Effects of exercise and dietary epigallocatechin gallate and β-alanine on skeletal muscle in aged mice


Abstract: Aging leads to sarcopenia and loss of physical function. We examined whether voluntary wheel running, when combined with dietary supplementation with (-)-epigallocatechin-3-gallate (EGCG) and β-alanine (β-ALA), could improve muscle function and alter gene expression in the gastrocnemius of aged mice. Seventeen-month-old BALB/cByJ mice were given access to a running wheel or remained sedentary for 41 days while receiving either AIN-93M (standard feed) or AIN-93M containing 1.5 mg kg⁻¹ EGCG and 3.43 mg kg⁻¹ β-ALA. Mice underwent tests over 11 days from day 29 to day 39 of the study period, including muscle function testing (grip strength, treadmill exhaustive fatigue, rotarod). Following a rest day, mice were euthanized and gastrocnemii were collected for analysis of gene expression by quantitative PCR. Voluntary wheel running (VWR) improved rotarod and treadmill exhaustive fatigue performance and maintained grip strength in aged mice, while dietary intervention had no effect. VWR increased gastrocnemius expression of several genes, including those encoding interleukin-6 (Il6, p = 0.001), superoxide dismutase 1 (Sod1, p = 0.046), peroxisome proliferator-activated receptor gamma coactivator 1α (Ppargc1a, p = 0.013), forhead box protein O3 (Foxo3, p = 0.005), and brain-derived neurotrophic factor (Bdnf, p = 0.008), while reducing gastrocnemius levels of the lipid peroxidation marker 4-hydroxynonenal (p = 0.019). Dietary intervention alone increased gastrocnemius expression of Ppargc1a (p = 0.033) and genes encoding NAD-dependent protein deacetylase sirtuin-1 (Sirt1, p = 0.039), insulin-like growth factor 1 (Igf1, p = 0.003), and macrophage marker CD11b (Itgam, p = 0.016). Exercise and a diet containing β-ALA and EGCG differentially regulated gene expression in the gastrocnemius of aged mice, while VWR but not dietary intervention improved muscle function. We found no synergistic effects between dietary intervention and VWR.

Key words: voluntary wheel running, gene expression, muscle function, animal model, supplementation, aging.

Résumé : Le vieillissement aboutit à la sarcopénie et à la perte de fonctions physiques. Nous vérifions si la course volatile dans une roue d’exercice combinée à une supplémentation alimentaire en (-)épigallocatechine-3-gallate (- EGCG) et en β-alanine (β-ALA) améliore la fonction musculaire et modifie l’expression génique dans le muscle gastrocnemius de la souris âgée. On donne à des souris BALB/cByJ âgées de 17 mois l’accès à une roue d’exercice ou on les maintient sédentaires durant 41 jours tout en leur procurant un régime de AIN-93M (nourriture standard) ou de AIN-93M contenant 1,5 mg kg⁻¹ EGCG et 3,43 mg kg⁻¹ β-ALA. Les souris sont soumises durant 11 jours (jour 29 – jour 39) à des tests incluant l’évaluation des fonctions musculaires (force de préhension, épuisement sur tapis roulant, tige tournante). Après une journée de repos, on euthanise les souris et on les retire au gastrocnemius pour l’analyse de l’expression génique au moyen de la PCR quantitative. La course volatile sur tapis roulant (VWR) améliore la performance au test de la tige tournante et de l’épuisement sur tapis roulant et maintient la force de préhension des souris âgées, alors que l’intervention alimentaire n’a aucun effet. La VWR améliore l’expression de plusieurs gènes dans le gastrocnemius, notamment ceux encodant l’interleukine-6 (Il6, p = 0.001), la superoxyde dismutase 1 (Sod1, p = 0.046), le coactivateur 1α du récepteur gamma activé de la prolifération des peroxysomes (- Ppargc1α) (-Ppargc1a, p = 0.013), forhead box 03 (Foxo3, p = 0.005), le facteur neurotrophique dérivé du cerveau (Bdnf, p = 0.008), mais diminue dans le gastrocnemius le taux de 4-hydroxynonenal, marqueur de la peroxydation des lipides (p = 0.019). L’intervention sur le plan alimentaire seul augmente l’expression des gènes du gastrocnemius encodant PGC-1α (p = 0.033), la déacétylase sirtuine 1 (Sirt1, p = 0.039), du facteur de
Introduction

Physiological aging includes a number of morphological and functional changes to many organs, including skeletal muscle. Total muscle mass decreases in humans starting at approximately age 30 (Short et al. 2003), and this change is associated with deficits in muscle strength. Mechanistically, a number of studies have noted an age-related decrease in the rate of protein synthesis in skeletal muscle with resistance exercise (Yarasheski et al. 2002), although the rate of protein synthesis in young versus old is thought to be unchanged at rest (Fry and Rasmussen 2011; Volpi et al. 2001). Additionally, skeletal muscle mitochondria become increasingly dysfunctional with advancing age (Rooyackers et al. 1996). Age-related loss of muscle mass and function (sarcopenia) is associated with impairments in mobility and locomotion (Morley et al. 2011). Whether these impairments are the consequence or the cause of sarcopenia is a matter of debate. Age-associated mitochondrial dysfunction in skeletal muscle (Rooyackers et al. 1996) is associated with an age-related increase in oxidative stress (Capel et al. 2005), and these findings are thought to be causally linked. Indeed, overexpression of the antioxidant enzyme catalase has been shown to prevent age-associated metabolic impairments in mice (Lee et al. 2010), and mitochondrial DNA deletions have been shown to co-localize with oxidative damage in sarcopenic muscle fibers isolated from aged rats (Wanagat et al. 2001). However, the exact link between oxidative stress and mitochondrial dysfunction in aging remains under investigation.

