Aerobic Exercise Training Increases Muscle Water Content in Obese Middle-Age Men

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ABSTRACT

MORA-RODRIGUEZ, R. C., A. SANCHEZ-RONCERO, V. E. FERNÁNDEZ-ELÍAS, A. GUADALUPE-GRAU, J. F. ORTEGA, F. DELA, and J. W. HELGE. Aerobic Exercise Training Increases Muscle Water Content in Obese Middle-Age Men. Med. Sci. Sports Exerc., Vol. 48, No. 5, pp. 00–00, 2016. Purpose: The objective of this study is to determine whether muscle water content (H2Omuscle) expands with training in deconditioned middle-age men and the effects of this expansion in other muscle metabolites. Methods: Eighteen obese (BMI = 33 ± 3 kg·m⁻²) untrained (VO2peak = 29 ± 7 mL·kg⁻¹·min⁻¹) metabolic syndrome men completed a 4-month aerobic cycling training program. Vastus lateralis muscle biopsies were collected before and 72 h after the completion of the last training bout. Water content, total protein, glycogen concentration, and citrate synthase activity were measured in biopsy tissue. Body composition was assessed using dual-energy X-ray absorptiometry, and cardiometabolic fitness was measured during an incremental cycling test. Results: Body weight and fat mass were reduced −1.9% and −5.4%, respectively (P < 0.05), whereas leg fat free mass increased with training (1.8%, P = 0.023). Cardiorespiratory fitness (i.e., VO2peak), maximal fat oxidation (i.e., FOmax), and maximal cycling power (i.e., Wmax) improved with training (11%, 33%, and 10%, respectively; P < 0.05). After 4 months of training, H2Omuscle increased from 783 ± 18 to 799 ± 24 g·kg⁻¹ wet weight (ww) (2%, P = 0.011), whereas muscle protein concentration decreased 11% (145 ± 15 to 129 ± 13 g·kg⁻¹ ww, P = 0.007). Citrate synthase activity (proxy for mitochondrial density) increased by 31% (17 ± 5 to 22 ± 5 mmol·min⁻¹·kg⁻¹·ww, P = 0.024). Muscle glycogen concentration increased by 14% (22 ± 7 to 25 ± 7 g·kg⁻¹ ww) although without reaching statistical significance as expressed as per kilogram of wet weight (P = 0.15). Conclusions: Our findings suggest that aerobic cycling training increases quadriceps muscle water although reduces muscle protein concentration in obese metabolic syndrome men. Reduced protein concentration coexists with increased leg lean mass suggestive of a water dilution effect that however does impair increased cycling leg power with training. Key Words: EXERCISE TRAINING, MUSCLE WATER CONTENT, MUSCLE HYPERTROPHY, AGING

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in resting healthy young subjects coincides across studies (i.e., −78% ± 3%) (6,9,13,14,20,22,29,30).

The water-to-FFM constant originates from experiments where animals or human cadavers were thoroughly homogenized, fat was extracted, and water content was determined by weighing homogenate aliquots before and after drying by evaporation or sublimation (i.e., freeze-drying). Similar processing of a muscle biopsy sample allows the measurement of muscle water content in vivo. Using this technique, it has been shown that congestive heart failure patients increase leg muscle water content in comparison with age-matched healthy controls (2) likely because of reduced venous return provoking edema. In healthy men, short intense exercise increases muscle water content (30) likely because of increased transcapillary pressure. Conversely, submaximal prolonged exercise in a hot environment results in whole body dehydration and losses of muscle water content (6). We have recently shown that whole body dehydration by 4.2% reduces muscle water content when 1 h of recovery is allowed to re-equilibrate among body fluids spaces (20).

