L-arginine supplementation and resistance training promotes increase of maximum strength and protection from DNA damage in rats

Abstract

L-arginine (Arg) is a precursor of several substances with remarkable physiological functions, such as nitric oxide (NO) and the amino acid creatine. Arg supplementation has been associated with increased strength in high-intensity exercise. However, there is no clear evidence of association of supplementation with Resistance Training (RT), on their antioxidant and ergogenic effects, specifically on their genotoxic protection. Therefore, we evaluated the effect of supplementation of Arg associated with RT 8weeks about gaining maximum strength and DNA damage in rats. Ten Wistar rats (220-270g, 90days old) were randomly divided into 3 groups: sedentary (SED, n=4), Resistance Training (RT, n=2) and Resistance Training + L-Arginine (RT+Arg, n=4). Trained rats were submitted to the protocol of RT in a squat apparatus adapted for rats (4 sets of 10-12 reps with 90s interval, 4times/week, 65-75% of One Maximum Repetition (1MR) for 8weeks). The supplemented group received Arg (500mg/kg) daily for 8weeks. After 8weeks, whole blood was collected from animals to perform the comet assay. It was used CASP software (CASPLabs®) for quantification of DNA damage. To quantify the maximum strength the 1MR test was performed at baseline and after the RT protocol. For comparisons among groups it was performed One-way ANOVA followed by post hoc test of Student-Newman-Keuls. For associations, it was used the Pearson correlation test. We considered the level of significance of 5%. The RT was able to protect DNA damage in rats. However, the association with Arg supplementation was able to promote greater genotoxic protection, compared to sedentary rats. These results indicate remarkable ergogenic action and genotoxic protection of Arg supplementation in association with RT.

Keywords: arginine, resistance training, supplementation, exercise, DNA damage, genotoxicity, rats

Introduction

The supplementation of L-arginine (Arg) has been showed to improve performance in high intensity and short duration exercises, such as weightlifting, possibly due to its vasodilating action. In this reaction mediated by the conversion of Arg to citrulline, the production of nitric oxide (NO) occurs by the enzymatic catalysis of nitric oxide synthase (NOS). The NO is a Reactive Oxygen and Nitrogen Specie (RONS), found in the form of molecular gas. The responsible enzyme to produce NO is NOS, which may be expressed constitutively in endothelial isoform (eNOS), neuronal (nNOS) and inducible form (iNOS). The decoupling of NOS constitutive forms lead to the overproduction of two oxidants in endothelium: the superoxide anion-radical (O$_2^-$) and peroxynitrite (ONOO-). These both free radicals are associated with the development of endothelial disfunction.

Under normal conditions, endothelium produces NO by Arg in the presence of oxygen catalyzed by eNOS. The NO diffuses into the adjacent smooth muscle, where it is modulated by guanylate cyclase. This enzyme has the function of converting guanosine triphosphate (GTP) into cyclic guanosine monophosphate (cGMP), thereby causing relaxation of vascular wall. In situations where it is established a dysfunction of vascular endothelium, production of NO occurs in the same manner, together with high concentrations of O$_2^-$, which is caused by the decoupling of electrons in the reaction of eNOS. In this case, the O$_2^-$ can be converted into hydrogen peroxide.

The NO may react with O$_2^-$ and form ONOO-, which is a highly cytotoxic compound. Free radicals have high reactivity to adjacent biomolecules. It is well established that reactive species leads to the impairment of important cellular structures such as lipid peroxidation (mainly in plasmatic membranes), damage in proteins (signaling proteins, enzymes and other proteins) and genomic damage (oxidation of nucleic acids).

The Resistance Training (RT) has been widely utilized by the general population as a very effective method in promoting gains of muscle strength, increase power performance in sports, preventing muscle lesions and increased quality of life. Despite of the scientific literature demonstrate benefits of Arg supplementation in exercised conditions, there are few evidence of associating of supplementation with RT on their ergogenic effects, as well as on genotoxic protection. For that reason, we evaluated the effect of supplementation of Arg associated with RT 8weeks about gaining maximum strength and DNA damage in rats.

Methodology

Ten Wistar rats (220-270g, 90days old) were randomly divided into 3 groups: Sedentary (SED, n=4), Resistance Training (RT, n=2) and Resistance Training + L-Arginine (RT+Arg, n=4). The animals were housed under standard conditions (food and water ad libitum, ...
temperature between 22 and 24°C, light - dark cycle of 12 hours). This study was approved by CEUA/UFCSPA, under the protocol number 114/13.

