Combined Effect of AMPK/PPAR Agonists and Exercise Training in *mdx* Mice Functional Performance

Carlos R. Bueno Júnior, Lucas C. Pantaleão, Vanessa A. Voltarelli, Luiz Henrique M. Bozi, Patricia Chakur Brum, Mayana Zatz

1 Human Genome Research Center - Institute of Biosciences, University of São Paulo, São Paulo, Brazil, 2 Department of Physiology and Biophysics - Institute of Biomedical Sciences, University of São Paulo, São Paulo, Brazil, 3 School of Physical Education and Sport, University of São Paulo, São Paulo, Brazil

Abstract

The present investigation was undertaken to test whether exercise training (ET) associated with AMPK/PPAR agonists (EM) would improve skeletal muscle function in *mdx* mice. These drugs have the potential to improve oxidative metabolism. This is of particular interest because oxidative muscle fibers are less affected in the course of the disease than glycolytic counterparts. Therefore, a cohort of 34 male congenic C57Bl/10J *mdx* mice included in this study was randomly assigned into four groups: vehicle solution (V), EM [AICAR (AMPK agonist, 50 mg/Kg·day−1, ip) and GW 1516 (PPARγ agonist, 2.5 mg/Kg·day−1, gavage)], ET (voluntary running on activity wheel) and EM+ET. Functional performance (grip meter and rotares), aerobic capacity (running test), muscle histopathology, serum creatine kinase (CK), levels of ubiquitinated proteins, oxidative metabolism protein expression (AMPK, PPAR, myoglobin and SCD) and intracellular calcium handling (DHPR, SERCA and NCX) protein expression were analyzed. Treatments started when the animals were two months old and were maintained for one month. A significant functional improvement (p<0.05) was observed in animals submitted to the combination of ET and EM. CK levels were decreased and the expression of proteins related to oxidative metabolism was increased in this group. There were no differences among the groups in the intracellular calcium handling protein expression. To our knowledge, this is the first study that tested the association of ET with EM in an experimental model of muscular dystrophy. Our results suggest that the association of ET and EM should be further tested as a potential therapeutic approach in muscular dystrophies.

Introduction

Duchenne muscular dystrophy (DMD) is an X-linked lethal genetic disease caused by the absence of the protein dystrophin [1]. This subsarcolemmal protein transmits the tension from the contractile proteins to the extracellular matrix and maintains the stability of the plasma membrane, preventing the extrusion of intracellular constituents, including creatine kinase to blood serum, which is hallmark of the disease [2]. All patients have grossly increased serum CK levels [3–4] since birth, which decrease with the progression of the dystrophic process, reaching almost normal levels in the late stages. Affected boys are usually wheelchair-bound around age 10–12 and assisted ventilation is required in order to prolong survival after adolescence. The mdx mouse also lack muscle dystrophin and is the most widely used animal model for DMD [5].

Many investigators suggest that exercise training is beneficial in muscular dystrophies, since it results in better intracellular calcium handling, activation of compensatory or antagonistic signaling pathways, increased antioxidant capacity and angiogenesis, improved ability to blunt alpha-adrenergic vasoconstriction, increased iNOS activity and membrane protective effects [6–12]. However, others claim it has detrimental effects, which could be caused by the increased susceptibility of dystrophic muscle to exercise-induced injury. It is also known that these contradictory results are mainly related to type and intensity of exercise [13–15].

AMPK (AMP-activated protein kinase) and PPAR (peroxisome proliferator-activated receptor) agonists are exercise mimetics. In 2009, Miura et al. demonstrated that a PPAR agonist reduced the number of skeletal muscle fiber membrane lesions and decreased force drop due to eccentric contractions in mdx mice [16]. It has been demonstrated that these drugs are able to improve the oxidative metabolism - they induce mitochondrial biogenesis and fatty acid oxidation [17]. The improvement of the oxidative metabolism in muscular dystrophies is critical since it has been demonstrated, both in experimental models and humans, that slow oxidative muscles display reduced damage when compared to the glycolytic counterparts [18–19].

As exercise training and its mimetics present both coincident and different beneficial effects [17], the aim of the present study was to test the hypothesis that the association of exercise training with AMPK and PPAR agonists can improve mdx functional performance, reducing the impact of muscle dystrophin deficiency. In order to address this question, several parameters, such as functional tests, histopathology, skeletal muscle protein expression...
related to oxidative metabolism and calcium handling, renal function, and fat in the carcass, were analyzed in animals submitted or not to exercise.

