Comparison of training and detraining on redox state of rats: gender specific differences

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Abstract. Given the fact that oxidative stress response induced by training/detraining has still not been clarified and may be influenced by gender, the aim of our investigation was to compare the effects of swimming training and detraining on oxidative and antioxidative parameters in rats, with a special focus on sex differences. Wistar albino rats (n = 64) were divided into 4 groups: control, trained group, groups exposed to 2 and 4 weeks of detraining. Each group included two subgroups: males and females. After sacrificing, hearts were isolated and retrogradely perfused according to Langendorff technique. Levels of superoxide anion radical, hydrogen peroxide, nitrites and thio-barbituric acid reactive substances were measured in plasma and coronary venous effluent, while reduced glutathione, activities of superoxide dismutase and catalase were measured in erythrocytes. Our results indicate that swimming training doesn’t promote oxidative damage, nor act protectively within the heart. However, 2 and 4 weeks of detraining led to a partial lost in exercise-induced adaptation. It seems that moderate-intensity physical exercise of sufficient duration leads to beneficial adaptations, which may be partially lost during detraining period. Positive antioxidative effects of training remained longer in males. Findings of present study may help in elucidation of training and detraining effects on modulation of redox homeostasis, especially from aspect of gender differences.

Introduction

Under normal physiological conditions, pro-oxidants are continuously produced as a result of essential metabolic processes and environmental factors (Pham-Huy et al. 2008). Aerobic organisms possess antioxidant system which consists of variety of enzymatic and nonenzymatic antioxidants that serve to counterbalance the effects of oxidants (Birben et al. 2012). Oxidative stress occurs when there is an imbalance between oxidants and antioxidants in favour of oxidants. Since pro-oxidants are very reactive molecules, they can cause tissue damage interacting in a destructive manner with practically every cellular component (Birben et al. 2012; Rahal et al. 2014).

Regular physical exercise has been shown to exert a myriad of beneficial effects for health, such as promotion of health and lifespan, improvement of quality of life and decrease the incidence of a life-style related diseases (Macera et al. 2003; Vina et al. 2012). Groundbreaking research which gave the first data about the association between exercise and oxidative stress was conducted by Dillardi et al. (1978) approximately 4 decades ago. This finding stimulated further curiosity of many authors regarding the role of reactive oxygen species (ROS) and reactive nitrogen species (RNS) in skeletal muscle and other metabolically active organ during physical exercise (Davies et al. 1982; Ammeren et al. 1992; Gomez-Cabrera et al. 2008). Numerous papers have been published proving that low physiological levels of pro-oxidants produced in the muscles have an important role in maintenance its normal tone and

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contractility. On the contrary, excessive production of ROS leads to a contractile dysfunction which is followed by muscle weakness and fatigue (Powers and Jackson 2008; Rahal et al. 2014). It has been established that swimming training causes changes in antioxidant enzymes, alters muscle gene expression, thus contributing to exercise-induced adaptations to skeletal muscle (Venditti and Di Meo 1997; Elikov 2016; Ruzicic et al. 2016). However it should be taken into consideration that various factors influence the oxidative stress response to swimming training, such as type of exercise, intensity, duration, gender and age of athletes etc. (Ruzicic et al. 2016).

Interestingly, prolonged cessation of training stimulus, known as detraining, may abolish training-induced adaptations in response to oxidative stress and antioxidant status markers (Fatouros et al. 2004; Agarwal et al. 2012). Nevertheless there is a lack of data referring to the oxidative stress response to detraining.

Gender differences in response to exercise-induced oxidative stress have gained increased attention despite controversial results (Liu et al. 2000; Balci and Pepe 2012; Farhat et al. 2017). To our best knowledge there are just a few studies which explored the potential gender difference in oxidative stress markers and antioxidant defense system in the period of detraining.

Given the fact that oxidative stress response induced by detraining has still not been clarified and may be influenced by gender, the aim of our investigation was to compare the effects of swimming training and detraining on oxidative stress parameters and parameters of antioxidant defense system in rats, with a special focus on sex differences.

Materials and Methods

The study was performed in the laboratory for cardiovascular physiology of the Faculty of Medical Sciences, University of Kragujevac, Serbia. It was approved and performed in accordance with the regulatives of the Faculty’s Ethical committee for the welfare of laboratory animals and principles of the Good laboratory practice and European Council Directive (86/609/EEC).

Subjects

Sixty four Wistar albino rats (males and females, eight weeks old at the beginning of the experiment, body weight 200 ± 50 g) were included in the study. The animals were housed at temperature of 22 ± 1°C, with 12 hours of automatic illumination daily. They consumed commercial rat food (20% protein rat food, Veterinary institute Subotica, Serbia) and water ad libitum.