β-alanine (β-ALA) is the non-proteinogenic β form of the amino acid alanine and is a precursor for carnosine, a dipeptide of β-ALA and β-histidine. In muscle, carnosine acts as a pH buffer and functions as an antioxidant (Boldyrev et al. 1993), which suggests a potential role for carnosine in reversing or limiting the effects of oxidative stress and cellular senescence. Carnosine additionally increases Ca²⁺ sensitivity and aids Ca²⁺ release from the sarcoplasmic reticulum in muscle (Dutka et al. 2012), thus directly enhancing muscle contractility and performance. Thus, supplementation with β-ALA has been widely examined for its effects on muscle carnosine content and physical function. Supplementation with β-ALA or carnosine is a strategy commonly used by athletes, as β-ALA-induced increases in muscle carnosine have been shown to decrease fatigue and improve muscular function (Derave et al. 2010). Recently, β-ALA has been investigated for its effects on skeletal muscle function in the elderly. β-ALA supplementation for 12 weeks increased muscle carnosine content and improved time-to-exhaustion in several tests of physical capacity in 60- to 80-year-olds (del Favero et al. 2012). Furthermore, there was a positive correlation between improved physical capacity and increases in skeletal muscle carnosine content. Because muscle contents of carnosine and its methylated analogue anserine naturally decrease with age (Stuerenburg and Kunze 1999), including in rodents (Derave et al. 2008; Johnson and Hammer 1992), β-ALA supplementation is an attractive strategy to minimize or reverse age-associated deficits in muscle functional capacity. However, recent data indicate that β-ALA supplementation has no effect on skeletal muscle function after a 6-week interval training program (Cochran et al. 2015), no effect on Wingate performance in women (Kresta et al. 2014), and no effect on measures of muscle function in young mice after a 4-week voluntary wheel running intervention (Bhattacharya et al. 2015). Thus, the value of β-ALA supplementation for improving muscle function remains somewhat controversial.

In addition to β-ALA, we used an abundant biologically active catechin, (-)-epigallocatechin-3-gallate (EGCG), because of its well-established antioxidant action. EGCG is the major polyphenolic catechin constituent of green tea and has been widely investigated for its antioxidant properties, as this is the major health benefit attributed to green tea (Yang 1999). EGCG acts as a scavenger of a number of free radicals and has been shown to exert anti-inflammatory effects (Aktas et al. 2004). Thus, there is considerable interest in the potential effects of EGCG in ameliorating a variety of oxidative and pro-inflammatory conditions, including normal aging. In contrast with β-ALA, there is limited information available on the role of EGCG in muscle, especially from an aging standpoint. In aged rats, EGCG increased skeletal muscle activity of several antioxidant enzymes (including superoxide dismutase, catalase, glutathione peroxidase, and glutathione reductase) and increased skeletal muscle levels of non-enzymatic antioxidants such as ascorbate and α-tocopherol (Senthil Kumaran et al. 2008). Several studies have reported that green tea extract or EGCG supplementation reduces muscle mass and (or) enhances muscle recovery during or after hind-limb suspension-induced unloading in aged rats (Alway et al. 2014, 2015). However, the effect of EGCG on skeletal muscle dysfunction resulting from normal physiological aging has yet to be adequately examined.

Exercise training has long been utilized as an intervention to improve muscle function in aged subjects. Although resistance exercise training is commonly used in this population, aerobic exercise training also has benefits in aged skeletal muscle. In the aged, aerobic exercise training increases skeletal muscle mitochondrial number and activity, enhances muscle protein synthesis, improves insulin sensitivity, and increases oxidative capacity (Short et al. 2003). Because these exercise-induced improvements are related to decreased oxidative stress and increased exercise capacity in aging (Leeuwenburgh and Heinecke 2001), aerobic exercise is a powerful modality by which to improve physical function in the elderly. An exercise-induced increase in mitochondrial number (mitochondriobiogenesis) is thought to be important in many of the above effects of exercise. Mitochondrial biogenesis increases owing to enhanced activity of peroxisome proliferator-activated receptor gamma coactivator 1α (PGC-1α, encoded by Ppargc1a), which coordinates the transcriptional activity of several factors related to mitochondrial biogenesis, is increased by exercise, and decreases in skeletal muscle with aging (Pugsley and Spiegelman 2003). Thus, it is thought that aerobic exercise mediates its positive effects on skeletal muscle through increasing the activity of PGC-1α. However, training adaptations are still reduced in aged individuals compared with young (Lanza et al. 2008); thus, adjunct strategies (e.g., nutritional supplementation) that increase the training response in aged individuals are of interest.

Given the potential benefits of EGCG and β-ALA to muscle function, the purpose of our study was to examine the effects of dietary supplementation with β-ALA and EGCG in combination with a voluntary exercise intervention on skeletal muscle function and body composition in aged mice. We hypothesized that in addition to having independent beneficial effects, dietary supplementation with β-ALA and EGCG would either additively or synergistically improve muscle function, body composition, markers of mitochondrial biogenesis, and gene expression of antioxidant enzymes in response to voluntary exercise in aged mice.
Materials and methods

Animals

Male BALB/cByJ retired breeder mice (8–10 months old) were purchased from The Jackson Laboratory (Bar Harbor, Maine, USA) and maintained until 17 months of age. Mice were then randomized to treatments as described below and began the feeding portion of the study. Mice were maintained in an AAALAC-accredited animal care facility and allowed ad libitum access to water and food. The mice were individually housed on a 12-h reversed light-dark cycle (dark period 1000 to 2200 h) at a constant temperature of 24 °C. All procedures used were approved by the Institutional Animal Care and Use Committee at the University of Illinois Urbana-Champaign.