Materials and Methods

Ethics Statement
This study was conducted in accordance with the ethical principles in animal research adopted by the Brazilian College of Animal Experimentation (www.cobea.org.br) and was approved by the University of São Paulo, Institute of Biosciences Ethical Committee (087/2009).

Study Population
A cohort of 34 two-months old male congenic C57Bl/10J mdx mice were randomly assigned into four groups: vehicle solution (V), exercise mimetics (EM), exercise training (ET) and EM+ET. Analysis started when the animals were three months-old. During all the month before the EM mice received AICAR (100 mg.Kg⁻¹.d⁻¹), ip, Cayman Chemical, catalog number 10010241, Ann Arbor, MI, USA) and GW 1516 (5 mg.Kg⁻¹.ip, Cayman Chemical, catalog number 10010241, Ann Arbor, MI, USA) detected by autoradiography. Quantification analysis of chemiluminescence (Amersham Biosciences; Piscataway, NJ, USA) was performed using peroxidase-conjugated primary antibody was detected using peroxidase-conjugated secondary antibody (anti-mouse or anti-rabbit, 1:100) for at least 30 minutes. The intensity of exercise was increased by 3 m/min (6–33 m/min) every 3 min at 0% grade until exhaustion.

Muscle Histopathology
After the tests, the mice were killed and triceps brachialis muscles were harvested, immediately frozen in melting isopentane and stored in liquid nitrogen. The frozen muscles were cut into 10-μm cross sections from the proximal to distal region using a cryostat (Cristostat Micron HM505E, Walldorf, Germany). Sections of muscle were then stained with haematoxylin and cosin [25] or used to perform histochemical myosin ATPase [26], as previously described. The cross-sectional area of the fibers was evaluated at 200x magnification and further analyzed on a digitizing unit connected to a computer (Image Pro-plus, Media Cybernetic, Silver Spring, MD, USA). The cross-sectional area and the percentage of interstitium and type I fibers in the muscle were analyzed in blind test. In addition, serum creatine kinase was analyzed when the animals were three months old, when normal values for control animals are reported to be low (around 100 U/l) [27].

Skeletal Muscle Protein Expression
In order to evaluate aspects related to muscular structure, oxidative metabolism and calcium handling, immunoblot of triceps brachialis muscle homogenates were performed according to Bacurau et al. [26]. Briefly, frozen muscles in liquid nitrogen were homogenized in a buffer containing 1 mM EDTA, 1 mM EGTA, 2 mM MgCl₂, 5 nM KCl, 25 mM HEPES (pH 7.5), 100 μM PMSF, 2 mM DTT, 1% Triton X-100 and protease inhibitor cocktail (1:100, Sigma-Aldrich, MO, USA). Samples were loaded and subjected to SDS-PAGE in polyacrylamide gels. After electrophoresis, proteins were electro-transferred to nitrocellulose membrane (Amersham Biosciences, NJ, USA). Equal loading of samples (25 μg) and even transfer efficiency were monitored using 0.5% Ponceau S staining of the blotted membrane. The blotted membrane was then incubated in a blocking buffer (5% nonfat dry milk, 10 mM Tris-HCl, pH 7.6, 150 mM NaCl, and 0.1% Tween 20) for 2 h at room temperature and then incubated overnight at 4°C with the following primary antibodies: ubiquitinated proteins (1:1000) from Santa Cruz Biotechnology; AMPKα (1:1000), p-AMPKα (Thr172, 1:1000), ACC (1:1000) and p-ACC (Ser79, 1:1000) from Cell Signaling Technology; PPARδ (1:1000), Myoglobin (1:1000) and SCD (1:1000) from Abcam; and DHPR-α (1:1000), SERCA 1 (1:1000) and NCX (1:1000) from ABR Incorporation. Binding of the primary antibody was detected using peroxidase-conjugated secondary antibodies (anti-mouse or anti-rabbit, 1:1000, for 90 minutes at room temperature) and developed using enhanced chemiluminescence (Amersham Biosciences; Piscataway, NJ, USA) detected by autoradiography. Quantification analysis of blots was performed using Scion Image software (Scion Corporation based on NIH image).