Exercise training protocol

Rats were divided into 4 groups, while each group consisted of 2 subgroups, males (M) and females (F). The first group was control group (C), subgroups CM and CF \( (n = 8 \text{ for each subgroup}) \). The second group was trained group (T), subgroups TM and TF \( (n = 8 \text{ for each subgroup}) \). The third group included 2 weeks detrained animals (D2), i.e., animals who were subjected to training, followed by 2 weeks of detraining period, subgroups DM2 and DF2 \( (n = 8 \text{ for each subgroup}) \). The forth group consisted of 4 weeks detrained animals (D4), i.e., animals who were subjected to training followed by 4 weeks of detraining, subgroups DM4 and DF4 \( (n = 8 \text{ for each subgroup}) \). Rats from the control group were placed in the pool 5 times a week for 3 minutes to achieve water induced-stress (Lima et al. 2013; Stojanovic Tosic et al. 2015). Trained group included rats who were subjected to moderate intensity exercise, such as swimming training (8 weeks, 5 days/week, 60 min/day) in a specially designed pool according to the protocol in

![Figure 1. Experimental protocol]
Figure 1A. Week before the experiment, rats were gradually exposed to swimming training from 5 to 15 minutes, in order to familiarize them to the swimming exercise. Subsequently they started with 8 weeks training process. Rats were sacrificed a day after accomplishing training process. On the same day, rats (the same age as in T group) from C groups were sacrificed as well. Animals from DM2, DF2 and DM4, DF4 groups were sacrificed after 2 and 4 weeks of cessation, respectively (Figure 1).

**Isolated rat heart preparation**

After a short-term ketamine/xylazine-induced narcosis rats were sacrificed by decapitation. The chest was then opened via midline thoracotomy. The hearts were immediately removed and immersed in cold saline and were then mounted on a stainless steel cannula of the Langendorff perfusion apparatus to provide retrograde perfusion under gradually increasing coronary perfusion pressure (CPP from 40 to 120 cmH₂O). Krebs–Henseleit buffer was used for retrograde perfusion (in mmol/l: NaCl 118, KCl 4.7, CaCl₂×2H₂O 2.5, MgSO₄×7H₂O 1.7, NaHCO₃ 25, KH₂PO₄ 1.2, glucose 11, and pyruvate 2). The buffer was balanced with 95% O₂ and 5% CO₂, with a pH of 7.4 and a temperature of 37°C. Following the establishment of heart perfusion, the preparations were stabilized within 30 min with a basal coronary perfusion pressure of 70 cmH₂O. After the stabilization period, the perfusion pressure was reduced to 50 and 40 cmH₂O and then gradually increased to 60, 80, 100 and 120 cmH₂O to establish coronary autoregulation.

**Biochemical analysis**

In order to test systemic oxidative stress response to training, after animal sacrifice blood samples for biochemical analysis were collected from jugular vein. After centrifugation of heparinised venous blood, plasma and erythrocytes were separated. In plasma the following parameters of redox balance were determined: the levels of superoxide anion radical (O₂⁻), nitrites (NO₂⁻), hydrogen peroxide (H₂O₂) and index of lipid peroxidation (measured as thiobarbituric acid reactive substances–TBARS). Superoxide dismutase (SOD), catalase (CAT) and reduced glutathione (GSH) were determined in erythrocytes samples. During each CPP coronary venous effluent was collected for the purpose of determination of levels of pro-oxidants such as O₂, NO₂⁻, H₂O₂ and TBARS. Analyses of the pro-oxidants were performed using the same methods as when analyzing plasma samples and coronary venous effluent except NO₂⁻ where the protocol differs. Biochemical parameters were measured spectrophotometrically, using UV-1800 Shimadzu UV spectrophotometer, Japan.

**Superoxide anion radical determination (O₂⁻)**

The level of superoxide anion radical (O₂⁻) was measured using nitro blue tetrazolium (NBT) reaction in TRIS-buffer combined with plasma sample or coronary venous effluent. The measurement was performed at a wavelength of 530 nm. For O₂⁻ determination in coronary venous effluent the Krebs-Henseleit solvent was used as the blank control, while in case of plasma samples distilled water served as a blank control (Auclair and Voisin 1980).

**Hydrogen peroxide determination (H₂O₂)**

The protocol for measurement of hydrogen peroxide (H₂O₂) is based on oxidation of phenol red in the presence of horseradish peroxidase. 200 µl sample with 800 µl PRS (phenol red solution) and 10 µl POD (horseradish peroxidase) were combined (1:20). The level of H₂O₂ was measured at 610 nm. For H₂O₂ determination in coronary venous effluent the Krebs–Henseleit solvent was used as the blank control, while in case of plasma samples distilled water served as a blank control (Pick and Keisari 1980).

