 334 Degree of Intermittency

The estimates of the degree of intermittency described by the model are reported in Figure 7. There was a main effect of direction (F = 11.15, p < 0.0001), muscle (F = 10.33, p < 0.0001) and MU subgroup (F = 38.37, p < 0.001) on the degree of intermittency, but no effect of LoS condition. The muscle effect was dependent on the common units indicated by the muscle x

339 MU subgroup interaction (F = 3.67, p = 0.02).

340	A detailed description of the post hoc test results (including 95% CI and p-values) are
341	reported in Table 4. Post hoc tests for the leaning direction showed that in the unique units for all
342	three muscles there was a decrease in the degree of intermittency compared to the $0^{\circ}$ and $120^{\circ}$
343	leaning directions during the 30° (0°: M = 0.16; 120°: M = 0.21), 60° (0°: M = 0.13; 120°: M
344	0.18) and 90° (0°: $M = 0.18$ ; 120°: $M = 0.21$ ) leaning directions. In addition, the post hoc test for
345	the MU subgroup showed that the unique units had more intermittent activity compared to the
346	common units during the $0^{\circ}$ (M = 0.13) and $120^{\circ}$ (M = 0.28) leaning directions. Finally, the post
347	hoc test for the muscle interaction showed that the MG common units had more intermittent
348	recruitment compared to the SOL common units during all leaning directions ( $M = 0.07$ ).
349	
350	Discussion
351	This study examined the motor control of two sub-populations of MUs (common and
352	unique MUs) within the ankle plantarflexors to meet the force requirements in different
353	directions during a voluntary multidirectional leaning task. The AFR of both common and
354	unique units was higher during the leaning directions that required higher forces. However, even
355	though the 80% LoS condition had greater force requirements, the AFR did not differ between
356	LoS conditions, relying on recruitment instead. The two MU subgroups displayed different
357	patterns of MU variability. The directional effect of $\text{CoV}_{\text{ISI}}$ was dependent on the two sub-
358	populations of MUs. The unique units had higher $\text{CoV}_{\text{ISI}}$ , especially in the 30-90° directions,
359	whereas the common units $\text{CoV}_{\text{ISI}}$ remained relatively constant, while intermittent firing

360 behaviour was higher across the leaning directions for the unique units.

361

362 Differential behaviour of motor unit subgroups

363	Across the leaning directions, the unique and common units discharged at higher AFRs
364	when force requirements were largest (during 30°-90° directions). When analyzing the
365	differences between the MU subgroups, the unique units discharged at higher AFR compared to
366	the common units during all leaning directions (Figure 5). The distributions of the AFR (Figure
367	4) provide a better indication of the differences seen. There was a shift towards higher firing
368	frequencies in the distribution of AFR for the unique units during the 30°, 60°, and 90° leaning
369	directions that was not observed in the common units. The shift (or lack thereof) in the AFR
370	depending on the MU subgroups may suggest differential control of the distinct subgroups.
371	Further evidence supporting differential control is the strikingly different patterns of the $CoV_{ISI}$
372	between the two MU subgroups (Figure 6). The directional effect on $\text{CoV}_{\text{ISI}}$ was dependent on
373	the MU subgroups; with unique units modulating their $\text{CoV}_{\text{ISI}}$ while the common remained
374	relatively constant. Further, the unique units had larger $\text{CoV}_{\text{ISI}}$ when force requirements were
375	largest, whereas the common units fired with a lower $\text{CoV}_{\text{ISI}}$ across the directions.
376	The differences in the AFR and $\text{CoV}_{\text{ISI}}$ between the unique and common units suggest a
377	degree of selectivity of synaptic inputs. If the synaptic input was distributed uniformly across
378	common and unique units, we would have expected a similar pattern of firing. A recent study by
379	Hug et al. (38) provides support for selectivity of synaptic inputs controlling firing behaviours.
380	During volitional contractions, the modulations in the firing rates of concurrently active MUs
381	have been shown to be correlated and are believed to reflect a common drive (12, 39). However,
382	Hug and colleagues have recently shown that muscles from the same anatomical muscle group
383	(e.g. the ankle plantarflexors) do not share the same common drive during isometric and standing
384	heel-raise contractions (38). It is therefore possible that there are selective synaptic inputs that
385	can control separate motoneuron pools with the ankle plantarflexors.

