INTRINSIC MOTOR NEURONE EXCITABILITY IS REDUCED IN SOLEUS AND TIBIALIS ANTERIOR OF OLDER ADULTS

Running title: Intrinsic motor neurone excitability is reduced with ageing.

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ABSTRACT

Age-related deterioration within both motor neurones and monoaminergic systems should theoretically reduce neuromodulation by weakening motor neuronal persistent inward current (PIC) strength. However, this assumption remains untested. Surface electromyographic signals were collected using two 32-channel electrode matrices placed on soleus and tibialis anterior of 25 older adults (70±4 years) and 17 young adults (29±5 years) to investigate motor unit discharge behaviours. Participants performed triangular-shaped plantar and dorsiflexion contractions to 20% of maximum torque at a rise-decline rate of 2%/s of each participant’s maximal torque. Pairwise and composite paired-motor unit analyses were adopted to calculate delta frequency (ΔF) and estimate PIC amplitudes. ΔF has been used to differentiate between the effects of synaptic excitation and intrinsic motor neuronal properties and is assumed to be proportional to PIC amplitude. The results show that soleus and tibialis anterior motor units in older adults had lower ΔFs when calculated with the pairwise (-0.99 and -1.29 pps, respectively) or composite (-1.65 and -2.26 pps, respectively) methods. Older adults’ motor units discharged at lower rates (-2.14 and -2.03 pps, respectively) and were recruited at lower torque levels (-1.50 and -2.06% of maximum, respectively) than young adults. These results demonstrate reduced intrinsic motor neurone excitability during low-force contractions in older adults, likely mediated by decreases in the strength of persistent inward currents. Our findings might be explained by deterioration in the motor neurones or monoaminergic systems, and could contribute to the decline in motor function during ageing; these assumptions should be explicitly tested in future investigations.

Keywords: Persistent inward current; Ageing; Motor unit; HD-EMG; Motoneuron.
KEY POINTS

- The persistent inward current (PIC) is an intrinsic motor neurone excitability property responsible for amplifying excitatory synaptic input into an appropriate motor output. This study compared estimates of PICs in soleus and tibialis anterior between young and older adults.

- Motor units of older adults presented lower estimates of PICs for both soleus and tibialis anterior. Older adults also presented lower motor unit discharge rates and they were recruited at lower torque levels than young adults.

- These results indicate a reduced intrinsic motor neurone excitability in soleus and tibialis anterior motor units, likely mediated by decreases in the strength of persistent inward currents.

- The reduced PIC strength in older adults may result from deterioration within both motor neurones and monoaminergic systems.

- These novel findings contribute to our understanding of the factors underpinning the declines in force production and motor function during ageing.
INTRODUCTION

The age-related loss of force production has been comprehensively described in the literature (Vandervoort, 2002; Orssatto et al., 2018; Suetta et al., 2019), with the physiological alterations affecting force production including changes in several pathways within the nervous system (Aagaard et al., 2010; Manini et al., 2013; Orssatto et al., 2018; Larsson et al., 2019). The motor neurone is an important component of the nervous system affected by ageing as it is responsible for integrating and amplifying excitatory synaptic input into an appropriate motor output (Heckman & Enoka, 2012). An essential intrinsic property of the motor neurone is its capacity to set up persistent inward currents (PICs), which are depolarising currents generated by voltage-sensitive sodium and calcium channels that increase cell excitability by amplifying and prolonging synaptic input (Gorassini et al., 2002; Heckman et al., 2005). Importantly, increases in the concentration of the monoamines serotonin and noradrenaline facilitate PIC development. Under conditions of high monoaminergic drive, synaptic input can be amplified by at least five-fold, suggesting that this amplification is a critical determinant of the motor neurone’s ability to achieve the discharge rates observed during normal motor behaviour (Hounsgaard & Kiehn, 1993; Lee & Heckman, 1999, 2000). Thus, potential physiological alterations in motor neurone intrinsic properties, or in the monoaminergic input to the motor neurone, might reduce the motor neurone’s ability to discharge at higher rates, thus reducing the ability to produce high muscle forces.

During ageing, several changes are observed in both the motor neurones and the monoaminergic system that might potentially reduce PIC strength in older adults, including lower discharge rates (Orssatto et al., 2020; Kirk et al., 2021), reduced incidence of doublet discharges (Christie & Kamen, 2006), and an increased afterhyperpolarisation duration (Piotrkiewicz et al., 2007). These changes are consistent with the lower motor neurone excitability that is also observed in aged rat models (Morales et al., 1987). With respect to the monoaminergic system, research using both human and animal models suggests that ageing is associated with reduced noradrenaline and serotonin secretions and thus input onto the motor neurones (Johnson et al., 1993; Ko et al., 1997; Míguez et al., 1999; Shibata et al., 2006; Liu et al., 2020), which might theoretically underpin PIC reduction with ageing. These findings indicate the possibility that PIC strength might be reduced in older adults; however, this hypothesis remains to be tested.