Creatinine Clearance and Fat in the Carcass
In order to evaluate mouse renal function, we evaluated creatinine clearance [28]. Carcass fat was determined by its basic parameters.

| Parameter                        | V (9)     | EM (8)    | ET (9)    | EM+ET (8) |
|----------------------------------|-----------|-----------|-----------|-----------|
| Body weight (g)                  | 31.4±1.1  | 29.2±1.1  | 29.4±0.7  | 29.9±1.2  |
| Food intake (g/day)              | 3.5±0.2   | 4±0.1     | 3.5±0.2   | 3.6±0.1   |
| Feces (g/day)                    | 2.4±0.1   | 2.7±0.2   | 2.1±0.1   | 2.2±0.1   |
| Water intake (g/day)             | 3.2±0.4   | 2.9±0.3   | 3.1±0.2   | 2.9±0.2   |
| Urine (g/day)                    | 2±0.3     | 1.7±0.2   | 1.5±0.1   | 1.9±0.1   |
| Glucose (mg/dl)                  | 90.8±4.7  | 100.7±5.1 | 83.5±6.6  | 97±7.3    |
| Triglycerides (mg/dl)            | 140±19    | 170±13    | 172±22    | 231±25*   |
| Cholesterol (mg/dl)              | 110.3±5.6 | 104.5±4.4 | 117.8±7.7 | 115.9±4.8 |

The number of animals in each group is shown in parentheses and the data are presented as mean ± SE. *P<0.05 versus vehicle group (V) after two-way ANOVA and Duncan post-hoc test.
doi:10.1371/journal.pone.0045699.t001
chemical analysis. Initially, carcasses stayed in a ventilated oven (70°C) for 7 days. Then, whole dry carcass was chopped up and wrapped in gauze and filter paper for determination of body fat by the solvent extraction technique using a Soxhlet apparatus and ethyl ether as solvent [29].

Statistical Analysis
All values are presented as means ± SE. Two-way analysis of variance (ANOVA; exercise mimetics and exercise training as variables) and Duncan post-hoc test (Statistica software, StatSoft, Inc., Tulsa, OK, USA) were used to compare the groups. Statistical significance was considered as p ≤ 0.05.

Results

Basic Parameters
As presented in the Table 1, there was no statistical difference among the groups in body weight, food intake, feces excretion, water intake and urine excretion. In addition, the data of serum glucose and cholesterol after night fasting were similar in the groups. Only triglycerides levels were increased in the group that received exercise mimetics and exercise training when compared to the values of the vehicle group.

Functional Capacity
Figure 1 shows that the group submitted to both exercise mimetics and exercise training presented improved functional performance (grip meter - Figure 1A, p = 0.05; and rotarod - Figure 1B, p = 0.009) as well as aerobic capacity (running test - Figure 1C, p < 0.001), which was statistically significant when compared to the vehicle group. Exercise mimetics only ameliorated significantly grip force (Figure 1A, p = 0.026) and exercise training only improved significantly aerobic capacity (Figure 1C, p < 0.001). Of interest, during the entire month of treatment activity wheel was moved 2.9 ± 0.4 hours per day in a speed of 9.6 ± 0.6 m/min by the exercise training group. These mean values were 2.8 ± 0.5 hours and 5.8 ± 0.6 m/min, respectively, for the group that received both interventions (there is a statistical difference between 9.6 ± 0.6 m/min and 5.8 ± 0.6 m/min, p = 0.004).

Skeletal Muscle Histology, Serum Creatine Kinase and Ubiquitined Proteins
In animals that had a better functional capacity (exercise mimetics + training), skeletal muscle showed statistically increased cross sectional area (Figure 2A, p = 0.024) associated with reduced interstitium (Figure 2B, p = 0.005), serum creatine kinase (Figure 2C, p = 0.048) and ubiquitinated proteins (Figure 2D, p = 0.029) as compared to the vehicle group. Exercise training
alone led to significantly augmented fibers (Figure 2A, p = 0.023) and reduced percentage of interstitium and ubiquitinated proteins levels (Figure 2B and D; p = 0.001 and 0.05, respectively), but increased levels of blood creatine kinase (Figure 2C, p = 0.05). Exercise mimetics alone, on the other hand, statistically improved cross sectional area (Figure 2A, p = 0.028), percentage of interstitium (Figure 2B, 0.015), percentage of type I fibers (Figure 2C, p = 0.003) and levels of ubiquitinated proteins (Figure 2D, p = 0.04), with no change on creatine kinase levels (Figure 2C).