17

386 In our previous study (19), we postulated that common units are recruited to produce a 387 baseline tonic level of force production. Once the task exceeded the capabilities of the common 388 units, recruitment of unique units with fibres grouped in different spatial locations were utilized 389 to meet the force requirements of the task. It is interesting to note that there were no statistically 390 significant differences in AFR between the LoS conditions. This may indicate that to achieve the 391 force required for the 80% LoS, recruitment of additional MUs is utilized over rate coding. The 392 results of our study provide indirect evidence for this hypothesis with more MUs identified 393 during the 80% LoS condition compared to the 60% condition (Table 1). 394 At first glance, it may appear that this organization is violating the size principle that has 395 been very well established in previous studies (40-46). However, the organization we describe 396 exists in harmony with the size principle. Henneman's size principle states that recruitment will 397 depend on the size of motor neurons receiving the same amount of synaptic input (47). This 398 implies that when a common input is sent to a motoneuron pool, recruitment will be governed by 399 the size of the units. However, when separate synaptic inputs are sent to different motoneuron 400 pools, the size principle will be governed separately within the pools. In fact, it has been 401 previously shown that different motoneuron pools may be controlled differentially when the 402 muscle contributes to different actions (48-51). Although the triceps surae are normally 403 considered to strictly plantarflex, there is evidence suggesting that MG and LG are also 404 responsible for providing inversion and eversion torques, respectively (18, 52, 53). Further, we 405 have previously demonstrated there is an uneven distribution of activation in the ankle 406 plantarflexors during postural control (18, 19). As the synaptic input to the ankle plantarflexors is 407 nonuniform and the muscles can contribute to different actions, it follows that the organization 408 we describe exists within the parameters of the size principle.

409 While selective synaptic inputs to the distinct motoneuron pools is supported in the 410 literature (18, 19, 38, 54-56), there is the possibility that the differential behaviour of the MU 411 subgroups may derive from the interaction between spinal mechanisms and muscle mechanics. 412 Neither spinal mechanisms (57) nor mechanical features alone can maintain upright standing in 413 humans (25, 58). However, the *combination* of spinal mechanisms and mechanical features of 414 the ankle plantarflexors without cortical input have been demonstrated in simulations to be 415 adequate for the maintenance of upright stance (57). The changing mechanical properties of the 416 muscles during the different leaning directions (i.e. nonuniform variations in length across the 417 different parts of the muscles) may interact with spinal proprioceptive activity (e.g. muscle 418 spindles, Golgi tendon organs, and interneurons) distributing nonuniformly on the motoneuron 419 pools. It should be noted that selective synaptic inputs, spinal mechanisms, and mechanical 420 properties may all interact to produce the differential behaviour output. Future research is 421 required to determine whether cortical, subcortical, and/or spinal contributions mediate the 422 differential behaviour observed.

423 Shifts in the distribution of the AFR

424 The distribution of AFRs in unique units showed peaks at lower frequency during the 0° 425 and 120° leaning directions and peaks at higher frequency during the 60°-90° (Figure 4). The 426 common units displayed lower frequency peaks for all leaning directions (Figure 4). To our 427 knowledge, this is the first study to demonstrate different AFR distributions for separate sub-428 populations of MUs within the same muscle in humans. Previous studies have investigated 429 similar characteristics in animal models that may help explain our findings. Tansey and 430 Botterman (59) assessed MU firing rate behaviours during muscle contractions evoked by 431 stimulation of the mesencephalic locomotor region in cats. The distribution of instantaneous

firing rates between slow-twitch and fast-twitch units was different. Both types of MUs displayed unimodal distribution patterns, but at opposite ends of the distribution pattern; slowtwitch units had distribution peaks near 20 Hz, whereas fast-twitch had distribution peaks near 40 Hz. In our study, it is possible that the common units were comprised of mostly slow-twitch units, whereas the unique units were both slow and fast-twitch units. Given that slow-twitch MUs tend to fire more tonically, this explanation fits well with the idea that the common units are first recruited to create baseline, stable torque (see above).