Although it is clear that muscle water content can transitorily change in response to a bout of exercise, the effects of chronic exercise training on muscle water content are not well
defined. To our knowledge, only four studies in humans report the effects of training on muscle water changes. All of them report water content as a secondary finding to the study of muscle hypertrophy. One study uses nuclear magnetic resonance to estimate leg water changes after leg extension training in young subjects finding no effects (17). Three of them use direct measurement of water content in leg muscle biopsies (13,14,29). One is a cross-sectional comparison between competitive and recreational runners and thus subjected to the effects of different diets, muscle fiber composition, genetics, and hormonal milieu between groups (29). The two remaining studies follow a repeated-measures design in the same group of subjects before and after 12 wk of cycle ergometer training. However, although the first study conducted in old women finds increases in muscle water content with training (10), the subsequent study conducted in old and young men does not (11).

Despite the utmost relevance of water for muscle energetics during and after exercise (9,10), the effects of chronic training on muscle water content remain unclear. A few aerobic training exercise sessions expand intravascular water (i.e., plasma volume expansion) allowing better cardiac function, cutaneous blood flow, and sweat gland fluid supply (4). It is however unclear if water within muscles is also expanded with aerobic training to improve muscle energetics and/or contractile function. To study muscle water changes in a sedentary population initially seems pertinent to unveil if muscle conditioning includes muscle water expansion and how does it relate to other known muscle training adaptations (e.g., increase in mitochondrial density or glycogen content). We hypothesized that a health-oriented fitness-training program using aerobic interval exercise will increase muscle water content in initially sedentary metabolic syndrome men. We will relate the changes in muscle water content to muscle variables known to change with training (mitochondrial and glycogen content) in an attempt to establish possible links.

METHODS

Participants. Eighteen obese men between 34 and 64 yr old (mean, 54 ± 8 yr old) completed the study. Participants were enrolled based on fulfilling the ≥3 METs criteria as per harmonized definition (1) using population Europid waist circumference cut-points. Exclusion criteria included cardiovascular or renal disease, peripheral vascular disease, and any disease associated with exercise intolerance. Body weight stability in the last 6 months (i.e., changes below 1% of initial body weight) was also a requirement. Participants reported not being engaged in regular physical activity beyond walking less than 30 min d⁻¹ in the past 6 months. The local hospital’s research ethics committee approved the study procedures and the informed consent documents. Subjects were informed of the purpose and risks involved in the study before signing the written consent. The study fulfilled the latest version of the Declaration of Helsinki.

Exercise training and dietary records. Subjects underwent supervised aerobic interval training (AIT) with a frequency of three times per week for 4 months. Training consisted of pedaling for 10 min as a warm-up at 70% HRmax followed by 4 × 4-min intervals at 90% of HRmax interspersed with 3-min active recovery at 70% HRmax and a 5-min cool-down period for a total of 43 min. Exercise intensity was increased as training adaptations developed to maintain the target HR (Accurex coted; Polar, Finland). Participants were required to attend at least 85% of all the exercise sessions. Subjects were instructed to maintain their dietary patterns during the duration of the study. A 3-d dietary log was collected monthly and analyzed for caloric intake and macronutrient composition.

Cardiorespiratory and metabolic fitness assessment. Before and after 4 months of training, we tested all subjects for weight, waist circumference, exercise maximal fat oxidation (FOmax), peak oxygen consumption (VO2peak), and body composition. All tests were scheduled before and at least 72 h after the last exercise training session to avoid measuring the acute effects of the last exercise bout rather than the chronic effects of the exercise training program. In testing days, subjects arrived at the laboratory after an overnight fast. Upon arrival, subjects voided and their body weight was assessed (Hawk, Mettler Toledo). Urine was analyzed for specific gravity (Usg, Uricon-NE; Atago, Japan) to ensure that subjects were euhydrated (Usg < 1.015). Subjects rested in a stretcher for 20 min while the resting ECG was being examined (Cosmed T12, Italy). Then, exercise testing started in an electromagnetically braked cycle ergometer (Cardiotest 100; Seca, Germany) with three to eight submaximal 4-min stages to assess maximal fat oxidation (FOmax). During the FOmax test, oxygen consumption and carbon dioxide production were analyzed in a breath-by-breath mode (Quark b2, Cosmed). The test was discontinued when the RER exceeded 1. The last minute of each stage was averaged to calculate the nonprotein respiratory quotient and fat oxidation rate (11). Next, subjects recovered for 40 min while 250 mL of juice was being ingested (125 kcal). Then, a graded exercise test (GXT) was conducted to volitional fatigue to determine the subject’s peak aerobic power (VO2peak). After a 5-min warm-up at 100 W, participants began cycling at 125 W with increments of 25 W each minute. Gas exchange data were collected using an automated breath-by-breath system and averaged every 15 s. A physician visually inspected the ECG tracing during the GXT.

