Metabolic and Morphometric Alterations Inherent to Neuromuscular Electric Stimulation in the Antagonist Muscle Submitted to Ankle Joint Immobilization

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ABSTRACT

The aim of this work was to investigate the effect of the neuromuscular electrical stimulation (ES) on the metabolic and morphometric profile of the tibialis anterior muscle, antagonist to the soleus muscle which was stimulated, under the joint immobilization condition of the ankle for 7 and 15 days. The immobilization promoted the reduction in the muscle mass (I7: 17.36%, I15: 20.83%), in the glycogen content (I7: 48%, I15: 48%), in the muscle fibers areas (I7: 27%, I15: 40%) and increase in intramuscular connective tissue density (I7: 122%, I15: 206%). The EE didn’t promote significant alterations in the mass of the immobilized groups, however, promoted increase in the glycogen (IEE7: 31.25%; IEE15: 56.25%), reduction in the muscle fibers areas (IEE7: 14%, IEE15: 24.69%) and reduction in the connective tissue density (IEE7: 25.63%, IEE15: 49.09%) when compared with the respective immobilized groups.

Key words: Immobilization, muscle disuse, metabolic, morphometric, neuromuscular electrical stimulation, rehabilitation

INTRODUCTION

Several studies have demonstrated that the muscle disuse induces a series of harmful effects, such as the proliferation of intramuscular connective tissue, muscular hypotrophy, reduction in glycogen content, capillary density, serial sarcomeres, as well as force and fatigue resistance (Józsa et al., 1990; Reardon et al., 2001). With the intention of minimizing the events set off by the muscle disuse, several techniques have been used, in an endeavor to improve the homeostatic conditions of the muscle fibers, with neuromuscular electrical stimulation (ES) (Silva et al., 1999). It is worth pointing out that ES has been used directly in the disused muscle. However, some studies were not limited to analyzing its effect on the stimulated muscle only, but also on the antagonist and contralateral muscles. In this context, Lieber et al. (1988) described the effectiveness of the resource in minimizing the hypotrophy in the directly stimulated muscle (tibialis anterior), and also its antagonist (soleus), since the authors considered that the latter underwent passive stretching during the ES. Ashmore and Summers (1981) described a direct relationship between the passive stretching and the increase in the muscle fiber diameter.

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Furthermore, it was also observed that the static stretching was more efficient for the muscle protein synthesis than the contraction induced by ES (Goldspink et al., 1995). Previous work have demonstrated that the ES minimized some of the harmful effects inherent to joint immobilization on the soleus muscle, such as the reduction of the muscle mass, fiber area, glycogen content, as well as an increase in the intramuscular connective tissue (Cancelliero et al., 2005; Durigan et al., 2006; Silva et al., 2006). These aspects have, however, not yet been characterized in the antagonistic muscles to those receiving the direct electrical stimulation, during the joint immobilization period.

Considering these observations, the aim of this study was to investigate the effect of the ES on the metabolic and morphometric profile of the tibialis anterior muscle, antagonist to the soleus muscle, which was stimulated under the condition of ankle immobilization in the neutral position for 7 and 15 days.

**MATERIAL AND METHODOS**

Male *Wistar* rats (3 to 4 months old; 250-300g) were kept under the controlled vivarium conditions, with free access to the food and water and treated in accordance with the recommendations in the Guide on the care and use of the laboratory animals (National Research Council, 1996). The size of the box containing three animals was 40 x 30cm and it was covered with the newspaper, in order to prevent the sawdust from entering the internal compartment of the orthoses and causing cutaneous lesions. The animals were divided into five groups (n=5): control (C), immobilized for 7 days (I7), immobilized for 15 days (I15), immobilized associated with the ES for 7 days (IES7) and immobilized associated with the ES for 15 days (IES15). For the immobilization, the rats were anesthetized with the sodic pentobarbital (50 mg/Kg weight) and the left hindlimb was tricotomized and ES was realized in daily sessions of 20 minutes for 7 and 15 days, initiating immediately after the immobilization. An electrode (1cm²) was placed in the inguinal region and the other (1cm²) in the triceps sural muscle in the immobilized limb so that this second electrode was connected inside of orthoses. The ES parameters were the frequency of 10Hz, pulselwidth of 0.4ms, two-phase quadratic pulse, intensity of 5.0mA, from the visualization of the muscle contraction, increased 1.0 mA after each 5 minutes to keep the contraction level during the all stimulation period. After the experimental period, the analyses performed in the tibialis anterior were glycogen content, weight, fiber area and intramuscular connective tissue.

