Original Article

Chronic training status affects muscle excitation of the vastus lateralis during repeated contractions

Sunggun Jeon a, Stephanie A. Sontag a, Trent J. Herda b, Michael A. Trevino a,∗

a Applied Neuromuscular Physiology Laboratory, Department of Kinesiology, Applied Health, and Recreation, Oklahoma State University, Stillwater, OK, USA
b Neuromechanics Laboratory, Department of Health, Sport, and Exercise Sciences, The University of Kansas, Lawrence, KS, USA

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ABSTRACT

This study examined electromyographic amplitude (EMG RMS)-force relationships during repeated submaximal knee extensor muscle actions among chronic aerobically-(AT), resistance-trained (RT), and sedentary (SED) individuals. Fifteen adults (5/group) attempted 20 isometric trapezoidal muscle actions at 50% of maximal strength. Surface electromyography (EMG) was recorded from vastus lateralis (VL) during the muscle actions. For the first and last successfully completed contractions, linear regression models were fit to the log-transformed EMG RMS-force relationships during the linearly increasing and decreasing segments, and the b terms (slope) and a terms (antilog of y-intercept) were calculated. EMG RMS was averaged during steady force. Only the AT completed all 20 muscle actions. During the first contraction, the b terms for RT (1.301 ± 0.197) were greater than AT (0.910 ± 0.123; p = 0.008) and SED (0.912 ± 0.162; p = 0.008) during the linearly increasing segment, and in comparison to the linearly decreasing segment (1.018 ± 0.139; p = 0.014), respectively. For the last contraction, the b terms for RT were greater than AT during the linearly increasing (RT = 1.373 ± 0.353; AT = 0.883 ± 0.123; p = 0.018) and decreasing (RT = 1.526 ± 0.328; AT = 0.970 ± 0.223; p = 0.010) segments. In addition, the b terms for SED increased from the linearly increasing (0.968 ± 0.144) to decreasing segment (1.268 ± 0.126; p = 0.015). There were no training, segment, or contraction differences for the a terms. EMG RMS during steady force increased from the first- ([64.08 ± 51.68] μV) to last-contraction ([86.73 ± 49.55] μV; p = 0.001) collapsed across training statuses. The b terms differentiated the rate of change for EMG RMS with increments in force among training groups, indicating greater muscle excitation to the motoneuron pool was necessary for the RT than AT during the linearly increasing and decreasing segments of a repetitive task.

INTRODUCTION

Surface electromyography (EMG) is the recording of myoelectric signals from the skin overlying active motor units (MUs),1,2 The time domain (amplitude) of the EMG signal is influenced by both the recruitment of MUs and their firing rates.3 Thus, surface EMG can provide information regarding MU control strategies.

It is well documented training elicits adaptations that are dependent on the mode of exercise.4–7 Numerous studies have reported acute and chronic aerobic- (AT) and resistance-training (RT) alters MU: firing rates,5–8 synchronization,9,10 recruitment thresholds,11,12 excitability,12,14 and MU pool output13,15 during submaximal voluntary contractions. Therefore, it is plausible that EMG amplitude (EMG RMS) would be sensitive to alterations in MU control strategies as a function of chronic training status; however, previous literature has been mixed. For example, Herda et al.16 reported no differences in the b terms (slopes) calculated from the natural log-transformed EMG RMS-force relationships during a linearly increasing muscle action up to 90% maximal voluntary contraction (MVC) among chronic training statuses (AT, RT, sedentary [SED]) with known fiber area differences for the vastus lateralis (VL). Conversely, Trevino and Herda17 reported b term differences among chronic training statuses for an isometric trapezoidal muscle action at 60% MVC that included a linearly increasing, steady force, and linearly decreasing segment. Therefore, the ability to differentiate muscle excitation to the motoneuron pool among chronic training statuses may depend on the targeted intensity and muscle action being examined.

Chronic training also elicits structural alterations; such as greater type I- and type IIA-heavy chain (MHC) expression for AT16,18 and RT-individuals,19,20 respectively, which would influence the

* Corresponding author. Applied Neuromuscular Physiology Laboratory, Department of Health and Human Performance, Oklahoma State University, 191 Colvin Recreation Center, Stillwater, OK, 74074, USA.
E-mail address: michael.a.trevino@okstate.edu (M.A. Trevino).