Body composition and biopsy tissue collection. Before the above-described test, subjects arrived after an overnight fast before and after training for body composition assessment and muscle biopsy collection. Percent body fat and right leg lean soft tissue mass were determined by dual-energy X-ray absorptiometry (DXA Hologic Serie Discovery Wi QDR; Hologic, Bedford, MA). Muscle biopsies were obtained before and at least 3 d after training from the vastus lateralis using a suction-modified Bergstrom technique (33). Skin was prepared with povidone-iodine (Betadine; MEDA, Paris, France), followed by injection of 2% lidocaine without epinephrine (Braun 2%; Braun Medical, Spain). Then, the skin and underlying
tissues were surgically opened (scalpel blade number 10; Braun, Germany) and muscle tissue was obtained using a 4-mm internal diameter Bergstrom biopsy needle. Upon collection, muscle samples were immediately cleaned of connective tissue, divided into two pieces, and rapidly frozen in liquid nitrogen for subsequent analysis of water and metabolite contents. The incision was closed using adhesive strips (Steri-Strip™, 3M) covered with an adhesive dressing pad (Tegaderm™+Pad, 3M) and compressive dressing (IcoVenda; Novico Medica, Spain).

Muscle water content measurement. All the samples from a given subject were analyzed in the same assay batch. Frozen samples were weighed on an electronic balance with a sensitivity of 0.1 µg (XB220A; Precisa, Switzerland). Elapsed time from sample removal from the freezer until weighing was recorded to permit correction for tissue water evaporation. Samples were freeze-dried in a thermoelectric freeze dryer (Cryodos-50; Telstar, Spain) for 6 h at −50°C and at a vacuum of 10⁻² Torr. In brief, this apparatus freezes the liquid in the sample to −50°C to then sublimate it with a potent vacuum pump at a high flux rate (83 L min⁻¹). Samples were then reweighed in the same precision scale to measure water content. Pilot data in our laboratory in fresh pig leg muscle indicated that H₂O_muscle measurement was highly reproducible (i.e., 6% CV). Data in one subject that underwent 10 resting biopsies within 8 wk in a euhydrated state (Usg < 1.020 and body weight ± 0.25 kg) confirmed the high reproducibility of this technique in our laboratory in human muscle (i.e., 5% CV).

Muscle metabolites analysis. Glycogen concentration was determined from the measurement of glucose after acid hydrolysis analysis (26). Briefly, muscle samples (~20 mg) were homogenized using a glass-on-glass system on ice with deionized water. Then, samples were hydrolyzed in 2 N hydrochloric acid and heated for 2 h at 100°C (Temibloc; JP Selecta, Spain). Finally, samples were neutralized to pH 6.5–7.5 with 1 N sodium hydroxide, and glucose concentration was analyzed by colorimetric assay (enzymatic glucose reagent; Thermo Scientific, Waltham, MA). Muscle protein was assayed after tissue homogenization using a modified Lowry technique with bicinechonic acid (16). Citrate synthase activity was measured from an approximately 10-mg portion of muscle through the reduction of 5,5'-dithiobis(2-nitrobenzoic acid) by the release of CoA-SH in the cleaving of acetyl-CoA (32).

Statistical analysis. Data are presented as mean ± SD. Sample size was calculated based on muscle water increases in three pilot subjects undergoing a similar training program. Power test revealed that at least 10 subjects were needed to reach significance for a statistical power at 80% (α = 0.05) (4). Normally distributed data were analyzed using Student’s two-tailed paired t-test (pre- to posttraining comparison). A Pearson product–moment correlation coefficient was used to establish linear correlations (dependence) between the changes with training in muscle variables and muscle water. Level of significance was set at P < 0.05. Cohen’s formula for effect size (ES) (4) was used, and the results were based on the following criteria: >0.70 large effect; 0.30–0.69 moderate effect; ≤0.30 small effect. Confidence intervals (95%) are also presented. Data analysis was performed using SPSS software for windows (v.18, IBM).