Study design

The study design is diagrammed in Supplementary Fig. S1.1 Mice were switched to either a control diet (Ctrl, N = 28) or an experimental diet containing β-ALA and EGCG (EGCG+β-ALA, N = 30) for 41 days. Body weight and food and water intake (by disappearance) were recorded twice weekly. Mice were randomized to have access to a running wheel (voluntary wheel running, VWR: Ctrl, N = 15; EGCG+β-ALA, N = 15) or were maintained in standard housing (sedentary, SED: Ctrl, N = 13; EGCG+β-ALA, N = 15) during this period. Thus, mice were randomly assigned to 1 of 4 treatment groups (Ctrl-SED, Ctrl-VWR, EGCG+β-ALA-SED, EGCG+β-ALA-VWR).

Beginning on day 29, mice underwent a battery of behavioral and functional tests. These tests included 7 days of Morris water maze (days 29–35) and 2 days of contextual fear conditioning (days 38–39), the results of which are reported elsewhere (Gibbons et al. 2014). On experimental day 36, mice underwent functional muscle testing, including grip strength, rotarod, and treadmill test to exhaustion. Only grip strength and rotarod were performed on day 37. Following the cessation of behavioral testing on day 39, mice were given a rest day on day 40 and were euthanized for tissue collection on day 41 prior to the onset of the dark cycle. Mice were euthanized by CO2 asphyxiation. Muscle samples were excised from both hind limbs, and the gastrocnemius complex was separated from the soleus, frozen on dry ice, and stored at −80 °C for analysis. The gastrocnemius was chosen for its importance in running movements and for its mixed fiber type composition, which we suggest gives a more representative picture of skeletal muscle adaptations to exercise than would the use of a predominantly oxidative (soleus) or glycolytic (quadriceps) muscle group (Jackaman et al. 2007).

Diets

Ctrl and EGCG+β-ALA diets were purchased from Research Diets Inc. (New Brunswick, N.J., USA) and were based on the purified AIN-93M mature rodent diet. Mice receiving Ctrl were given ad libitum access to the AIN-93M diet throughout the study. EGCG+β-ALA was manufactured by Research Diets by mixing 1.7 mg of Teavigo (90% EGCG, DSM Nutritional Products, Basel, Switzerland) and 3.43 mg of β-ALA (NutraBio, Middlesex, N.J., USA) per gram of AIN-93M diet, which was then pelleted to match the consistency and appearance of Ctrl. Dietary constituents have been previously published (Gibbons et al. 2014). If the EGCG+β-ALA had yielded positive effects, the individual dietary components would have been examined to determine their contribution. Since this was not the case, we did not test the individual components, as they were unlikely to have positive effects where the combined diet did not, as was shown in our previous study in young mice (Bhattacharya et al. 2015).

The diets were independently assayed by Covance Inc. (Princeton, N.J., USA), and the experimental diet was found to contain 1.49 mg of EGCG per gram of diet (99.3% of expected) and 3.34 mg of β-ALA per gram of diet (97.4% of expected). The control diet, AIN-93M, was found to be free of both EGCG and β-ALA. Mice consumed an average of 182 mg·kg−1·day−1 and 417 mg·kg−1·day−1 of EGCG and β-ALA, respectively, based on average daily diet disappearance and mean body weight. Target EGCG dosage was determined based on previous studies demonstrating beneficial cognitive effects of EGCG in mice (Li et al. 2009). Because studies examining β-ALA supplementation and its effects on cognition or muscle function in mice are lacking, we calculated β-ALA dosage using the 2.4 g·day−1 dose that has been shown to improve physical work capacity in humans (Stout et al. 2008). For a 70-kg human, this equates to 34 mg·kg−1·day−1 and was adjusted for mice using the FDA-recommended conversion factor of 12.3 (US Department of Health and Human Services 2005). Thus, the target dose for β-ALA in this study was 418 mg·kg−1·day−1.

Wheel running

VWR mice were given constant access to a running wheel (Respironics, Bend, Ore., USA), while SED mice were housed in standard shoebox cages. Both VWR and SED mice remained in their respective conditions throughout the entirety of the study period. Wheel turns were monitored by computerized software (VitalView, Respironics) in 1-h increments, and resulting data were analyzed as wheel distance (km) covered in a 24-h period. VWR and SED mice were subjected to similar handling throughout the experiment.

Body composition

Body composition was analyzed prior to and on the final day of the pretesting portion of the study (day 0 and day 28, respectively) by small animal magnetic resonance imaging (EchoMRI, Houston, Tex., USA). Total fat mass and lean mass were assessed for each mouse at each time point by this method. Data were expressed as percent fat mass or percent lean mass by dividing fat mass or lean mass by total body mass at each time point.

Muscle function testing

Forelimb grip strength was assessed using a commercially available force gauge (Columbus Instruments, Columbus, Ohio, USA) in 5 separate trials per day over 2 consecutive days both at the beginning of the study (days 1 and 2) and during the muscle function testing period (days 36 and 37) by the same investigator. Grip strength was quantified as the average of the highest grip force recorded on each testing day at each time point and as the percent change in maximal grip strength from the beginning to the end of the study in each group. Grip strength was expressed as peak force in newtons.

An exhaustive treadmill test was performed to assess fatigability (day 36). Mice ran on an inclined (5%), motorized treadmill (JOG A DOG, Ottawa Lake, Mich., USA) using an incremental running velocity protocol as previously described (Martin et al. 2013). Fatigue was defined as an inability to continue running despite gentle prodding for at least 10 s. The test ended at 120 min if mice had not reached fatigue. No electric shock was used. Data were expressed as time-to-exhaustion (min).