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fatigability and twitch forces of the MU pool. During fatiguing contractions, increases in MU recruitment and firing rates in conjunction with decreases in recruitment thresholds have been observed. In addition, there is evidence suggesting that muscle excitation to the motoneuron pool will adjust in response to MU twitch forces when producing a desired force output. Thus, examining the EMGRMS patterns of responses among AT, RT, and SED during a fatigue inducing task may indicate chronic training related specific adjustments in MU control strategies (MU recruitment and/or firing rates) when maintaining a targeted force. In addition, investigating MU control strategy responses during fatigue as a result of specific chronic training may provide information for strength and conditioning coaches and clinicians that allows for better exercise and rehabilitation programming, respectively. Furthermore, investigating a series of isometric trapezoidal muscle actions that includes a linearly increasing, steady force, and decreasing segment, mimics the MU activation, force maintenance, and MU deactivation-strategies performed during the cyclical movement patterns that humans perform during activities of daily living more so than a single, sustained contraction to failure. However, we are aware of only one study that has examined the influence of chronic training status on the EMGRMS patterns of response for the VL during a fatiguing isometric muscle action. Beck et al. reported no differences for absolute and normalized EMGRMS between chronic AT and RT individuals with known MHC expression differences during a 30 s step contraction of the knee extensors at 50% MVC. Recently, our group reported greater fatigability for chronic RT and SED individuals in comparison to AT, which was associated with differences in mechanical behavior (mechanomyographic amplitude patterns) of the VL during a series of repetitive muscle actions. Subsequently, based on the findings of Trevino and Herda, and Olmos et al., a series of isometric trapezoidal muscle actions may be sensitive to adjustments in the electrophysiological behavior during fatigue among chronic training statuses. This has yet to be examined and warrants further investigation.

The patterns of response for EMGRMS-force relationships during linearly increasing muscle actions have typically been examined with polynomial regression or analysis of variance (ANOVA) models of composite EMGRMS values at discrete %MVC levels. However, due to large variability among individuals, it has been suggested that linearly varying muscle actions should be examined on a subject-by-subject basis to better describe the individual patterns of response. Subsequently, Herda et al. proposed log-transforming the EMGRMS and force values and calculating b terms to investigate possible changes in the individual patterns. In addition, the 95% confidence intervals (CI) calculated around the b terms provide insight on the linearity of the relationship. For example, if the b term is equal to 1 or the 95% CI include 1, the relationship between EMGRMS and force is linear. If the b term is greater than 1 and the 95% CI do not include 1, the relationship accelerates across the force spectrum as the rate of change is greater for the Y variable (EMGRMS) than the X variable (force). Furthermore, the y-intercepts (a terms) of the log-transformed relationships reflect upward or downward shifting of the overall exponential relationship without changes to the EMGRMS patterns. Previously, the b terms calculated from EMGRMS-force relationships have identified MU activation- and muscle action-related differences among chronic training statuses during a 60% MVC. Thus, examining the b and a terms during repetitive isometric trapezoidal muscle actions may provide insight on differences in MU control strategies among chronically trained individuals during fatigue.

Therefore, the purpose of this study was to examine EMGRMS-force relationships during repetitive muscle actions at 50% MVC that contained linearly increasing, steady force, and linearly decreasing segments for AT, RT, and SED individuals. Based on the findings of Trevino and Herda, we hypothesized greater b terms for the RT compared to the AT and SED during the linearly increasing segment of the first contraction. In addition, we hypothesized muscle action-related differences for the first contraction, such as greater b terms for the RT during the increasing-in comparison to the decreasing-segment and the converse for the AT. During fatiguing contractions, it has been reported that as MU twitch forces decline, MU firing rates increase, MU recruitment thresholds decrease, and muscle excitation from the central nervous system to motoneuron pool increases to maintain the target force level. Thus, for the last contraction, we hypothesized greater b terms for the RT compared to the AT during the linearly increasing and decreasing segments due to the accumulation of fatigue. Based on the findings of Beck et al. and Trevino and Herda, we hypothesized EMGRMS from the steady force segment would not differentiate training statuses.

Material and methods

Subjects

Fifteen healthy adults (mean ± standard deviation [SD]; age = (21.80 ± 3.67) years [yr]; body weight = (73.59 ± 22.79) kg; height = (172.85 ± 11.71) cm) participated in this study. Based on training status, participants were categorized as AT (five participants; age = [19.20 ± 0.45] yr; body weight = [59.02 ± 11.98] kg; height = [171.89 ± 15.81] cm, RT (five participants; age = [25 ± 4.53] yr; body weight = [99.22 ± 17.87] kg; height = [178.74 ± 8.09] cm) or SED (five participants; age = [21.20 ± 2.17] yr; body weight = [62.52 ± 10.69] kg; height = [167.90 ± 9.45] cm) for further statistical analysis. Individuals in the AT group participated in a structured running program for at least 3 years prior to the study, completing an average of (61 ± 15) miles per week for 7–10 h per week, and none of them reported engaging in resistance training. RT individuals reported engaging in a structured resistance training program for at least 4 years prior to the study and performed 4–8 h per week of resistance training without engaging in any type of aerobic activity and self-reported a one-repetition back squat of at least twice their body mass. SED individuals reported no participation in any form of structured physical activity or exercise for 3 years prior to this study. According to Herda et al. and Fry et al., differences in %MHC isoform expression for the VL have been reported among individuals with similar training.