RESULTS

The group was quite homogenous regarding body weight, BMI, and waist circumference (Table 1, pretraining column). Although their initial cardiorespiratory fitness (VO₂peak, Table 1) had a coefficient of variation of 24%, they were all largely untrained according to the normative values for their age and gender (38).

Cardiometabolic fitness and exercise capacity. VO₂peak increased significantly after 4 months of AIT by 3 mL kg⁻¹ min⁻¹ (95% CI, 5–2 mL kg⁻¹ min⁻¹; ES = 0.44; P = 0.001). In turn, FO_max increased by 0.08 g min⁻¹ (i.e., 33%) after 4 months of training (95% CI, 0.12–0.04 g min⁻¹; ES = 0.78; P = 0.003). During the GXT, cycling power output increased by 23 W (i.e., 10% from 232 ± 67 to 255 ± 75 W; Table 1) at the end of the training program (95% CI, 32–14 W; ES = 0.32; P = 0.001).

Anthropometry and body composition. Although subjects did not undergo a hypocaloric diet, body weight was reduced by 1.85 kg (1.9%; from 95.2 ± 9.7 to 93.4 ± 9.6 kg) after 4 months of training (95% CI, −0.4 to −3.3 kg; ES = 0.19; P = 0.022). Furthermore, macronutrient distribution in diet remained constant throughout the 4 months of the study, with 41% ± 2% of energy intake from carbohydrates, 38% ± 1% from fat (40% saturated), and 21% ± 1% from protein. Because of the body weight losses, BMI was also decreased by a similar magnitude (from 33.4 ± 2.9 to 32.7 kg m⁻²; 95% CI, −0.2 to −1.3 kg m⁻²; ES = 0.30; P = 0.013).Fat mass measured by DXA was also significantly reduced with training (−1.7 kg; 95% CI, −3.28 kg; ES = 0.37; P = 0.044), whereas whole body FFM did not significantly change. However, sectional DXA analysis of the right leg revealed a significant increase of 1.8% in FFMM (0.2 kg; 95% CI, 0.31–0.05 kg; ES = 0.10; P = 0.023; Table 1).

Muscle water content. Before exercise, vastus lateralis contained 363 ± 36 mL of water per 100-g dry weight (dw) muscle. After 4 months of exercise, muscle water content (i.e., H₂O_muscle) increased by 12% to 404 ± 69 mL per 100-g dw muscle (P = 0.019). When the changes in water were expressed

### TABLE 1. Exercise and anthropometric changes with 4 months of AIT.

<table>
<thead>
<tr>
<th></th>
<th>Pretraining</th>
<th>Posttraining</th>
<th>% Change</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>VO₂peak (mLO₂ kg⁻¹ min⁻¹)</td>
<td>28.8 ± 7</td>
<td>32.1 ± 8</td>
<td>11%</td>
<td>0.001*</td>
</tr>
<tr>
<td>FO_max (g min⁻¹)</td>
<td>0.04 ± 0.09</td>
<td>0.02 ± 0.11</td>
<td>33%</td>
<td>0.003*</td>
</tr>
<tr>
<td>Workload max (W_max)</td>
<td>232 ± 67</td>
<td>255 ± 75</td>
<td>10%</td>
<td>0.001*</td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>95.2 ± 10</td>
<td>90.26 ± 10</td>
<td>-1.9%</td>
<td>0.023*</td>
</tr>
<tr>
<td>BMI (kg cm⁻²)</td>
<td>33.4 ± 3</td>
<td>32.7 ± 2</td>
<td>-2.2%</td>
<td>0.013*</td>
</tr>
<tr>
<td>Waist circumference (cm)</td>
<td>109 ± 5</td>
<td>107 ± 5</td>
<td>-1.8%</td>
<td>0.016*</td>
</tr>
<tr>
<td>Fat mass (kg)</td>
<td>31.9 ± 4.3</td>
<td>30.1 ± 5.1</td>
<td>-5.4%</td>
<td>0.044*</td>
</tr>
<tr>
<td>FFM (kg)</td>
<td>59.9 ± 9</td>
<td>60.3 ± 8</td>
<td>0.5%</td>
<td>0.477</td>
</tr>
<tr>
<td>Right leg FFM (kg)</td>
<td>9.84 ± 1.84</td>
<td>10.02 ± 1.85</td>
<td>1.8%</td>
<td>0.023*</td>
</tr>
</tbody>
</table>

