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61 Abstract

Individuals with multiple sclerosis (MS) experience progressive declines in movement 62 capabilities, especially walking performance. The purpose of our study was to compare the 63 amount of variance in walking performance that could be explained by the functional capabilities 64 of lower leg muscles in persons with MS and a sex- and age-matched control group. Participants 65 performed two walking tests (6-min walk and 25-ft walk), strength tests for the plantar flexor and 66 dorsiflexor muscles, and steady submaximal (10% and 20% maximum) isometric contractions. 67 High-density electromyography (EMG) was recorded during the steady contractions and the 68 signals were decomposed to identify the discharge times of concurrently active motor units. 69 70 There were significant differences between the two groups in the force fluctuations during the steady contractions (force steadiness), the strength of the plantar flexor and dorsiflexor muscles, 71 72 and the discharge characteristics during the steady contractions. Performance on the two walking tests by the MS group was moderately associated with force steadiness of the plantar 73 flexor and dorsiflexor muscles; worse force steadiness was associated with poorer walking 74 75 performance. In contrast, the performance of the control group was associated with muscle 76 strength (25-ft test) and force steadiness of the dorsiflexors and variance in common input of motor units to the plantar flexors (6-min test). These findings indicate that a reduction in the 77 78 ability to maintain a steady force during submaximal isometric contractions is moderately 79 associated with walking performance of persons with MS.

New & Noteworthy. The variance in walking endurance and walking speed was associated with
force control of the lower leg muscles during submaximal isometric contractions in individuals
with MS. In contrast, the fast walking speed of a sex- and age-matched control group was
associated with the strength of lower leg muscles. These findings indicate that moderate

- 84 *declines in the walking performance of persons with MS are more associated with impairments*
- 85 *in force control rather than decreases in muscle strength.*
- 86 Keywords. Multiple sclerosis, walking, force steadiness, common drive, motor units, high-
- 87 density EMG

88 Introduction

Multiple sclerosis (MS) is a demyelinating neurological disorder that compromises function 89 within the central and peripheral nervous systems. The damage caused by the disease invariably 90 leads to increases in self-reported levels of fatigue, worse walking and balance, and decreased 91 quality of life. When walking at a preferred speed, for example, individuals with MS take 92 shorter, wider, and more variable strides and spend a greater percentage of the gait cycle in the 93 double-support phase (Benedetti et al. 1999; Comber et al. 2017; Galea et al. 2017; Gutierrez et 94 al. 2005; Sosnoff et al. 2012). These declines in gait kinematics are associated with a greater 95 energetic cost of walking for persons with MS (Motl et al. 2012; Sebastião et ak, 2018), which 96 may contribute to the elevated levels of fatigue they report (Hadjimichael et al. 2008; Kluger et 97 al. 2012; Loy et al. 2017; Zijdewind et al. 2016). 98

99 The mobility-limiting symptoms experienced by persons with MS arise in part from the progressive decline in sensorimotor function (Cabib et al. 2015; Newsome et al. 2011; 100 Zackowski et al. 2015). The disease compromises functional connectivity within the central 101 102 nervous system (Fling et al. 2015, Fritzer et al. 2017; Schlaeger et al. 2015), including the activation signal sent from the spinal cord to muscle (DeLuca et al. 2013; Kearney et al. 2015; 103 Petrova et al. 2018). These changes result in a decrease in the maximal capacity to activate 104 105 muscle (Andreasen et al. 2009; Ng et al. 2004), a reduction in the number of functioning motor units (Vogt et al. 2009), and a lesser range of rate coding for single motor units (Rice et al. 106 1992). Consistent with these observations, we have found that the normalized amplitude of the 107 force fluctuations and the mean interspike interval of motor unit action potentials in plantar 108 flexor muscles during steady submaximal contractions can explain moderate, but significant, 109

amounts of the variance in performance of individuals with MS on tests of maximal walkingspeed and walking endurance (Almuklass et al. 2018).