The strength of PICs that are initiated close to the recruitment threshold of motor neurones can be estimated in humans using the paired motor unit technique (Gorassini et al.,
2002; Powers et al., 2008; Stephenson & Maluf, 2011), with data obtained using high-density surface electromyography (Holobar & Zazula, 2007; Enoka, 2019). This technique requires the pairing of the discharge rates of a low-threshold (control unit) to a higher-threshold (test unit) motor unit, obtained during a slowly-increasing and decreasing triangular-shaped contraction (Gorassini et al., 2002; Powers et al., 2008; Orssatto et al., 2021). Subsequently, the difference in discharge rate of the control unit between the time of recruitment and de-recruitment of the test unit is computed as the change in frequency ($\Delta F$). $\Delta F$ has been used to differentiate between the effects of synaptic excitation and motor neuronal intrinsic properties and is assumed to be proportional to PIC strength (Afsharipour et al., 2020). With these techniques, it is now possible to estimate and compare the strengths of these PICs in motor units of young and older adults.

The present study compared $\Delta F$s to estimate differences in PIC strength of soleus and tibialis anterior motor units between young and older adults. Additionally, we explored the relationship between $\Delta F$ and the peak discharge rates. We hypothesised that there would be a reduction in PIC in both soleus and tibialis anterior, and that $\Delta F$ would be strongly associated with motor unit peak discharge rate. Soleus and tibialis anterior were selected for study because the control and timing of their activation are both critical to the performance of daily activities such as standing and walking in older adults (Polcyn et al., 1998; Laughton et al., 2003).

METHODS

Participants and ethical procedures

Forty-four participants volunteered for the study, including 18 young adults and 26 older adults (participant characteristics are documented in Table 1). More older adults were recruited because it was expected that fewer motor units would be identified during decomposition and that some participants may not be able to perform the triangle-shaped contractions with the necessary torque rise and fall accuracy. One participant per group was excluded from the study because no motor units were identified in either soleus or tibialis anterior. To participate, volunteers had to be free from neurological and lower limb musculoskeletal injuries and disorders. Participants were excluded if they were on medications that could influence the monoaminergic system, such as serotonin or noradrenaline (e.g., beta-blockers and serotonin reuptake inhibitors) or consumed caffèinated foods (e.g., coffee). They were also excluded from the analyses if: a) no usable motor units were identified by the decomposition algorithm, or b) if it was not possible to achieve all the
assumptions required in the paired motor unit analysis (as described below) for either *soleus* or *tibialis anterior*. The study was approved by the University Human Research Ethics Committee, and all participants gave written informed consent before participating. Data collection was conducted during the COVID-19 pandemic and all safety procedures followed the local state government policies.

**Table 1.** Participant characteristics.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Young Adults (n = 17)</th>
<th>Older Adults (n = 25)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>29 ± 5</td>
<td>70 ± 4</td>
</tr>
<tr>
<td>Sex (n, %)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td>9 (53%)</td>
<td>11 (44%)</td>
</tr>
<tr>
<td>Women</td>
<td>8 (47%)</td>
<td>14 (56%)</td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td>71.9 ± 13.6</td>
<td>77.8 ± 19.3</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>174 ± 11</td>
<td>163 ± 17</td>
</tr>
<tr>
<td>Peak torque (N·m)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plantar flexion</td>
<td>156 ± 47</td>
<td>85 ± 32</td>
</tr>
<tr>
<td>Dorsiflexion</td>
<td>41 ± 14</td>
<td>29 ± 7</td>
</tr>
<tr>
<td>Physical activity level (MET min/week)</td>
<td>2541 (1902–3039)</td>
<td>2994 (1386–5139)</td>
</tr>
</tbody>
</table>

Note: Data are presented as mean ± standard deviation, except for Physical activity level, which is presented as median with interquartile range.

