

LJMU Research Online

Conceicao, MS, Traina Chacon-Mikahil, MP, Telles, GD, Libardi, CA, Mendes Junior, EM, Vechin, FC, Lugnani De Andrade, AL, Gaspari, AF, Brum, PC, Cavaglieri, CR, Serag, S, Spiegelman, BM, Hawley, JA and Camera, DM

Attenuated PGC-1 alpha Isoforms following Endurance Exercise with Blood Flow Restriction

http://researchonline.ljmu.ac.uk/id/eprint/5434/

Article

Citation (please note it is advisable to refer to the publisher's version if you intend to cite from this work)

Conceicao, MS, Traina Chacon-Mikahil, MP, Telles, GD, Libardi, CA, Mendes Junior, EM, Vechin, FC, Lugnani De Andrade, AL, Gaspari, AF, Brum, PC, Cavaglieri, CR, Serag, S, Spiegelman, BM, Hawley, JA and Camera, DM (2016) Attenuated PGC-1 alpha Isoforms following Endurance Exercise with

LJMU has developed LJMU Research Online for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact researchonline@ljmu.ac.uk

http://researchonline.ljmu.ac.uk/

http://researchonline.ljmu.ac.uk/

- 1 Running Title: PGC-1α isoform and blood flow restriction
- 2

3

4	
5	Miguel S. Conceição ¹ ; Mara P.T Chacon-Mikahil ¹ ; Guilherme D. Telles ¹ ; Cleiton A.
6	Libardi ² ; Edson M. M. Junior ¹ ; Felipe C. Vechin ³ ; André L. Andrade ¹ ; Arthur F.
7	Gáspari ¹ ; Patrícia C. Brum ³ ; Cláudia R. Cavaglieri ¹ ; Sara Serag ⁴ ; Bruce M.
8	Spiegelman ⁴ ; John A. Hawley ^{5,6} ; Donny M. Camera ⁵ .
9	
10	¹ Faculty of Physical Education, University of Campinas – Campinas/Brazil; ² Laboratory
11	of Neuromuscular Adaptations to Resistance Training, Department of Physical
12	Education, Federal University of São Carlos – São Carlos/Brazil; ³ School of Physical
13	Education and Sport, University of São Paulo, São Paulo/Brazil; ⁴ Department of Cell
14	Biology, Dana-Farber Cancer Institute, Harvard Medical School, Boston/USA; ⁵ Mary
15	MacKillop Institute for Health Research, Centre for Exercise and Nutrition, Australian
16	Catholic University, Melbourne, Australia; ⁶ Research Institute for Sport and Exercise
17	Sciences, Liverpool John Moores University, Liverpool, United Kingdom
18	
19	Author for correspondence

Attenuated PGC-1a isoforms following endurance exercise with blood flow restriction

- 20 Miguel S. Conceição
- 21 Faculty of Physical Education, University of Campinas
- 22 Av. Érico Veríssimo, 701, Cidade Universitária "Zeferino Vaz" Barão Geraldo,
- 23 Campinas, São Paulo, Brazil. CEP: 13083-851
- 24 Email: conceicao.miguel0106@gmail.com
- 25 Phone: + 55 19 35216625

26 ABSTRACT

27 Introduction: Exercise performed with blood flow restriction simultaneously enhances the acute responses to both myogenic and mitochondrial pathways with roles in training 28 29 adaptation. We investigated isoform-specific gene expression of the peroxisome proliferator-activated receptor gamma (PPARG) coactivator 1 and selected target genes 30 and proteins regulating skeletal muscle training adaptation. Methods: 9 healthy, 31 32 untrained males participated in a randomized, counter-balanced, cross-over design in which each subject completed a bout of low-intensity endurance exercise performed 33 with blood flow restriction (15 min cycling at 40% of VO_{2peak}, BFR-EE), endurance 34 exercise (30 min cycling at 70% of VO_{2peak}, EE) or resistance exercise (4 x 10 35 repetitions of leg press at 70% of 1-repetition maximum, RE) separated by at least one-36 week recovery. A single resting muscle biopsy (vastus lateralis) was obtained two 37 38 weeks before the first exercise trial (rest) and 3 h after each boat. Results: Total PGC-1a mRNA abundance, along with all four isoforms, increased above rest with EE only 39 40 (P<0.05) being higher than BFR-EE (P<0.05). PGC-1a1, 2 and 4 were higher after EE compared to RE (P<0.05). EE also increased VEGF, Hif-1a and MuRF-1 mRNA 41 abundance above rest (P<0.05) while COXIV mRNA expression increased with EE 42 compared to BFR-EE (P<0.05). Conclusion: The attenuated expression of all four PGC-43 1α isoforms when endurance exercise is performed with blood flow restriction suggests 44 this type of exercise provides an insufficient stimulus to activate the signaling pathways 45 governing mitochondrial and angiogenesis responses observed with moderate- to high 46 47 intensity endurance exercise.

48

Key words: mitochondrial biogenesis; cell signalling; skeletal muscle; adaptation;angiogenesis; high intensity exercise

51 Introduction

52 Skeletal muscle is a highly malleable tissue that can alter its phenotype according to the contractile stimulus imposed (39). For instance, moderate-intensity 53 54 (i.e., <65% of peak oxygen uptake [VO_{2peak}]) endurance exercise training enhances whole-body $\dot{V}O_{2peak}$ (4), increases the maximal activities of oxidative enzymes, and 55 56 shifts patterns of substrate selection from carbohydrate- to fat-based fuels (18). In 57 contrast, strenuous (80% of one repetition maximum [1-RM]) resistance exercise has little or no effects on whole-body VO_{2peak} and oxidative enzyme profiles (11) but 58 increases myofibrillar protein accretion and muscle cross-sectional area (CSA) (30). 59

60

While resistance and endurance exercise could be considered at opposite ends of 61 62 the 'adaptation continuum' by virtue of their divergent biochemical and morphological 63 phenotypes, blood flow restriction during low-intensity endurance exercise (BFR-EE) improves both VO_{2peak}, muscle strength and CSA (1, 2). Abe and co-workers (1) 64 65 reported increased isometric muscle strength, muscle CSA and VO_{2peak} following 8 weeks (24 training sessions) of low-intensity cycle exercise (15 min at 40% VO_{2peak}) 66 performed with BFR-EE compared to same exercise undertaken without BFR. While 67 68 these adaptation responses are considerably lower in magnitude relative to conventional endurance and resistance training performed at higher intensities, the local hypoxia 69 induced by BFR appears to induce an additive 'metabolic stressor' that perturbs cellular 70 homeostasis (17) and concomitantly enhances both anabolic and oxidative adaptations. 71