112 Technological advances in electromyography (EMG) now make it possible to evaluate more 113 thoroughly the influence of the impairments in muscle activation on the capacity of an individual to control muscle force (Farina et al. 2016). In particular, multi-channel electrode systems 114 115 enable the decoding of the neural drive to muscle by identifying the discharge times of several concurrently active motor units from surface EMG recordings (Farina et al. 2010). In a seminal 116 117 study, for example, Negro et al. (2009) demonstrated that the first principal component of the smoothed discharge rates of ≥ 4 motor units in a hand muscle was strongly correlated (72 ± 13%) 118 with the fluctuations in abductor force applied by the little finger during steady submaximal 119 contractions. They found a similar association $(58 \pm 14\%)$ for motor units in tibialis anterior 120 when the dorsiflexors performed a steady, isometric contraction at 10% maximal voluntary 121 contraction (MVC) force. Subsequently, Thompson et al. (2018) demonstrated with an in vivo 122 cat model that the cumulative spike trains of ≥ 9 motor units were strongly correlated with both 123 the total force (99.2 \pm 0.2%) and the higher frequency force fluctuations (83.5 \pm 1.3%) evoked in 124 the soleus muscle. 125

The normalized amplitude of the force fluctuations (coefficient of variation for force) during steady submaximal contractions provides a measure of force steadiness (Galganski et al. 1993). Force steadiness is a sensitive marker of fine motor control as it varies with the target force for the steady contraction (Galganski et al. 1993; Laidlaw et al. 2000; Tracy and Enoka, 2002), the test muscle (Grunte et al. 2010; Tracy et al. 2007), the difficulty of the task (Almuklass et al. 2016; Holmes et al. 2015), the level of physiological arousal (Christou et al. 2004; Pereira et al. 2015; Rice et al. 2015), the age of the performer (Castronovo et al. 2018; Hamilton et al. 2017;

133 Marmon et al. 2011), and the neurological health of the individual (Bilodeau et al. 2000; Carville

et al. 2007; Gould et al. 2018). Critically, force steadiness has consistently emerges as a

statistically significant explanatory variable for the variance in performance on a number of

dynamic tasks, such as walking (Almuklass et al. 2018, Mani et al. 2018), manual dexterity

137 (Almuklass et al. 2016, Feeney et al. 2018; Hamilton et al. 2017; Marmon et al. 2011), risk of

falls in older adults (Carville et al. 2007), and postural sway (Davis et al. 2020).

The purpose of our study was to compare the amount of variance in walking performance that 139 could be explained by the functional capabilities of lower leg muscles in persons with MS and an 140 age- and sex-matched control group. Walking performance was quantified as fast walking speed 141 (25-ft test) and walking endurance (6-min test), whereas muscle function was characterized with 142 measures of muscle strength, force steadiness during submaximal isometric contractions, and the 143 discharge characteristics of single motor units during the steady contractions. Due to the 144 explanatory power of force steadiness for the performance of older adults on tests of motor 145 function, we hypothesized that significant amounts of the variance in walking performance for 146 individuals with MS would be explained by differences in force steadiness, whereas we expected 147 measurements of muscle strength to be the dominant explanatory variable for the control group. 148

149 METHODS

150 *Subjects*

151 Data from 18 participants with relapsing-remitting MS who participated in a previous study (52

 ± 8 yrs, 7 men, 3.6 ± 0.96 PDDS; Almuklass et al., 2017, Almuklass et al., 2018) were compared

- with 18 sex- and age-matched (\pm 5 yrs) control subjects (52 \pm 9 yrs) who reported no
- 154 neurological diseases. The control subjects were explicitly recruited for our current report. The
- 155 MS participants provided written informed consent (IRB#: 13-0720) for the previous study. For

156 the current study, a waiver-of-consent document was approved in accordance with the exception to the regulations of the Office of Human Research Protections in that there was no more than 157 minimal risk of harm to subjects and it involved no procedures for which written consent is 158 normally required outside the research context. The MS subjects were sent an approved e-mail 159 message that requested permission to use their data by replying either 'yes, I consent for you to 160 use my data' or 'no, I do not consent for you to use my data.' The control group provided 161 written informed consent to participate in a protocol (IRB#: 16-0396) that was approved by the 162 Institutional Review Board at the University of Colorado. 163

The MS participants completed a baseline evaluation over a 2-day period that included 164 measurements of muscle strength, force steadiness, walking endurance, and fast walking speed 165 (Almuklass et al., 2017). They then performed a 6-wk intervention in which neuromuscular 166 electrical stimulation was applied to their plantar flexor and dorsiflexor muscles 3x/wk for a total 167 of 18 treatment sessions. The current report includes only those data that were obtained from the 168 169 MS participants during the baseline evaluation. These baseline data were compared with those for the control group that was collected using the same protocol performed by the MS group, but 170 within a single session. 171