#### 439 Coefficient of Variation of the Interspike Interval and Degree of Intermittency

440 There are subtle but important differences between the CoV<sub>ISI</sub> and the degree of intermittency. CoV<sub>ISI</sub> is a measure of the variability of the MU firing instants and is associated 441 442 with the steadiness of the MU firings with a higher CoV<sub>ISI</sub> being indicative of fluctuating 443 (unsteady) firing intervals. The CoV<sub>ISI</sub> has been utilized and investigated extensively and is 444 generally used to describe the modulation of MU discharge times and force output steadiness 445 (60-63). The degree of intermittency is a standardized value measuring the extent to which the 446 firing rate has dropped below a physiological threshold of 4 Hz and determines whether a unit is 447 recruited sporadically. A low degree of intermittency is indicative of less MU derecruitment 448 occurrences, and therefore less sporadic (continuous) activity. The degree of intermittency has 449 been investigated to a lesser extent compared to CoV<sub>ISI</sub> but is believed to be due to the 450 physiological properties of the muscle (e.g. recruitment thresholds), the time constant of the 451 standing body (22, 23), the neural drive delivered to the motoneuron pool, or a combination of 452 the three (22, 23, 57, 64).

Unique and common units were observed to have different patterns of CoV<sub>ISI</sub>, but similar
 patterns of the degree of intermittency across directions. Interestingly, during the leaning task,

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455 the unique units CoV<sub>ISI</sub> increased across the directions and the degree of intermittency decreased. 456 This observation demonstrates that the unique units had higher firing variability but are behaving 457 less sporadically as the leaning directions shifted towards 90°. Contrastingly, common units 458 CoV<sub>ISI</sub> remained constant across directions, while their degree of intermittency decreased, 459 however this decrease was not found to be statistically significant. This is suggestive of a 460 steadier firing pattern and a more continuous activation compared to the unique units. These 461 observations may illustrate two things. First, it is possible that the higher CoV<sub>ISI</sub> may be a 462 response to the fluctuations in participant's CoP during the leaning task. As the unique units are 463 recruited in optimal locations (see above), the CNS may be modulating the activity of the units 464 producing the most efficient torques. Second, it may provide evidence of different mechanisms 465 or synaptic inputs controlling the two constructs. One might assume if the CoV<sub>ISI</sub> and degree of 466 intermittency were similarly controlled, they would follow the same trend (as MU firings 467 become more variable, they also become more intermittent) and they would have the same effect 468 in both MU subgroups. We observed an opposite trend (more variable firings and less 469 intermittency), and different effects in the unique and common units CoV<sub>ISI</sub>. While speculative, 470 this may suggest that the constructs are controlled by different mechanisms (e.g., physiological 471 organization vs synaptic inputs) or by a certain degree of independent synaptic inputs. Future 472 studies are required to confirm these potential explanations. 473 The degree of intermittency of both MU subgroups had a similar response to the

directions, however the unique units were more intermittent compared to the common units. It
has been reported that an intermittent control pattern allows for precise control (65) while
minimizing the neural cost compared to a constant signal (64). This efficiency allows the CNS to
reduce the degrees of freedom of standing balance (64). While these models were derived during

quiet standing, our results provide more evidence that intermittent control could be used toreduce the degrees of freedom during more demanding tasks.

480 The intermittent behaviour in the unique units is modulated based on direction. As the 481 leaning directions shifted towards the 90° target angle (directly forward movement), the degree 482 of intermittency of the ankle plantarflexors decreased. This is in agreement with other studies in 483 which intermittency has been observed to decrease during forward sway (12, 23). However, 484 previous studies have only investigated standing balance where directional information is 485 minimal. As mentioned, the 90° target angle requires maintaining the CoP position far forward 486 from its average position during quiet standing and thus imposing greater demands for 487 plantarflexion ankle torque. Given that leaning directly forward is controlled mostly by the ankle 488 plantarflexors with little help from other musculature (66, 67), it is not surprising that the degree 489 of intermittency decreased at 90° leaning direction. At this forward CoP position, fluctuations in 490 ankle torque necessary to compensate for the bodily sways would be superimposed on high, 491 average plantarflexion torque. As the leaning directions shift clockwise and counter clockwise, 492 more hip musculature may be recruited (67) that increases the feedback from group II muscle 493 spindles of the hip musculature received by CNS. Group II afferents can evoke opposite effects 494 in motor neurons that innervate flexor and extensor muscles in heteronymous connections (68, 495 69). This feedback system may allow the ankle plantarflexor to become more intermittent when 496 the leaning task is being assisted by other muscles to maintain the low metabolic cost of 497 standing.