72

The cellular mechanisms mediating adaptation responses to exercise are complex involving the cross talk of several intracellular signaling systems that ultimately form the basis for specific phenotypic responses with divergent contractile 76 modes (17). The transcriptional co-activator Peroxisome proliferator-activated receptor 77 gamma (PPARG) coactivator 1 alpha (PGC-1a) is a 'master regulator' of many endurance exercise-induced adaptations by virtue of its central role in promoting 78 79 mitochondrial biogenesis, angiogenesis, and inflammatory proteins (17). Transcription of the PGC-1 α gene has been shown to be under the control of several promoter regions 80 with activation of the alternative PGC-1 α 1 promoter resulting in the transcription of 81 three additional isoforms: PGC-1 α 2, - α 3 and- α 4. Ruas and colleagues (32) recently 82 demonstrated a preferential increase in the PGC1-a4 isoform following resistance 83 exercise in human skeletal muscle. However, little is known about the regulation of the 84 85 α^2 and α^3 isoforms and, to date, no studies have investigated the expression of all four PGC-1a isoforms to diverse contractile stimuli such as resistance and endurance 86 exercise, or following BFR, in humans. Accordingly, the aim of the present study was 87 88 to compare the acute molecular responses mediated by the different PGC-1 α isoforms following low intensity endurance exercise (BFR-EE), resistance exercise (RE) and 89 90 moderate endurance exercise (EE). As BFR-EE can promote both endurance capacity 91 and muscle hypertrophy responses, we hypothesised EE and RE would selectively increase the expression of the PGC-1 α 1 and α 4 isoforms, respectively. In contrast, we 92 hypothesized that BFR-EE would upregulate a molecular signature involving the 93 increase of both isoforms and their respective anabolic and mitochondrial gene targets. 94

95

96 METHODS

97 Subjects

Nine untrained, healthy male subjects [age 22.4 \pm 3.0 yr, body mass (BM) 73.5 \pm 99 9.7 kg, height 1.79 \pm 0.05 m, maximal oxygen uptake test (VO_{2peak}) 36.8 \pm 4.8 mLlkg⁻¹ 100 ¹·min⁻¹, leg press one repetition maximum (1-RM) 266 \pm 66 kg; values are mean \pm SD] voluntarily participated in this study. The experimental procedures and possible risks
associated with the study were explained to all subjects, who provided written informed
consent before participation. The study was approved by the local University's Ethics
Committee and conducted in conformity with the policy statement regarding the use of
human subjects according to the latest revision of the *Declaration of Helsinki*.

106

107 Experimental Design

108 The study employed a randomized counter-balanced, cross-over design in which each subject completed a bout of either resistance exercise (RE), endurance cycling 109 exercise (EE) or low-intensity cycling exercise combined with blood flow restriction 110 (BFR-EE). Two weeks prior to the first exercise session, a resting muscle biopsy was 111 obtained before participants underwent VO_{2peak} and one-repetition maximum (1-RM) 112 113 testing, and exercise familiarization. Exercise trials were separated by a one-week 114 recovery period during which time subjects maintained their habitual diet and physical activity patterns. 115

116

117 Preliminary Testing

 VO_{2peak} . Participants performed a maximum graded exercise test on a cycle ergometer 118 119 with electromagnetic braking (Quinton modelo: Corival 400, Lode BV, Groningen, 120 Netherlands) based on a protocol used in previously published paper that investigated BFR-EE (1). Briefly, after resting on the bike for 5 min, participants commenced the 121 incremental test protocol. Briefly, subjects commenced cycling at an initial load of 50 122 W for 1 min and the workload was increased by 15 W/min until a workload of 200 W 123 124 was reached, after which further increases were 10 W/min increments. The test continued until voluntary exhaustion, defined by two of the three following criteria: 125

 VO_{2peak} plateau (< 2.1 mL.kg⁻¹.min⁻¹ of variation), > 1.10 respiratory exchange ratio, 126 and/or heart rate higher than 90% of maximum estimated from age (19). Gas exchange 127 data were collected continuously using an automated breath-by-breath metabolic system 128 129 (CPX, Medical Graphics, St. Paul, Minnesota, USA) and the highest oxygen consumption value was defined as the peak oxygen consumption (VO_{2peak}) over any 30 130 131 sec period. To confirm the appropriateness of this protocol for this study we performed 132 a pilot study to verify repeatability in VO_{2peak} measures and observed a strong repeatability in VO_{2peak} (3.0%), power (1.9%), respiratory exchange ratio (RER) (5.6%), 133 and time to exhaustion (1.6%) measures. 134

135

136 Maximal Strength

137 The one-repetition maximum (1-RM) test was performed on a leg press machine 138 (45° leg press, G3-PL70; Matrix, São Paulo, Brazil) as previously described (8). Briefly, 139 participants performed a 5 min warm-up on a cycle ergometer riding at 25 W. 140 Participants then undertook 1 x 10 repetitions at 50% of their estimated 1-RM, followed 141 by 1 x 3 repetitions at 70% of the estimated 1-RM with 1-min rest between sets. Participants then performed a series of single repetitions until the maximum load (1-142 RM) lifted was established with fully eccentric-concentric movement with 90° range of 143 144 motion. Repetitions were separated by a 3-min recovery and were used to establish the 145 maximum load/weight that could be moved through the full range of motion once, but 146 not a second time.

147

148 Diet/Exercise Control

Before each experimental trial (described subsequently), subjects were instructedto refrain from exercise training and vigorous physical activity, and alcohol and caffeine

151 consumption for a minimum of 48 h. Subjects were provided with standardized 152 prepacked meals that consisted of 3 g carbohydrate/kg body mass (BM), 0.5 g 153 protein/kg BM, and 0.3 g fat/kg body mass consumed as the final caloric intake the 154 evening before reporting for an experimental trial.