**Study design and neuromuscular testing procedures**

Participants visited the laboratory on a single occasion in which they were familiarised with the testing procedures and data were collected. Initially, participants signed the informed consent and completed the International Physical Activity Questionnaire (Hagströmer *et al.*, 2006). After electrode placement on *soleus* and *tibialis anterior*, the participants were seated upright in the chair of an isokinetic dynamometer (Biodex System 4, Biodex Medical system, Shirley, NY) with the knee fully extended (0°) and ankle in the anatomical position (0°). A warm-up consisting of six 5-s submaximal voluntary isometric plantar and dorsiflexion contractions (2 × 30%, 2 × 60%, and 2 × 80% of perceived maximal effort) was performed, followed by three maximal voluntary contractions of ~3-s with 30-s rest intervals. The maximum torque achieved was recorded as the maximal voluntary
contraction peak torque. Subsequently, participants were familiarised with the triangular-shaped contractions to 20% of their maximum voluntary torque level. Triangular contractions to 20% of maximal torque have been extensively used for ΔF calculations using the paired motor unit technique (Gorassini et al., 2002; Afsharipour et al., 2020; Hassan et al., 2020; Kim et al., 2020; Trajano et al., 2020), and this force was considered similar to the maximum torques developed during daily activities such as standing and walking. All contractions had a duration of 20 s (10-s up and 10-s down) and were performed at a rate of torque increase and decrease of ~2%/s. Participants were instructed to follow the torque path provided in real time on a 58-cm computer monitor during each contraction. Data collection commenced 5 min after the end of familiarisation (usually requiring ~3-10 × 20% triangular contractions with 30-s rest), during which the participants then performed four triangular contractions with 60-s rest intervals. When an abrupt increase or decrease in torque was observed (i.e. the torque trajectory was not closely followed), the trial was excluded and repeated. The maximum voluntary isometric torque and order of triangular contraction completion was randomised between soleus and tibialis anterior.

**Surface electromyography**

Surface electromyograms (sEMG) were recorded during the 20% triangular contractions using four semi-disposable 32-channel electrode grids with a 10-mm interelectrode distance (ELSCH032NM6, OTBioelettronica, Torino, Italy). After skin shaving, abrasion, and cleansing with 70% isopropyl alcohol, two electrode grids were placed over the medial and lateral portions of soleus (either side of the Achilles tendon) and another two electrode grids were placed over the superior and inferior aspect of tibialis anterior using a bi-adhesive foam layer and conductive paste (Ten20, Weaver and Company, Colorado, USA). A strap electrode (WS2, OTBioelettronica, Torino, Italy) was dampened and positioned around the ankle joint as a ground electrode. The sEMG signals were acquired in monopolar mode, amplified (256×), band-pass filtered (10–500 Hz), and converted to a digital signal at 2048 Hz by a 16-bit wireless amplifier (Sessantaquattro, OTBioelettronica, Torino, Italy) using OTBioLab+ software (version 1.3.0., OTBioelettronica, Torino, Italy) before being stored for offline analysis.

**Motor unit analyses**

*Motor unit identification*
The recorded data were processed offline using the DEMUSE software (Holobar & Zazula, 2007). For each contraction intensity, only the triangular contraction yielding the lowest deviation from the torque trajectory was analysed. If both contractions presented a similar torque trajectory, the contraction with the highest number of identified motor units was analysed. High-density sEMG signals were band-pass filtered (20-500 Hz) with a second-order, zero-lag Butterworth filter. Thereafter, a blind source separation method, the convolutive kernel compensation (CKC) method, was used for signal decomposition (Holobar & Zazula, 2007; Holobar et al., 2014) from each triangular contraction. CKC yields the filters of individual motor units (so-called motor unit filters) that, when applied to high-density sEMG signals, estimate the motor unit spike trains (Holobar & Zazula, 2007; Holobar et al., 2014). After decomposition, a trained investigator manually inspected motor unit spike trains and edited the discharge patterns of the motor units. Only the motor units with a pulse-to-noise ratio equal to or greater than 30 dB were kept for further analysis (Holobar et al., 2014).

*Estimation of PIC amplitude (ΔF) and peak discharge rate*

The observed discharge events for each motor unit were converted into instantaneous discharge rates and fitted into a 5th-order polynomial function. The maximum value obtained from the polynomial curve was considered the peak discharge rate. Thereafter, PIC amplitude was estimated using the paired motor unit analysis (Gorassini et al., 2002), referred through the manuscript as pairwise method. Motor units with a low recruitment threshold (i.e., control units) were paired with higher recruitment threshold motor units (i.e., test units). ΔF was calculated as the change in discharge rates of the control motor unit from the moment of recruitment to the moment of de-recruitment of the test unit (Gorassini et al., 2002; Heckman et al., 2005). In order to produce motor unit pairs, the following criteria were adopted: 1) rate-to-rate correlations between the smoothed discharge rate polynomials of the test and control units was $r \geq 0.7$; 2) test units were recruited at least 1.0 s after the control units; and 3) the control unit did not show discharge rate saturation after the moment of test unit recruitment (>0.5 pps) (Gorassini et al., 2002; Udina et al., 2010; Vandenberk & Kalmar, 2014; Binder et al., 2020; Hassan et al., 2020). ΔFs obtained for each pair were averaged to obtain a single ΔF for each motor unit.