Abbreviations:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>a term</td>
<td>antilog of the y-intercept</td>
</tr>
<tr>
<td>ANOVA</td>
<td>Analysis of Variance</td>
</tr>
<tr>
<td>AT</td>
<td>Aerobically Trained</td>
</tr>
<tr>
<td>b term</td>
<td>slope</td>
</tr>
<tr>
<td>CI</td>
<td>Confidence Interval</td>
</tr>
<tr>
<td>cm</td>
<td>Centimeter</td>
</tr>
<tr>
<td>EMG</td>
<td>Electromyography</td>
</tr>
<tr>
<td>EMGRMS</td>
<td>Electromyographic Amplitude</td>
</tr>
<tr>
<td>h</td>
<td>Hour</td>
</tr>
<tr>
<td>Hz</td>
<td>Hertz</td>
</tr>
<tr>
<td>Kg</td>
<td>Kilogram</td>
</tr>
<tr>
<td>kHz</td>
<td>Kilohertz</td>
</tr>
<tr>
<td>MHC</td>
<td>Myosin Heavy Chain</td>
</tr>
<tr>
<td>MU</td>
<td>Motor Unit</td>
</tr>
<tr>
<td>MUAP</td>
<td>Motor Unit Action Potential</td>
</tr>
<tr>
<td>MVC</td>
<td>Maximal Voluntary Contraction</td>
</tr>
<tr>
<td>RMS</td>
<td>Root-Mean-Square</td>
</tr>
<tr>
<td>RT</td>
<td>Resistance Trained</td>
</tr>
<tr>
<td>SD</td>
<td>Standard Deviation</td>
</tr>
<tr>
<td>SED</td>
<td>Sedentary</td>
</tr>
<tr>
<td>SPSS</td>
<td>Statistical Package for the Social Sciences</td>
</tr>
<tr>
<td>VL</td>
<td>Vastus Lateralis</td>
</tr>
</tbody>
</table>
histories, such as a greater type I %MHC isoform expression for AT than the RT and SED. The sample size (n = 5 for each group) was based on previous investigations with similar study designs that examined the log transformed EMG_RMS-force relationships of the VL for AT, RT, and SED groups. In addition, power calculation software (G^power 3.1.9.7, Heinrich-Heine-Universität Düsseldorf, Düsseldorf, Germany) indicated that 5 participants per group were sufficient for detecting training status related differences. Prior to experimental testing, participants completed an informed consent and health and exercise status questionnaire. No participants reported any current or previous neuromuscular diseases or musculoskeletal injuries specific to the ankle, knee, or hip joints. This study was approved by the University Institutional Review Board for human subject research. This study was performed in line with the principles of the Declaration of Helsinki. Approval was granted by the Ethics Committee of University of Kansas (10-30-2012/HSCL #20495).

Isometric testing

Each participant was seated on a Biodex dynamometer (Biodex Medical System, Inc., Shirley, NY, USA) with restraining straps over the pelvis, trunk, and left thigh. The right femur was aligned with the input axis of the Biodex and all isometric knee extensor strength assessments were performed on the right leg at a knee joint angle of 90° (Biodex Pro Manual, Application/Operations, 1998). The force output of knee extensors was measured using a load cell (LC402, Omegadyne, Inc., Sunbury, OH, USA) that was fitted to the Biodex System 3 isokinetic dynamometer.

Participants visited the laboratory for one experimental visit. During experimental testing, participants performed three isometric MVCs for each of the knee extensors with 3 min rest between muscle actions. Following a 5-min rest period, participants were asked to complete 20 repetitive submaximal isometric trapezoidal muscle actions at 50% MVC. The trapezoidal trajectory contained a 5 s baseline, a linearly increasing segment from baseline at a rate of 10% MVC/s, a 12 s steady force segment at the targeted %MVC, a linearly decreasing segment to baseline at a rate of 10% MVC/s, and a 3 s baseline (Fig. 1). Thus, participants were given an 8–9 s rest period between contractions. Participants repeatedly performed the muscle actions until they completed the 20 repetitions or the average force decreased by > 5% MVC from the 50% MVC target. Prior to the repeated trapezoidal muscle actions at 50% MVC, participants practiced the isometric trapezoidal muscle actions at 20% MVC. A computer monitor was provided to display the target force template and real-time force output, and participants were asked to maintain their force as close as possible to target force template during the testing.