172 Strength and force steadiness.

Both groups performed measurements of muscle strength and force steadiness with the
dorsiflexor and plantar flexor muscles. Participants lay in a supine position with the hip and
knee joints at neutral angles and the ankle at 90°. A strap was placed around the widest section
of the forefoot and connected to a strain-gauge transducer (MLP-300, Transducer Techniques,
Temecula, CA). The force signal was sampled at 1 kHz (Power 1401, Cambridge Electronic
Design, Cambridge, UK) and displayed on a monitor that was placed ~1 m in front of the

participants. They performed maximal voluntary contractions (MVC) with the ankle-spanning dorsiflexor and plantar flexor muscles; the task was to increase muscle force gradually (~30% MVC/s) up to maximum and then sustain it for ~ 3 s. They were verbally encouraged to provide a maximal effort during each MVC. They performed 2-5 MVCs until the peak forces in 2 trials were within 10% of each other, the greater of which was taken as the MVC force. The moment arm for the muscle torque calculation was measured as the distance from the lateral malleolus to the center of the strap around the forefoot.

Participants then performed submaximal isometric contractions for ~30 s while matching target 186 187 forces of 10% or 20% MVC force (Figure 1. A and B). The submaximal contractions were performed with the dominant leg (Waterloo footedness questionnaire) by the control group and 188 in the self-reported leg with fewer symptoms (less-affected leg) by the MS group. The task was 189 to increase the force from an initial resting value up to the target force displayed on a monitor 190 (visual angle $\sim 0.25^{\circ}$) and then to maintain a steady contraction for ~ 30 s. Participants were not 191 asked to achieve a specific a rate of increase in force, but typically reached the target force ≥ 2 s 192 $(\sim 5 - 10\% \text{ MVC/s})$. Participants performed two trials at each target force for both muscle 193 194 groups.

The force signals were amplified (Coulbourn Instruments, Allentown, PA) and then recorded on a computer with Spike2 data acquisition software (Version 6.17, Cambridge Electronic Design, Cambridge, UK). The force signals were subsequently low-pass filtered with a cut-off frequency of 20 Hz and the fluctuations in force during the steady contractions (force steadiness) were quantified as the coefficient of variation for force (Galganski et al., 1993).

200 High-density surface EMG.

Three high-density surface electrodes (4x8 detection points, 10 mm interelectrode distance) were placed longitudinally over the tibialis anterior, medial gastrocnemius, and lateral portion of the soleus muscles during the steady isometric contractions (Figure 1.A; plantar flexors or dorsiflexors x two target forces x two trials at each target force). A custom MATLAB script was used to decompose motor unit activity from the EMG signals (Figure 1, C and D; Holobar & Zazula, 2007).

207

[Figure 1]

The interspike intervals (ISI) for each motor unit were calculated as the difference between 208 consecutive discharge times. Due to limitations in decomposition algorithms, the signals likely 209 contained both motor unit signals as well as non-physiological waveforms. To control for 210 211 aberrant waveforms, the decomposed signal was deemed not to be a motor unit when all four of the following criteria were met: 1) 25 < ISI > 400 ms; 2) 8% < coefficient of variation for ISI >212 55%; 3) if the skewness of the ISI histogram was between -0.5 and 0.5; and 4) if the coefficient 213 of variation for ISI was <10% and the skewness of the ISI histogram was <1 (Almuklass et al., 214 2018). 215

When the discharge times of ≥ 5 motor units were identified for a given trial, the data were 216 entered into a state-space model (Feeney et al., 2017, Macke et al., 2011) to estimate the variance 217 in the common synaptic input to the motor neuron pools during the steady contractions. Briefly, 218 219 the unobserved common input at a given time, x(t), is related to a linear approximation of the common input (A) at the previous time point, the influence of the intended target force on 220 discharge rate (bf(t)), and the effects of synaptic noise on signal transmission as represented as a 221 222 Gaussian noise vector (E(t)). The target force is represented by f(t), which is influenced by a vector representing discharge rate saturation in low threshold motor units (b) 223

$$\boldsymbol{x}(t) = \boldsymbol{A}\boldsymbol{x}(t-1) + \boldsymbol{b}\boldsymbol{f}(t) + \boldsymbol{E}(t)$$

The common synaptic input varies over the contraction and is modeled with Poisson lineardynamics.

$$x_i \sim N(x_o, Q_o)$$
$$x_t | x_{t1} \sim N(Ax_{t1}, bf_t, Q)$$

The term Q represents the variance in the state-space trajectory, which corresponds to the fluctuations in the common input signal. Q is assumed to remain constant during each steady contraction.