To determine the muscle glycogen, the muscle samples were digested in 30% KOH at 100°C and the glycogen was precipitated by ethanol, after which, the sample was centrifuged at 3000 rpm for 15 minutes. The precipitated glycogen was submitted to the acid hydrolysis in the presence of phenol. The values were expressed in mg/100 mg of wet weight and the method used was of Siu et al. (1970).

For the morphometric analysis, the ventral segment of the tibialis anterior was fixed in buffered 10% formol solution. The tissue was embedded in the paraffin, cut into several non-serry 7 µm thick cross-sections and stained with hematoxylin-eosin. The image analysis system used consisted of the Image Pro-plus 4.0 software (Media Cybernetics, Silver Spring, MD, USA) and a digital camera (JVC® manufacturer, Lawrenceville, GA, USA) coupled to a microscope (Zeiss, Narberth, PA, USA) and connected to a microcomputer. All the images were captured at 10X magnification. The areas of 375 tibialis anterior muscle fibers from each animal were analyzed as follows; 15 fibers per area in five areas per section, and a total of five sections per animal. A square reticulum was used for randomly choosing 15 fibers per cuts that coincided with the straight intersections.

A planimetry system was used for the analysis of the intramuscular connective tissue density by scoring the points by means of a reticulum with 2500µm² squares containing 56 straight line intersections (Mathieu et al., 1981). The coincident points in the endomysium and perimysium in five areas per section in five sections per animal corresponded to a total of 1400 points per animal. The relative area of connective tissue (area density) was calculated by dividing the sum of the number of the coincident points in the straight line intersected
intersections in the connective tissue (endomysium and perimysium) by the total number of points. The statistical analysis was initially performed by the Kolmogorov-Smirnov normality test and for the homoscedasticity test (Bartlett criterion). After the observation that the variables had contemplated the parametric methodology, ANOVA test, was performed and when the presented difference was significant, the Tukey test was performed for the multiple comparisons. The level of significance was set at P < 0.05 for all the analyses.

RESULTS

Significant reduction was observed in the muscle mass (mg) as follows: 17.36% (p<0.05) in I7 group; and 20.83% (p<0.05) in I15 group, when compared with C, and no significant difference was observed between I7 and I15. Moreover, there was a reduction of 48% (p<0.05) in the glycogen content in both the immobilized groups. In the morphometric analysis, I7 and I15 groups demonstrated significant reduction in the fiber area when compared with C, represented by 27% and 40% (p<0.05), respectively, in addition to presenting significant differences among them. The intramuscular connective tissue density demonstrated a significant increase of 122% in I7 and of 206% in I15, in relation to C (p<0.05), and I7 presented significant difference when compared with I15.

The ES did not promote significant alterations (p>0.05) in the muscle mass in the IES7 and IES15 groups in relation to the respective immobilized groups. On the other hand, there was a significant increase of 31.25% in the glycogen content in the IES7 (p<0.05) and 56.25% in IES15 (p<0.05), when compared with the respective immobilized groups. Moreover, IES7 did not show significant difference when compared with IES15. The muscle fiber area presented a reduction of 14% in IES7 and 24.69% in IES15 (p<0.05) when compared with the respective immobilized groups, whereas IES7 and IES15 presented no significant difference. The resource also promoted a significant reduction in intramuscular connective tissue density of 25.63% in the IES7 (p <0.05) and 49.09% in the IES15 (p <0.05) when compared with the respective immobilized groups. Furthermore, the IES7 presented no significant difference when compared with the IES15. (Table 1 and Fig. 1).

DISCUSSION

There are several factors that influence the results mentioned in the studies involving immobilization, with emphasis on the joint position, which could interfere in the development of the harmful effects inherent to the muscular disuse. In general, the shortened position reduces the muscle mass, fiber area, number of serial sarcomeres and connective tissue proliferation. The immobilization in the stretched position leads to the increase of the muscular mass, protein synthesis, additional serial sarcomeres and less connective tissue (Fournier et al., 1983; Williams and Goldspink, 1984; Appell, 1986; Williams et al., 1988; Savolainen et al., 1988). This analyzed the joint immobilization in the neutral position, because in most of the studies it has been done in flexion or extension, either by the plaster cast, resin, steel mesh, or even by the hindlimb suspension. There are few studies that used the neutral ankle position (90º), showing harmful effects on the skeletal muscle system, which were associated with knee and hip joint immobilization at 90º (Qin et al., 1997; Hirose et al., 2000). The acrylic resin orthoses model used in this study, however, differed from the others, because it kept the knee and hip joints free (Silva et al., 2006). Not only the joint position interferes in the results, but also the disuse period. Some studies suggest that most of the changes take place in the first seven days of the muscular disuse (Williams and Goldspink, 1984; Okita et al., 2004). Other authors also observed alterations in the period of 15 days (Bodine et al., 2001; Picquet and Falempin, 2003). Thus, in the present study the immobilization periods of 7 and 15 days were opted for.
Table 1 - Mean±sd of tibialis anterior mass (mg), muscle fibers area (µm$^2$), connective tissue area density (%), glycogen content (mg/100 mg) of control group (C), immobilized for 7 days (I7), immobilized for 15 days (I15), immobilized treated with ES for 7 days (IES7) and immobilized treated with ES for 15 days (IES15) n=5, p<0.05, * compared with control group, † compared with respective immobilized group, # compared with I7.