We also conducted an additional analysis using the composite paired motor unit method to calculate ΔF values for each motor unit (Afsharipour et al., 2020). This method is characterised by the overlay of 3 lower-threshold motor units to construct a single composite...
control unit profile to be paired with the test units. The composite method has been suggested to address some of the limitations observed with the pairwise method, such as underestimation and overestimation of $\Delta F$ values, reducing its variability. However, strict assumptions are made for eligible motor units to be included in the analysis, which allows its use only in muscles in which it is possible to identify many motor units (e.g., *tibialis anterior*) but not in those in which fewer motor units are identified using the decomposition method (e.g., *soleus*). Moreover, this method does not allow calculation of $\Delta F$ values for lower-threshold motor units since they are used to construct the composite control unit. In summary, the control unit is the overlay of at least three motor units recruited at <3% of the maximum voluntary torque and presenting a similar discharge profile. It was assumed that lower-threshold motor units had their discharge rate profile more linearly related to the synaptic input profile because PICs were almost fully activated at the time of recruitment; therefore, it would be more appropriate to use the composite unit as control. Also, this method requires the removal of the acceleration phase of the discharge rates (i.e., secondary range), making the polynomial ascending-to-descending slope ratio near 1, which is important when measuring PIC amplitudes to avoid any rate-dependent effects on motor unit recruitment or derecruitment (Desmedt & Godaux, 1977). The test units should start discharging in the tertiary range (i.e., after the secondary range and before the descending phase). This method provides a single $\Delta F$ for each motor unit, which was then used in the data analysis. Figure 1 illustrates the pairwise and composited paired motor unit analysis methods on *tibialis anterior* motor units for one participant per group. Panels C and D display the test units used for both methods. Panels E and F display the control units for the pairwise method and panels G and H display the control units for the composite method.
Figure 1. Data from a single participant for each group showing torque during triangle-shaped contractions and delta frequency (ΔF) calculation in tibialis anterior for both the pairwise and composite paired motor unit analyses. Data from a young adult is displayed in the left panels and from an older adult in the right panels. Panels A and B on the first row show the torque traces for contractions with 20% of the participant’s maximal voluntary
torque. The participants’ test units are displayed on panels C and D (purple motor unit),
control units for the pairwise method on panels E and F (red motor unit), and control units for
the composite method on panels G and H (red, green, and blue motor units). The black
continuous lines are the 5th-order polynomial fits for the control units. Note that for the
composite method the polynomial curve starts from the tertiary range. The gray-shaded areas
represent the ΔF amplitude for each participant and analysis.

Data analysis

All analyses were undertaken in R (version 4.0.3) using RStudio environment
(version 1.3.1093). Models were fitted using the lmerTest package (Kuznetsova et al., 2017).
A linear mixed-effects model was used to compare estimates of ΔF of soleus and tibialis
anterior motor units between young and older adults. The model included: age group, muscle
type, recruitment threshold, age group by muscle type, and muscle type by recruitment
threshold as fixed effects. A random intercept and slope (recruitment threshold by muscle
type) was included for each participant in the study, to account for the correlation between
repeated observations on each individual. This model was selected from a series of candidate
models (Supplement 1), based on the smallest Bayesian Information Criteria value. The
recruitment threshold was standardised (mean = 0, SD = 1) before analysis.

Separate linear mixed-effects models were used to analyse peak discharge rate and
recruitment threshold data. These models included: age group, muscle type and age group by
muscle type as fixed factors; and a random intercept and slope (muscle group) for each
participant. The estimated marginal mean difference and 90% and 95% confidence intervals
(CI) in ΔF, peak discharge rate, and recruitment threshold between young and older adults,
were determined using the emmeans package (Lenth et al., 2021). The standardised
difference, denoted \(d\), was also calculated using the population SD from each respective
linear mixed-effects model as the denominator (Lenth et al., 2021).

The association between ΔF and peak discharge rate was determined by fitting two
models: (1) a base model including: age group and muscle type as fixed factors, and a
random intercept and slope (muscle type) for each participant, and (2) a second model
including all covariates from the base model, in addition to ΔF (standardised; mean 0, SD 1).
The difference in the variance explained by the fixed effects (R²) between the models was
calculated (Nakagawa et al., 2017) to isolate the contribution of ΔF. The models were also
compared to determine whether there was a reduction in the residual sum of squares with the
inclusion of $\Delta F$, which is an indication of a better fitting model (Chambers, 1992). Differences between young and older adults in peak plantar flexion and dorsiflexion torque, and physical activity levels, were determined using independent t-tests. The $\alpha$ level for all tests was 5%. The dataset and R code can be found at https://github.com/orssatto/PICs-ageing.

RESULTS

Effects of age and muscle group on $\Delta F$, peak discharge rate, and recruitment threshold

Motor units of older adults had lower $\Delta$Fs, discharged at lower rates, and were recruited at lower torque (muscle force) levels than young adults in both soleus and tibialis anterior. Also, $\Delta F$ levels and peak discharge rates were lower in soleus than tibialis anterior, independent of age.