An automated rotarod unit (Accuscan, Columbus, Ohio, USA) with a 30 mm diameter rotating dowel and a 63 cm fall height was utilized. Mice were placed on the dowel, and rotation started at 0 r/min with constant acceleration to a maximum of 60 r/min. Timing was controlled by photobeam, and timing for each mouse was stopped automatically by the system when the falling mouse broke the plane of the photobeam. Mice underwent 4 consecutive trials per day at the initiation of the dietary/exercise intervention (days 1 and 2) and after intervention (days 37 and 38). Data were expressed as the average performance across all 8 trials at pre-
intervention (pre) and the average performance across all 8 trials at post-intervention (post).

**Gastrocnemius gene expression**

Total RNA was isolated from frozen gastrocnemius samples by TRizol (Invitrogen, Carlsbad, Calif., USA) according to the manufacturer’s instructions. RNA was reverse-transcribed to cDNA using a commercially available high-capacity cDNA reverse transcription kit (Applied Biosystems, Carlsbad, Calif., USA). The cDNA was stored at −20 °C until gene expression analysis.

Gene expression analysis was performed by a Taqman Low Density Array (TLDA, Applied Biosystems, Carlsbad, Calif., USA) according to the manufacturer’s instructions. A total of 1000 ng of cDNA was loaded per sample. Commercially available, prevalidated primers (Applied Biosystems) were used for TLDA analysis (Supplementary Table S1†). TLDA cards were run on a high-throughput real-time polymerase chain reaction system (7900HT, Applied Biosystems), and threshold cycle (Ct) values were determined using the SDS 2.4 and RQ Manager 1.2.1 software packages (Applied Biosystems). All samples were run in duplicate, and gene expression was expressed relative to the housekeeping gene (Gapdh) as fold change from the referent Ctrl-SED group by the 2−ΔΔCt method.

**4-hydroxynonenal assay**

The level of 4-hydroxynonenal (4-HNE) in gastrocnemius extracts was analyzed by enzyme-linked immunosorbant assay (ELISA) according to the manufacturer’s instructions (OxiSelect HNE Adduct ELISA Kit, Cell Biolabs, San Diego, Calif., USA). Gastrocnemius samples were homogenized in sterile phosphate-buffered saline (PBS) supplemented with 5 μL of protease inhibitor cocktail (Sigma–Aldrich, St. Louis, Mo., USA) and stored at −80 °C until analysis. Total protein content in homogenized samples was quantified by DC Protein Assay (Bio-Rad, Hercules, Calif., USA), and samples were diluted in PBS to a concentration of 10 μg protein·mL−1 for use in the ELISA kit. All samples assayed for 4-HNE were tested in duplicate. Color development was read at 450 nm on a spectrophotometric plate reader (BioTek, Winooski, VT, USA).

**Sample preparation for muscle carnosine/β-alanine detection**

All reagents used, except those specifically mentioned, were obtained from Sigma–Aldrich. Ethanol (200 proof) was from Decon Laboratories (King of Prussia, Penn., USA).

Muscle samples were prechilled with liquid nitrogen and pulverized using a tissue pulverizer (BioSpec Products, Bartlesville, Okla., USA). Analytes were extracted using a multistage extraction procedure. During the first stage, 200 μL of absolute ethanol containing 25 mg/L butylated hydroxytoluene (BHT) was added to the tube containing the pulverized, frozen, and weighed tissue. Tissue samples were manually stirred/homogenized with 10–15 strokes of pestle, incubated for 15 min on ice, and centrifuged at 21 000g using an Eppendorf 5810R refrigerated benchtop centrifuge (Eppendorf, Hauppauge, N.Y., USA) for 10 min at 4 °C. Supernatant was collected and 100 μL of water containing 25 mg/L BHT was added to the pellet. A 15-min incubation was followed by 10 min of centrifugation at 21 000g. Supernatant was collected and combined with supernatant from the first extraction. The last step of analyte extraction from the remaining pellet resembles the first step except that only 50 μL of ethanol containing 25 mg/L BHT was added. All 3 supernatants were combined and analyzed by liquid chromatography–mass spectrometry (LC-MS) within 12 h. Unless otherwise indicated, each procedure was performed either on ice or at 4 °C.

**Quantitative measurement of analytes using multiple reaction monitoring (MRM)**

An MRM assay was performed using an atmospheric pressure ionization ultrahigh performance liquid chromatography–triple quadrupole–mass spectrometry (UHPLC-TQ-MS) system from Bruker (Bruker Daltonics, Billerica, Mass., USA) consisting of an Advance UHPLC module and an EVOQ Elite triple quadrupole-mass spectrometer. A Kinetex 2.6 μm HILIC 50 × 2.1 mm internal diameter column (Phenomenex, Torrance, Calif., USA) was used for LC analyte separation. Mobile phase A was composed of 50 mmol/L ammonium acetate (pH 4) and acetonitrile (5/95, v/v). Mobile phase B consisted of acetonitrile, water, and 50 mmol/L ammonium acetate (pH 4) at a ratio of 50/40/10 (v/v). The flow rate was set to 250 μL/min. The LC conditions were as follows (time in minutes/mobile phase A %/mobile phase B %): 0.00/100/0, 0.05/100/0, 1.10/5/95, 5.30/5/95, 5.40/100/0.

An atmospheric-pressure chemical ionization source was used in MS/MS analysis. The analysis was performed using positive ionization polarity. Monitored MRM transitions for β-ALA were 90→72.2, 90→45.2, and 90→43.2. Transitions for carnosine were 227→110.1, 227→156.1, and 227→122. The spray current was 15 μA. Temperatures of cone and heated probe were set to 150 °C and 450 °C. Gas flows were 10 units on cone, 15 units on probe, and 60 units on nebulizer.