**Nitrite determination (NO₂⁻)**

Nitric oxide (NO) decomposes rapidly to form stable metabolite nitrite/nitrate products. Nitrite (NO₂⁻) was determined as an index of nitric oxide production with Griess reagent. For NO₂⁻ determination in plasma 0.1 ml 3 N PCA (perchloric acid), 0.4 ml 20 mM ethylenediaminetetraacetic acid (EDTA), and 0.2 ml plasma were put on ice for 15 min, then centrifuged 15 min at 6,000 rpm. After pouring off the supernatant, 220 µl K₂CO₃ was added. Nitrites were measured at 550 nm. Distilled water was used as a blank probe.

For NO₂⁻ determination in coronary venous effluent, 0.5 ml of the perfusate was precipitated with 200 µl of 30% sulfosalicylic acid, mixed for 30 min and centrifuged at 3000 × g. Equal volumes of the supernatant and Griess reagent were mixed and stabilized for 10 min in the dark, and then the sample was measured spectrophotometrically at a wavelength of 543 nm. The nitrite concentrations were determined using sodium nitrite as the standard (Green et al. 1982).

**Determination of the index of lipid peroxidation measured as TBARS**

The degree of lipid peroxidation in the sample (plasma and coronary venous effluent) was estimated by measuring of TBARS using 1% TBA (thiobarbituric acid) in 0.05 NaOH, incubated with sample at 100°C for 15 min and read at 530 nm. TBA extract was obtained by combining 0.8 ml
sample and 0.4 ml trichloro acetic acid (TCA), then samples were put on ice for 10 min, and centrifuged for 15 min at 6,000 rpm. The Krebs-Henselheit solvent was used as a blank control when TBARS was determined in coronary venous effluent, while in case of plasma sample distilled water was used (Ohkawa et al. 1979).

**Determination of antioxidant enzymes (CAT, SOD)**

Isolated RBCs were washed three times with three volumes of ice-cold 0.9 mmol/l NaCl and hemolysates containing about 50 g Hb/l (prepared according to McCord and Fridovich 1969) were used for the determination of CAT activity (Beutler 1982). Then 50 μl CAT buffer, 100 μl sample, and 1 ml 10 mM H2O2 were added to the samples. Detection was performed at 360 nm. SOD activity was determined by the epinephrine method (Misra and Fridovich 1972). A 100 μl lysate and 1 ml carbonate buffer were mixed, and then 100 μl of epinephrine was added. Detection was performed at 470 nm. Distilled water was used as a blank probe.

**Determination of reduced glutathione (GSH)**

Level of GSH was determined spectrophotometrically, and it is based on GSH oxidation via 5,5-dithiobis-6,2-nitrobenzoic acid. GSH extract was obtained by combining 0.1 ml 0.1% EDTA, 400 μl haemolysate, and 750 μl precipitation solution (containing 1.67 g metaphosphoric acid, 0.2 g EDTA, 30 g NaCl, and filled with distilled water until 100 ml; the solution is stable for 3 weeks at +4°C). After mixing in the vortex machine and extraction on cold ice (15 min), it was centrifuged on 4000 rpm (10 min). Distilled water was used as a blank probe. Measuring was performed at 420 nm. The concentration is expressed as nanomoles per milliliter of RBCs (Beutler 1975).

**Statistical analysis**

IBM SPSS Statistics 20.0 for Windows was used for statistical analysis. Values were expressed as mean ± standard deviation (SD). Descriptive statistics were used to calculate arithmetic mean with dispersion measures (standard deviation SD and standard error SE). Distribution of data was checked by Shapiro-Wilk test. Where distribution between groups was normal, statistical comparisons were performed using the one-way ANOVA tests with a Tukey’s post hoc test for multiple comparisons. Kruskal-Wallis was used for comparison between groups where the distribution of data was different than normal. Values of $p < 0.05$ were considered to be statistically significant, while values of $p < 0.01$ were considered to be statistically high significant.