Resistance training protocol

Trained rats were submitted to a RT protocol in a squat apparatus adapted for rats10-12 (4 sets of 10-12 reps with 90s interval, 4 times/week, 65-75% of One Maximum Repetition (1MR) for 8 weeks). An electrical stimulus (4-5mA, 0.3 seconds long, with a 3 second interval between each repetition)10 was applied in the rat’s tail using a surface electrode, in order to provoke the extension movement of the lower limbs of the rat.

To quantify the maximum strength, it was performed the 1MR test at baseline and after the RT protocol. To determine the maximum lifted load in one repetition, the One Maximum Repetition (1MR) was utilized. From the obtained value, the load percentage required to perform the training protocol were determined. In response to training, it is expected strength gains along the training, making the realization of retests every two weeks necessary, in order to adjust the training load.

L-Arginine supplementation

It was utilized L-Arginine (presentation form: powder, with purity of 99.9%, Sigma-Aldrich®; Brazil). The supplementation was given by gavage solution, as this resembles human oral consumption and ensures that the desired dosage is achieved. The supplemented group received Arg (500mg/kg) daily for 8 weeks. The animals received the supplement every day before training for the entire period of the protocol (including the days on which they did not train).

Comet assay (alkaline version)

After 8 weeks, whole blood was collected from animals to perform the comet assay. The alkaline Comet Assay in peripheral blood was performed as described in the literature2-13 with minor changes. It was used 20μL of whole peripheral blood embedded in 90μL of 0.75% low-melting point agarose at 37°C. This mixture was placed into a slide previously coated with 1.5% of normal melting point agarose processed at 60°C. A cover glass was placed on top and the agar allowed to set at 4°C and, after gel solidifying, the cover glass were removed. Then, the slides were immersed in iced-cold lysis solution (2.5 M NaCl, 100 mM EDTA and 10mM Tris, pH 10.0; containing freshly added 1% (v/v) Triton X-100 and 10% (v/v) dimethylsulfoxide (DMSO) at 4°C in dark for a minimum of 1h. Afterwards, to allow DNA unwinding, slides were incubated in a freshly made alkaline electrophoresis buffer (0.3MNaOH and 1mM EDTA; pH >13) at 4°C for 5min in a horizontal electrophoresis box. The alkaline electrophoresis was carried out for 15min at 25 V and 300mA. Every step was carried in dark. After electrophoresis, slides were washed three times in a neutralization buffer (0.4M Tris; pH 7.5) for 5min, rinsed twice in distilled water, and left to dry overnight at room temperature. Then, the slides were fixed for 10min in trichloroacetic acid 15% w/v, zinc sulfate 5% w/v, glycerol 5% w/v, rinsed three times in distilled water, and dried for 2h at 37°C. Finally, the slides were stained with silver nitrate as previously described by Nadin et al.,14

For DNA damage quantification, it was used CASP software (CASPLabs®).15 The software can detect three different types of markers of genotoxic damage, such as percentage of tail DNA, tail moment (represents the distance of DNA migration from the head of the comet) and olive tail moment (represents the product of the tail length and the fraction of DNA in the tail). For image analysis it was scored 100 comets per slide.

Statistical analysis

The results are expressed in mean±SD. For comparisons among groups it was performed One-way ANOVA followed by post hoc test of Student-Newman-Keuls. For associations, it was used the Pearson correlation test. We considered the level of significance of 5%.

Results

All of the groups showed no differences at the baseline of resistance training protocol (P>0.05). After 8 weeks of training, it was observed higher maximum strength in the trained groups, when compared to the sedentary group (P<0.05). However, the RT + Arg group showed greater maximum strength than the other groups (P<0.05) (Figure 1).

Regarding damage to DNA, % Tail DNA was lower in RT + Arg, compared with RT and SED groups (P<0.05) (Figure 2) as well as in other genotoxic markers: Tail Moment (TM) (P<0.05) (Figure 3) and Olive Tail Moment (OTM) (P<0.05) (Figure 4).

It was observed a strong inversely proportional correlation between different markers of DNA damage with maximum strength. Percentage of Tail DNA with 1MR (r=-0.9719, P<0.05), TM with 1MR (r=-0.9046, P<0.05) and OTM with 1MR (r=-0.9079, P<0.05).

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**Conclusion**

The RT was able to protect DNA damage in rats. However, the association with Arg supplementation was able to promote greater genotoxic protection, compared to sedentary rats. Also, the association of RT with Arg supplementation increase maximum strength after the training protocol. These results indicate remarkable ergogenic action and genotoxic protection of Arg supplementation in association with RT.

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**Conflict of interest**

Author declares that there is no conflict of interest.

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