*Stock solutions containing standards of interest — β-ALA and carnosine — were prepared in ethanol. The stock solutions were diluted with mobile phase A (described above), creating a set of samples with concentrations of the analyte standards ranging from ppb to ppm. Absolute concentrations of the analytes were determined using the standard addition approach to reduce the effects of sample matrix on the measured values without the use of stable isotope labeled standards. Briefly, equal volumes of various concentrations of standards were spiked into sets of aliquots of 2 muscle samples, which were measured along with one aliquot of each sample that was spiked with only mobile phase A. Concentrations of 2 ppm, 5 ppm, 10 ppm, 20 ppm, and 50 ppm for the carnosine standard and 100 ppb, 200 ppb, 500 ppb, 1 ppm, and 2 ppm for the β-ALA standard were spiked into the aliquots. For each aliquot, 2 technical replicates were acquired.*

**Data analysis**

Body weight, body composition, and grip strength were analyzed by repeated measures analysis of variance (ANOVA) in a 2 × 2 × 2 design (time × diet × activity). Food and water disappearance as well as running distance were analyzed by factorial ANOVA using a 2 × 2 (diet × activity) design. Grip strength ratios and rotarod performance scores were also analyzed by factorial ANOVA using a 2 × 2 (diet × activity) design. Treadmill time to fatigue was analyzed by 2 × 2 (diet × activity) ANOVA on rank-transformed data owing to non-normality of the data and the failure of standard transformations to transform the data to normal.

MRM data were processed using Bruker MS software (Bruker MS Workstation 8.1 for EVOQ). The software was used to calculate and process peak areas and plot them on calibration curves, and final statistics were calculated in Excel. The concentrations of β-ALA and carnosine in each sample were normalized to the sample’s wet weight.

For data gene expression and 4-HNE levels in the gastrocnemius were analyzed by factorial ANOVA using a 2 × 2 (diet × activity) design. Mice displaying splenomegaly indicative of immune activation at tissue collection were excluded from these analyses (N = 2, one each from Ctrl-SED and EGCG+β-ALA-SED). For all ANOVA analyses in this study, post hoc mean separation was performed by Tukey’s HSD in the event of a significant main effect or interaction. Because no significant interactions were detected in our data, we have reported only post hoc tests for main effects in the Results section. All data analysis was performed using IBM SPSS Statistics for
Table 1. Descriptive data.

<table>
<thead>
<tr>
<th></th>
<th>Ctrl-SED</th>
<th>EGCG+β-ALA-SED</th>
<th>Ctrl-VWR</th>
<th>EGCG+β-ALA-VWR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1 body weight (g)</td>
<td>31.1±0.5</td>
<td>30.3±0.6</td>
<td>30.5±0.6</td>
<td>30.7±0.5</td>
</tr>
<tr>
<td>Day 41 body weight (g)</td>
<td>30.0±0.6</td>
<td>29.1±0.6</td>
<td>27.0±0.5</td>
<td>27.5±0.5*</td>
</tr>
<tr>
<td>Food disappearance (g·day⁻¹)</td>
<td>3.4±0.1</td>
<td>3.5±0.1</td>
<td>3.6±0.1</td>
<td>3.6±0.1</td>
</tr>
<tr>
<td>Water disappearance (g·day⁻¹)</td>
<td>2.7±0.2</td>
<td>3.4±0.2</td>
<td>3.1±0.2</td>
<td>3.3±0.2</td>
</tr>
<tr>
<td>Running distance (km·day⁻¹)</td>
<td>—</td>
<td>—</td>
<td>4.8±0.8</td>
<td>4.5±0.4†</td>
</tr>
<tr>
<td>Lean mass pre-intervention (% body weight)</td>
<td>83.7±0.8</td>
<td>83.1±0.8</td>
<td>83.6±0.8</td>
<td>82.8±0.7</td>
</tr>
<tr>
<td>Lean mass post-intervention (% body weight)</td>
<td>83.9±1.0</td>
<td>85.3±0.9</td>
<td>84.0±0.6</td>
<td>84.0±0.7</td>
</tr>
<tr>
<td>Fat mass pre-intervention (% body weight)</td>
<td>10.9±0.8</td>
<td>10.4±0.8</td>
<td>11.2±0.8</td>
<td>11.4±0.7</td>
</tr>
<tr>
<td>Fat mass post-intervention (% body weight)</td>
<td>9.3±1.0</td>
<td>9.0±0.7</td>
<td>7.8±0.8</td>
<td>8.3±0.7</td>
</tr>
</tbody>
</table>

Note: Ctrl, control diet; EGCG+β-ALA, control diet supplemented with (−)-epigallocatechin-3-gallate and β-alanine; SED, sedentary; VWR, voluntary wheel running. N = 10–15/group for all variables. Data are means ± SE.

Results

Descriptive data and muscle carnosine/β-ALA

Lean and fat mass and other descriptive data have been published previously (Gibbons et al. 2014) and are summarized here. There were no differences in body weights between groups prior to intervention. VWR reduced body weights from pre- to post-intervention (F[1,52] = 4.691, p = 0.035, not shown). Post hoc analysis revealed that body weights were decreased from pre- to post-intervention in both Ctrl-VWR and EGCG+β-ALA-VWR groups (p < 0.05, not shown). Table 1 shows lean and fat mass pre- and post-intervention. For percent fat mass, there was a significant time × VWR interaction (F[1,52] = 6.062, p = 0.017). Post hoc analysis indicated a near-significant reduction in fat mass percentage from pre- to post-intervention in the Ctrl-VWR group (p = 0.057). Percent lean mass did not differ between groups at either time point. The dietary treatment increased muscle concentration of carnosine at sacrifice, but there was no significant effect of exercise (Ctrl-SED, 2.45 ± 0.15 µg/mg; EGCG+β-ALA-SED, 2.85 ± 0.21 µg/mg; Ctrl-VWR, 1.76 ± 0.32 µg/mg; EGCG+β-ALA-VWR, 2.91 ± 0.15 µg/mg; diet main effect, F[1,140] = 12.498, p = 0.003, N = 5/group). Gastrocnemius β-ALA did not differ between groups.