Electromyography

Surface EMG signals were recorded from the VL during the trapezoidal muscle action with a 5 pin array sensor (Delsys, Boston, MA, USA). Each pin had a 0.5 mm diameter, with 4 pins located at the corners of a 5 × 5 mm square and the 5th pin at the center of the square. Before sensor placement, the surface of the skin was shaved, adhesive tape was used to remove superficial dead skin, and the skin was sterilized with alcohol. The sensor was placed on the muscle belly of the VL at half the distance between the greater trochanter and lateral condyle of the femur with adhesive tape. A reference electrode was placed over the left patella after superficial skin was shaved, cleaned, and sterilized. The EMG signals from the 4 pins of the sensor were differentially amplified and filtered with a bandwidth of 20 Hz to 9.5 kHz. The EMG signal recorded from channel 1 was used for all subsequent analyses and statistical comparisons.

Signal processing

Both EMG (µV) and force (N) signals were sampled at 20 kHz with a Delsys data acquisition system (Bagnoli-16 channel EMG system, Delsys, Inc., Boston, MA) during each muscle action. All subsequent signals were stored and processed off-line with a customized LabVIEW program (LabVIEW, version 11; National Instruments, Austin, TX). The EMG signals were bandpass filtered (fourth-order Butterworth) at 10–500 Hz. The force and EMG signals were analyzed with consecutive, non-overlapping 0.25 s epochs during the submaximal isometric trapezoidal muscle actions. The amplitude of the EMG signal was calculated as the root-mean-square (RMS).

Skinfold thickness

Skinfold thickness measurements were taken at the location of the EMG sensor placement for the VL. An experienced investigator performed the measurements with a calibrated Harpenden caliper (John Bull, UK) in accordance to the recommendations of Jackson and Pollock. Three measurements were recorded and the average was defined as the representative skinfold thickness for each participant. It has previously been suggested that subcutaneous fat may low-pass filter the EMG signals.

Statistical analyses

For the linearly increasing (Fig. 1-A) and decreasing (Fig. 1-C) segments of the trapezoid, simple linear regression models were fit to log-transformed EMG_RMS-force relationships. The equations were represented as:

\[ \ln(Y) = b \ln(X) + \ln(a) \]  

Where \( \ln(Y) \) = the natural log of the EMG_RMS values, \( \ln(X) \) = the natural log of the force values, \( b \) = slope, and \( \ln(a) \) = the natural log of the y-
intercept. This can also be expressed as an exponential equation after antilog transformation:

\[ Y = ax^b \]  

(2)

Where \( Y \) is the predicted EMGRMS values, \( X = \) force, \( b = \) slope of equation (1), and \( a = \) the antilog of the y-intercept from equation (1). Slopes \((b)\) were calculated using Microsoft Excel (Microsoft Excel, version 2010; Microsoft, Inc., Redmond, WA).

For the steady force segment of the trapezoid (Fig. 1–B), EMGRMS was calculated by averaging the values for each 0.25 s epoch from the entire 12 s targeted \%MVC. Two separate three-way mixed factorial ANOVAs (training status [AT vs. RT vs. SED] \(\times\) segment [linear increase vs. linear decrease] \(\times\) contraction [first vs. last]) was used to examine differences in the \(b\) and \(a\) terms from the log-transformed EMGRMS-force relationships during linear increasing and linear decreasing segments of submaximal isometric trapezoid muscle actions. A two-way mixed factorial ANOVA (training status [AT vs. RT vs. SED] \(\times\) contraction [first vs. last]) was used to examine possible differences in EMGRMS among training statuses during the steady force segment of the isometric trapezoid muscle action. In addition, Pearson's product moment correlation coefficients were calculated comparing skinfold thicknesses among the \(a\) and \(b\) terms and EMGRMS during the steady force segments. When appropriate, follow-up tests included one-way ANOVAs and paired sample t-tests for the \(b\) terms of log-transformed EMGRMS-force relationships with Bonferroni corrections. The partial \(\eta^2\) statistics were calculated with values of 0.01, 0.06, and 0.14 corresponding to small, medium, and large effect size, respectively. In addition, Hedge's \(g\) was calculated for paired comparisons, with 0.2, 0.5, and 0.8 corresponding to small, medium, and large effect size, respectively. All statistical analyses were performed using Statistical Package for the Social Sciences (SPSS) (version 24, IBM Corporation, Armonk, New York, USA) with alpha set at 0.05.