There were effects of age ($\beta = -0.99$, 95% CI = -1.49, -0.50; $p < .001$) and muscle ($\beta = 1.29$, 95% CI = 0.65, 1.91; $p < .001$) but no age group by muscle effect ($\beta = -0.29$, 95% CI = -1.08, 0.55; $p = .45$), on $\Delta F$ when calculated using the pairwise paired motor unit method. $\Delta F$ was lower in older adults (Figure 2A) in both soleus ($d = -1.50$; Figure 3A) and tibialis anterior ($d = -1.42$; Figure 3A). There were effects of age ($\beta = -2.14$, 95% CI = -2.98, -1.33; $p < .001$) and muscle ($\beta = 3.71$, 95% CI = 2.64, 4.83; $p < .001$) on peak discharge rate, but there was no age by muscle effect on peak discharge rate ($\beta = 0.11$, 95% CI = -1.37, 1.56; $p = .88$). Peak discharge rate was lower in older adults (Figure 2B) in both soleus ($d = -0.98$; Figure 3B) and tibialis anterior ($d = -1.27$; Figure 3B). There was an age effect on recruitment threshold, with thresholds lower in older adults ($\beta = -1.50$, 95% CI = -2.89, -0.12; $p = .040$; Figure 3C). There was no evidence of muscle ($\beta = 1.12$, 95% CI = -0.35, 2.65; $p = .15$) or age by muscle ($\beta = 1.12$, 95% CI = -2.59, 1.38; $p = .58$) effects on recruitment threshold.

The results of the additional, composite method, analysis revealed there was effects of age ($\beta = 1.65$, 95% CI = -2.54, 0.77; $p < .001$) and muscle ($\beta = 0.82$, 95% CI = 0.22, 1.42; $p = .008$) on $\Delta F$, but no age by muscle effect on $\Delta F$ ($\beta = 0.61$, 95% CI = -1.43, 0.21; $p = .150$). $\Delta$Fs were lower in older adults in both soleus ($d = -2.14$) and tibialis anterior ($d = -2.93$), as shown in Figure 4.
Figure 2. \(\Delta F\) calculated with the pairwise paired motor unit method (A), peak discharge rate (B), and recruitment threshold (C) in *soleus* and *tibialis anterior* in young and older adults.
The mean (circle) and 95% confidence interval are offset to the right, with individual data points coloured by participant. pps = peaks per second.
Figure 3. The marginal mean difference with 90% (thick line) and 95% (thick line) confidence intervals between young and older adults for \( \Delta F \) calculated with the pairwise paired motor unit method (A), peak discharge rate (B), and recruitment threshold (RT) (C) in soleus and tibialis anterior. Negative values on all panels indicate the lower responses in older adults. There was an effect of age on \( \Delta F \), peak discharge rate, and recruitment threshold, with all variables lower in older adults. There was no age by muscle effect on \( \Delta F \), peak discharge rate, or recruitment threshold—indicating that age-dependent differences in these variables were similar for both muscles.
Figure 4. ΔF calculated with the composite paired motor unit method (A), and the marginal mean difference with 90% (thick line) and 95% (thick line) confidence intervals between young and older adults for ΔF (B). In panel A, the mean (circle) and 95% confidence interval are offset to the right, with individual data points coloured by participant. In panel B, negative values indicate ΔF was lower responses in older adults. pps = peaks per second.
Contribution of ΔF to peak discharge rate

There was a small, significant improvement in model fit when ΔF was included as a covariate in the prediction of peak discharge rate \( (X^2 = 10.444, \ p = .001) \). The inclusion of ΔF increased the \( R^2 \) by 0.017, from 0.445 (i.e., base model without ΔF) to 0.462. There was a statistical effect of ΔF on peak discharge rate \( (\beta = 0.28, 95\% \ CI = 0.11, 0.45, \ p = .001) \), in addition to effects of age \( (\beta = -1.90, 95\% \ CI = -2.68, -1.14, \ p < .001) \) and muscle \( (\beta = 3.54, 95\% \ CI = 2.83, 4.28, \ p < .001) \).

Peak torque and physical activity levels

Peak torques and physical activity levels are shown in Table 1. Peak torque was lower in older adults for both plantar flexion (mean difference = -70 N·m, 95% CI = -95, -46; \( p < .001; \ d = -1.83 \)) and dorsiflexion (mean difference = -12 N·m, 95% CI = -18, -5; \( p < .001; \ d = -1.18 \)). Physical activity levels were not statistically different between young and older adults (mean difference = 594 MET min/week, 95% CI = -1269, 2457; \( p = .52; \ d = 0.20 \)).