Muscle function tests

There was a time × VWR (F[1,52] = 7.360, p = 0.009) and a near-significant EGCG+β-ALA × VWR (F[1,52] = 3.753, p = 0.058) interaction for grip strength from pre- to post-intervention (Fig. 1A). Post hoc analysis revealed no significant differences when grip strength was expressed as absolute force. Normalization of grip strength to lean mass for each individual mouse at each time point did not alter interpretation of these data (not shown). When forelimb grip strength was analyzed on a relative basis by calculating the ratio of post-intervention to pre-intervention values (Fig. 1B), there was an effect of VWR (F[1,52] = 6.761, p = 0.012). VWR maintained maximal grip strength from pre- to post-intervention compared with SED. EGCG+β-ALA alone or in combination with VWR had no effect on grip strength from pre- to post-intervention. Pre-intervention absolute grip strength was not significantly different between groups (one-way ANOVA, F[3,63] = 0.612, p = 0.610, not shown).

In the treadmill test to exhaustion, there was a main effect of VWR (F[1,52] = 12.932, p = 0.001) such that VWR mice ran longer than their SED counterparts (Fig. 2A). EGCG+β-ALA had no effect on performance on the treadmill test in either VWR or SED groups, nor was there synergy between EGCG+β-ALA and VWR.

In the rotarod test, there was a main effect of VWR (F[1,52] = 6.115, p = 0.017) and a trend toward an effect for EGCG+β-ALA (F[1,52] = 2.814, p = 0.099) such that VWR increased and EGCG+β-ALA
decreased average rotarod performance compared with the respective SED or Ctrl counterpart groups (Fig. 2B).

**Gastrocnemius gene expression**

VWR increased expression of Il6, the gene encoding interleukin (IL)-6 ($F_{[1,49]} = 13.410, p = 0.001$, Fig. 3A) within the gastrocnemius muscle. There were no significant main effects or interactions for other cytokine genes (Fig. 3A) including Il1b (IL-1β) and Tnf (tumor necrosis factor). Expression of Il10 was negligible in muscles from all treatment groups (mean $C_t > 35$, not shown). Interestingly, EGCG+β-ALA increased expression of Il10 in the gastrocnemius and macrophage marker Cd11b ($F_{[1,49]} = 6.268, p = 0.016$, Fig. 3A), but there were no significant main effects or interactions for expression of other macrophage marker genes (not shown), including Itgam (CD11c), Mrc1 (CD206), and Retnla (cysteine-rich secreted protein FIZZ1). Neither EGCG+β-ALA nor VWR affected gene expression of chemokines for macrophages (C2c, monocyte chemotactrant protein 1) or neutrophils (Cxcl1, keratinocyte chemotactrant) in the gastrocnemius (data not shown).

For genes associated with oxidative stress (Fig. 3B), there was a main effect of VWR for Sod1, the gene encoding superoxide dismutase (SOD) 1 ($F_{[1,49]} = 4.193, p = 0.046$), as well as a near-significant main effect of VWR for Gpx1, which encodes glutathione peroxidase 1 ($F_{[1,49]} = 3.750, p = 0.059$), in which VWR increased expression of both of these genes. There was a trend for EGCG+β-ALA to increase expression of Sod2 (SOD2), although this effect did not reach significance ($F_{[1,49]} = 3.423, p = 0.070$). Neither EGCG+β-ALA nor VWR affected expression of Nos2, the gene encoding nitric oxide synthase 2.

Both VWR and EGCG+β-ALA affected expression of several genes associated with important intracellular signaling pathways (Fig. 3C). There were significant main effects for both EGCG+β-ALA ($F_{[1,49]} = 4.802, p = 0.033$) and VWR ($F_{[1,49]} = 6.672, p = 0.013$) for Ppargc1a, the gene encoding PGC-1α, such that both EGCG+β-ALA and VWR increased expression of this gene compared with normal chow-fed or sedentary mice, respectively. Likewise, EGCG+β-ALA increased expression of Sirt1 (NAD-dependent protein deacetylase sirtuin-1; $F_{[1,49]} = 4.481, p = 0.039$), while there was a near-significant effect of VWR ($F_{[1,49]} = 3.805, p = 0.057$) in increasing expression of this gene. VWR also increased expression of Foxx3 (forkhead box protein O3, FOXO3; $F_{[1,49]} = 8.446, p = 0.005$). There was no significant main effect or interaction with respect to expression of Prkg2, a gene encoding the gamma-2 subunit of 5′-AMP-activated protein kinase, although EGCG+β-ALA tended to increase expression of this gene ($F_{[1,49]} = 2.849, p = 0.098$).

Finally, VWR increased expression of Bdnf, the gene encoding brain-derived neurotrophic factor (BDNF; $F_{[1,49]} = 7.568, p = 0.008$, Fig. 3D). There was also a significant main effect of EGCG+β-ALA for Igl1 expression (insulin-like growth factor (IGF) I), such that EGCG+β-ALA increased expression of this gene in the gastrocnemius ($F_{[1,49]} = 9.451, p = 0.003$, Fig. 3D). Neither EGCG+β-ALA nor VWR affected expression of Vegfa, the gene encoding vascular endothelial growth factor A (data not shown).