Protein Expression Related to Oxidative Metabolism and Calcium Handling
Since an improvement in oxidative metabolism is a proposed mechanism to explain the beneficial effects of both exercise mimetics and exercise training, we evaluated the expression of proteins related to oxidative metabolism. As observed in Figure 3, the association of both approaches increased the expression of key proteins (p = 0.046 for AMPKa, p = 0.003 for p-AMPKa, p = 0.009 for p-ACC/ACC ratio, p = 0.031 for PPARδ and p = 0.05 for myoglobin) and the activity of one of them (AMPKa, p = 0.003), assessed by phosphorylation levels (Figure 3A–E).

Figure 2. When associated with its mimetics, exercise training does not impair skeletal muscle structure in mdx mice. Cross sectional area (A), intersticium (B), percentage of type I muscle fibers (C), blood creatine kinase (D) and ubiquitinated proteins (E) after one month of vehicle solution (V), exercise mimetics (EM), exercise training (ET) and EM+ET. The number of animals in each group is shown in parentheses and the data are presented as mean ± SE. *P<0.05 versus vehicle group (V) after two-way ANOVA and Duncan post-hoc test.

doi:10.1371/journal.pone.0045699.g002
Exercise Training and Its Mimetics in mdx Mice

A

B

C

D

E

F

G

AMPK (V % of V)

p-AMPK (Thr172) (V % of V)

p-ACC/ACC ratio (V % of V)

PPARδ (V % of V)

Myoglobin (V % of V)

SCD (V % of V)

Ponceau

V (4) EM (3) ET (3) EM+ET (4)

V (8) EM (6) ET (5) EM+ET (6)

V (5) EM (5) ET (5) EM+ET (5)

V (7) EM (6) ET (6) EM+ET (7)

V (4) EM (3) ET (3) EM+ET (4)

V (7) EM (6) ET (6) EM+ET (7)

AMPKα 62 kDa

p-AMPKα (Thr172) 62 kDa

p-ACC 280 kDa

PPARδ 50 kDa

Myoglobin 17 kDa

SCD 41 kDa

Ponceau 195 kDa
Interestingly, in this group the expression of SCD, a pivotal enzyme in lipogenesis, was decreased (Figure 3F, p = 0.05) when compared to the control group. However, when we analyzed exercise training and its mimetics isolatedly, there was a significant increase only in the level of AMPK\(^\alpha\) phosphorylation in the exercise training group (Figure 3B, p = 0.05) and in the p-ACC/ACC ratio in both groups (Figure 3C, p = 0.018 for exercise mimetics group and p = 0.044 for wheel activity group).

Since impaired transsarcolemmal calcium flux and sarcoplasmic reticulum calcium release/reuptake has been identified as main contributors to skeletal muscle functional and structural abnormalities [30], we further investigated whether exercise mimetics and exercise training would change the expression of DHPR, SERCA and NCX. However, as shown in Figure 4, there was no statistical difference among the groups studied.

Renal Function and Fat in the Carcass

As a potential adverse effect of different drugs can be an impaired renal function, we analyzed creatinine clearance. Interestingly, the exercise mimetics group displayed better renal function than the other groups (Figure 5A, p = 0.033). There was no difference among the other groups (Figure 5A).

Finally, as both aerobic exercise training and AMPK/PPAR agonists accelerate the oxidative metabolism, an expected consequence would be a reduction in the amount of fat in the body. In order to test this hypothesis, we analyzed the carcass fat of the animals. The results are presented in the Figure 5B. When compared to the vehicle group, all three approaches reduced the amount of fat in mice (p = 0.024 for the exercise mimetics group and p<0.001 for the exercise training and combined therapy groups), but the group that received both exercise mimetics and exercise training showed a statistically significant decrease when compared to the groups submitted to the isolated approaches (p = 0.007 for the exercise mimetics group and p = 0.005 for the exercise training group).