229 Motor unit discharge times were sorted into 10-ms bins and arranged into a binary matrix where the occurrence of an action potential was represented by 1. The matrix was entered into an open 230 231 source Expectation-Maximization algorithm (Macke et al., 2011; Buesing et al., 2012) to estimate the parameters of the state-space model. The Q values were subsequently low-pass 232 filtered to account for the response characteristics of muscle. The unfiltered Q values were 233 interpreted as the variance in the common synaptic input received by the motor neuron pool, 234 whereas the filtered Q values estimated the variance in the activation signal received by the 235 muscles. 236

237 Walking tests.

Each participant performed two walking tests: 25-ft walk (fast walking speed) and the 6-min
walk (walking endurance). Fast walking speed was represented as the average of two 25-ft tests.
Participants stood ~1 m before the start line and a timer was started as soon as one foot stepped
over the line and stopped when one foot stepped over the end line. The 6-min test required
participants to walk around an indoor track for 6 minutes and the distance covered was

243 measured. The instruction for both tests was to walk as quickly but as safely as possible. Due to 244 the influence of fatigue on the performance of persons with MS, they performed the two walking 245 tests on one day and the tests of muscle function on another day. The control subjects performed 246 all the tests on the same day.

247 Questionnaires.

The disability status of the MS participants was quantified with three questionnaires: patient determined disease steps (PDDS), MS walking scale-12 (MSWS-12), and modified fatigue impact score (MFIS). Only the PDDS scores are reported here.

251 Data Analysis.

Descriptive statistics for both pairwise and group values were calculated and are reported as 252 253 mean \pm standard deviation. Pairwise differences across groups were calculated as the absolute difference between paired subjects for each variable. When the 95% confidence interval of the 254 differences included 0, the difference was deemed not to be statistically significant. Conversely, 255 the absence of 0 in the 95% confidence interval corresponded to a statistically significant 256 pairwise difference. Group differences were examined with linear regression analysis for each 257 variable (P < 0.05). Muscle strength was compared between 1) the less-affected leg from the MS 258 group to the dominant leg from the control group, and 2) the more-affected leg of the MS group 259 and the non-dominant leg of the control group. Due to the variable numbers of motor units 260 across subjects, we could only examine group differences for both the filtered and unfiltered Q 261 values. 262

Pearson-product correlations were used to compare performance on the walking tests (25-ft test and 6-min test) with muscle strength, force steadiness, and the variability in common input (Q) for each group of participants. Significantly correlated variables were entered in a backward

linear regression analysis to identify the measures of muscle function that were most stronglyassociated with the variability in walking performance for each group.

All statistical procedures were performed in R (Version 3.5.1) with an α level of P<0.05 (with

269 Bonferroni correction for multiple comparisons) for statistical significance.

270 **RESULTS**

The results comprise an assessment of the associations between walking performance and measures of muscle function and motor unit activity in lower leg muscles of persons with MS and an age- and sex-matched control group.

274 Strength, force steadiness, and walking.

Table 1 contains the pairwise and group differences for the measurements of force steadiness and 275 muscle strength. In contrast to the control group, force steadiness (coefficient of variation for 276 force) for both the plantar flexors and dorsiflexors was not less at the greater target force (20% 277 278 MVC) than the lower target force (10% MVC) for the MS group. However, force steadiness for the dorsiflexors of both groups was greater at the two target forces than that for the plantar 279 flexors. Nonetheless, there were no significant differences between groups in force steadiness of 280 281 the plantar flexors at either target force (10% or 20% MVC) and in the force steadiness for the dorsiflexors at 10% MVC. At the greater target force (20% MVC) for the dorsiflexors, however, 282 283 the MS participants exhibited significantly worse force steadiness (greater force fluctuations) 284 than the control group.