**Oxidative stress**

Analysis of 4-HNE as a measure of oxidative stress in gastrocnemius samples revealed a significant EGCG+β-ALA × VWR interaction ($F_{[1,52]} = 5.583, p = 0.022$) and a significant main effect of VWR ($F_{[1,52]} = 5.815, p = 0.019$) such that VWR reduced 4-HNE in the gastrocnemius muscle compared with SED controls (Fig. 4). Post hoc mean separation by Tukey’s HSD revealed that Ctrl-VWR mice had reduced 4-HNE compared with Ctrl-SED mice ($p = 0.009$).

**Discussion**

We have demonstrated an effect of voluntary wheel running in improving muscle function and increasing expression of antioxidant genes. Although the results are perhaps unsurprising given the extant literature on exercise and aging, they are relatively novel in the context of aging and VWR. A recent study examining the impact of 4 weeks of VWR in extremely old (28–30 months) C57BL/6j mice showed similar VWR-mediated improvements in grip strength, endurance, and rotarod performance [Graber et al. 2015]. However, the effects of exercise on muscle function tests can be highly strain-dependent [Merritt and Rhodes 2015], and we believe that the current study is the first to demonstrate VWR-mediated improvements in rotarod and grip strength in aged BALB/cByJ mice.

In addition to muscle function tests, we examined gene expression for markers of a number of pathways that might affect muscle (dys)function in aged animals. For ease of presentation, we have grouped these into 4 categories: inflammation, oxidative stress, cellular signaling, and growth factors. Both inflammation [Kalinkovich and Livshits 2015] and oxidative stress [Wohlgemuth et al. 2010] are known to be associated with impaired muscle function in aging, and treatments that reduce inflammation and oxidative stress have been shown to improve muscle growth and function.
function (Marzani et al. 2008; Mourkioti et al. 2006). Likewise, growth factors such as IGF-I are important for promotion of muscle growth and have been shown to decline with age (Kalinkovich and Livshits 2015; Sharples et al. 2015). Finally, both mitochondrial function (Marzetti et al. 2013) and autophagy (Wohlgemuth et al. 2010) have been shown to be dysregulated in skeletal muscle during aging, and we examined expression of markers related to these pathways as well.

Aerobic exercise training has been previously shown to reduce oxidative stress (Leeuwenburgh and Heinecke 2001), and in this study we have demonstrated a significant increase in expression of the antioxidant gene \textit{Sod1} as well as a significant decrease in gastrocnemius levels of 4-HNE, a product of lipid peroxidation used here as a tissue marker of overall oxidative stress. \textit{Sod2} was unaffected by these treatments, although there was a trend for exercise to upregulate the expression of this gene. Both \textit{Sod1} and \textit{Sod2} were examined because \textit{Sod1} is abundant in the cytosol, while \textit{Sod2} is primarily present in the mitochondria (Fukui and Zhu 2010).

Fig. 3. Gastrocnemius gene expression of (A) inflammatory markers, (B) oxidative stress markers, (C) intracellular signaling molecules, and (D) growth factors. Ctrl, control diet; EGCG+\textbeta-ALA, control diet supplemented with (−)-epigallocatechin-3-gallate and \textbeta-alanine; SED, sedentary; VWR, voluntary wheel running. †, Significant main effect of VWR (\(p < 0.05\)); ‡, significant main effect of EGCG+\textbeta-ALA (\(p < 0.05\)). \(N = 12\text{–}14\)/treatment combination. Data are means ± SE.

Fig. 4. Gastrocnemius 4-hydroxynonenal (4-HNE). Ctrl, control diet; EGCG+\textbeta-ALA, control diet supplemented with (−)-epigallocatechin-3-gallate and \textbeta-alanine; SED, sedentary; VWR, voluntary wheel running. †, Significant main effect of VWR (\(p = 0.019\)). \(N = 12\text{–}15\)/group. Data are means ± SE.
Aerobic exercise is also well known to increase mitochondrial biogenesis (Pilegaard et al. 2003), and in this study we demonstrated a significant increase in expression of Pparacl in the gastrocnemius of VWR mice. We additionally noted increased expression of Bdnf, the gene encoding BDNF, which acts to increase neurogenesis and angiogenesis (Kermani and Hempstead 2007). BDNF and II-6, gene expression of which was also increased in our study, additionally increase myogenes and fat oxidation in muscle (Pedersen 2013), and both have been routinely shown to be upregulated with exercise (Pedersen 2013). However, Il6 is generally upregulated only after acute exercise (Pedersen 2013); thus, the finding of an upregulation of Il6 expression 12 h after the last exercise bout is surprising. The less than 2-fold upregulation of Il6 in this study, while still significant, is far less striking than that induced by acute exercise, and this may potentially be explained by increased cage activity in VWR mice (although this is speculative, as cage activity was not monitored).

Interestingly, VWR also increased expression of Foxo3, the gene encoding FOXO3. FOXO3 is a transcription factor that has been implicated in a number of physiological processes. For example, FOXO3 has been demonstrated to promote DNA repair and antioxidant enzyme function (Kops et al. 2002) and to upregulate gluconeogenesis (Barthel et al. 2005). Additionally, FOXO3 enhances autophagy in atrophying muscle cells (Zhao et al. 2007), and stress-induced autophagy has been shown to be essential to muscle protein and glucose homeostasis during exercise (He et al. 2012).