Data are mean ± SD for 18 subjects. *Significantly different from pretraining.
as per wet weight (i.e., physiological muscle condition), the increases in \( \text{H}_2\text{O}_{\text{muscle}} \) were of 16 g kg\(^{-1}\), increasing from 783 ± 18 to 799 ± 24 g kg\(^{-1}\) ww (95% CI, 27–5 kg; ES = 0.76; \( P = 0.011 \); Fig. 1).

**Muscle metabolites.** Glycogen concentration increased by 19% as a result of training (546 ± 146 to 650 ± 158 mmol kg\(^{-1}\) dw) when expressed as per dry weight (95% CI, 180–27 mmol kg\(^{-1}\) dw; ES = 0.68; \( P = 0.017 \)). However, when glycogen concentrations were expressed as per wet weight, the increases were of 14% and did not reach statistical significance (from 22 ± 7 to 25 ± 7 g kg\(^{-1}\) ww; 95% CI, 7 to −1 g kg\(^{-1}\) ww; ES = 0.43; \( P = 0.154 \); Fig. 2). Total protein concentration was significantly reduced after 4 months of training by 11% from 145 ± 15 to 129 ± 13 g kg\(^{-1}\) ww (95% CI, −6 to −26 g kg\(^{-1}\) ww; ES = 1.17; \( P = 0.007 \); Fig. 2). Citrate synthase activity increased by 31% from 17 ± 5 to 22 ± 5 mmol min\(^{-1}\) kg\(^{-1}\) ww (95% CI, 8–3 mmol min\(^{-1}\) kg\(^{-1}\) ww; ES = 1.08; \( P = 0.001 \); Fig. 2).

**Metabolic syndrome components.** After 16 wk of training, three out of the five components of metabolic syndrome significantly improved. Waist circumference (index of abdominal obesity) was reduced by 2% (i.e., 2 cm), fasting blood glucose by 7% (0.5 mmol \( \text{L}^{-1} \)), and mean arterial pressure by 7% (8 mm Hg). However, blood triglyceride and HDL cholesterol concentrations did not improve with training.

**Correlations of muscle tissue analysis.** Pearson correlation coefficient analyses were performed in selected variables using wet weight as the physiological muscle conditions for expressing concentrations (glycogen and protein), content (water), and enzyme activity (citrate synthase (\( \text{CS}_{\text{activity}} \))). The increase in \( \text{H}_2\text{O}_{\text{muscle}} \) was associated with reductions in protein, muscle glycogen concentration, and citrate synthase activity (Table 2). Conversely, the changes in protein concentration were positively associated with the changes in \( \text{CS}_{\text{activity}} \).

**DISCUSSION**

We trained 18 obese metabolic syndrome patients for 4 months (i.e., 48 sessions) using an intense AIT program (34) and obtained the typical cardiometabolic and body composition improvements previously reported in the literature (21). Our subjects lost body weight and fat mass (−1.9% and −5.4%, respectively; Table 1). They also increased their cardiorespiratory fitness, exercise maximal fat oxidation, and cycling peak power by 11%, 33%, and 10%, respectively (Table 1). The novel finding of our study is that skeletal muscle (i.e., \( \text{vastus lateralis} \)) water content (i.e., \( \text{H}_2\text{O}_{\text{muscle}} \))
increased, whereas, surprisingly, muscle total protein concentration decreased after 4 months of aerobic training. DXA analysis reflected an increase in leg FFM suggestive of muscle hypertrophy despite reductions in muscle protein concentration. Other investigators have shown hypertrophic response to similar cycling endurance training programs in older women and men using microscopic determination of the myofiber diameter and magnetic resonance imaging of the leg (12,14). Our data suggest that increases in muscle water content notably participates in the hypertrophic response to aerobic training in middle-age (54 ± 8 yr old) initially untrained metabolic syndrome men. It also suggests that the mild hypertrophic response to aerobic training detected in our individuals is not due to muscle protein accretion. Furthermore, correlations (Table 2) suggest that the reduction in protein concentration is related to the gain in muscle water probably through a dilution effect.