Motor unit identification

The number of motor units identified in soleus was 113 for the young group and 211 for the older group. The median (IQR) number per younger participant was 7 (5–8) and per older participant 8 (6–9). For tibialis anterior, 293 motor units were identified in the young group and 411 in the older group. The median number per younger participant was 19 (16–23) and per older participant 20 (16–27). For the pairwise paired motor unit analysis, it was possible to obtain ΔF values from 16 young adults and 23 older adults in soleus, and from 16 young adults and 19 older adults in tibialis anterior. The number of test unit ΔFs for soleus in the young group was 70 and in the older group was 117. The median number per young participant was 5 (3–6) and per older participant was 4 (3–6). The number of test unit ΔFs for tibialis anterior in the young group was 185 and in the older group was 257. The median number per younger participant was 11 (8–14) and per older participant was 14 (9–18).

For the composite paired motor unit analysis, it was possible to obtain ΔF values from 4 young adults and 7 older adults for soleus and from 15 young adults and 18 older adults for tibialis anterior. The number of test unit ΔFs for soleus in the young group was 12 and in the older group was 17, with the median number per younger participant being 3 (3–4) and per older participant being 2 (1–4). The number of test unit ΔFs for tibialis anterior in the young
group was 93 and in the older group was 111, with the median number per young participant being 5 (4–8) and per older participant being 6 (3–8). No motor units were identified in one young participant and one older participant in soleus, and in one young participant and six older participants in tibialis anterior, respectively. These participants were not included in the analyses for the respective muscles.

DISCUSSION

The present study compared strengths of motor neuronal PICs, estimated using the paired motor unit technique (ΔF), between young and older adults. The main findings indicate that ΔF values are considerably lower in both soleus and tibialis anterior in older adults. These reductions are accompanied by lower peak discharge rates and the recruitment of motor units at lower torque levels in the older than younger adults in both muscles. As an exploratory analysis, a model was developed that showed a small contribution of ΔF to the between-subject variance in peak discharge rates after accounting for the effects of age, muscle. These findings point to a meaningful reduction in the intrinsic motor neurone excitability in older adults that is at least partly associated with a reduced amplitude of persistent inward currents triggered by muscle recruitment.

Estimates of persistent inward currents (ΔF)

The ΔF values obtained from soleus and tibialis anterior in young adults are similar to those obtained in previous studies using triangular-shaped contractions with a 2%/s force increase-decrease rate (Afsharipour et al., 2020; Kim et al., 2020; Trajano et al., 2020) and indicate that ΔFs from older adults are substantially lower than in young adults in both soleus and tibialis anterior. These differences can be a result of the detrimental changes within the monoaminergic systems, which indicate lower release of serotonin and noradrenaline with ageing. Locus coeruleus and the dorsal raphe nucleus are the major central sources of noradrenaline and serotonin, respectively (Cassano et al., 2009). Diminution of structural integrity (Liu et al., 2019, 2020) and in neuromelanin (a product of noradrenaline synthesis) content in noradrenergic neurones emanating from locus coeruleus (Shibata et al., 2006) indicate impaired noradrenaline secretion in older adults. Moreover, noradrenaline and serotonin concentration decrements have been observed in the brains of aged rats (Ko et al., 1997; Míguez et al., 1999), and degeneration of serotonergic axons projecting to the ventral horn of the lumbar segment of the spinal cord (where motor neurones innervating lower limb muscles emanate) have been detected (Johnson et al., 1993). Also, serotonin receptors are
affected by expanded circulation of cytokines, resulting in increased re-uptake of serotonin (Michaud et al., 2013). Thus, older adults might speculatively present reduced noradrenaline and serotonin secretions, and hence input onto motor neurones, which would then impair the initiation and modulation of PICs in this population. Changes in motor neurone integrity might also partly underpin reductions in intrinsic excitability. During ageing, axonal demyelination due to reduced expression of proteins responsible for myelination (Pannese, 2011) as well as axonal atrophy and degeneration have been observed, possibly subsequent to deregulated Ca\(^{2+}\) homeostasis (Nikoletopoulou & Tavernarakis, 2012) and to toxic, metabolic, or infectious injury sustained throughout the lifespan, or due to high levels of chronic inflammation and oxidative stress (Misgeld, 2011; Selman et al., 2012). Motor neuronal death, especially in higher-threshold motor axons, leading to denervation of motor units has also been documented (Gordon et al., 2004). In these cases, denervated motor units may remodel through reinnervation by nearby lower-threshold motor neurones (Manini et al., 2013; Piasecki et al., 2016), which may explain the reduced recruitment threshold observed in the older adults (as discussed below). These detrimental alterations in motor neurone structure are associated with reduced Ca\(^{2+}\)-mediated plateau potential durations in striatal neurones (from aged rats) (Dunia et al., 1996), slower conduction velocity of efferent motor axons (Di Iorio et al., 2006), lower incidence of doublet discharges, slower maximum discharge rates (Klass et al., 2008; Orssatto et al., 2020; Kirk et al., 2021) alongside an increased afterhyperpolarisation duration (Piotrkiewicz et al., 2007), and, as evidenced in the present study, lower PIC strength in the motor neurones. It is a logical hypothesis that the changes within the monoaminergic system and motor neurone structural integrity would possibly explain the reduced $\Delta$Fs observed in the present study. However, our findings indicate the need for further study of the cause-effect relationship between these mechanisms and reduced PICs in humans.