155

156 Experimental Testing Sessions

On the morning of an experimental trial, subjects reported to the laboratory after a ~10-157 158 h overnight fast. After resting in the supine position for ~15 min and under local anaesthesia (2-3 mL of 1% Xylocaine), a resting biopsy was obtained from the vastus 159 160 lateralis using a 5-mm Bergstrom needle modified with suction (7). Approximately 100 mg of muscle was removed, dissected free from blood and connective tissue and snap 161 frozen in liquid nitrogen before being stored at -80° C until subsequent analyses. Due to 162 163 ethical constraints regarding the total number of muscle biopsies allowed, this single resting biopsy was used as a basal control for all subsequent exercise trials. Two weeks 164 165 later participants returned to the laboratory having (after the same pre-trial diet and 166 exercise control) to undertake the first of three randomly assigned exercise sessions (described below). Each exercise trial was separated by a one week wash out. Following 167 the completion of each exercise session, subjects rested for 180 min after which time a 168 169 muscle biopsy was obtained. Subsequent incisions were performed 3 cm proximal to each other. Blood samples were collected before each exercise session and immediately, 170 1, 2 hr and 3 hr post exercise. Blood samples were immediately placed in microtubes 171 172 containing 1% sodium fluoride and then centrifuged at 3000 rpm for 5 min to separate the plasma before being aliquoted and frozen in liquid nitrogen and stored at -80°C. 173

174

175 *Resistance Exercise (RE)*

After a standardized warm-up on a cycle ergometer consisting of 5 min light 176 cycling at 25 W, subjects performed 4 sets of 10 repetitions leg press exercise (45° leg 177 press machine; G3-PL70; Matrix) at 70% of 1-RM. Each set was separated by a 1 min 178 179 recovery period during which time subjects remained seated on the leg press machine. Complete concentric/eccentric movements were performed with 90° of range of motion 180 and strong verbal encouragement was provided during each set. The volume and 181 182 intensity of this session was based on the recommendations of American College Sports 183 Medicine (ACSM) (3). All participants completed every repetition from each respective 184 set.

185

186 Endurance Exercise (EE)

Following a standardized warm up (described previously), subjects performed 30 min of continuous cycling at a power output that elicited ~at 70% of individual VO_{2peak}. Subjects were fan-cooled and provided visual feedback for pedal frequency, power output, and elapsed time were provided to subjects. The volume and intensity of this session were based on the recommendations of ACSM (4). All participants completed the full 30 min session.

193

194 *Low Intensity Blood Flow Restriction* (BFR-EE)

Subjects performed 15 min continuous cycling with a cuff strapped over the thigh at a power output that elicited at 40% of VO_{2peak} , as previously reported (1). An 18-cm wide cuff was placed on the proximal portion of the thigh (inguinal fold region) over the tibial artery and once in position, was inflated until an absence of auditory blood pulse detected through auscultation with a vascular Doppler probe (DV-600; Marted, São Paulo, Brazil). Pressure was then slowly released until the first arterial pulse was detected which was considered the systolic pressure at the tibial artery. Cuff pressure was set at 80% of the maximum tibial arterial pressure and the cuff was inflated through-out the entire exercise session (22).

204

205 Analytical Procedures

206 Blood Lactate

Plasma lactate concentration was measured on a spectrophotometer (ELx800,
Biotek, Winooski, USA) using a commercial kit (Biotecnica, Varginha, Brazil)
according to the manufacturer's protocol.

210

211 RNA Extraction and Quantification

Approximately 20 mg of skeletal muscle was homogenized in TRIzol with chloroform added to form an aqueous RNA phase. This RNA phase was then precipitated by mixing with ice-cold isopropanol alcohol and the resulting pellet was washed and re-suspended in 40 µl of RNase-free water. Extracted RNA was quantified using a NanoDrop 1000 spectrophotometer (Nanodrop Technologies, Wilmington, USA) by measuring absorbance at 260 nm and 280 nm.

218

219 *Reverse Transcription*

First-strand complementary DNA (cDNA) synthesis was performed using commercially available TaqMan Reverse Transcription Reagents (Invitrogen, Melbourne, Australia) in a final reaction volume of 20 μ L. All RNA and negative control samples were reverse transcribed to cDNA in a single run from the same reverse transcription master mix. Serial dilutions of a template human skeletal muscle RNA (AMBION; Cat No AM7982) was included to ensure efficiency of reverse transcription and for calculation of a standard curve for real-time quantitative polymerase chainreaction (RT-PCR).

228

229 *Real-Time PCR*

Quantification (in duplicate) of mRNA was performed using a CFX96 Touch[™] 230 Real-Time PCR Detection System (Bio Rad, California, USA). Tagman-FAM-labelled 231 primer/probes for MuRF-1 (Cat No. Hs00822397_m1), COXIV (Cat 232 No. 233 Hs00971639_m1), IL-6 (Cat No. Hs00985639_m1), PGC-1a (Cat No. Hs01016719_m1), HIF-1a (Cat No. Hs00153153 m1), Myostatin (Hs00976237 m1), 234 IGF-1 (Hs01547656_m1) and VEGF (Cat No. Hs00900055_m1) were used in a final 235 reaction volume of 20 µL. PCR treatments were 2 min at 50 °C for UNG activation, 10 236 min at 95 °C then 40 cycles of 95 °C for 15 s and 60 °C for 60s. Glyceraldehyde-3-237 238 phosphate dehydrogenase (GAPDH) (Cat No Hs02758991_g1) was used as a 239 housekeeping gene and was stably expressed between exercise interventions (data not 240 shown). The relative amounts of mRNAs were calculated using the relative 241 quantification ($\Delta\Delta$ CT) method (24). All Taqman-based PCR experiments were performed in the Centre for Exercise and Nutrition laboratory at the Australian Catholic 242 University. 243

244

245 *Quantification of PGC-1α isoforms*

RNA was extracted from a separate piece of snap frozen muscle (~20 mg) using TRIzol
(Invitrogen) and purified using QIAGEN RNeasy mini-columns. Reverse transcription
was performed using a High Capacity cDNA Reverse Transcription kit (Applied Biosystems). Real-Time Quantitative PCR was carried out in a SYBR Green ER PCR
Master Mix (Invitrogen)/ 384-well format using an ABI PRISM 7900HT (Applied

251 Biosystems). Relative mRNA levels were calculated using the comparative CT method 252 and normalized to cyclophilin mRNA. Primer sequences are as follows: Cyclophilin (forward: GGAGATGGCACAGGAGGAA; reverse: GCCCGTAGTGC TTCAGTTT), 253 PGC1a1 (forward: ATG GAG TGA CAT CGA GTG TGC T; reverse: GAG TCC ACC 254 CAG AAA GCT GT), PGC1a2 (forward: AGT CCA CCC AGA AAG CTG TCT; 255 reverse: ATG AAT GAC ACA CAT GTT GGG), PGC1a3 (forward: CTG CAC CTA 256 GGA GGC TTT ATG C; reverse: CAA TCC ACC CAG AAA GCT GTC T), and 257 258 PGC1a4 (forward: TCA CAC CAA ACC CAC AGA GA; reverse: CTG GAA GAT ATG GCA CAT). All SYBR Green-based PCR experiments were performed in the 259 260 Department of Cell Biology laboratory at the Dana-Farber Cancer Institute, Harvard 261 Medical School (USA).