498

499 Possible Limitations

500	While impossible to utilize with our methodology, the ability to record recruitment
501	thresholds for different MUs would have aided our interpretation of the findings. While other
502	studies provide information regarding slow twitch and fast twitch MUs, being able to record
503	recruitment thresholds would help confirm our hypothesis. In this study, the recruitment of motor
504	units occurs during a dynamic movement to the target location. Currently, decomposition of
505	HDs-EMG during dynamic movements has not been validated (70). Further development of
506	dynamic decomposition of HDs-EMG needs to be completed before we can utilize this
507	technique.

508 The contraction during the leaning task was not completely isometric; this caused slight 509 changes in muscle length during maintenance of the CoP within the targets. We nevertheless do 510 not expect these changes, which have been reported to be smaller than 50 µm (22), to have 511 affected the identification of firing instants. Indeed, the pulse-to-noise ratio, a metric indicating 512 the quality of EMG decomposition (71), was consistently high across all MUs analysed (>28dB). 513 Changes in muscle length across leaning directions were however presumably greater than those 514 occurring during each leaning direction. For this reason, we raised the error threshold for 515 matching units to 10% (19, 20), compensating for shape changes of common MUs waveform 516 during different leaning directions.

517

#### 518 Conclusion

This study provides evidence that there is differential behaviour of distinct motoneuron pools within the ankle plantarflexors utilized during postural control. Further research is necessary to determine the precise mechanism of the differential behaviour of distinct motoneuron subpopulations. 523

#### 524 **Conflict of interest**

525 The authors have no conflicts of interest to declare

526

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- 529
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- 708
- 709 Figure Legends
- 710 Figure 1
- 711 Schematic of the experimental set-up for the multi-directional leaning task. A. A participant
- standing on a force platform observing their centre of pressure movements towards the leaning
- targets in real time. B. The screen displaying participant's centre of pressure, leaning targets and
- the 60% (broken line) and 80% (solid line) limits of stability (LoS). The grey circles represent
- 715 the 5 different leaning directions  $(0^{\circ}, 30^{\circ}, 60^{\circ}, 90^{\circ}, \text{and } 120^{\circ})$  placed along the ellipses
- 716 calculated from the participant's limit of stability measures. Only one target was displayed at a
- time during the experiment. C. Placement of the high-density surface EMG sensors over the
- 718 medial (MG) and lateral (LG) gastrocnemius (two 32 electrode grids) and the soleus (SOL; a
- 519 single 64 electrode grid). D. Center of pressure (CoP) of a single participant during the multi-
- 720 directional leaning task.
- 721
- Figure 2
- Examples of motor unit action potentials from each plantarflexor muscle of a single participant
- during the 30° leaning direction. Medial and lateral gastrocnemius (MG and LG) are displayed
   on the top. Soleus (SOL) is displayed on the bottom. The vertical and horizontal axes represent
- the channel rows (vertical axes) and columns (horizontal axes) from the high-density surface
- 727 EMG.
- 728
- Figure 3
- 730 Example of motor unit pairs that were matched (A), and two motor unit (MU) pairs that were
- 731 untracked (B and C). The reference MU is depicted in red, and the paired MU in black. A
- displays a motor unit that was considered common, with the mean square error (MSE) = 6%. B
- and C display motor unit pairs that were considered unique, with MSE close to the 10%
- threshold (B: MSE = 16%) and very far from the threshold (C: MSE = 80%). Note that in C the
- 735 locations of the motor units are different. The vertical and horizontal axes represent the channel
- rows and columns numbers from the high-density surface EMG, respectively.
- 737
- 738 Figure 4
- 739 Violin plots for each leaning direction of the average firing rate (AFR) distribution of the motor
- vunits (MUs) for all participants combined, identified during both 60% and 80% limits of stability