<table>
<thead>
<tr>
<th></th>
<th>C</th>
<th>I7</th>
<th>IES7</th>
<th>I15</th>
<th>IES15</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass (mg)</td>
<td>432.6±51</td>
<td>357.2±37.5*</td>
<td>341±55*</td>
<td>342.8±44.1*</td>
<td>324.9±23.4*</td>
</tr>
<tr>
<td>Area (µm$^2$)</td>
<td>2046±436</td>
<td>1488±391*</td>
<td>1697±330*†</td>
<td>1211±297*#</td>
<td>1510±395*†</td>
</tr>
<tr>
<td>Connective (%)</td>
<td>7.78±3.3</td>
<td>17.28±4.27*</td>
<td>12.85±5.15* †</td>
<td>23.85±6.06*#</td>
<td>12.14±4.08*†</td>
</tr>
<tr>
<td>Glycogen (mg/100 mg)</td>
<td>0.31±0.03</td>
<td>0.16±0.04*</td>
<td>0.21±0.02*†</td>
<td>0.16±0.03*#</td>
<td>0.25±0.05*†</td>
</tr>
</tbody>
</table>

Figure 1 - Tibialis anterior muscle fibers of C group (A), muscle fibers of I7 group (B), muscle fibers of IES7 group (C), muscle fibers of I15 group (D), muscle fibers of IES15 group (E). A reduction of muscle fiber area (*) and an increase of intramuscular connective tissue (arrows) can be seen in immobilized groups with compared to control group. An increase of muscle fiber area (*) and a decrease of intramuscular connective tissue (arrows) can be seen in immobilized associated to ES groups with compared to respective immobilized groups.
The immobilization for 7 and 15 days in the neutral position promoted significant reduction in the tibialis anterior muscle mass, fiber area, increase of intramuscular connective tissue, in addition to reduction of the glycogen content, showing the interrelation of contractile activity with energy homeostasis and muscle fiber morphology, pointing towards a condition of muscle hypotrophy. According to Qin et al. (1997), the immobilization for different periods results in the hypotrophy varying from 15% to 70%, depending on the animals used and on the results in the hypotrophy varying from 15% to 70%, depending on the animals used and on the type of fibers assessed. Savolainen et al. (1988) reported reduction in the muscle mass to the order of 35.14% in the shortened position and of 16.31% in the stretched position in the rat tibialis anterior muscles submitted to the plaster cast ankle immobilization for 7 days. The immobilization for 7 days also promoted significant increase in the connective tissue density to the order of 122% and 168% on the fifteenth day, when compared with the control group. These results are in agreement with the literature, since for Józsa et al. (1990), irrespective of the muscular disuse model studied (immobilization, tenotomy or denervation) and of the muscle type, the amount of connective tissue in the endomysium and perimysium increased significantly, varying from 50% to 700%.

The muscle disuse caused by the conditions of long periods in the bed, ortheses or limb fixations and microgravity, induces resistance to the insulin and a catabolic state in the affected skeletal muscles of the humans (Ferrando et al., 1996). Hirose et al. (2000), in rats that had the left hindlimb immobilized in the neutral position for 7 days, observed reduction in the transduction of the intracellular signal stimulated by the insulin, suggesting deficit in the insulin receptor activation and in the molecules activated by this, indicating that the insulin resistance could also be set off during the immobilization. This condition of resistance to insulin could explain the compromised glycogen content as a result of the immobilization of the tibialis anterior muscle analyzed in the above mentioned study.