### Results

Table 1 contains the individual values for EMGRMS at steady-force and the \(a\) and \(b\) terms from the log-transformed EMGRMS-force relationships during the linearly increasing and decreasing segments of the first and last successfully completed isometric trapezoidal contraction.

#### Linearly increasing and decreasing segments EMGRMS-force relationships

For the \(b\) terms, the analyses indicated a significant three-way interaction (training status \(\times\) segment \(\times\) contraction; \(F = 4.926, p = 0.027\), partial \(\eta^2 = 0.451\)). The \(b\) terms for RT during the linearly increasing segment of the first contraction (1.301 \(\pm\) 0.197) were greater than the AT (0.910 \(\pm\) 0.123; \(p = 0.008, g = 2.381\)) and SED (0.912 \(\pm\) 0.162; \(p = 0.008, g = 2.157\)), the \(b\) terms for the RT during the first contraction were greater during the linearly increasing segment than decreasing segment (1.018 \(\pm\) 0.139; \(p = 0.014, g = 1.660\)), the \(b\) terms for the RT during the linearly increasing (1.373 \(\pm\) 0.353) and decreasing segment (1.526 \(\pm\) 0.328) of the last contraction were greater than the AT (linear increase = 0.883 \(\pm\) 0.129, \(p = 0.018, g = 1.844\); linear decrease = 0.970 \(\pm\) 0.223, \(p = 0.010, g = 1.983\)), and the \(b\) terms for SED during the last contraction were less during the linearly increasing (0.968 \(\pm\) 0.144) than decreasing segment (1.268 \(\pm\) 0.126; \(p = 0.015, g = 2.217\) (Fig. 2)). There were no other differences reported among training statuses, between segments, or between repetitions. Fig. 3 illustrates the mean EMGRMS Patterns for the AT, RT, and SED during the linearly increasing and decreasing segments of the first and last contractions.

**Table 1** Electromyographic amplitude (EMGRMS) during steady-force and the \(a\) and \(b\) terms calculated from the log-transformed EMGRMS-force relationship during linearly increasing and decreasing segments for the first and last successfully completed isometric trapezoidal contraction for the aerobically- (AT), resistance-trained (RT) and sedentary (SED) individuals.

<table>
<thead>
<tr>
<th>Group</th>
<th>Subject</th>
<th>First contraction</th>
<th>Last contraction</th>
<th>Coefficients</th>
<th>First contraction</th>
<th>Last contraction</th>
</tr>
</thead>
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<tr>
<td></td>
<td></td>
<td>EMGRMS ((\mu V))</td>
<td>EMGRMS ((\mu V))</td>
<td>Increase</td>
<td>Decrease</td>
<td>Increase</td>
</tr>
<tr>
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<td>0.270</td>
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<td></td>
<td>2</td>
<td>51.05</td>
<td>60.44</td>
<td>1.013^a</td>
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<tr>
<td></td>
<td>3</td>
<td>80.65</td>
<td>98.14</td>
<td>0.530</td>
<td>0.542</td>
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<tr>
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<td>4</td>
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<td>78.65</td>
<td>0.838^a</td>
<td>0.855^a</td>
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<tr>
<td></td>
<td>5</td>
<td>36.02</td>
<td>37.27</td>
<td>0.634</td>
<td>0.240</td>
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<tr>
<td>RT</td>
<td>6</td>
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<td>47.47</td>
<td>0.172</td>
<td>0.126</td>
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<tr>
<td></td>
<td>7</td>
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<td>88.54</td>
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<td>1.165^a</td>
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<td>8</td>
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<td></td>
<td>9</td>
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<td>133.40</td>
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<td>0.866^a</td>
<td>0.748^a</td>
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<tr>
<td></td>
<td>10</td>
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<tr>
<td>SED</td>
<td>11</td>
<td>27.95</td>
<td>60.70</td>
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<td>0.141</td>
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<tr>
<td></td>
<td>12</td>
<td>26.48</td>
<td>39.39</td>
<td>0.863^a</td>
<td>0.863^a</td>
<td>1.333^a</td>
</tr>
<tr>
<td></td>
<td>13</td>
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<td>32.80</td>
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<td>0.042</td>
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<tr>
<td></td>
<td>14</td>
<td>234.04</td>
<td>215.42</td>
<td>1.249^a</td>
<td>1.105^b</td>
<td>0.806^a</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>82.59</td>
<td>149.52</td>
<td>0.004</td>
<td>0.022</td>
<td>0.110</td>
</tr>
</tbody>
</table>