Discussion

The main result of the present study was that voluntary exercise training combined with AMPK/PPAR agonists improves the functional performance and the aerobic capacity of mdx mice. Of particular interest is the fact that these improvements might be associated to reduced muscle degeneration (increased muscle fiber cross sectional area and diminished ubiquitinated proteins expression) and improved efficiency of the aerobic metabolism. To our knowledge, this is the first study that tested the association of

Figure 3. Exercise training associated with its mimetics improves protein expression related to the oxidative metabolism. Muscular protein expression of AMPK\(\alpha\) (A), p-AMPK\(\alpha\)(Thr172) (B), p-ACC/ACC ratio (C), PPAR\(\delta\) (D), Myoglobin (E) and SCD (F) after one month of vehicle solution (V), exercise mimetics (EM), exercise training (ET) and EM+ET. The number of animals in each group is shown in parentheses and the data are presented as mean ± SE. *\(P<0.05\) versus vehicle group (V) after two-way ANOVA and Duncan post-hoc test.

doi:10.1371/journal.pone.0045699.g003

Figure 4. Exercise training and its mimetics do not change calcium handling protein expression. DHPR\(\alpha1\) (A), SERCA 1 (B) and NCX (C) protein expression after one month of vehicle solution (V), exercise mimetics (EM), exercise training (ET) and EM+ET. The number of animals in each group is shown in parentheses and the data are presented as mean ± SE. *\(P<0.05\) versus vehicle group (V) after two-way ANOVA and Duncan post-hoc test.

doi:10.1371/journal.pone.0045699.g004
exercise training with AMPK/PPAR agonists in a disease murine model.

Here we show that the association of exercise training and AMPK/PPAR agonists resulted in functional improvements related to grip meter, rotarod and aerobic capacity (running test). The animals submitted only to the exercise training showed improved aerobic capacity, supporting the efficiency of the training, and the animals that received only the drugs had an improvement in the grip meter test. Our data in the running test corroborated Narkar et al. [17], who reported that the association of AICAR and exercise training improved the running capacity in control mice.

Interestingly, the exercise training group presented increased values of training intensity in wheel activity when compared to the animals that also received exercise mimetics - the time of activity per day was similar in both groups. This can help explain, at least in part, the results where the combined therapy did not result in a better outcome as compared to the isolated approaches. Although different intensities between the groups can be a limitation of the present study, we did the running test before the treatment to certify that both groups presented similar aerobic capacity (data not shown). Furthermore, the absence of exercise intensity control is inherent to the activity wheel method and this kind of voluntary training was chosen because it has been demonstrated that it avoid excessive overload to the dystrophic muscle. Exercise intensity is a prominent factor to induce muscle adaptations, but dystrophic patients are not able to perform intense exercises because their muscles are more susceptible to mechanical stress [31].

Although strength training is more efficient to induce hypertrophic response, aerobic exercise training also can increase muscle mass, particularly in animals with skeletal muscle abnormalities [26,32]. In addition, related to the exercise mimetics group, it is possible that AMPK/PPAR agonists might increase protein synthesis through higher metabolism efficiency and reducing the need of proteolysis to generate energy by gluconeogenesis. In fact, both groups that received AMPK/PPAR agonists had decreased ubiquitinated proteins - the ubiquitination of proteins has a direct relationship with the activity of the ubiquitin-proteasome system, the main mechanism of protein breakdown in the cell [33].

As stated before, glycolytic muscle fibers are more affected by the dystrophy when compared to the oxidative counterparts and only the exercise mimetics group presented increased percentage of type I fibers than the control group. This effect of drugs is in accordance with Narkar et al. [17]. Furthermore, the observation that exercise training is unable to increase the percentage of oxidative fibers corroborated other studies [34]. One possible explanation is that the intensity of the exercise was excessive for the dystrophic mice, which would also explain the higher levels of serum CK in the exercise training group. It was possible to observe that the animals performed high intensity exercises to move the wheel, which continued in movement by inertia. Although one month can be insufficient to result in changes in the type fiber profile, serum CK may increase rapidly after exercise. Finally, in the animals that received both treatments the high intensity exercise could have caused a negative interference in the effect of the exercise mimetics, which resemble aerobic exercise [35].

We have no explanation for the apparently decreased activity in serum CK in the mimetics group. It could be due to the small sample size or to the natural fluctuation that is seen in this enzyme which might not be directly related to the dystrophic process [36].