It seems contradictory that aerobic training would reduce muscle protein concentration although improve leg muscle power (i.e., peak cycling power; Table 1). It could be argued that the weight loss experienced by our subjects set them in a catabolic state, preventing muscle protein anabolism or muscle protein accretion. However, our subjects neither followed a hypocaloric diet nor reduced the percent of protein ingested in the diet (i.e., 21% of energy intake). Thus, the moderate weight loss experienced during training (i.e., 1.85-kg weight loss or 1.9% of body mass) was most likely induced by a lack of compensatory increase in calorie intake to match energy expenditure during training. In fact, most of the weight loss (93%; i.e., 1.73 out of 1.85 kg) could be accounted for by reduction in fat mass tissue (Table 1) in the abdominal region as judged by the reduction in waist perimeter. Harber et al. (12) subjected old women (71 ± 2 yr old) to 12 wk of aerobic training and also found increased skeletal muscle water content and reductions in muscle protein concentration despite null losses of body weight. Furthermore, they measured isolated fiber contractile function and found increased unloaded contraction velocity and absolute power in type I fibers (12). Our data amount to theirs to suggest that aerobic training improves leg peak power ($W_{max}$; Table 1) despite reducing muscle protein concentration.

Although H$_2$O$_{muscle}$ gains have been reported as an acute response during the first stages of moderate (5) and intense exercise (30), we are the first to report an increased H$_2$O$_{muscle}$ 72 h after the final session of exercise of a 4-month-long training program. We interpret this to suggest that gain in muscle water is another of the adaptations to endurance training. Aerobic or resistance training seems to reduce specific force (the ratio between isometric force and fiber cross-sectional area) in both young (18) and old subjects (12,14). Reduction in specific force evidences that the increase in size due to aerobic training is not entirely functional. Others have suggested that the reduced specific force is due to edema or swelling. Increased in H$_2$O$_{muscle}$ diluting myofibrillar protein could explain the reduced specific force after aerobic training reported in the literature. Our measurements of increased H$_2$O$_{muscle}$ support this view.

The gain in muscle water that we observed may have remodeled vastus lateralis fibers’ contractile performance. Dehydration of rat skinned isolated single fibers by bathing the muscle preparation in a hyperosmotic fluid (10% dextran) reduces the space between myofilaments (myosin and actin) and lowers the maximal shortening velocity. The authors suggest that filament lattice compression likely affects the rate constant for cross-bridge detachment (19). In humans, 17 d of bed rest increased the type I fiber unloaded shortening velocity, although they reduced their peak isometric force (39). On the other hand, resistance training can increase the muscle strength and cross-sectional area without changes in myofilament spacing (3) when a reduction in this spacing was expected because of myofilament packing (27). It is then possible that the gain in muscle water reported presently constitutes a training adaptation geared to increase the space between thin and thick filaments allowing faster cross-bridge cycling. Vastus lateralis importantly contributes to cycling mechanics, and the increase in cycling power in our data suggests improvement in the rate of force applied to the pedals from this muscle. Based on the current literature, it is reasonable to hypothesize that the gains in muscle water content increased filament spacing allowing higher muscle power.