- conditions overlaid by the MU AFR for the unique (left column; A, B) and common (right
- column; C, D) units in the medial and lateral gastrocnemius (MG: white circles, LG: black
- circles; A, C) and soleus (SOL: white circles; B, D). The number of MUs for each direction are
- indicated under each violin plot. A small amount of swarm was added to the x-axis for clarity of
- display. The horizontal location of the individual MU AFR does not have an effect on the
- distributions of the violin plots. There is a shift toward higher frequency in the AFR distribution
- for the unique units during the  $30^{\circ}$ - $90^{\circ}$  leaning directions.
- 748
- Figure 5
- Average firing rates (AFR) for unique (circles) and common (triangles) motor units during 60%
- 751 (grey) and 80% (black) limits of stability (LoS) reported for the medial gastrocnemius (MG),
- 752 lateral gastrocnemius (LG), and soleus (SOL). The data are estimated means and 95%
- confidence intervals. The estimates are calculated from the linear mixed model, with subjects as
- random intercepts, and fixed effects of leaning direction, muscle, limit of stability, and motor
- unit subgroup. Main effects are reported, no interactions were observed. The AFR was
   significantly higher for both unique and common units in medial and lateral gastrocnemius as
- significantly higher for both unique and common units in medial and lateral gastrocnemius as
   compared to soleus (see Table 2). \* Indicates significantly different from the 0° and 120° leaning
- directions for both 60% and 80% LoS (p < 0.001). † Indicates significantly different between
- 759 unique and common units (p < 0.001).
- 760
- 761 Figure 6
- 762 Interspike intervals coefficient of variation ( $CoV_{ISI}$ ) for unique (circles) and common (triangles) 763 motor units during 60% (grey) and 80% (black) limits of stability (LoS) reported for the medial
- 764 gastrocnemius (MG), lateral gastrocnemius (LG), and soleus (SOL). The data are estimated
- means and 95% confidence intervals. The estimates are calculated from the linear mixed models,
- with a participant as a random intercept, and fixed effects of leaning direction, muscle, limit of
- stability and motor unit subgroup. The CoV<sub>ISI</sub> was significantly higher for both unique and
- common units in the medial and lateral gastrocnemius as compared to soleus (see Table 3). × and
- <sup>769</sup> \* indicate significantly different from the 0° and 120° leaning direction, respectively, for both
- 60% and 80% LoS. † Indicates significantly different CoV<sub>ISI</sub> between unique and common units.
   † Indicates significantly different CoV<sub>ISI</sub> between 60% and 80% LoS for both unique and
- 772 common units.
- 773
- 774 Figure 7
- 775 Degree of intermittency for unique (circles) and common (triangles) motor units. Only the 60%
- (grey) limits of stability (LoS) are reported for the medial gastrocnemius (MG), lateral
- gastrocnemius (LG), and soleus (SOL). The 80% LoS had near identical values, and has been
- omitted for clarity. The data are estimated means and 95% confidence intervals for the log
- transformed values. The estimates are calculated from the linear mixed model with participants
- as a random intercept, and fixed effects of leaning direction, muscle, limit of stability and motor
- via unit subgroup. \* Indicates significantly different from the  $0^{\circ}$  and  $120^{\circ}$  leaning directions for all
- muscles (p < 0.001). † Indicates significantly different degree of intermittency between unique
- and common units. The degree of intermittency was significantly higher for the common units in
- the medial gastrocnemius as compared to soleus (see Table 4).



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55 X \* MG 50 45 Τ T T τŢ 40 Ŧ١ τľ T Y Y 35 30 × \* 55 LG 50 45 t Т 40 Ī Unique 80% LoS 35 Unique 60% LoS Common 80% LoS Common 60% Los 30 55 SOL × \* 50 45 40 T ΤI Т Ý 35 30 60 0 30 90 120

Coefficient of Variation (%)

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Degree of intermittency

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Muscle	Medial Gastrocnemius	Lateral Gastrocnemius	Soleus
Total	60%: 649 units, $44 \pm 27$ (24-59) 80%: 684 units, $47 \pm 28$ (25-63)	$60\%: 324 \text{ units}, 20 \pm 17$ (6-48) 80\%: 402 units, $25 \pm 19$ (10-52)	60%: 616 units, $42 \pm 25$ (26-52) 80%: 894 units, $56 \pm 30$ (28-82)
Unique	60%: 468 units, $31 \pm 20$	60%: 227 units, $15 \pm 10$	60%: 448 units, $32 \pm 19$
	(13-52)	(4-32)	(23-48)
	80%: 520 units, $36 \pm 21$	80%: 299 units, $21 \pm 14$	80%: 715 units, $51 \pm 32$
	(13-58)	(5-43)	(33-78)
Common	60%: 181 units, $13 \pm 11$	60%: 97 units, $5 \pm 3$	60%: 168 units, $12 \pm 6$
	(4-30)	(1-12)	(5-23)
	80%: 164 units, $11 \pm 10$	80%: 103 units, $5 \pm 5$	80%: 179 units, $14 \pm 8$
	(5-22)	(1-13)	(6-24)

Table 1. Number of identified motor units

Values are group totals, mean  $\pm$  standard deviation and (minimum-maximum range) across participants. 60% and 80% correspond to the limit of stability condition.