262

263 Western Blots

Approximately 30 mg of muscle was homogenized in buffer containing 50 mM 264 265 Tris·HCl pH 7.5, 1 mM EDTA, 1 mM EGTA, 10% glycerol, 1% Triton X-100, 50 mM 266 NaF, 5 mM sodium pyrophosphate, 1 mM DTT, 10% µg/ml trypsin inhibitor, 2 µg/ml aprotinin, 1mM benzamidine, and 1 mM PMSF. After determination of protein 267 concentration (Pierce, Rockford, IL), lysate was resuspended in Laemmli sample buffer. 268 269 Lysate was then re-suspended in Laemmli sample buffer with 40 µg of protein loaded 270 onto 4–20% Mini-PROTEAN TGX Stain-Free[™] Gels (Bio Rad, California, USA). Post electrophoresis gels were activated according to the manufacturer's details (Chemidoc, 271 Bio-Rad, Gladesville, Australia) and then transferred to polyvinylidine fluoride (PVDF) 272 membranes. After transfer, a Stain-Free image of the PVDF membranes for total protein 273 274 normalization was obtained before membranes were rinsed briefly in distilled water and blocked with 5% non-fat milk, washed with 10 mM of Tris-HCl, 100 mM of NaCl, and 275

0.02% Tween 20, and incubated with primary antibody (1:1000) overnight at 4 °C. 276 Membranes were incubated with secondary antibody (1:2,000), and proteins were 277 detected via chemiluminescence (Amersham Biosciences, Buckinghamshire, UK; 278 Pierce Biotechnology, Rockford, IL) and quantified by densitometry. All sample time 279 points for each subject were run on the same gel. Polyclonal anti-phospho-mTOR^{Ser2448} 280 (no. 2971), -p70 S6K^{Thr389} (no. 9206), - adenosine monophosphate kinase (AMPK)^{Thr172} 281 (no. 2531), eukaryotic translation initiation factor 4E-binding protein 1 (4E-BP1) 4E-282 BP1^{Thr37/46} (no. 9459), eEF2 eukaryotic translation elongation factor 2 (eEF2) eEF2^{Thr56} 283 (no. 2331) and p53^{Ser15} (no. 9284) were purchased from Cell Signaling Technology 284 285 (Danvers, MA, USA). Volume density of each target protein band was normalized to 286 the total protein loaded into each lane using stain-free technology (15) with data expressed in arbitrary units. Due to low sample availability, phosphorylated proteins 287 288 were unable to be normalized to their respective total protein content and were therefore 289 also normalized to stain-free protein levels.

290

291 Statistical analysis

Statistical analysis was performed using SAS version 9.3 for Windows (SAS 292 Institute Inc., Cary, NC, USA). Data normality and variance equality were assessed 293 294 through the Shapiro-Wilk and Levene tests. One-way ANOVA with repeated measures 295 (factor: condition) was performed for gene and protein expression analyses. A mixed 296 model ANOVA, assuming group and time as fixed factors and subjects as a random factor, was performed for blood lactate data. Tukey post hoc analysis was used for 297 multiple comparison purposes when significant F-values were found. The significance 298 299 level was set at $P \le 0.05$. Data are presented as Mean \pm Standard Deviation (SD).

300

302 **mRNA expression**

303 Total PGC1-α and isoforms

Total PGC-1α mRNA (Figure 1A) increased with EE above rest (P<0.0001), RE 304 (P=0.0013) and BFR-EE (P>0.0001). There was a significant increase in PGC-1 α 1 305 mRNA with EE above rest (P=0.0450), RE (P=0.0069) and BFR-EE (P=0.0349) 306 (Figure 1B). There was also a significant increase in PGC-1 α 2 mRNA (Figure 1C) with 307 308 EE above rest (P<0.0001), RE (P=0.0003) and BFR-EE (P<0.0001). PGC-1α3 mRNA (Figure 1D) increased with EE above rest (P=0.0389). There was also increases PGC-309 310 1α4 mRNA (Figure 1E) with EE above rest (P=0.0035), RE (P=0.0469) and BFR-EE (P=0.0140). 311 312 313 **Figure 1 here** 314 315 VEGF, COXIV, HIF-1a 316 There was a significant increase in VEGF mRNA (Figure 2A) with EE above rest (P=0.0180) and RE (P=0.0069). COXIV mRNA expression increased with EE 317

above BFR-EE (P=0.0550) (Figure 2B). There was a significant increase in HIF-1a

abundance with EE above Rest (P=0.0530) (Figure 2C).

320

321 ******Figure 2 here**

322

323 IL-6 – IGF-1 - Myostatin - MurRF1

324 IL-6, IGF-1 and Myostatin mRNA expression were unchanged post-exercise
325 (Figure 3 A, B, C). There was a post-exercise increase in MuRF1 mRNA abundance

326	with EE above Rest (P=0.0003), RE (P=0.0256) and BFR-EE (P=0.0007) (Figure 3D).
327	
328	**Figure 3 here**
329	
330	Cell Signaling
331	mTOR -p7086K -4E-BP1 -eEF2
332	There were no changes in mTOR ^{Ser2448} , p70S6K ^{Thr389} , 4E-BP1 ^{Thr37/46} or eEF2 ^{Thr56}
333	phosphorylation post-exercise or between exercise groups (Figure 4).
334	
335	**Figure 4 here**
336	
337	АМРК -р53
220	AMDE The 156 and p52 Sec 15 phosphory lation were upshaped post everying (Figure
220	AMPK and p35 phosphorylation were unchanged post-exercise (Figure
339	5).
340	
2.44	<u></u>
341	**Figure 5 here**
342	
343	Plasma lactate concentration
344	Lactate concentration increased above rest immediately post-exercise for all
345	interventions (P<.0001 for all comparisons; Table 1). Lactate concentration remained
346	elevated at 1 h, 2 h and 3 h post-exercise for EE and RE, and 1 h and 2 h for BFR-EE
347	(P<.0001 for all comparisons).
348	