As regards the use of the ES, the resource was shown to be effective in minimizing the increase of the connective tissue density, as well as reducing the muscle fiber area of the tibialis anterior submitted to immobilization for 7 and 15 days. These results corroborate the statement of Avramidis et al. (2003) who described the importance of the ES to recovering the muscle force, reducing the connective tissue, reducing the time of rehabilitation and preventing the muscle atrophy due to the immobilization. Qin et al. (1997) used the ES with a frequency of 50Hz applied daily for 30 minutes, five times the week for three weeks in the rabbit tibialis anterior muscles and observed that the resource was effective for preventing the muscular hypotrophy, minimizing the fiber area reduction, interstitial fibroses and blood supply deficiency.

According to Goodyear et al. (1992), the increase in the glycogen content was due to the larger glucose uptake by the GLUT4 population that was insensitive to the insulin, externalized and resulted from the activation of the enzymatic cytosolic systems in the glycogenesis. Other studies also mentioned the ES as a resource that increased the glucose uptake and metabolism, and cellular metabolic pathway activities (Silva et al., 1999; Guirro et al., 2004). Etgen et al. (1993) evaluated the GLUT 4 content in the rat plantar muscle after the chronic ES and observed an increase of 82% in its content. Longer periods of the ES, 30 to 40 and 60 to 90 days, only showed a tendency towards increase in GLUT4 content, reaching a plateau around 30 to 40 days. An important result of the study of Hamada et al. (2003) was that the corporal glucose uptake in the rats was acutely increased in response to 20 minutes of the ES, and this increase lasted for at least 90 minutes after the use of the resource ended. The results of the present study demonstrated that the muscular glycogen remained elevated even after 24 h of the ES.

Other results add support to the concept that passive muscular tension is an important factor in the attempt to minimize the harmful effects on the skeletal muscle system during the disuse. Lieber et al. (1988) studied the effects of the ES applied in the directly stimulated tibialis anterior muscle and in the soleus muscle (antagonist), associated with the joint immobilization of the rat ankles in a neutral position for four weeks. The following parameters were used: frequencies of 10 and 50Hz, pulsewidth of 0.3ms, 1 hour a day, applied five times a week. The resource minimized the fiber area and muscle mass reduction, both of the tibialis anterior and the soleus, in comparison with the control group, possibly due to the passive tension imposed on the soleus because of activating the tibialis anterior, as well as reflex...
activation through the antidromic transport.
The results of the present study corroborated these findings, since the direct stimulation of the soleus muscle minimized the reduction of the fiber area, glycogen content and the increase in the connective tissue density in the tibialis anterior muscle. These findings cannot, however, be attributed to the passive stretching, since the immobilization promoted by the acrylic resin orthoses fixed the ankle in a neutral position that disabled the stretching of the tibialis anterior muscle. Moreover the current density used in that study was 5 mA/cm², being extremely high, considering that for the first flexors muscle contraction elicited by the ES in the human, a current density of between 0.5 and 2 mA/cm² was used. (Wang et al., 2004)
The high current density applied in the animals might have generated an electrical field broad enough to stimulate both the posterior compartment (directly stimulated soleus muscle), and the anterior (tibialis anterior). The proximity of the soleus muscles and tibialis anterior, as well as the good electrical current transport by the muscular tissue should also be considered. Thus, the tibialis anterior could have been activated by the action of the electrical field, minimizing the harmful effects promoted by the immobilization.
The ES applied directly in the soleus muscle minimized the reduction of the glycogen content, fiber area and increase in the connective tissue density in the tibialis anterior muscle during the joint immobilization condition in the periods of 7 and 15 days. Thus, the importance of the physical therapy intervention with the ES during this period could be important, not only to minimize the muscle skeletal alterations in the directly stimulated muscle, but also in its antagonist, since the ES was a resource of extreme importance in the rehabilitation process.

RESUMO
O objetivo desse trabalho foi investigar o efeito da estimulação elétrica neuromuscular (EE) sobre o perfil metabólico e morfométrico do músculo tibial anterior, antagonista ao músculo sóleo, o qual foi estimulado, sob a condição de immobilização articular de tornozelo durante 7 e 15 dias. A immobilização promoveu redução na massa muscular (I7: 17,36%, I15: 20,83%), no conteúdo de glicogênio (I7: 48%, I15: 48%), na área das fibras musculares (I7: 27%, I15: 40%) e aumento na densidade do tecido conjuntivo intramuscular (I7: 122%, I15: 206%). A EE não promoveu alterações significativas na massa muscular dos grupos imobilizados, porém, promoveu aumento no glicogênio (IEE7: 31,25%; IEE15: 56,25%), redução na área das fibras musculares (IEE7: 14%, IEE15: 24,69%) e redução na densidade do tecido conjuntivo de (IEE7: 25,63%, IEE15: 49,09%) quando comparado aos respectivos grupos imobilizados.

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