285

[Table 1]

There were no statistically significant differences in MVC torque between legs for the plantar
flexor muscles of each group of participants and for the dorsiflexor muscles in the control group.

In contrast, dorsiflexor MVC torque for the more-affected leg was significantly less for the lessaffected leg in the MS group. Further, the MS group was consistently weaker than the control
group for both the plantar flexor and dorsiflexor muscles of each leg.

The difference in time to complete the 25-ft walk test between the MS and control groups was not statistically significant despite substantially different group means. This was due to the large standard deviation for the MS group (Table 2). In contrast, the control group walked further during the 6-min walk test at each recorded timepoint (1 min, 2 min, 4 min, and 6 min) than the MS group. Furthermore, the absolute difference in distance walked between participants matched across groups increased progressively during the 6-min test (Table 2).

297

[Table 2]

298 Motor units.

Motor unit data were recorded for the tibialis anterior, medial gastrocnemius, and soleus muscles 299 during the submaximal contractions at the two target forces. Our analysis required ≥ 5 identified 300 301 motor units in each trial in order to calculate Q (Feeney et al., 2017). As the 19 participants in each group performed 2 trials for every condition, there was the possibility of recording usable 302 data from 38 trials for each muscle group at each target force. However, the number of viable 303 304 trials was less than 38 for both the MS and control groups, respectively: tibialis anterior (10%) MVC: 24 and 32 trials; 20% MVC: 30 and 36 trials), soleus (10% MVC: 13 and 20 trials; 20% 305 MVC: 18 and 25 trials), and medial gastrocnemius (10% MVC: 6 and 20 trials; 20% MVC: 14 306 307 and 21 trials). Thus, it was more difficult to identify a sufficient number of motor units in the MS participants than the control group and in the plantar flexor muscles than the dorsiflexor 308 309 muscle. Due to the shared contribution of medial gastrocnemius and soleus to the net plantar

flexor torque, the discharge times of motor units recorded from the medial gastrocnemius and 310 soleus were pooled and the variance in the common input (Q) was calculated for the pooled data. 311 312 When performing a steady isometric contraction at 10% MVC force with the dorsiflexor 313 muscles, the mean and coefficient of variation for interspike interval for motor units in tibialis anterior were greater and the unfiltered Q value was lower for the MS group than the control 314 315 group (Table 3). At the higher target force (20% MVC), only the coefficient of variation for 316 interspike interval of motor units in tibialis anterior was greater for the MS group than the 317 control group (Table 3).

There were fewer group differences in motor unit activity during the steady contractions 318 performed with the plantar flexor muscles. The most consistent difference between the groups 319 320 was the lower unfiltered Q values (medial gastrocnemius + soleus) for the MS group compared with the control group at both target forces (10% & 20% MVC; Table 5). Also, the mean 321 interspike interval for motor units in medial gastrocnemius was greater at the 20% target force 322 for the MS group than the control group. Notably, there was only one statistically significant 323 difference for either group of participants in the mean and coefficient of variation for interspike 324 intervals across target forces (Table 4). Mean interspike interval for motor units in the tibialis 325 anterior of the control group were significantly lower at 20% MVC than at 10% MVC. 326

327

The filtered Q values for tibialis anterior were correlated with force steadiness in the dorsiflexor muscles of the MS group. When the target forces (10 and 20% MVC) were pooled, the correlation coefficient was 0.32 (P = 0.04) for the MS group and 0.23 (P = 0.06) for the control group. The association between the filtered Q values from the tibialis anterior and force steadiness of the dorsiflexors when the MS group maintained a steady contraction at 10% MVC

was not statistically significant when the data were not collapsed across target forces (r = 0.37, *P* = 0.10). The filtered Q values from tibialis anterior and force steadiness for the dorsiflexors at 20% MVC were significantly correlated (r = 0.45, *P* = 0.008) in the control group. There were no significant correlations between the filtered Q values and force steadiness of the plantar flexors.

338 Walking correlations and regression analyses.

To assess the relative explanatory power of the outcome variables on the two tests of walking 339 performance, we first identified the variables that showed statistically significant correlations 340 with each walking test and then compared the relative influence of these correlations by 341 developing regression models that retained the most influential variables. Table 6 shows that 342 343 there were a greater number of significant Pearson-product correlations between muscle function (force steadiness, strength, and motor unit discharge characteristics) and walking performance 344 345 for the MS group than for the control group. The distance walked in 6 min by the MS group was 346 correlated with three of the four measures of force steadiness, with the filtered Q values for the motor units in the plantar flexors, and with the strength of the dorsiflexors in the more-affected 347 348 leg. The distance walked in 6 min by the control group was correlated with force steadiness for 349 the dorsiflexors at the lower target force and the filtered Q value for the plantar flexors.