**Results**

**Cardiac oxidative stress markers**

**Levels of superoxide anion radical (O$_2^-$)**

There was a statistically significant difference between groups in the level of O$_2^-$ as determined by one-way ANOVA ($F(7,121) = 7.362$, $p = 0.00$ for CPP 40 cmH$_2$O, $F(7,121) = 3.155$, $p = 0.01$ for CPP 60 cmH$_2$O, $F(7,121) = 9.229$, $p = 0.00$ for CPP 80 cmH$_2$O, $F(7,121) = 9.769$, $p = 0.00$ for CPP 100 cmH$_2$O, $F(7,121) = 11.322$, $p = 0.00$ for CPP 120 cmH$_2$O). A Tukey post hoc test revealed significant decrease in the level of O$_2^-$ in trained females compared to their controls at all CPP ($p = 0.009$ at CPP 40 cmH$_2$O, $p = 0.015$ at CPP 60 cmH$_2$O, $p = 0.018$ at CPP 80 cmH$_2$O, $p = 0.006$ at CPP 100 cmH$_2$O, $p = 0.002$ at CPP 120 cmH$_2$O). The same trend was revealed in trained males compared to their controls at all CPP values ($p = 0.004$ at CPP 40 cmH$_2$O, $p = 0.008$ at CPP 60 cmH$_2$O, $p = 0.005$ at CPP 80 cmH$_2$O, $p = 0.003$ at CPP 100 cmH$_2$O, $p = 0.002$ at CPP 120 cmH$_2$O (Figure 2A). This parameter was significantly higher in 4 weeks detrained females than in 2 weeks detrained females at CPP 80–120 cmH$_2$O ($p = 0.039$ at CPP 80 cmH$_2$O, $p = 0.032$ at CPP 100 cmH$_2$O, $p = 0.029$ at CPP 120 cmH$_2$O). On the other hand, level of O$_2^-$ in 4 weeks detrained males was lower than in 2 weeks detrained males at CPP 40–80 cmH$_2$O ($p = 0.029$ at CPP 40 cmH$_2$O, $p = 0.021$ at CPP 60 cmH$_2$O, $p = 0.034$ at CPP 80 cmH$_2$O). Significantly lower level of O$_2^-$ was revealed in 4 weeks detrained males compared to 4 weeks detrained females at all CPP values ($p = 0.031$ at CPP 40 cmH$_2$O, $p = 0.007$ at CPP 60 cmH$_2$O, $p = 0.002$ at CPP 80 cmH$_2$O, $p = 0.004$ at CPP 100 cmH$_2$O, $p = 0.006$ at CPP 120 cmH$_2$O) (Figure 2B). There was a decrease in the level of this parameter in trained females compared to 2 weeks detrained females at all CPP ($p = 0.023$ at CPP 40, $p = 0.005$ at CPP 60, $p = 0.007$ at CPP 80 cmH$_2$O, $p = 0.014$ at CPP 100 cmH$_2$O, $p = 0.034$ at CPP 120 cmH$_2$O) and 4 weeks detrained females at all CPP ($p = 0.020$ at CPP 40 cmH$_2$O, $p = 0.006$ at CPP 60 cmH$_2$O, $p = 0.002$ at CPP 80 cmH$_2$O, $p = 0.003$ at CPP 100 cmH$_2$O, $p = 0.015$ at CPP 120 cmH$_2$O) (Figure 2C). Training led to a decrease in the level of O$_2^-$ in males compared to 2 weeks detrained males at all CPP values ($p = 0.015$ at CPP 40 cmH$_2$O, $p = 0.011$ at CPP 60 cmH$_2$O, $p = 0.017$ at CPP 80 cmH$_2$O, $p = 0.019$ at CPP 100 cmH$_2$O, $p = 0.015$ at CPP 120 cmH$_2$O). In 4 weeks detrained males values were higher than in trained males at 100 and 120 cmH$_2$O ($p = 0.036$ at CPP 100 cmH$_2$O, $p = 0.033$ at CPP 120 cmH$_2$O) (Figure 2D).

**Levels of nitrites (NO$_2^-$)**

One-way ANOVA revealed there was a significant difference between groups in the level of O$_2^-$ ($F(7,121) = 2.112$, $p = 0.03$
for CPP = 40 cmH$_2$O, F(7,121) = 2,267, p = 0.02 for CPP = 60 cmH$_2$O, F(7,121) = 3,569, p = 0.00 for CPP = 80 cmH$_2$O, F(7,121) = 6,782, p = 0.00 for CPP = 120 cmH$_2$O). There was a decrease in the level of this parameter in trained females compared to control females at 100 and 120 cmH$_2$O (p = 0.023 at CPP 100 cmH$_2$O, p = 0.021 at CPP 120 cmH$_2$O) (Figure 3A).

In addition level of NO$_2^-$ observed in trained females in comparison to 2 (p = 0.032) and 4 weeks detrained females were lower at CPP = 120 cmH$_2$O (p = 0.039) (Figure 3C).

**Levels of hydrogen peroxide (H$_2$O$_2$)**

A significant difference between groups in the level of H$_2$O$_2$ was found (F(7,121) = 2,122, p = 0.002 for CPP = 40 cmH$_2$O, F(7,121) = 2,346, p = 0.00 for CPP = 60 cmH$_2$O, F(7,121) = 8,791, p = 0.00 for CPP = 80 cmH$_2$O, F(7,121) = 7,562, p = 0.00 for CPP = 100 cmH$_2$O, F(7,121) = 10,289, p = 0.00 for CPP = 120 cmH$_2$O). After the training, level of H$_2$O$_2$ was significantly decreased compared to the control group at male rats (p = 0.030 at CPP 40 cmH$_2$O, p = 0.032 at CPP 60 cmH$_2$O, p = 0.031 at CPP 80 cmH$_2$O, p = 0.026 at CPP 100 cmH$_2$O, p = 0.029 at CPP 120 cmH$_2$O). The same trend was noticed in female rats (p = 0.043 at CPP 40 cmH$_2$O, p = 0.035 at CPP 60 cmH$_2$O, p = 0.033 at CPP 80 cmH$_2$O, p = 0.029 at CPP 100 cmH$_2$O, p = 0.026 at CPP 120 cmH$_2$O) (Figure 4A). Gender difference in the level of H$_2$O$_2$ in 2 weeks detrained females compared to 2 weeks detrained males was observed, where levels were lower in female rats (p = 0.028 at CPP 100 cmH$_2$O, p = 0.022 at CPP 120 cmH$_2$O) (Figure 4A).