^a Indicates relationship was linear.  
^b Indicates relationship was curvilinear.
For the terms, the analyses indicated no significant three-way interaction (training status × segment × contraction; \( F = 0.715, p = 0.509 \), partial \( \eta^2 = 0.107 \)), no two-way interactions (segment × contraction, \( F = 2.529, p = 0.138 \), partial \( \eta^2 = 0.174 \); training status × segment, \( F = 1.601, p = 0.242 \), partial \( \eta^2 = 0.211 \); training status × contraction, \( F = 1.432, p = 0.277 \), partial \( \eta^2 = 0.193 \)) or main effects for training status (\( F = 3.677, p = 0.057 \), partial \( \eta^2 = 0.380 \)), segment (\( F = 2.860, p = 0.117 \), partial \( \eta^2 = 0.192 \)), or contraction (\( F = 1.985, p = 0.184 \), partial \( \eta^2 = 0.142 \)).

For the terms, the analyses indicated no significant three-way interaction (training status × segment × contraction; \( F = 0.715, p = 0.509 \), partial \( \eta^2 = 0.107 \)), no two-way interactions (segment × contraction, \( F = 2.529, p = 0.138 \), partial \( \eta^2 = 0.174 \); training status × segment, \( F = 1.601, p = 0.242 \), partial \( \eta^2 = 0.211 \); training status × contraction, \( F = 1.432, p = 0.277 \), partial \( \eta^2 = 0.193 \)) or main effects for training status (\( F = 3.677, p = 0.057 \), partial \( \eta^2 = 0.380 \)), segment (\( F = 2.860, p = 0.117 \), partial \( \eta^2 = 0.192 \)), or contraction (\( F = 1.985, p = 0.184 \), partial \( \eta^2 = 0.142 \)).

**Steady force segments**

For the steady force segments, the analyses indicated neither a two-way interaction (training status × contraction; \( F = 1.463, p = 0.270 \), partial \( \eta^2 = 0.196 \)) nor a main effect for training status (\( F = 0.325, p = 0.729 \), partial \( \eta^2 = 0.051 \)). There was a main effect for contraction (\( F = 16.787, p = 0.001 \), partial \( \eta^2 = 0.583 \)). EMGRMS during the steady force segment was greater for the last contraction (\( 86.73 \pm 49.55 \) μV) than the first contraction (\( 64.08 \pm 51.68 \) μV) when collapsed across training status (Fig. 4).

**Correlations**

Pearson's product moment correlations were not significant for skinfold thickness with the terms from the linearly increasing and decreasing segments for the first and last contractions (\( p = 0.114–0.733 \), \( r = -0.425 \) to 0.327). In addition, there were no correlations for skinfold thickness among the terms for the linearly increasing or decreasing segments for the first contraction (\( p = 0.114–0.855 \), \( r = -0.052 \) to -0.379), or the linearly decreasing segment for the last contraction (\( p = 0.018 \)).
Skinfold thickness was correlated with the $a$ terms during the linearly increasing segment of the last contraction ($p = 0.025$, $r = -0.576$). Therefore, only 1 of 8 (12.5%) correlations were significant for skinfold thickness among the coefficients, which is in agreement with previous examinations\(^{17,28}\) and provides further confidence that training status related differences for the $b$ terms were not influenced by skinfold thicknesses. Skinfold thickness was correlated with EMGRMS from the steady force segments for the first ($p = 0.014$, $r = -0.617$) and the last contractions ($p = 0.040$, $r = -0.535$).

**Discussion**

All 60 log-transformed EMGRMS-force relationships were significant for the linearly increasing ($p < 0.05$; $r$ range = 0.808–0.988) and decreasing segments ($p < 0.05$; $r$ range = 0.899–0.988) of the first and last completed isometric trapezoidal muscle action. As previously reported by our group,\(^{20}\) only the AT were able to successfully complete all 20 isometric trapezoidal muscle actions, as well as maintain maximal strength following the repeated muscle actions. Additionally, during the first isometric trapezoidal muscle action, the RT exhibited greater $b$ terms than the AT and SED during the linearly increasing segment, and only the RT displayed muscle action-related differences.\(^{13}\) Significant and varied findings during the last contraction in response to fatigue include greater $b$ terms for the RT during the linearly increasing and decreasing segments in comparison to the AT, and muscle action-related differences for the SED where the $b$ terms were greater during the linearly decreasing in comparison to the linearly increasing segment.