350 The time it took the MS group to walk 25 ft was correlated with the same set of outcome

variables as for the 6-min walk, except not with the strength of the dorsiflexors in the more

affected leg. The time it took the control group to walk 25 ft was correlated with one measure of

353 force steadiness (plantar flexors at 10% MVC) and the strength of plantar flexors and

dorsiflexors of the left and right legs.

355 Four regression models were developed (2 groups x 2 tests of walking performance); each model involved up to three variables with the strongest correlations to walking performance. The 356 regression model for the 6-min distance of the MS group explained 54% of the variance with two 357 predictor variables (Figure 2A): force steadiness of the plantar flexors at 20% MVC force (partial 358 r = -0.52) and force steadiness for the dorsiflexors at 10% MVC (partial r = -0.36). The 359 regression model for the 6-min distance of the control group explained 36% of the variance with 360 two predictor variables (Figure 2B): force steadiness of the dorsiflexors at 20% MVC force; 361 partial r = -0.54) and the filtered O values for the plantar flexors (partial r = -0.19). 362 The regression model for the time it took the MS group to walk 25 ft explained 34% of the 363 variance with two predictor variables (Figure 2C): force steadiness of the plantar flexors at 20% 364 MVC force (partial r = -0.47) and force steadiness for the dorsiflexors at 10% MVC (partial r =365 0.48). The regression model for the time it took the control group to walk 25 ft explained 56% 366 of the variance with two predictor variables (Figure 2D): MVC torque for the plantar flexors of 367 the non-dominant leg (partial r = -0.49) and MVC torque for the dorsiflexors of the dominant leg 368 (partial r = -0.39). 369

370

[Table 6 & Figure 2]

371 **DISCUSSION**

The main finding of our study was that moderate amounts of the variance in two tests of walking performance for persons with MS and control participants were explained by measurements derived from submaximal, isometric contractions with the plantar flexor and dorsiflexor muscles. The exception to this general conclusion was that the explanatory variables for the fast walking speed (25-ft test) performed by the control group were two measurements of muscle strength. We hypothesized that significant amounts of the variance in walking performance would be

explained by differences in force steadiness for individuals with MS, whereas we expectedmuscle strength to be the dominant explanatory variable for the control group.

380 Walking, strength, and force steadiness.

We found that greater coefficients of variation in force (worse force steadiness) corresponded to 381 a shorter distance walked in 6 min in both groups of participants and to a longer time to walk 25 382 ft in the MS participants. These associations were stronger for the MS group than the control 383 group. The functional significance of force steadiness is that it represents an individual's 384 accuracy in producing and maintaining a force trajectory (Christou et al. 2003). Within the last 5 385 years, our lab has found that force steadiness, a measurement derived from submaximal 386 isometric contractions, consistently emerges as a significant explanatory variable in regression 387 388 models to explain the variance in performance of a dynamic task (Almuklass et al. 2016, Almuklass et al. 2018, Davis et al. 2020, Feeney et al. 2018, Mani et al. 2018). 389 Although the direction of the correlations (positive or negative) has differed across studies, the 390 correlation coefficients and partial-r values have consistently comprised moderate-to-strong 391 values. Of particular note, when both force-steadiness values and muscle strength (MVC 392 torques) have both been entered into the regression models, force steadiness frequently has the 393 greater explanatory power for the variance in the outcome variables (Almuklass et al. 2016; 394 Hamilton et al. 2017; Justice et al. 2014; Mani et al. 2018; Marmon et al. 2011). Taken together 395 with previous research, these results suggest that an individual's ability to sustain a constant 396 force during a steady isometric contraction at a submaximal target force is more often 397 functionally relevant than maximum muscle strength for most activities of daily living. The 398 399 stronger correlations between walking performance and force steadiness for the MS group than the control group suggests that the negative impact of the disease on the ability to sustain steady 400

submaximal contractions has functional consequences. Moreover, the underlying mechanisms
presumably involve the variance in the common input received by the activated motor neurons
(Feeney et al. 2017; Negro et al. 2009; Thompson et al. 2018).