350

351 **Discussion**

352 Low intensity (<50% of VO_{2peak}) endurance training with blood flow restriction has been shown to concomitantly promote isometric muscle strength, muscle CSA and 353 VO_{2peak} (1, 2). While these enhanced adaptation responses are considerably lower in 354 magnitude compared to conventional resistance or endurance exercise performed 355 356 without any blood flow restriction, the underlying molecular mechanisms mediating these responses remain largely undefined. For the first time we report that low intensity 357 358 endurance cycling exercise performed with blood flow restriction failed to increase PGC-1 α expression to that commonly observed with 'conventional' endurance exercise. 359 360 Moreover, we show isoform-specific post-exercise increases in the α 4 isoform along 361 with Hif-1 α and VEGF mRNA expression following higher intensity endurance 362 exercise without blood flow restriction. Taken collectively, our novel findings suggest 363 that cycle exercise undertaken with blood flow restriction is unable to provoke the 364 perturbations to cellular homeostasis necessary to induce activation of the cell signaling events regulating mitochondrial biogenesis and angiogenesis that take place with higher 365 366 intensity endurance exercise without blood flow restriction.

367

A growing body of evidence suggests that exercise undertaken with blood flow restriction can enhance exercise adaptation. A recent meta-analysis reported both low load/intensity resistance (20–30% 1 RM) and aerobic walking exercise performed with blood flow restriction can induce increases in muscle strength and hypertrophy, although with smaller gains compared to high intensity resistance exercise alone (35). However, little is known about the molecular mechanisms mediating these responses

349

when low intensity endurance exercise is undertaken with blood flow restriction. As such, we compared the expression of key gene and protein targets implicated in a range of exercise adaptation responses such as hypertrophy, mitochondrial biogenesis, muscle proteolysis, substrate metabolism and angiogenesis between BFR-EE, and conventional bouts of RE and EE. We particularly focused on the four different full-length PGC-1 α isoforms putatively implicated in anabolic and mitochondrial-related adaptation responses.

381

In agreement with previous studies (5, 23, 29), we observed significant increases in total 382 PGC-1a mRNA following continuous endurance exercise performed at 70% of VO_{2peak}. 383 384 This increase in PGC-1a mRNA was concomitant with greater abundance of VEGF, a target of PGC-1 α (37). However, in contrast to our original hypothesis, this response 385 386 was absent following a bout of low-intensity endurance exercise (40% VO_{2peak}) performed with blood flow restriction. In an attempt to identify possible mechanisms 387 388 responsible for this attenuated PGC-1a response, we investigated IL-6 expression to 389 determine whether an increase in the muscular inflammatory program was implicated in the blunted response. This hypothesis was based on previous data showing an inverse 390 relationship between skeletal muscle PGC-1 α and IL-6 expression (16). However, IL-6 391 392 mRNA expression post-exercise was unchanged in all exercise groups suggesting any 393 acute increase in muscle inflammation caused by BFR-EE was not responsible for the 394 reduced PGC-1a1 expression observed. We also investigated other cellular markers implicated in exercise adaptation responses that can regulate PGC-1 α expression. 395 AMPK is an intracellular 'fuel gauge' that can phosphorylate PGC-1a and increase its 396 397 transcriptional activity (36) while the apoptogenic protein p53 has emerged as another 398 signaling regulator of skeletal muscle exercise-induced mitochondrial biogenesis and

399 substrate metabolism that can translocate to the nucleus upon activation and induce PGC-1 α expression (17). Phosphorylation of either of these protein targets was 400 unaltered post-exercise suggesting other molecular markers and/or physiological 401 402 mechanisms may be responsible for the upregulation of PGC-1 α with high intensity endurance exercise. One plausible explanation for these discrepant findings may be the 403 level of glycogen utilization between exercise sessions in our untrained subjects. We 404 (10) and others (6, 31) have shown greater post-exercise PGC-1 α expression with low-405 406 compared to normal or high glycogen concentration and although we did not measure muscle glycogen use in the current study due to limited muscle tissue availability, the 407 408 longer duration and higher intensity exercise bout is likely to have induced greater glycogen depletion compared to the endurance exercise session performed with blood 409 410 flow restriction.

411

412 Another possible explanation for the discrepancy in PGC-1 α 1 expression between the 413 two endurance-based exercise bouts is the large differences in estimated energy 414 expenditure. Exercise energy expenditure after BFR-EE was ~4 fold less compared to the EE protocol with total energy expenditure positively associated with PGC-1a 415 expression (r=0.73, P=0.039). Increased PGC-1a mRNA expression has been observed 416 417 after 30 min running compared to bouts of 20 and 10 min (37). Thus, total exerciseinduced energy expenditure may be an overriding determinant of PGC-1 α expression 418 responses post-exercise. 419

420

Low intensity endurance exercise with BFR was also unable to induce the expression of
PGC-1α4 compared to higher intensity endurance exercise without blood flow
restriction. The PGC-1α4 isoform has been proposed to promote muscle hypertrophy by

inducing IGF-1 expression and reducing the expression of myostatin, a negative 424 425 regulator of muscle growth (32). The increase in PGC-1a4 mRNA expression with EE in the current study was mirrored by a small, non-significant, increase and decrease in 426 IGF1 and myostatin expression, respectively. Ruas and colleagues were the first to 427 show a selective increase in PGC-1a4 expression (concomitant with decreased 428 myostatin abundance) with resistance compared to endurance exercise in human skeletal 429 muscle (32). However, this expression pattern was observed following 8 weeks whole-430 431 body resistance training. Thus, a limitation of our study is that we only incorporated a single bout of isolated leg press suggesting longer training programs/ exercise stimulus 432 433 may be required to induce this selective PGC-1 α 4 response. Nonetheless, another recent publication reported increased truncated and non-truncated PGC-1a transcripts from 434 both alternative and proximal promoter sites 2 hours following an acute bout of 435 436 resistance exercise that incorporated the same volume and intensity as our study (40). 437 This indicates the resistance exercise bout performed in our study was likely sufficient 438 to induce the appropriate signal to increase the expression of this isoform however 439 potential differences in post-exercise biopsy timing between this study and ours (2 h vs. 3 h) may explain why we did not observe this increase with resistance exercise. 440