A limitation of this study was the measurement of gene expression at a single time point. This provides only a snapshot of overall physiological activity and may not completely reflect downstream protein synthesis or functional activity. Additionally, mRNA levels likely vary considerably within subjects across the duration of the treatment. However, measurement of mRNA in mice requires sufficient tissue mass so as to necessitate euthanasia, and the need to maintain statistical power for behavioral testing precluded euthanizing mice for gene expression analysis at multiple time points in this study.

In addition to the described alterations in gastrocnemius gene expression, VWR also improved physical function as defined by increased performance by VWR mice in forelimb grip strength, the rotarod test, and a treadmill test to fatigue. Although these findings are not novel, the combination of VWR-induced improvements in muscle function and increases in gene expression does serve to underscore the utility of VWR as an appropriate model for aerobic exercise training. There has long been debate, both formal and informal, about the propriety of VWR as an exercise modality, given the nature of the activity in mice. Consistent with our study, mice routinely run multiple kilometers in a 24-h period, while forced treadmill-trained mice generally run less than 400 m per day.

It is often argued that forced treadmill training offers a better model of human exercise, given the greater volume and reduced intensity of VWR. Despite this, VWR still yields many of the same positive changes seen with forced treadmill exercise, including adaptations in cardiac and skeletal muscle (Allen et al. 2001) consistent with other endurance exercise modalities. Our findings reinforce the need for careful selection of an exercise modality in rodent exercise studies, as recent reports have demonstrated differential effects of VWR and forced treadmill running (Cook et al. 2013), despite these strategies sharing many of the same adaptive benefits.

In contrast to VWR, dietary supplementation of EGCG and β-ALA had little effect on the parameters measured in this study. The dietary intervention increased expression of 2 signaling molecule genes, Pparacl and Sirt1, indicating that our dietary intervention may increase mitochondrial biogenesis in aged muscle irrespective of exercise training. EGCG+β-ALA had no effect on expression of antioxidant genes or on gastrocnemius oxidative stress as measured by 4-HNE.

Perhaps the most interesting effect of our dietary intervention is the increase in expression of Ilgam, the gene encoding CD11b, a common marker for macrophages as well as other leukocytes (Solovyov et al. 2005). Although gene expression by itself is insufficient to demonstrate an increase in gastrocnemius macrophage content, it does suggest that our dietary intervention may act to increase either macrophage influx or proliferation in skeletal muscle. Because this hypothesis is tangential to our main aims in this study, we did not pursue the finding further, but it offers an interesting potential avenue for future exploration in this area. The observed increased expression of Ilgam may also partially explain the dietary intervention’s effect of increasing expression of ifg, as macrophages have previously been demonstrated to produce IGF-I and to be involved in muscle injury repair (Lu et al. 2011).

Although we noted that the combined intervention (EGCG+β-ALA-VWR) resulted in the greatest expression of several genes in absolute terms (most notably Pparacl, Sirt1, and Foxo3), the lack of statistically significant activity × diet interactions for these genes as well as the relatively low number of significant main effects of diet give us no evidence to support our hypothesis that the combined VWR and EGCG+β-ALA diet treatment acted synergistically to improve outcomes in this study. Thus, we conclude that given the parameters of our study design, a combination diet including supplemental β-ALA and EGCG is ineffective in improving muscle function in aged mice.

Alternatively, studies may show different results. In rodent studies examining the impact of EGCG on skeletal muscle, doses range from 1 mg kg\(^{-1}\)day\(^{-1}\) (Chen et al. 2009) to 1500 mg kg\(^{-1}\)day\(^{-1}\) (Friedrich et al. 2012). Additionally, delivery methods vary widely, with strategies such as oral gavage (Senthil Kumaran et al. 2008) and supplementation in the drinking water (Chen et al. 2009) being most common. The number of studies that have incorporated EGCG into the diet is relatively limited, especially in relation to the effects of EGCG on skeletal muscle. However, successful studies using diet-incorporated EGCG that target skeletal muscle have reported EGCG concentrations (\(\mu g/g\)) in the diet of 0.32% (Sae-Tan et al. 2011) and 0.5% and 1% (Friedrich et al. 2012), higher than the concentration of 0.135% used in this study. Moreover, these studies used high-fat diet feeding, which may also explain the efficacy of EGCG.

Likewise, despite the vast literature examining β-ALA and muscle, surprisingly little information is available on β-ALA supplementation in rodents. A recent study supplemented mice with 0.6% or 1.2% β-ALA (\(\mu g/\mu l\)) in drinking water for 8 weeks and demonstrated increased muscle carnosine and enhanced fatigue resistance at the higher dose (Eiveraert et al. 2013). This represents a dosage approximately 3.5-fold greater than that used in our study (~1420 mg kg\(^{-1}\)day\(^{-1}\) vs. 414 mg kg\(^{-1}\)day\(^{-1}\)), and the dietary treatment was applied for twice as long (8 weeks vs. 4 weeks). While we noted no deficit in treadmill or rotarod performance due to dietary intervention in either SED or VWR mice, it is possible that paresthesia (“pins and needles”) obscured a potential β-ALA benefit to performance on these tests. It is unclear to what extent mice are affected by this phenomenon, but β-ALA supplementation in drinking water at a dose of 40 mg kg\(^{-1}\) was sufficient to induce itching behavior in mice in a recent study (Liu et al. 2012).

Conclusions

In this study, we demonstrated significant positive effects of voluntary wheel running on several muscle function measures in aged mice. Additionally, wheel running enhanced gastrocnemius expression of a variety of genes known to be related to antioxidant and biogenic mechanisms in skeletal muscle. In contrast, a diet containing epigallocatechin gallate and β-alanine had no effect on muscle function and caused only a slight increase in expression of several genes related to mitochondrial biogenesis.
Furthermore, we found no synergistic effects when dietary and exercise interventions were combined.

Conflict of interest statement
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