The only case in which muscle strength emerged as a significant explanatory variable was for the 25-ft test performed by the control group. Stronger lower leg muscles were associated with faster times to walk 25-ft (faster). As the goal of this test is to walk as quickly as possible for a relatively short distance, the same relative muscle strength produces greater absolute forces and faster gait cycles (Bohanan, 1997; Broekmans et al. 2012; Clark et al. 2013). Although muscle strength did not emerge as an explanatory variable for the MS participants, dorsiflexor MVC torque for the more-affected leg was negatively correlated with their walking endurance.

411 *Motor units*.

Previously, Almuklass et al. (2018) found that the walking performance of individuals with MS (n = 27) was associated with the mean interspike intervals of single motor units in medial gastrocnemius and soleus muscles during steady contractions (10% MVC force) averaged across 2 to 3 time points. The current study comprised the data obtained from a subset of these participants (n = 18) in only one of the experimental sessions. In this reduced data set, we did not find any significant associations between either the mean or the coefficient of variation for interspike interval with performance on the two walking tests.

Instead of estimating the cumulative influence of motor unit activity by collapsing data across experimental sessions (Almuklass et al. 2018), we derived Q values from the discharge times of many motor units during steady submaximal contractions to infer the activity of the motor unit pool. We found that lower filtered Q values for the plantar flexors of individuals with MS

corresponded to a longer time to complete the 25-ft test and a shorter distance walked in 6 min. 423 In contrast, the filtered Q value for the plantar flexors of the control group was negatively 424 associated with walking endurance, but not with fast-walking speed. These findings suggest that 425 greater variance in the common input to motor neurons, once it had been low-pass filtered by the 426 muscle, was associated with better walking endurance and faster walking speed in individuals 427 428 with MS, but with worse walking endurance in healthy adults. Although we hypothesized that variance in the common input would be associated with better walking performance, we 429 expected the association to be for the dorsiflexors and not the plantar flexors. 430 The current study appears to be the first to examine the common input to the motor unit pool in 431 individuals with MS. The moderate correlations between the variance in the common 432 modulation of motor units within the tibialis anterior and force steadiness of the dorsiflexors are 433 consistent with previous work reported from our lab on the wrist extensor muscles (Feeney et al. 434 2018). In that study, they found that the unfiltered Q values derived from the discharge times of 435 motor units during a steady contraction at 20% MVC force were significantly correlated with 436 force steadiness for both young ($r^2 = 0.31$) and older ($r^2 = 0.39$) adults. However, only the 437 correlation for the older adults had a functional influence on the time it took older adults to 438 439 complete a pegboard test of manual dexterity. Perhaps these findings suggest that the functional significance of the variance in common input is most evident in individuals with a reduced 440 integrity of the sensorimotor system. 441

A key issue that needs to be addressed in this field is the discrepancy in the magnitude of the
correlation between the variance in common input and force steadiness (fluctuations in force) for
the classic studies (Negro et al. 2009; Thompson et al. 2018) and our work (Feeney et al. 2018;
current study). One likely confounding factor is the number of muscles that contribute to the

applied force. Negro et al. (2009) studied the sole contributor to the abduction force exerted by 446 the little finger (abductor digiti minimi), whereas Thompson et al. (2018) examined the isolated 447 soleus muscle in a decerebrate cat preparation. In these two studies, 72% and 84%, respectively, 448 of the force fluctuations during steady contractions were explained by the estimated variance on 449 the common input signal of the activated motor units. In contrast, the current study assessed the 450 association between the net force and motor units in just a few of the involved muscles. For 451 example, there are nine muscles that can contribute to the net plantar flexor force with each 452 muscle varying in architecture and lines of action. Although some evidence suggests that the 453 454 motor neuron pools of synergist muscles share most of their synaptic input (Laine et al. 2015), the generalizability of this conclusion needs to be examined in a broader array of synergistic 455 muscles. Even among the dorsiflexor muscles where tibialis anterior contributes approximately 456 60% to the applied force (Andreassen & Arendt-Nielsen, 1987), the Q value calculated from the 457 discharge times of those motor units was only moderately associated with the force fluctuations 458 (current study and Negro et al., 2009). Moreover, it is the fluctuations in the net force rather than 459 the activity of a few selected motor units that is more strongly associated with motor 460 performance. 461