During the repetitive submaximal isometric trapezoidal muscle actions, there were numerous differences as a function of training status and muscle action. For the linearly increasing segment of the first contraction, the $b$ terms for the RT were greater than the AT and SED. Previously, Trevino and Herda\(^{17}\) reported similar findings during an isometric trapezoidal contraction performed at 60% MVC. It is suggested that EMGRMS-force relationships reflect changes in MU recruitment and firing rates to modulate force production.\(^{1,2}\) Thus, it was hypothesized the larger $b$ terms for the RT indicated a greater amount of MU activation were necessary to match the targeted force compared to the AT and SED.\(^{17}\) However, it has recently been reported that EMGRMS is primarily explained by the size of the motor unit action potentials (MUAPs).\(^{39}\) Numerous studies have reported chronic resistance-training increases muscle cross-sectional area\(^{40,41}\) particularly due to the hypertrophy of type II muscle fibers.\(^{40}\) In addition, MUAPs are positively correlated to muscle fiber size.\(^{32,41}\) Therefore, the greater $b$ terms for the RT compared to the AT and SED during the linearly increasing segment of the first contraction is likely due to the recruitment of MU comprised of larger muscle fibers, rather than greater MU activation as fatigue was unlikely accumulating during the 5 s linearly increasing portion of the isometric trapezoidal contraction.\(^{39}\)

During the first contraction, the $b$ terms for RT were greater during the linearly increasing than decreasing segment, which is agreement with Trevino and Herda.\(^{17}\) The differences in MU activation and deactivation strategies for the RT may be a result of the 12 s steady force segment between the linearly increasing and decreasing segments. MU potentiation of the VL was previously reported for chronic RT individuals during the linearly increasing and decreasing segments of the last contraction,\(^{4,41}\) MU potentiation would increase MU twitch force output during the contraction, allowing potentiated MUs to be derecruited at higher force levels during the linearly decreasing segment than which they were initially recruited at during the increasing segment.\(^{4,41,45}\) Thus, the amount of muscle excitation to the motoneuron pool and the size of the largest activated MUs would be less at the same relative force level (%MVC) during the decreasing segment, which is supported by significant decrease in $b$ terms for the RT. Conversely, the AT and SED displayed no significant differences in the $b$ terms between increasing and decreasing segments. Mettler and Griffin\(^{26}\) and Herda et al.\(^{44}\) reported a lack of MU potentiation for the adductor pollicis and VL following endurance training and for chronic AT individuals, respectively. In addition, no differences in $b$ terms from EMGRMS-force relationships were previously reported for SED between linearly increasing and decreasing muscle actions.\(^{17}\) Thus, the findings suggest alterations in MU potentiation may require chronic resistance training.

There were additional chronic training status related differences during the last contraction, such as greater $b$ terms for the RT during the linearly increasing and decreasing segments in comparison to the AT. The chronic training-related differences in MU activation and deactivation strategies may be due to differences in fatigue resistance. During muscular fatigue, it is well understood that MU twitch forces decrease.\(^{53}\) Consequently, the central nervous system increases the amount of excitation to the motoneuron pool and the recruitment thresholds decrease for higher-threshold MUs to compensate for the decline in twitch forces of the fatiguing MUs.\(^{24}\) Aerobic training elicits relatively higher amounts of type I % MHC expression,\(^{17,45}\) whereas the converse is true for resistance training.\(^{40}\) Thus, the VL for the RT is likely comprised of a greater percentage of MUs that possess lower fatigue resistance than the AT, and the larger $b$ terms indicate a greater increase in MU recruitment and/or firing rates to match the same %MVC during the linearly increasing and decreasing segments of the last contraction. Conversely, the SED showed no significant differences in the $b$ terms during the linearly increasing and decreasing segments compared with the RT and AT, although it has been reported the proportion of total type II% MHC for SED-is similar with the RT-individuals.\(^{16,34}\) Therefore, it would be expected the RT and SED would possess similar fatigability, which has previously been reported in this cohort of participants.\(^{26}\) However, the lack of differences for the SED in comparison to the AT and RT may suggest that neuromuscular and muscle structure training-related adaptions are both necessary for the $b$ terms to differentiate MU activation and deactivation strategies among chronic training status during fatigue. There were also muscle action related differences for the SED during the last contraction, such as greater $b$ terms during the linearly decreasing in comparison to the linearly increasing segment. It is well documented that muscle excitation and descending drive from central nervous system increases with muscular fatigue.\(^{39}\) Thus, the greater $b$ terms for the SED during the linearly decreasing indicates increased muscle excitation in comparison to the linearly increasing segment when trying to produce force. Nonetheless, the findings suggest the log-transformed EMGRMS-force relationships may be sensitive to chronic training induced alterations in motor control strategies during fatigue.