PICs are highly sensitive to synaptic inhibition, and both reciprocal and recurrent inhibition directly influence intrinsic motor neurone excitability, being effective PIC deactivators by opposing the facilitatory effects on PICs of descending brainstem neuromodulatory systems (Kuo et al., 2003; Hyngstrom et al., 2007, 2008; Bui et al., 2008a, 2008b). It would therefore be expected that the reduced PIC strength estimates observed in the older adults in the present study might result from an age-dependent increase in reciprocal and/or recurrent inhibition. However, older adults present reduced reciprocal inhibition from the common peroneal nerve onto soleus and from the tibial nerve onto the tibialis anterior (Kido et al., 2004). Also, recurrent inhibition onto the soleus motor neurone pool is suspected
to be relatively unaffected by ageing (Chalmers & Knutzen, 2004). Therefore, a reasonable supposition is that the behaviour of spinal inhibitory/excitatory systems (i.e., reduced reciprocal inhibition and preserved recurrent inhibition) of older adults are unlikely to be responsible for the decreases in PIC strength.

**Recruitment threshold**

Motor units that were identified in the older adults showed a lower recruitment threshold and were thus recruited at a lower torque level than in young adults. This behaviour may result from a compression in the range of firing rates observed previously in older adults motor units (Barry et al., 2007; Girts et al., 2020). Since motor unit discharge rate modulation in response to force changes is impaired in older adults, additional motor units must be recruited earlier in a triangular-shaped contraction to continually increase muscle force (Barry et al., 2007; Enoka & Duchateau, 2017). Additionally, the observed lower recruitment thresholds might reflect the recruitment of a greater relative number of smaller motor neurones during the task. Older adults have a greater proportion of lower-threshold motor units as a result of motor unit remodelling subsequent to motor neuronal denervation, so motor units previously innervated by higher-threshold motor units become reinnervated by lower-threshold motor units (Piasecki et al., 2016), as described above. In summary, our data indicate that aged motor neurones have a constrained capacity to amplify the excitatory synaptic input, consequently demanding an earlier recruitment of additional motor units to achieve the required motor output.

It is important to note that Ca\textsuperscript{2+} PIC channels can be activated below the action potential threshold (i.e., subthreshold PICs), strongly influencing motor unit recruitment (Li et al., 2004). The possibility exists that the lower recruitment threshold of motor units of the older adults observed in the present study reflects a higher motor neurone excitability (i.e. enhanced motor neurone recruitment for a given input) because of stronger subthreshold PIC activation. An increase in the proportion of subthreshold PICs would reduce the recruitment threshold while also decreasing $\Delta F$ values. Subthreshold PICs may be stronger in smaller motor neurones, resulting in a larger overall (sub- plus supra-threshold) PIC strength (Afsharipour et al., 2020). However, the $\Delta F$ method only estimates the suprathreshold contribution of the PICs to the discharge behaviour of motor units. Consequently, it is not possible to ascertain the behaviour of subthreshold PICs using the $\Delta F$ method. Our analytical approach accounted for any effect of recruitment threshold (and possibly motor neurone size).
on $\Delta F$, that may otherwise confound any age-related effect, by including recruitment threshold in our modelling of $\Delta F$. Further, the reanalysis of $\Delta F$ using the composite paired motor unit method appeared to remove the effect of recruitment threshold on $\Delta F$ values (from $\beta = 0.27$ to $\beta = -0.05$), which is an expected characteristic of the method (Afsharipour et al., 2020). Therefore, the reduced $\Delta F$s observed in older adults in the present study is not likely to have been an artefact of differences in motor neurone recruitment thresholds.