The variance in the common input in the current study was quantified using a state-space model, which Feeney et al. (2017) used to model the variance of the common input signal to motor neurons. This parameter differs from the cumulative spike train (CST) estimate, which explains a large part of the fluctuations in the motor output from the spinal cord during isometric contractions (Negro et al., 2009). To enable a more direct comparison with the estimate of the variance derived from the CST, the Q values were filtered to mimic the low-pass filtering influence of muscle on the variance in the common synaptic input to motor neurons (Farina et

469 al., 2014). Consistent with the classic finding of a significant association between CST values and force fluctuations (Negro et al., 2009; Thompson et al., 2018), we found that it was the 470 filtered O values that were significantly correlated for three of the four walking-test results 471 (Table 6). Moreover, the filtered Q values for the plantar flexors emerged as one of the two 472 significant explanatory variables for the distance walked in 6 min by the control group. 473 474 Although the unfiltered Q values for the motor units in the plantar flexors during the two steady contractions (10% and 20% MVC) were greater for the control group, this difference did not 475 explain the variance in performance on the walking tests for either group of participants. Taken 476 477 together, these findings reveal some of the differences between the CST and state-space model measures of variance in common input. It appears that the state-space model may approximate 478 the common input to the motor neurons, whereas the CST more accurately represents the 479 common output of the motor pool. 480

481 Conclusion

482 The time it took individuals with MS to walk 25 ft and the distance they could walk in 6 min were most strongly associated with the fluctuations in force during steady, submaximal 483 484 contractions (force steadiness) with the plantar flexors and the dorsiflexors. In contrast, the 25-ft 485 time for the control group was associated with the strength (MVC torque) of the plantar flexors, 486 whereas their 6-min distance was explained by force steadiness for the dorsiflexors and the 487 variance in common input to the plantar flexors during the steady contractions. These findings 488 indicate that the ability to maintain a constant force during a submaximal, isometric contraction has a greater influence than the strength of lower leg muscles on the walking performance of 489 persons with MS. 490

491

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497 **Disclosures**

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- 499

500 <u>Author contributions</u>

- 501 L.A.D., A.M.A. and R.M.E. conceived and designed research; L.A.D., A.M.A., and M.S.A.
- 502 performed experiments; L.A.D, A.M.A, M.S.A, D.F.F, T.V., A.B., R.M.E interpreted results of
- 503 experiments; L.A.D, M.S.A, and R.M.E. prepared figures; L.A.D. and R.M.E. drafted
- 504 manuscript; L.A.D, A.M.A, M.S.A, D.F.F, T.V., A.B., R.M.E edited and revised manuscript;
- 505 L.A.D, A.M.A, M.S.A, D.F.F, T.V., A.B., R.M.E approved final version of manuscript.
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510 **References**

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688 Figure Legends

Figure 1. A. The experimental setup for measuring muscle strength and force steadiness for the

- 690 dorsiflexors (Above) and the plantar flexors (Below) of the right leg. High-density grid electrodes were
- 691 placed over the tibialis anterior, medial gastrocnemius, and soleus muscles. B. Participant's maintained a
- steady contraction for 30 s at 10% and 20% MVC force. C. Monopolar signals were recorded during
- 693 submaximal, isometric contractions. The high-density grid recordings resulted in bipolar
- 694 electromyographic recordings that were subsequently decomposed into motor unit discharge times. D.
- 695 The discharge times of 9 motor units during a steady, isometric contraction.
- 696 Figure 2. Regression models for fast-walking speed (A, B) and walking endurance (C, D) for the
- 697 participants in the MS (n = 18; A, C) and control (n = 18; B, D) groups. The explanatory variables for
- each model were as follows: A, force steadiness of the plantar flexors at 20% MVC force and
- 699 force steadiness for the dorsiflexors at 10% MVC; B, force steadiness of the dorsiflexors at 20%
- 700 MVC force and the filtered Q values for the plantar flexors; C, force steadiness of the plantar
- flexors at 20% MVC force and force steadiness for the dorsiflexors at 10% MVC; D, MVC
- torque for the plantar flexors of the non-dominant leg and MVC torque for the dorsiflexors of the

703 dominant leg.

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