We also observed that in the group in which exercise training was associated with AMPK/PPAR agonists, there was an increased expression of AMPK, phosphorylated AMPK and PPARβ. The exercise training group also presented increased AMPK activity evaluated by phosphorylation and these results corroborate the running test data since both groups with highest running performance presented increased AMPK activity. Additionally, it is known that mice with dysfunction in the skeletal muscle AMPK signaling present reduced running capacity [37–38] and many studies have demonstrated that exercise training increases the AMPK activation [17,28,39]. Finally, the group submitted to both exercise training and its mimetics displayed increased myoglobin protein expression - a protein that has pivotal importance in the storage of oxygen in the muscle cell, especially in dystrophic muscle fiber, in which myoglobin can leak from the cell cytosol due to the disruption of plasma membrane [40]. This group also presented reduced levels of SCD, a key enzyme in the lipogenesis [41]. In fact, in this set of animals the levels of triglycerides were increased.

However, the high levels of triglycerides in this group do not represent a metabolic disorder - these animals do not present increased cholesterol and glucose values, for example. The result can be related to decreased lipogenesis (discussed before) and the time that blood was harvested for analysis. In addition, Narkar et al. [17] have found no correlation between serum CK and muscle damage, which is consistent with our results.
al. demonstrated that triglycerides levels may be increased by exercise [17].

Although they were not analyzed in previous studies with AMPK/PPAR agonists [16–17], potential adverse effects related to exercise mimetics should be monitored carefully. Interestingly, the exercise mimetics group showed decreased renal function by creatinine clearance test. Finally, we showed that all strategies reduced carcass fat, but with higher magnitude in the animals submitted to both exercise training and AMPK/PPAR agonists. This result suggests increased lipolysis and generation of energy by the oxidative metabolism, which is supported by the increased serum levels of triglycerides in this group. The significance of the reduction in the carcass fat is highlighted if we consider that there was no statistical difference among the groups in body weight.

We also demonstrated that the ratio between protein expressions of phosphorylated and total acetyl-CoA carboxylase (Ser79) was increased in the three groups submitted to the treatments, which indicate inhibition of carboxylation of acetyl-CoA to malonyl-CoA in the biosynthesis of fatty acids. These results might explain the results from the carcass fat. In addition, these data reinforce the effectiveness of the exercise mimetics treatment because an AMPK downstream activity is the phosphorylation of acetyl-CoA carboxylase [42].

In short, a beneficial effect of combination of exercise training and AMPK/PPAR agonists was observed in mdx mice. Although exercise training alone or alternatively its mimetics showed some improvement in skeletal muscle histology and aerobic capacity of mdx mice, the combination of both strategies seems more effective. In addition, our results suggest that a favorable protein turnover (represented by decreased levels of ubiquitinated proteins and increased cross sectional area) as well an improved efficiency of the oxidative metabolism might explain, at least in part, the observed functional improvements. It will be interesting to repeat these studies in other experimental models of muscular dystrophy. Although most dystrophic patients are not able to run, the animals in the present study are in the early stages of dystrophy, when many children would still be able to exercise. It would also be of interest to assess if the association of passive exercise and AMPK/PPAR agonists have a comparable beneficial effect.

**Author Contributions**

Conceived and designed the experiments: CRBJ PCB MZ. Performed the experiments: CRBJ LCP VAV LHMB. Analyzed the data: CRBJ LCP VAV LHMB PCB MZ. Contributed reagents/materials/analysis tools: CRBJ LCP VAV LHMB PCB MZ. Wrote the paper: CRBJ PCB MZ.

**References**

1. Worton RG, Thompson MW (1988) Genetics of Duchenne muscular dystrophy. Am Rev Genet 22: 601–629.

2. Mozzetta C, Minetti G, Puri PL (2009) Regenerative pharmacology in the treatment of genetic diseases: the paradigm of muscular dystrophy. Int J Biochem Cell Biol 41(4): 701–710.

3. Zatz M, Frota-Pessoa O, Levy JA, Peres CA (1996) Creatine-phosphokinase (CPK) activity in relatives of patients with x-linked muscular-dystrophies - A Brazilian study. Jornal de Genetique Humaine 24: 153–168.