**Peak discharge rates**

The lower $\Delta F$ values in the older adults were also accompanied by reduced motor neurone discharge rates. The PIC is an important modulator of discharge rate output (Gorassini et al., 2002; Orsatto et al., 2021) and the available monoamines, serotonin and noradrenaline, facilitate PICs to increase motor neuronal gain and alter the input-output relationship according to the required output (Heckman & Binder, 1991; Heckman, 1994; Powers et al., 2008; Powers & Heckman, 2015; Huh et al., 2017; Orssatto et al., 2021). PICs can amplify synaptic input by more than five-fold, and are thus a determinant mechanism influencing the capacity for motor neurones to achieve the necessary discharge rates to obtain very high muscle activation levels (Hounsgaard & Kiehn, 1993; Lee & Heckman, 1999, 2000). We have recently shown that increases in discharge rate during triangular-shaped contractions at different force levels were strongly associated with $\Delta F$ increases, and thus PIC amplitudes, using a within-subject design (Orssatto et al., 2021). Nonetheless, data from the current study showed that $\Delta F$ explained only 2% of the between-subject variance in discharge rate output. Therefore, it may be case that $\Delta F$ explains more of the within-subject than between-subject differences in motor unit discharge rates. It is important to mention that motor unit discharge rates are not only modulated by PICs. Evidence in both animal (Chase et al., 1985; Maxwell et al., 2018) and human models (Rowe et al., 2006) suggests that the synaptic input into the motor neurone decreases with ageing, which may result from increases in intracortical inhibition or reduced intracortical facilitation in older adults (Todd et al., 2003; McGinley et al., 2010; Opie et al., 2020).

**Strengths and limitations**

The main strength of our study was the use of two validated (Gorassini et al., 2002; Powers et al., 2008; Afsharipour et al., 2020) and a widely used methods to estimate PIC
strength in humans (Gorassini et al., 2002; Powers et al., 2008; Udina et al., 2010; Hassan et al., 2019, 2020; Trajano et al., 2020; Orssatto et al., 2021); however, both methods have limitations that should be pointed out. The pairwise method (Gorassini et al., 2002) allowed us to obtain several pairs of motor units, having a larger amount of test units per participant. On the other hand, this method present a higher variance as a result of under and overestimation of ΔFs as a consequence of adopting control units with varied recruitment thresholds (Afsharipour et al., 2020). Recently, Afsharipour et al., (2020) proposed the use of a composite control unit to reduce the ΔF variance present in the conventional paired motor unit analysis. However, this method requires some additional assumptions, such as the overlay of three lower threshold motor units with a similar discharge rate profile and with recruitment threshold below 3%. When following these assumptions, there was an important reduction of motor unit pairs, particularly for soleus. Therefore, we initially adopted the pairwise method as it permitted comparison between soleus and tibialis anterior ΔFs. We also ran the composite method as an additional analysis to examine whether reducing ΔF variability and removing the influence of the recruitment threshold on ΔF obtained with the pairwise method would affect our main outcomes. Using both methods allowed us to identify a large difference in ΔF between young and older adults, minimising the methodological limitations. Furthermore, ΔF values obtained in our study are derived from lower threshold soleus and tibialis anterior motor units recruited at a low force level (20% of peak torque); this is a commonly-used force target and is also similar to forces that might be expected in daily activities such as standing and walking. Motor neurones from distinct muscles depict different discharge behaviours during ageing (Kirk et al., 2021); thus, it is possible that ΔF behaviour might also differ. Furthermore, there is evidence that the function and structure of higher threshold motor neurones are more affected than lower threshold motor neurones (Orssatto et al., 2018). Therefore, we recommend that future studies investigate the effect of ageing on ΔF values from different muscles and at different contraction intensities.

**Conclusions**

The present study provides novel evidence of reduced intrinsic motor neurone excitability in humans by estimating PIC strengths using the paired-motor unit analyses. Older adults had substantially lower ΔFs, and presumably PIC strengths, in both soleus and tibialis anterior than young adults with matched physical activity level. This would likely influence the capacity of older individuals to activate the muscles, thus requiring a greater descending drive from cortical areas and hence level of volitional effort, and greater number
of recruited motor units to achieve the same force level (relative to maximum). We also identified a small contribution of $\Delta F$ to the between-subject variability in peak discharge rates. The present findings contribute to our understanding of the effects of ageing on motor neurone excitability, which is a potential mechanism underpinning motor functional loss during ageing; this hypothesis should be explicitly tested in future studies. A logical next step is to examine the effect of ageing on monoaminergic projections onto the motor neurones and their relationship with the reduced PIC strength observed in the present study.

ADDITIONAL INFORMATION

Data availability
The dataset and R code are available at https://github.com/orssatto/PICs-ageing.

Competing interests
The authors declare no competing interest related to this manuscript.

Author contributions
LBRO, AJS, AJB, and GST contributed with the conception and design of the work. LBRO acquired data. RLS developed the MATLAB script for $\Delta F$, peak discharge rate, and recruitment threshold calculation. LBRO conducted the biological signals data analyses, and DNB developed the R script and conducted statistical analyses. All authors interpreted and discussed the data, drafted the manuscript, and revised it critically providing important intellectual content.

All authors approved the final version of the manuscript; agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved; and qualify for authorship, and all those who qualify for authorship are listed.

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