441

Increased PGC-1 α 4 and VEGF expression has also been reported in primary myotubes treated under hypoxic conditions suggesting low oxygen conditions to be favorable for the activation of this isoform (38). In the current study, the transcription factor Hif-1 α , a key regulator of angiogenesis in situations of hypoxia (34), was unchanged following BFR-EE, while RE and EE induced 2-fold higher post-exercise changes in lactate compared to BFR-EE. While it is possible a greater metabolic and hypoxic stimulus may be required to increase PGC-1 α 4 signaling, others have reported unchanged blood

lactate following aerobic-based exercise with blood flow restriction (26). Moreover, the 449 450 same occlusion protocol (15 min cycle at 40% VO_{2peak}) has been shown to improve muscle volume and VO_{2peak}, during a chronic training intervention (1). Thus, it is 451 452 possible chronic exposure to this occlusion stimulus may be required to elicit increases in PGC-1 α 4 expression. As this is the first study to investigate changes in Hif-1 α 453 454 following endurance cycling exercise with BFR it is difficult to compare our results to 455 those of previous investigations incorporating resistance exercise and BFR. However, 456 we speculate that when performed with blood flow restriction, the lower contractile intensity associated with 'conventional' endurance compared to resistance (or sprint) 457 458 exercise, provides adequate blood flow to the exercising musculature and adjoining capillary beds in order to prevent tissue de-oxygenation. Further studies comparing 459 different low intensity endurance exercise protocols with resistance exercise that 460 461 incorporate blood flow restriction are required to corroborate this hypothesis.

462

463 Another novel finding from the current study was the post-exercise increases in the 464 PGC-1 α 2 and 3 isoforms. Similar to the α 1 and α 4 isoforms, both PGC-1 α 2 and α 3 increased above rest with higher intensity endurance exercise and were significantly 465 elevated compared to resistance exercise. Both isoforms are expressed in skeletal 466 muscle and brown adipose tissue although little is known about the regulatory targets of 467 these isoforms and their capacity to mediate exercise adaptation responses (27). Based 468 on the elevated response following endurance compared to resistance exercise, we 469 470 propose these isoforms to mediate physiological processes related to mitochondrial biogenesis and substrate metabolism. 471

472

Considering low load endurance exercise with BFR can increase muscle strength and 473 474 hypertrophy (35), we also investigated markers of translation initiation, elongation and muscle proteolysis. Previous studies have reported increases in mTOR and p70S6K 475 476 phosphorylation that have formed the basis for enhanced rates of muscle protein synthesis following resistance exercise with blood flow restriction (13, 14). 477 478 Nonetheless, the phosphorylation status of these proteins as well as 4E-BP1 and eEF2 479 were unchanged 3 h post-exercise in the current study. This is in agreement with the 480 results of Ozaki and colleagues (28) who observed no changes in Akt, mTOR or p70S6K phosphorylation following 20 min treadmill walking performed with blood 481 flow restriction despite a higher intensity exercise bout (55% VO_{2peak}) compared to our 482 protocol. While our study design was somewhat limited by only having the single post-483 484 exercise biopsy (9), this sampling time-point was specifically chosen based on previous 485 studies showing significant, and in some cases maximal, increases in PGC-1a mRNA 486 expression in response to an exercise challenge (5, 23). Future studies investigating 487 endurance exercise undertaken with BFR-EE should include a time-course of signaling 488 responses in order to determine the optimal 'window' for muscle sampling in subsequent investigations. 489

490

Several other factors including the width and pressure of cuff used during BFR must also be considered. Previous studies have reported smaller increases in muscle CSA when lower body resistance training is undertaken with BFR (compared to no BFR) at the site of the cuff (12, 20). While this indicates a narrow cuff may be advantageous for promoting anabolic adaptation responses due to compressing less muscle tissue, a recent study comparing the effects of a wide versus narrow cuff reported similar increases in maximum strength and muscle cross sectional area following 12 weeks of unilateral

elbow flexion performed at 20% of 1RM (21). Also, a recent study showed that there 498 499 was no difference in either muscle strength or hypertrophy between different occlusion pressures (25). Thus, the use of a wider cuff, as used in our protocol, appears unlikely to 500 501 attenuate chronic muscle anabolic responses. Regardless, these studies are currently only limited to BFR with resistance exercise. Future studies comparing these parameters 502 when endurance exercise is performed with BFR are required. Finally, MuRF-1 mRNA 503 expression increased post endurance exercise which resulted in a higher expression 504 505 above endurance exercise with BFR and resistance exercise. MuRF-1 mediates the ubiquitin proteasome system by 'labelling' cleaved myofibril segments for degradation 506 507 (33). It is unclear whether this increase in expression with high intensity endurance exercise represents general tissue remodeling, particularly considering our participants 508 were untrained and the unaccustomed contractile stimulus, or a greater induction of 509 510 protein degradation.

511

512 In summary, this is the first study to investigate the molecular mechanisms mediating 513 muscle adaptation responses to low intensity endurance cycling exercise with blood flow restriction. The attenuated expression of all four PGC-1a isoforms when endurance 514 515 exercise is performed with blood flow restriction suggests this type of exercise is unable 516 to induce the appropriate metabolic perturbation capable of activating the cell signaling 517 machinery responsible for mitochondrial biogenesis and angiogenesis responses with moderate-to-high intensity endurance exercise. Longer training programs incorporating 518 519 endurance exercise with BFR that correlate measurements of these molecular markers with functional adaptation responses such as changes in VO_{2peak} and cycle time to 520 521 fatigue will yield important information to the efficacy of this training method to

522 enhance training adaptation and subsequently improve health outcomes in populations523 that may be unable to perform, prolonged exercise.

524

525 Acknowledgments

526	The authors would like to express gratitude for the FAPESP (2014/00985-0) and
527	FAEPEX for financial support. The writing of this publication was also supported by a
528	Collaborative Research Networks (CRN) grant awarded to J. A. Hawley (2013000443).
529	The authors declare no declare no conflicts of interest. The results of the present study
530	do not constitute endorsement by ACSM.
531	
532	
533	
534	
535	
536	
537	
538	
539	
540	
541	
542	
543	
544	
545	
546	

547 **References**

- Abe T, Fujita S, Nakajima T et al. Effects of Low-Intensity Cycle Training with
 Restricted Leg Blood Flow on Thigh Muscle Volume and VO2MAX in Young
 Men. *J Sports Sci Med.* 2010;9(3):452-8.
- Abe T, Kearns CF, Sato Y. Muscle size and strength are increased following
 walk training with restricted venous blood flow from the leg muscle, Kaatsuwalk training. *J Appl Physiol (1985)*. 2006;100(5):1460-6.
- ACSM. American College of Sports Medicine position stand. Progression
 models in resistance training for healthy adults. *Med Sci Sports Exerc*.
 2009;41(3):687-708.
- ACSM. American College of Sports Medicine position stand. Quantity and
 quality of exercise for developing and maintaining cardiorespiratory,
 musculoskeletal, and neuromotor fitness in apparently healthy adults: guidance
 for prescribing exercise. *Med Sci Sports Exerc*. 2011;43(7):1334-59.
- 5. Bartlett JD, Hwa Joo C, Jeong TS et al. Matched work high-intensity interval
 and continuous running induce similar increases in PGC-1alpha mRNA, AMPK,
- p38, and p53 phosphorylation in human skeletal muscle. *J Appl Physiol (1985)*.