Our proxy marker of mitochondrial density (citrate synthase activity) increased by 31% with the completion of the 48 exercise training sessions (Fig. 2). Mitochondria constitutes approximately 4%–6% of muscle tissue (15), and its proliferation could be contributing to the modest leg hypertrophy currently reported (Table 1). However, the decrease in protein concentration is counterintuitive in the face of an increased mitochondrial density. Sarcomeres have more protein concentration than mitochondria and thus a superior mitochondrial proliferation with respect to sarcomere could explain the reduction in total protein concentration. We did not separate the different protein fractions, but Harber et al. (12) reported that 12 wk of aerobic training reduces myofibrillar protein although does not affect sarcoplasmic protein concentration. Thus, the large increase in citrate synthase suggests gains in mitochondria that may have contributed to the reduced protein concentration.

On the other hand, the increased H$_2$O$_{muscle}$ with aerobic training may be influenced by the 14% higher muscle glycogen storage after training. The literature suggest that muscle glycogen may require water for its storage (10,24). Other researchers have proposed that the combined increases in mitochondrial volume and glycogen stores could account for a significant portion of the increases in leg cross-sectional area with exercise training (23). Water and glycogen were the two components of muscle that increased after training. The increase in muscle water content was inversely associated with typically regarded adaptations to endurance training such as increased muscle glycogen, protein concentration, and citrate synthase activity (Table 2). This inverse association reveals the strong dilution effect that muscle water has in these muscle metabolites. In fact, the changes in glycogen, protein, and citrate synthase were
positively associated among them, probably driven by the dilution effect of water in all of them.

In contrast, other investigators have reported no changes in $H_2O_{\text{muscle}}$ after training young individuals using resistance training (17). Competitive runners display lower muscle water content than recreational runners (29), arguing against muscle water expansion as an adaptation to endurance training. However, the same authors found that training for a marathon reduces slow and fast twitch fiber size (35). A deficit in nutritional water during the extraneous training for the marathon may refrain the otherwise natural muscle water expansion. Of note, our findings of increased $H_2O_{\text{muscle}}$ with training might be specific to very untrained individuals because our subjects’ $\dot{V}O_{2\text{peak}}$ ranked in the lower 20% percentile according to the American College of Sports Medicine guidelines (38). Possibly, the initial muscle atrophy due to detraining and a suboptimal muscle water content may set the scenario for the increases in muscle water content with aerobic training. In fact, muscle water content before training in the current metabolic syndrome men was 8% lower than in a group of young endurance-trained cyclist recently tested in our laboratory (363 ± 9 vs 395 mL per 100-g dw) (9). Likewise, initial muscle water levels may explain why identical training increased muscle water content in old women (71.9% ± 1.0% of water in muscle) (13) but not in old men (77.8% ± 0.9% of water in muscle) (14). Our data suggest that muscle water expansion is not gender specific because our men increased muscle water content, but it is influenced by pretraining water content (73.8% of initial muscle water in muscle in our data).

In summary, we found increases in muscle water as a result of an endurance training program in much deconditioned ($\dot{V}O_{2\text{peak}}$, 28.8 ± 7 mL$O_2$·kg$^{-1}$·min$^{-1}$) obese metabolic syndrome men. The increases in muscle water were negatively associated with changes in protein and glycogen muscle concentration, suggesting a dilution effect. Nevertheless, glycogen concentration tended to increase while muscle protein concentration decreased. Interestingly, the increase in muscle water rather than protein accretion can explain the mild leg hypertrophy observed after the endurance training program (48 sessions). Two possible factors accounting for the increased cycling peak power despite reduced total muscle protein are improved neuromuscular function with training and/or myofilament remodeling linked to the gains in muscle water. Lastly, the diluting effect of muscle water expansion should be taken into account when expressing muscle metabolites or substrates as per unit of wet weight (i.e., original physiological state).

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REFERENCES


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AQ5 = Please provide the city and state/country of IBM.

AQ6 = In Table 2, should “ms” be changed to “muscle”?

AQ7 = The sentence beginning “In fact, most of the” has been altered for clarity, please check that the meaning is correct.

AQ8 = The sentence beginning “Sarcomeres have more” has been altered for clarity, please check that the meaning is correct.

AQ9 = References 12 and 13 contain almost the same data. Please check and renumber.

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