**Figure 2.** Effects of training/detraining on superoxide anion radical level determined in coronary venous effluent (O$_2^-$). A. Comparison between control and trained groups. B. Comparison between 2 and 4 weeks detrained groups. C. Comparison between trained, 2 and 4 weeks detrained females. D. Comparison between trained, 2 and 4 weeks detrained males. * p < 0.05, ** p < 0.01 TM vs. CM; * p < 0.05, ** p < 0.01 TF vs. CF; * p < 0.05, ** p < 0.01 DM2 vs. DM4; * p < 0.05, ** p < 0.01 TF vs. DF2, * p < 0.05, ** p < 0.01 TM vs. DM2; * p < 0.05, ** p < 0.01 TF vs. DF2; * p < 0.05, ** p < 0.01 TM vs. DM4; * p < 0.05, ** p < 0.01 TF vs. DF4; * p < 0.05, ** p < 0.01 DM2 vs. DF2, * p < 0.05, ** p < 0.01 DM4 vs. DF4. Data are means ± SD. CPP, coronary perfusion pressure; TM, trained males; DM2, 2 weeks detrained males; DM4, 4 weeks detrained males; CM, control males; TF, trained females; DF2, 2 weeks detrained females; DF4, 4 weeks detrained females; CF, control females.
Furthermore, level of this parameter was lower in 4 weeks detrained females compared to 4 weeks detrained males ($p = 0.013$ at CPP 100 cmH$_2$O, $p = 0.018$ at CPP 120 cmH$_2$O) (Figure 4B). In trained females, level of H$_2$O$_2$ was significantly lower than in 2 weeks detrained at CPP 60–120 cmH$_2$O ($p = 0.046$ at CPP 60 cmH$_2$O, $p = 0.038$ at CPP 80 cmH$_2$O, $p = 0.029$ at CPP 100 cmH$_2$O, $p = 0.035$ at CPP 120 cmH$_2$O). In 4 weeks detrained females, level of this parameter was higher in comparison to trained females ($p = 0.045$ at CPP 60 cmH$_2$O, $p = 0.039$ at CPP 80 cmH$_2$O, $p = 0.037$ at CPP 120 cmH$_2$O (Figure 4C). The same trend was revealed in males when compared trained group to 2 weeks detrained ($p = 0.035$ at CPP 40 cmH$_2$O, $p = 0.031$ at CPP 60 cmH$_2$O, $p = 0.022$ at CPP 80 cmH$_2$O, $p = 0.007$ at CPP 100 cmH$_2$O, $p = 0.006$ at CPP 120 cmH$_2$O). There was an increase after 4 weeks of detraining in males compared to training at all CPP values ($p = 0.029$ at CPP 40 cmH$_2$O, $p = 0.027$ at CPP 60 cmH$_2$O, $p = 0.018$ at CPP 80 cmH$_2$O, $p = 0.009$ at CPP 100 cmH$_2$O, $p = 0.005$ at CPP 120 cmH$_2$O) (Figure 4D).

Levels of TBARS

There was a significant difference between groups in the level of TBARS ($F(7,121) = 7.229$, $p = 0.00$ for CPP = 40 cmH$_2$O, $F(7,121) = 5.671$, $p = 0.00$ for CPP = 60 cmH$_2$O, $F(7,121) = 3.362$, $p = 0.000$ for CPP = 80 cmH$_2$O, $F(7,121) = 2.469$, $p = 0.004$ for CPP = 100 cmH$_2$O, $F(7,121) = 10.322$, $p = 0.00$ for CPP = 120 cmH$_2$O. In the trained group, a decrease in level of TBARS was noticed by comparing to the control group in male rats at all CPP ($p = 0.019$ at CPP 40 cmH$_2$O, $p = 0.009$ at CPP 60 cmH$_2$O, $p = 0.005$ at CPP 80 cmH$_2$O, $p = 0.008$ at CPP 100 cmH$_2$O, $p = 0.009$ at CPP 120 cmH$_2$O). Training led to the decrease in the level of this parameter in females at all CPP values compared to controls ($p = 0.013$ at CPP 40 cmH$_2$O, $p = 0.009$ at CPP 60 cmH$_2$O, $p = 0.007$ at CPP 80 cmH$_2$O, $p = 0.006$ at CPP 100 cmH$_2$O, $p = 0.002$ at CPP 120 cmH$_2$O) (Figure 5A). There was a significant increase in 2 weeks detrained females compared to 2 weeks detrained males at all CPP values ($p = 0.022$ at CPP 40 cmH$_2$O, $p = 0.018$ at CPP 60 cmH$_2$O, $p = 0.025$ at CPP
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80 cmH₂O, p = 0.031 at CPP 100 cmH₂O, p = 0.033 at CPP 120 cmH₂O (Figure 5B). In group of male rats, there was a decrease in values of this parameter in 2 weeks detrained compared to trained rats, but only in the lowest CPP (p = 0.029 at CPP 40 cmH₂O, p = 0.032 at CPP 60 cmH₂O). In 4 weeks detrained males levels were lower than in trained males at the highest CPP (p = 0.022 at CPP 80 cmH₂O, p = 0.027 at CPP 100 cmH₂O, p = 0.029 at CPP 120 cmH₂O) (Figure 5D).