564 2012;112(7):1135-43.

- 565 6. Bartlett JD, Louhelainen J, Iqbal Z et al. Reduced carbohydrate availability
 566 enhances exercise-induced p53 signaling in human skeletal muscle: implications
 567 for mitochondrial biogenesis. *Am J Physiol Regul Integr Comp Physiol.*568 2013;304(6):R450-8.
- 569 7. Bergstrom J. Percutaneous needle biopsy of skeletal muscle in physiological and
 570 clinical research. *Scandinavian journal of clinical and laboratory investigation*.
 571 1975;35(7):609-16.

- Brown LE, Weir JP. Procedures recommendation I: Accurate assessment of
 muscular strength and power. *Journal of Exercise Physiology Online*. 2001;4:121.
- 575 9. Camera DM, Edge J, Short MJ, Hawley JA, Coffey VG. Early time course of
 576 Akt phosphorylation after endurance and resistance exercise. *Med Sci Sports*577 *Exerc.* 2010;42(10):1843-52.
- Camera DM, Hawley JA, Coffey VG. Resistance exercise with low glycogen
 increases p53 phosphorylation and PGC-1alpha mRNA in skeletal muscle. *Eur J Appl Physiol.* 2015;115(6):1185-94.
- 11. Campos GE, Luecke TJ, Wendeln HK et al. Muscular adaptations in response to
 three different resistance-training regimens: specificity of repetition maximum
 training zones. *Eur J Appl Physiol*. 2002;88(1-2):50-60.
- Ellefsen S, Hammarstrom D, Strand TA et al. Blood flow-restricted strength
 training displays high functional and biological efficacy in women: a withinsubject comparison with high-load strength training. *Am J Physiol Regul Integr Comp Physiol.* 2015;309(7):R767-79.
- 588 13. Fry CS, Glynn EL, Drummond MJ et al. Blood flow restriction exercise
 589 stimulates mTORC1 signaling and muscle protein synthesis in older men. *J Appl*590 *Physiol.* 2010;108(5):1199-209.
- 591 14. Gundermann DM, Walker DK, Reidy PT et al. Activation of mTORC1 signaling 592 and protein synthesis in human muscle following blood flow restriction exercise 593 is inhibited by rapamycin. Am J**Physiol** Endocrinol Metab. 2014;306(10):E1198-204. 594
- 595 15. Gurtler A, Kunz N, Gomolka M et al. Stain-Free technology as a normalization
 596 tool in Western blot analysis. *Anal Biochem.* 2013;433(2):105-11.

- Handschin C, Choi CS, Chin S et al. Abnormal glucose homeostasis in skeletal
 muscle-specific PGC-1alpha knockout mice reveals skeletal muscle-pancreatic
 beta cell crosstalk. *The Journal of clinical investigation*. 2007;117(11):3463-74.
- Hawley JA, Hargreaves M, Joyner MJ, Zierath JR. Integrative biology of
 exercise. *Cell*. 2014;159(4):738-49.
- Holloszy JO, Booth FW. Biochemical adaptations to endurance exercise in
 muscle. *Annual review of physiology*. 1976;38:273-91.
- Howley ET, Bassett DR, Jr., Welch HG. Criteria for maximal oxygen uptake:
 review and commentary. *Med Sci Sports Exerc.* 1995;27(9):1292-301.
- Kacin A, Strazar K. Frequent low-load ischemic resistance exercise to failure
 enhances muscle oxygen delivery and endurance capacity. *Scand J Med Sci Sports*. 2011;21(6):e231-41.
- Laurentino GC, Loenneke JP, Teixeira EL, Nakajima E, Iared W, Tricoli V. The
 Effect of Cuff Width on Muscle Adaptations after Blood Flow Restriction
 Training. *Med Sci Sports Exerc.* 2015.
- Laurentino GC, Ugrinowitsch C, Roschel H et al. Strength training with blood
 flow restriction diminishes myostatin gene expression. *Med Sci Sports Exerc*.
 2012;44(3):406-12.
- Little JP, Safdar A, Bishop D, Tarnopolsky MA, Gibala MJ. An acute bout of
 high-intensity interval training increases the nuclear abundance of PGC-1alpha
 and activates mitochondrial biogenesis in human skeletal muscle. *Am J Physiol Regul Integr Comp Physiol.* 2011;300(6):R1303-10.
- Livak KJ, Schmittgen TD. Analysis of relative gene expression data using realtime quantitative PCR and the 2(-Delta Delta C(T)) Method. *Methods*.
 2001;25(4):402-8.

- Lixandrao ME, Ugrinowitsch C, Laurentino G et al. Effects of exercise intensity
 and occlusion pressure after 12 weeks of resistance training with blood-flow
 restriction. *Eur J Appl Physiol*. 2015.
- Loenneke JP, Thrower AD, Balapur A, Barnes JT, Pujol TJ. Blood flowrestricted walking does not result in an accumulation of metabolites. *Clin Physiol Funct Imaging*. 2012;32(1):80-2.
- 628 27. Martinez-Redondo V, Pettersson AT, Ruas JL. The hitchhiker's guide to PGC629 1alpha isoform structure and biological functions. *Diabetologia*.
 630 2015;58(9):1969-77.
- 631 28. Ozaki H, Kakigi R, Kobayashi H, Loenneke JP, Abe T, Naito H. Effects of
 632 walking combined with restricted leg blood flow on mTOR and MAPK
 633 signalling in young men. *Acta Physiol (Oxf)*. 2014;211(1):97-106.
- Perry CG, Lally J, Holloway GP, Heigenhauser GJ, Bonen A, Spriet LL.
 Repeated transient mRNA bursts precede increases in transcriptional and
 mitochondrial proteins during training in human skeletal muscle. *J Physiol.*2010;588(Pt 23):4795-810.
- 638 30. Phillips SM. Physiologic and molecular bases of muscle hypertrophy and
 639 atrophy: impact of resistance exercise on human skeletal muscle (protein and
 640 exercise dose effects). *Appl Physiol Nutr Metab.* 2009;34(3):403-10.
- 641 31. Psilander N, Frank P, Flockhart M, Sahlin K. Exercise with low glycogen
 642 increases PGC-1alpha gene expression in human skeletal muscle. *Eur J Appl*643 *Physiol.* 2013;113(4):951-63.
- Ruas JL, White JP, Rao RR et al. A PGC-1alpha isoform induced by resistance
 training regulates skeletal muscle hypertrophy. *Cell*. 2012;151(6):1319-31.