For the steady force segment, EMGRMS was greater during the last contraction in comparison to the first contraction when collapsed across training statuses. The amplitude of the EMG signal is influenced by the number of recruited MUs and their firing rates.\(^{1,2}\) Thus, the change in EMGRMS likely resulted from the increase in muscle excitation that occurs in response to muscular fatigue (e.g., increase in firing rates, accelerated recruitment of higher-threshold MUs).\(^{34}\) EMGRMS during steady force did not differentiate among groups ($p = 0.051$), which supports the findings of Beck et al.\(^{17}\) that reported no difference in EMGRMS of the VL between AT- and RT-individuals during a fatiguing 30 s submaximal isometric muscle action of the leg extensors at 50% MVC. These findings further support that utilizing the $b$ terms from the log-transformed EMGRMS-force relationships during linearly increasing and decreasing segments of an isometric trapezoidal contraction may provide more insight than EMGRMS recorded during steady force to elucidate neuromuscular adaptions resulting from chronic training.\(^{17,25}\) It should be noted that EMGRMS recorded during the 12 s steady force segment was negatively correlated with skinfold thickness, which can serve as a low-pass filter.\(^{15}\) However, it was previously reported that skinfold thickness differences did not exist among groups for this cohort of individuals,\(^{26}\) which may suggest low-pass filtering was not responsible for the lack of training-related differences in EMGRMS during the 12 s steady force segment. Nonetheless, EMGRMS at steady force was not able to differentiate among AT, RT, and SED during the first and last successfully completed muscle action.
We would like to acknowledge this study did not collect muscle biopsies, which is a limitation. Due to our strict inclusion criteria based of Herda et al.,10 Fry et al.,14 and Beck et al.,15 we were confident our participants would possess differences in MHC isoform expression. For example, 2 AT and 1 RT individual participated in another study for our laboratory that included MHC analysis of the VL. The participants exhibited training status related differences for MHC isoforms of the VL, such as the type I MHC isoform expression percentages for the AT individuals were 55% and 68%, respectively, whereas the type I MHC isoform expression percentage for the RT individual was 31%. However, since we did not collect muscle biopsies for every participant in our study, we cannot confirm that differences for type I MHC isoform expression existed between the chronic training groups.

In conclusion, the b terms from the log-transformed EMGRMS-force relationships during linearly increasing and decreasing muscle actions differentiated training statuses during pre- and late-fatigue isometric trapezoidal contractions, such as greater acceleration in EMGRMS throughout the force spectrum for the RT than AT. The greater acceleration in EMGRMS for RT during the first contraction may reflect the recruitment of larger sized MUs compared to the AT as fatigue was likely not present yet, whereas it is plausible the group differences during the last contraction is due to accelerated recruitment of additional higher-threshold MUs and/or greater firing rates of the recruited MUs as fatigue was accumulating to a greater extent for the RT. In addition, there were muscle-action related differences (linearly increasing vs. decreasing) among training statuses. The RT exhibited greater acceleration during the linearly increasing muscle action in comparison to the linearly decreasing for the first contraction, whereas the opposite was true for the SED during the last contraction. Conversely, there were no muscle-action related difference for the AT. Therefore, different modes of chronic training elicited specific EMGRMS patterns of response (i.e. muscle excitation to motoneuron pool) during a fatigueing-type task. The findings suggest strength coaches and practitioners should consider the athletic demands of their athletes and the goals of their patients when designing exercise programs, as solely engaging in resistance training resulted in a significantly greater amount of necessary muscle excitation to produce a relative targeted force during repetitive MU activation and deactivation tasks compared to individuals who strictly aerobically train. Thus, athletes or individuals who currently resistance train but have task demands lasting longer than a few seconds (repetitive or continuous) may see increased performance in the latter stages of the activity by incorporating some aerobic training. Future research should utilize EMG signal decomposition techniques to examine the influence of chronic training on MU control strategies (e.g., MU recruitment and firing rates) during fatigue.

Submission statement

The work described has not been published previously, that it is not under consideration for publication elsewhere, that its publication is approved by all authors and tacitly or explicitly by the responsible authorities where the work was carried out, and that, if accepted, it will not be published elsewhere including electronically in the same form, in English or any other language, without the written consent of the copyright-holder.

Authors’ contributions

All authors contributed to the study conception and design. Material preparation, data collection, and analysis were performed by SJ, SAS, TJH, and MAT. The first draft of the manuscript was written by SJ and MAT, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.