Predictors	Comparison	Estimated	95% CI	p-value
	-	difference (Hz)		-
Direction				
Unique & Common	0°-30°	1.65	1.19-2.12	< 0.0001
-	$0^{\circ}$ - $60^{\circ}$	1.52	1.10-2.67	< 0.0001
	0°-90°	1.27	0.98-2.67	0.003
	$120^{\circ}-30^{\circ}$	2.32	1.05-3.92	< 0.0001
	$120^{\circ}-60^{\circ}$	2.15	1.76-3.52	< 0.0001
	120°-90°	2.09	1.65-3.52	< 0.0001
Muscle				
Unique & Common	MG-SOL	1.44	0.98-2.05	< 0.0001
	LG-SOL	1.39	0.98-2.50	< 0.0001
MU Subgroup				
	Unique-Common	3.48	1.46-7.40	< 0.0001

Table 2: Post hoc results from the linear mixed model using average firing rate as the dependent variable.

Estimated difference = Mean estimated difference between comparisons in Hz. CI = confidence intervals. MU = Motor unit. P-values are adjusted using Bonferroni corrections. No interactions were observed; therefore, the motor unit subgroup marginal means are given equal weighting.

Predictors	Comparison	Estimated	95% CI	p-value
		difference (a.u.)		1
Direction				
Unique	$0^{\circ}-60^{\circ}$	0.02	0.01-0.04	0.002
1	$120^{\circ}-30^{\circ}$	0.04	0.02-0.06	< 0.0001
	$120^{\circ}-60^{\circ}$	0.05	0.03-0.07	< 0.0001
	$120^{\circ}-90^{\circ}$	0.04	0.02-0.06	< 0.0001
Muscle				
Unique	MG-SOL	0.03	0.02-0.05	< 0.0001
-	LG-SOL	0.04	0.02-0.06	< 0.0001
Common	MG-SOL	0.06	0.03-0.08	< 0.0001
	LG-SOL	0.04	0.02-0.06	0.003
MU Subgroup				
	Unique-Common	0.071	0.21-0.164	< 0.0001
<b>Direction x MU</b>				
Subgroup				
Unique-Common	$0^{\circ}-60^{\circ}$	0.015	0.01-0.03	0.04
-	$120^{\circ}-60^{\circ}$	0.04	0.02-0.06	0.009
LoS				
	60%-80%	0.02	0.01-0.04	< 0.0001
$\Gamma_{11}$ + 1.1.00	<b>N</b> (		1	1.

Table 3: Post hoc results from the linear mixed model using coefficient of variation of the interspike interval ( $CoV_{ISI}$ ) as the dependent variable

Estimated difference = Mean estimated difference between conditions. a.u. = arbitrary units. CI = confidence interval. MU = motor unit. LoS = limit of stability. P-values are adjusted using a Bonferroni correction.

Predictors	Comparison	Estimated	95% CI	p-value
	1	difference		
		$(\log(IO/s))$		
Direction				
Unique	0°-30°	0.16	0.027-0.29	0.003
-	$0^{\circ}-60^{\circ}$	0.13	0.007-0.26	0.006
	0°-90°	0.18	0.05-0.33	< 0.0001
	$120^{\circ}-30^{\circ}$	0.21	0.06-0.32	< 0.0001
	$120^{\circ}-60^{\circ}$	0.18	0.06-0.32	< 0.0001
	120°-90°	0.21	0.11-0.37	< 0.0001
MU Subgroup				
Unique-Common	$0^{\mathrm{o}}$	0.13	0.05-0.26	0.0024
	120°	0.28	0.13-0.35	< 0.0001
Muscle x MU				
Subgroup				
Common	MG-SOL	0.07	0.02-0.38	0.04

Table 4: Post hoc results from the linear mixed model using degree of intermittency as a dependent variable.

Estimated difference = Mean estimated difference between conditions. IO/s = intermittent occurrences/second. CI = confidence interval. LoS = limit of stability. P-values are adjusted using a Bonferroni correction.

# Differential control of distinct motoneuron pools in the ankle plantarflexors

## **METHODS**

## OUTCOME



## CONCLUSION

behaviour (average firing rate, coefficient of variation, intermittent behaviour) for MU active across multiple directions (common) and in one direction (unique)

There is differential behavior of distinct motoneuron pools rection (unique) within the triceps surae utilized during postural control Downloaded from journals.physiology.org/journal/jn at Univ Western Ontario (129.100.255.024) on December 12, 2022.