4. Zatz M, Rapaport D, Vainzof M, Rocha JML, Passos-Bueno MR, et al. (1991) Serum creatine-kinase (CK) and pyruvate-kinase (PK) activity as a function of clinical evolution in Duchenne (DMD) and Becker (BMD) muscular dystrophies. J Neurol Sci 102: 190–196.

5. Vainzof M, Ayub-Guerrieri D, Onofre PCG, Martins PCM, Lopes LF, et al. (2000) Animal models for genetic neuromuscular diseases. J Mol Neurosci 34: 241–248.

6. Bouchentouf M, Benabdallah BF, Mills P, Tremblay JP (2006) Exercise and Duchenne muscular dystrophy: a Brazilian study. J Neurol Sci 241–248.

7. Call JA, Voolker KA, Wolff AV (2008) Endurance capacity in maturing mdx mice is markedly enhanced by combined voluntary wheel running and green tea extract. J Appl Physiol 105: 923–932.

8. Carter GT, Wineinger MA, Walsh SA, Horasek SJ (1995) Effect of voluntary wheel-running exercise on muscles of the mdx mouse. Neuromuscul Disord 5: 323–332.

9. Dupont-Versteegden EE, McCarthy DJ, Kazis MS (1994) Voluntary exercise decreases progression of muscular dystrophy in diaphragm of mdx mice. J Appl Physiol 77: 1736–1741.

10. Hayes A, Williams DA (1996) Beneficial effects of voluntary wheel running on the properties of dystrophic mouse muscle. J Appl Physiol 80(2): 670–679.

11. Hayes A, Williams DA (1998) Contracture function and low-intensity exercise effects of old dystrophic (mdx) mice. Am J Physiol Cell Physiol 274: 1138–1144.

12. Markert CD, Ambrosio F, Call JA, Grange RW (2011) Exercise and Duchenne muscular dystrophy: toward evidence-based exercise prescription. Muscle Nerve 43(4): 464–478.

13. Carter GT, Abresch RT, Fowler WM Jr (2002) Adaptations to exercise training and contraction-induced muscle injury in animal models of muscular dystrophy. Am J Phys Med Rehabil 81: S151–S161.

14. Eagle M (2002) Report on muscular dystrophy campaign workshop: exercise in neuromuscular diseases. Neuromuscul Disord 12: 973–983.

15. Granchelli JA, Pollina C, Levy JA, Peres CA (1996) Creatine-phosphokinase (CPK) activity in relatives of patients with x-linked muscular-dystrophies - A Brazilian study. Jornal de Genetique Humaine 24: 153–168.

16. Miura P, Chakkalakal JV, Boudreault L, Belanger G, Hebert RL, et al. (2009) AMPK and PPARdelta agonists are exercise mimetics. Cell 134(3): 405–415.

17. Narkar VA, Downes M, Yu RT, Embler E, Wang YX, et al. (2008) AMPK and PPARDelta agonists are exercise mimetics. Cell 134(3): 405–415.

18. Moens P, Baatsen PH, Marechal G (1999) Increased susceptibility of EDL muscles from mdx mice to damage induced by contractions with stretch. J Muscle Res Cell Motil 14: 446–451.

19. Webster C, Sillerstein L, Hays AP, Blaa HM (1980) Fast muscle fibers are preferentially affected in Duchenne muscular dystrophy. Cell 52: 503–513.

20. Anderson KD, Abdul M, Steward O (2004) Quantitative assessment of deficits and recovery of forcible motor function after cervical spinal cord injury in mice. Exp Neurol 180(1): 184–191.

21. Bueno CR Jr, Ferreira JC, Pereira MG, Barcruz AV, Braga Jr, Bruno PM (2010) Aerobic exercise training improves skeletal muscle function and Ca2+ handling-related protein expression in sympathetic hyperactivity-induced heart failure. J Appl Physiol 109(3): 702–709.

22. Li ZF, Wu X, Jiang Y, Liu J, Wu C, et al. (2008) Non-pathogenic protein aggregates in skeletal muscle in MLF1 transgenic mice. J Neurol Sci 264(1-2): 77–86.

23. Turgeman T, Hagem Y, Hubner K, Jasaal DS, Anderson JE, et al. (2008) Prevention of muscle fibrosis and improvement in muscle performance in the mdx mouse by halofuginone. Neuromuscul Disord 18(11): 857–868.