**Systemic redox state**

**Levels of superoxide anion radical (O₂⁻)**

There was a significant difference between groups in the level of O₂⁻ (F(7,56) = 8.229, p = 0.00). Significantly lower levels were noticed in trained females compared to 2 weeks detrained females (p = 0.007) and trained males in comparison to 2 weeks (p = 0.008) and 4 weeks detrained males (p = 0.007). There was a decrease in 4 weeks detrained females compared to 2 weeks detrained females (p = 0.009). Level of O₂⁻ in 4 weeks detrained females was significantly lower than in 4 weeks detrained males (p = 0.006) (Figure 6A).

**Levels of nitrites (NO₂⁻)**

A significant difference between groups in the level of NO₂⁻ was noticed (F(7,56) = 9.125, p = 0.00). There was a decrease in the level of NO₂⁻ parameter in 2 weeks detrained females in comparison to trained females (p = 0.024) and 2 weeks detrained males in comparison to trained males (p = 0.029). Level of this parameter in 4 weeks detrained males was decreased compared to trained males (p = 0.028). In addition, level of NO₂⁻ in 4 weeks detrained females was increased compared to 2 weeks detrained females (p =
In a 4 weeks detrained group, levels of NO\textsubscript{2} was significantly higher in female rats than in male rats (0.026) (Figure 6B).

**Levels of hydrogen peroxide (H\textsubscript{2}O\textsubscript{2})**

According to the results of one-way ANOVA, there was a statistically significant difference between groups in the level of H\textsubscript{2}O\textsubscript{2} (F(7, 56) = 8.125, p = 0.00). A decrease in the level of H\textsubscript{2}O\textsubscript{2} in 2 weeks detrained males compared to trained males (p = 0.009) and 2 weeks detrained females group compared to trained females group was revealed (0.008). In addition, there was a significant decrease in the level of this parameter in 4 weeks detrained males compared to trained males (p = 0.007). In 4 weeks detrained females levels were increased compared to 2 weeks detrained females (p = 0.006). In a group of rats who were subjected to 4 weeks of detraining, levels of H\textsubscript{2}O\textsubscript{2} were higher in females (p = 0.004) (Figure 6C).

**Levels of TBARS**

There was a significant difference between groups in the level of TBARS (F(7, 56) = 5.125, p = 0.01). Level of TBARS in plasma increased in trained groups compared to control at females (p = 0.07) and males (p = 0.004). An increase in the level of this parameter in 4 weeks detrained females compared to trained females (p = 0.029) and 2 weeks detrained females (p = 0.034) was noticed as well (Figure 6D).

**Activity of superoxide dismutase (SOD)**

Activity of SOD differed between groups (F(7, 56) = 7.221, p = 0.00). A significant increase in the activity of this enzyme in trained group in males was observed in comparison to control males (p = 0.023). There was a decrease in the activity of SOD in 2 weeks detrained females (p = 0.021) and 4 weeks detrained females (p = 0.009) compared to trained females.
In 4 weeks detrained males activity of SOD was decreased in comparison to 2 weeks detrained males ($p = 0.006$) and trained males ($p = 0.008$). There was a statistically significant increase in SOD activity in male rats compared to female rats in the period of 2 weeks detraining ($0.003$) (Figure 7A).

**Activity of catalase (CAT)**

Statistically significant difference between groups in the activity of catalase ($F(7,56) = 10.221, p = 0.00$) was found. There was an increase in the activity of catalase (CAT) in trained group compared to the control group in females ($p = 0.008$), while in males the difference was not significant. In 2 weeks detrained females the activity was decreased compared to trained females ($p = 0.009$). A decrease in the activity of CAT was observed in 4 weeks detrained females compared to trained females ($p = 0.032$), and that value was increased compared to 2 weeks detrained females ($p = 0.029$). An increased activity was noticed in 4 weeks detrained males compared to trained males ($p = 0.009$) and 2 weeks detrained males ($p = 0.016$). The only difference in values between males and females was found in trained groups, where females had higher values of CAT activity ($p = 0.006$) (Figure 7B).