- Sanchez AM, Candau RB, Bernardi H. FoxO transcription factors: their roles in
 the maintenance of skeletal muscle homeostasis. *Cellular and molecular life sciences : CMLS*. 2014;71(9):1657-71.
- 649 34. Semenza GL. Regulation of physiological responses to continuous and
 650 intermittent hypoxia by hypoxia-inducible factor 1. *Experimental physiology*.
 651 2006;91(5):803-6.
- Slysz J, Stultz J, Burr JF. The efficacy of blood flow restricted exercise: A
 systematic review & meta-analysis. *J Sci Med Sport*. 2015.
- 36. Steinberg GR, Kemp BE. AMPK in Health and Disease. *Physiological reviews*.
 2009;89(3):1025-78.
- Taylor CW, Ingham SA, Ferguson RA. Acute and chronic effect of sprint
 interval training combined with postexercise blood-flow restriction in trained
 individuals. *Experimental physiology*. 2016;101(1):143-54.
- Thom R, Rowe GC, Jang C, Safdar A, Arany Z. Hypoxic induction of vascular
 endothelial growth factor (VEGF) and angiogenesis in muscle by truncated
 peroxisome proliferator-activated receptor gamma coactivator (PGC)-1alpha. J *Biol Chem.* 2014;289(13):8810-7.
- Wilkinson SB, Phillips SM, Atherton PJ et al. Differential effects of resistance
 and endurance exercise in the fed state on signalling molecule phosphorylation
 and protein synthesis in human muscle. *J Physiol.* 2008;586(Pt 15):3701-17.
- 40. Ydfors M, Fischer H, Mascher H, Blomstrand E, Norrbom J, Gustafsson T. The
 truncated splice variants, NT-PGC-1alpha and PGC-1alpha4, increase with both
 endurance and resistance exercise in human skeletal muscle. *Physiol Rep.*2013;1(6):e00140.

670

671	Figure 1. (A) Total Peroxisome proliferator-activated receptor- γ coactivator 1 α (PGC-
672	1a), (B) Peroxisome proliferator-activated receptor- γ coactivator 1a (PGC-1a1), (C)
673	Peroxisome proliferator-activated receptor- γ coactivator 1 α 2 (PGC-1 α 2), (D)
674	Peroxisome proliferator-activated receptor- γ coactivator 1 α 3 (PGC-1 α 3) and (E)
675	Peroxisome proliferator-activated receptor- γ coactivator 1 α 4 (PGC-1 α 4) mRNA
676	abundance at rest and 3 h post-exercise recovery following endurance exercise (EE),
677	resistance exercise (RE) or low-intensity associated with blood flow restriction (BFR-
678	EE). Values are expressed relative to GAPDH and presented in arbitrary units (mean \pm
679	SD, n=9). a= Significant different from Rest (P \leq 0.05); b= Significant different from
680	HI-RT ($P \le 0.05$); c= Significant different from BFR-EE ($P \le 0.05$).
681	
682	
683	
684	
685	
686	
687	
688	
689	
690	
691	
692	
693	
694	
695	

696	Figure 2. (A) Vascular endothelial growth factor (VEGF), (B) Cytochrome c oxidase
697	subunit 4 isoform 1 (COXIV) and (C) hypoxia-inducible factor-1 alpha (HIF-1 α)
698	mRNA abundance at rest and 3 h post-exercise recovery following endurance exercise
699	(EE), resistance exercise (RE) or low-intensity associated with blood flow restriction
700	(BFR-EE). Values are expressed relative to GAPDH and presented in arbitrary units
701	(mean \pm SD, n=9). a= Significant different from Rest (P \leq 0.05); b= Significant different
702	from HI-RT ($P \le 0.05$); c= Significant different from BFR-EE ($P \le 0.05$).
703	
704	
705	
706	
707	
708	
709	
710	
711	
712	
713	
714	
715	
716	
717	
718	
719	
720	

721	Figure 3. (A) Interleukin 6 (IL-6), (B) Insulin-like growth factor 1(IGF-1), (C) Muscle
722	RING finger 1 (MURF1) and (D) Myostatin mRNA abundance at rest and 3 h post-
723	exercise recovery following endurance exercise (EE), resistance exercise (RE) or low-
724	intensity associated with blood flow restriction (BFR-EE). Values are expressed relative
725	to GAPDH and presented in arbitrary units (mean \pm SD, n=9). a= Significant different
726	from Rest (P \leq 0.05); b= Significant different from HI-RT (P \leq 0.05); c= Significant
727	different from BFR-EE (P \leq 0.05).
728	
729	
730	
/ 30	
731	
732	
733	
734	
735	
736	
737	
738	
720	
/39	

740	Figure 4. (A) Mechanistic target of rapamycin (mTOR) ^{Ser2448} (B) p70S6K ^{Thr389} (C)
741	eukaryotic translation initiation factor 4E-binding protein 1 (4E-BP1) ^{Thr37/46} eukaryotic
742	elongation factor 2 (eEF2) ^{Thr56} phosphorylation in skeletal muscle at rest and after 3 h
743	post-exercise recovery following endurance exercise (EE), resistance exercise (RE) or
744	low-intensity associated with blood flow restriction (BFR-EE). Values are normalized
745	to total protein loaded determined by stain free technology in arbitrary units (mean \pm
746	SD, n=9).
747	
748	
749	
750	
751	
752	
753	
754	
755	
756	

758	Figure 5. (A) Adenosine Monophosphate-Activated Protein (AMPK) ^{Thr172} and (B)
759	p53 ^{Ser15} phosphorylation in skeletal muscle at rest and after 3 h post-exercise recovery
760	following endurance exercise (EE), resistance exercise (RE) or low-intensity associated
761	with blood flow restriction (BFR-EE). Values are normalized to total protein loaded
762	determined by stain free technology in arbitrary units (mean \pm SD, n=9).
763	
764	
765	
766	
767	