24. Ferreira JC, Barcruz AV, Bueno CR Jr, Cunha TC, Tanaka LY, et al. (2010) Aerobic exercise training improves Ca2+ handling and redox status of skeletal muscle in mice. Exp Biol Med 235(4): 497–503.

25. Dubowitz V (1985) Muscle biopsy: a practical approach. London: Bailliere Tindall, 385 p.

26. Barcruz AV, Jardim MA, Ferreira JC, Bechara LR, Bueno CR Jr, et al. (2009) Sympathetic hyperactivity differentially affects skeletal muscle mass in developing heart failure: role of exercise training. J Appl Physiol 105(6): 1631–1640.

27. Tanigui AP, Pertille A, Matsamura CY, Santo Neto H, Marques MJ (2011) Prevention of muscle fibrosis and myogenesis in mdx mice by sartamin, a TGF-β1 blocker. Muscle Nerve 43(1): 82–87.

28. Durante PE, Mustard JK, Park SH, Winder WW, Hardie DG (2002) Effects of endurance training on activity and expression of AMP-activated protein kinase isofoms in rat muscles. Am J Physiol Endocrinol Metab 283: E178–E186.

29. Donato JJ Jr, Pedrosa RG, Coutzi VF, Pires IS, Tiranegui J (2006) Effects of leucine supplementation on the body composition and protein status of rats submitted to food restriction. Nutri 22(3): 529–527.

30. Franco A Jr, Lansman JB (1990) Calcium entry through stretch-inactivated ion channels in mdx mouse. J Physiol 435(3): 647–678.

31. Baltgalvis KA, Call JA, Cochrane GD, Laker RC, Yan Z, et al. (2012) Exercise training improves plantarflexor muscle function in mdx mice. Med Sci Sports Exerc. In press.

32. Call JA, McKenzie PN, Novotny SA, Lose DA (2010) Progressive resistance voluntary wheel running in the mdx mouse. Muscle Nerve 42(6): 871–870.

33. Clague MJ, Urbé S (2010) Ubiquitin: same molecule, different degradation pathways. Cell 143(3): 602–605.

34. Landrich RM, Kose AM, Nelson SA, Balghavi KA, Lose DA (2008) Adaptive and nonadaptive responses to voluntary wheel running by mdx mouse. Muscle Nerve 38: 1290–1309.

35. Wilson JM, Marin PJ, Rheo MR, Wilson SM, Lorenne JP, et al. (2012) Concurrent exercise training: a meta analysis examining interference of aerobic and resistance exercise. J Strength Cond Res 26(10): 2293–2307.

36. Ozawa E, Hagiwara Y, Yoshida M (1999) Creatine kinase, cell membrane and redox status of skeletal muscles from mdx mice to damage induced by contractions with stretch. J Muscle Res Cell Motil 14: 446–451.
37. Mu J, Brozinick JT Jr, Valladares O, Bocan M, Birnbaum MJ (2001) A role for AMP-activated protein kinase in contraction- and hypoxia-regulated glucose transport in skeletal muscle. Mol Cell 7: 1085–1094.
38. Thomson DM, Porter BB, Tall JH, Kim HJ, Barrow JR, et al. (2007) Skeletal muscle and heart LKB1 deficiency causes decreased voluntary running and reduced muscle mitochondrial marker enzyme expression in mice. Am J Physiol Endocrinol Metab 292: E196–E202.
39. Frøsig C, Jørgensen SB, Hardie DG, Richter EA, Wojtaszewski JF (2004) 5′-AMP-activated protein kinase activity and protein expression are regulated by endurance training in human skeletal muscle. Am J Physiol Endocrinol Metab 286: E411–E417.
40. Garrood P, Eagle M, Jardine PE, Bushby K, Straub V (2008) Myoglobinuria in boys with Duchenne muscular dystrophy on corticosteroid therapy. Neurom Disord 18(1): 71–73.
41. Flowers MT, Ntambi JM (2009) Stearoyl-CoA desaturase and its relation to high-carbohydrate diets and obesity. Biochim Biophys Acta 1791(2): 85–91.
42. Kreuz S, Schoelch C, Thomas L, Rist W, Rippmann JF, et al. (2009) Acetyl-CoA carboxylase 1 and 2 show distinct expression patterns in rats and humans and alterations in obesity and diabetes. Diabetes Metab Res Rev 25(6): 577–586.