**Levels of reduced glutathione (GSH)**

There was a statistically significant difference between groups in the level of GSH ($F(7,56) = 4.298, p = 0.02$). Level of reduced glutathione (GSH) was significantly increased in trained groups compared to control groups in females ($p = 0.002$) and males ($p = 0.003$). In 2 weeks detrained males values were increased compared to trained males ($p = 0.027$). In 4 weeks detrained males group level of GSH was increased compared to trained males ($p = 0.022$). In addition, level of GSH was lower in 4 weeks detrained females than in 2 weeks detrained females ($p = 0.021$). Gender difference in 4 detrained females in comparison to the 4 weeks detrained males group was noticed, where values in female rats were lower than in male rats ($0.018$) (Figure 7C).

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**Figure 6.** Effects of training/detraining on level of pro-oxidants determined in plasma: $O_2^*$ (A), $NO_2^-$ (B), $H_2O_2$ (C), TBARS (D). * $p < 0.05$, ** $p < 0.01$ C vs. T vs. D2 vs. D4; * $p < 0.05$, ** $p < 0.01$ males vs. females. Data are means ± SD. C, control group; T, trained group; D2, 2 weeks detrained group; D4, 4 weeks detrained group.
Discussion

Potential source of oxidative stress during anaerobic training is an increased aerobic metabolism (Stankovic and Radovanovic 2012). ROS and RNS production during exercise follows the principal of hormesis and may represent an adaptive response of cells to stressors, such as physical activity. The responses of biological systems may be described with a bell-shaped curve whose two endpoints are inactivity and overtraining (Stojanovic Tosić et al. 2015; Radak et al. 2017).

Numerous investigations aimed to examine the influence of aerobic exercise on oxidative stress markers were mostly focused on treadmill or cycle ergometer (Powers et al. 2016). We’ve chosen to examine the effects of swimming training due to the fact that swimming is considered as a natural ability of rats and it has also been proposed as a convenient model for studying the physiological changes and stress response to training (Balci and Pepe 2012; Araujo et al. 2015).

Results of our study clearly show that 8 weeks of swimming training led to the decrease of almost all pro-oxidants measured in the heart both in male and female rats. Analysis of aforementioned parameters in the coronary venous effluent during coronary autoregulation refers to the oxidative stress in the endocardium of the left ventricle and endothelium of the coronary circulation as well. In order to complete the picture about the role of oxidative stress in physiology of effort, we investigated the systemic oxidative stress response to training and detraining. Generally viewed, there were no changes in the release of the measured pro-oxidants in plasma, thus suggesting that applied intensity and duration of swimming training may affect only local production of ROS (in the heart), while systemic response was not changed. In addition, having in mind that release of pro-oxidants was even decreased in the heart, it seems that swimming training of this type doesn’t promote oxidative damage, nor act protectively within the heart.

Regarding the components of antioxidant defense system, SOD, CAT and GSH, training led to the significant increase of GSH values at both sexes, CAT in females and SOD in males. It can be assumed that effects of training on these antioxidant parameters depend on their chemical characteristics. Also, results for these enzymes can not be interpreted independently of $O_2^–/H_2O_2$ dynamic. In female, unchanged activity of SOD in training induced less scavenging of $O_2^–$, leading to higher $H_2O_2$ values (compared to 2 weeks detrained group), which can induce enhanced CAT.
activity. Increased activity of antioxidants is in correlation with unaltered levels of pro-oxidants in the plasma and may explain these results.

Our results are in agreement with the study conducted by Balci et al. who found decreased malondialdehyde (MDA) levels in the heart of female rats at rest, however they didn't observe any change in MDA and NO levels in male rats in the heart (Balci and Pepe 2012). Neither did we reveal the difference in level of NO\textsuperscript{2−} in the heart when compared trained and untrained rats both sexes at rest. The potential explanation for this may be the interaction of NO with reactive oxygen species (ROS), particularly O\textsuperscript{2−}. Significantly lower levels of O\textsuperscript{2−} in the present study may be a consequence of this interaction resulting in generation of peroxynitrite (ONOO\textsuperscript{−}).

A group of researchers whose methodology differed from ours in case of duration of swimming training noticed that this type of activity reduced lipid peroxidation in the heart (Venditti and Di Meo 1997). It should be taken into consideration that pro-oxidants detected in blood plasma and erythrocytes reflect the redox state of all components which are included in the motor act during physical exercise. In that sense it's logical to expect the different values of parameters of oxidative stress response measured in heart and in plasma (Elikov 2016). Furthermore, Hu et al. (2000) observed no change in the lipid peroxidation level in heart, which may be explained by a shorter period of swimming training (7 days, 45 minutes per day) insufficient to establish positive adaptations to exercise.

When discussing antioxidant defense system, our results are not in accordance with the results of Balci and Pepe (2012) who revealed that 8 weeks of swimming training caused a decrease in SOD activity and didn’t affect total GSH levels in rat heart. Furthermore, Lima and co-workers proved increased reduced glutathione (GSH) content and reduced/oxidized (GSH/GSSG) ratio, higher superoxide dismutase activity in liver mitochondria after 6-week swimming training protocol (Lima et al. 2013). Others showed an increase in serum superoxide dismutase activity induced by swimming as well (Botezelli et al. 2011). We expected that gender differences in response to training exist, since it's been reported that female rats often show lower oxidative damage than males (Stankovic and Radovanovic 2012). However, values of all measured pro-oxidants were similar in female and male rats in training.

It's been known that the exercise-induced adaptive process is reversible in case of cardiovascular function and mitochondrial enzyme activity (Mujika and Padilla 2000). After 2 weeks of detraining, the release of O\textsubscript{2} and H\textsubscript{2}O\textsubscript{2} in the heart was increased compared to the values in training. Regarding the systemic response, we noticed the same trend in O\textsubscript{2} production, while levels of NO\textsuperscript{2−} and H\textsubscript{2}O\textsubscript{2} were decreased in comparison to the levels in training. The explanation for decreased plasma level in H\textsubscript{2}O\textsubscript{2} detraining period may be an increased catalase activity during training, which catalyzed the decomposition of hydrogen peroxide to water and oxygen. Furthermore, SOD and CAT activity were lower after 2 weeks of training cessation in females and level of GSH was higher in males in comparison with the values in training. The difference between gender that we noticed was an increased activity of SOD in males and higher TBARS production in heart of females.

We detected an increase in cardiac release of O\textsubscript{2} at both sexes and H\textsubscript{2}O\textsubscript{2} at males after four weeks of training cessation compared to the values in training. Interestingly, after 4 weeks of detraining levels of the most of measured pro-oxidants in females were similar to those in training. Regarding the parameters of antioxidant defense system, 4 weeks detraining led to the decrease in SOD activity at both sexes and CAT in females compared with training. On the contrary, GSH and CAT values in males remained increased in detraining. There is a data that increased total antioxidant capacity was associated with an increased circulating CD34+/VEGFR2\textsuperscript{+} cells in detraining (Witkowski et al. 2010).

Radak et al. examined the effects of 8 weeks of swimming and 8 weeks of detraining on the level of free radical species in the cerebellum. They proved that positive effects of training were maintained during detraining (Radak et al. 2006). Others showed that 16 week of walking/jogging at 50–80% of HR(max) decreased MDA levels and increased total antioxidant capacity (TAC) and glutathione peroxidase activity (GPX) (Fatouros et al. 2004). However after 4 months of training cessation those effects were eliminated. It has been previously reported that effects of treadmill training on paraventricular nucleus in hypertensive rats reversed after 2 weeks of training cessation (Agarwal et al. 2012).

Although estrogen 17β-estradiol and different levels of ferritin may be responsible for the higher antioxidant protection noticed in females compared to males, we revealed lower antioxidant protection in females during detraining (Català-Niell et al. 2008; Stankovic and Radovanovic 2012). Based on our results we may hypothesize that probably there are other mechanisms independent of the change in estrogen and iron metabolism, thus contributing to sex differences in oxidative-stress response to exercise cessation. Beside mentioned mechanisms involved in effects of training/detraining on diversity between male and female antioxidative status, a very recent study on rats have shown gender difference in mitochondrial function which can be affected by exercise or cessation of it (Farhat et al. 2017). It can be assumed that diminished mitochondrial activity can lead to depressed production of mitochondrial SOD and thus impair functioning of cellular antioxidant pathways (Macak-Safranko et al. 2011). Lower antioxidant values in our study support higher pro-oxidant levels after 4 weeks of detraining noticed in females.
We confirmed the fact that sexual dimorphism in oxidative capacity exists. These results suggest different dynamic of production of specific pro-oxidants among sexes during period of training cessation. One of limitations of our investigation was the absence of technique through which cellular mechanisms of obtained effects could be proved. Therefore further studies are necessary for better understanding the possible mechanisms underlying the gender differences in response to training and detraining.

Our results illustrated that moderate-intensity physical exercise of sufficient duration leads to the beneficial adaptations, manifested as improvement of antioxidant defense system. In addition, these results suggest that 2 and 4 weeks of training cessation may lead to a partial lost in exercise-induced adaptation. Positive antioxidative effects of training remained longer in males. Findings of the present study may help elucidation of training and detraining effects on modulation of redox homeostasis, especially from aspect of gender differences.

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