EXERCISE-INDUCED ALTERATIONS OF THE OXIDATIVE STRESS BIOMARKERS IN ERYTHROCYTES OF PONIES INVOLVED IN RECREATIONAL HORSEBACK RIDING

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The aim of the current study was the analysis of the oxidative stress biomarkers, as well as the osmotic-induced resistance of erythrocytes in mares and stallions of ponies involved in recreational horseback riding in Pomeranian regions. Ten healthy adult Hucul ponies (5 stallions and 5 mares), 5-11 years old, from Pomeranian regions in Poland (Ustka city, Pomeranian Voivodship, Poland) were used in our study. All horses participated in recreational horseback riding and were subjected to the resembling type of management. The training was continued 1 hour and included a ride of cross country by walking (10 min), trotting (15 min), walking (10 min), galloping (15 min), and walking (10 min). Blood was drawn from jugular veins of the animals in the morning, 90 minutes after feeding, and immediately after the exercise session. Marker of lipid peroxidation (2-Thiobarbituric acid reactive substances, TBARS), aldehydic and ketonic derivatives in the erythrocytes, total antioxidant capacity (TAC) level in the erythrocytes of both mares and stallions exhibited non-significant changes immediately after exercise as compared to the resting period. Both at the rest and after the training session, the levels of TBARS, aldehydic and ketonic derivatives in the erythrocyte suspensions of mares were non-significantly higher compared to stallions. In a like manner, differences of aldehydic and ketonic derivatives of oxidatively modified proteins between mares and stallions after training sessions were noted. Both at the rest and after the training session, the TAC level in the erythrocyte suspensions of stallions was non-significantly higher compared to mares. Among both mares and stallions, a non-significantly decreased erythrocyte hemolysis was observed after training sessions compared to the rest period. A comparison of erythrocyte hemolysis in mares and stallions at the rest period showed increased values of hemolysis in the stallions. After the training session, decreased hemolysis was observed in the stallions compared to mares. Efforts should be directed toward a thorough characterization of antioxidant defenses, as well as the correlation links between oxidative stress biomarkers and antioxidant defenses including age- and gender-related differences in the training programs of horses involved in the recreational horseback ridings.

Keywords: 2-Thiobarbituric acid reactive substances, aldehydic and ketonic derivatives, oxidatively modified proteins, total antioxidant capacity, erythrocyte hemolysis, training program

Exercise has been shown to increase the production of reactive oxygen species (ROS) to the point that it can exceed antioxidant defenses to cause oxidative stress [20]. Oxidants are essentially generated by metabolic enzymes, inflammatory cells, and mitochondrial electron leakage; they are indispensable for the cellular redox regulation and may, under certain conditions, have a pro-inflammatory stimulatory role [11]. High levels of oxidants and their by-products can compromise the athletic ability of a horse and
may lead to chronic diseases, i.e. fatigue, muscle damage, and reduced immune function, which can all affect exercise performance [7, 9-11]. Thus, there is a paradox between acute exercise, increasing oxidative stress, and regular exercise having beneficial effects on health and exercise performance [16-17]. However, exercise-induced oxidative stress and increased ROS generation may induce membrane lipid peroxidation and protein damage of erythrocytes and thereby decrease their membrane integrity [6].

Erythrocytes appear much more vulnerable to oxidative damage during intense exercise because of their continuous exposure to high oxygen fluxes and their high concentrations of polyunsaturated fatty acids and heme iron [14]. The major generators of reactive species located in blood during exercise could be erythrocytes (mainly due to their quantity) and leukocytes (mainly due to their drastic activation during exercise) [13]. In our previous studies, different responses of hematological parameters, oxidative stress biomarkers, as well as antioxidant defenses in the blood of well-trained horses and horses involved in the recreational horseback ridings were demonstrated [1-4, 18]. Recent studies of equine exercise physiology have focused mainly on the evaluation of particular parameters (hematological and biochemical) and changes in their reference ranges depend on the type of exercise [15].

The physical effort and training-induced stress in horses differ depending on equine sporting discipline, as well as on horse breeds. The proper training program aims to adapt a horse’s organism to the physiological changes occurring during intense physical effort [8, 15]. Necessary adaptive mechanisms are formed during that training cycle, giving the possibility of reaching a high level of efficiency with a concurrent decrease in the risk of injuries [8]. The adaptive changes include an increase in heart stroke volume and as a consequence an increase in oxygen utilization in muscles [15]. The cardiovascular system is exposed to modifications essential to ensure efficient transfer of oxygen and substrates to the muscles and allowing removal of metabolites from them. Information concerning physical effort is thus essential in making suitable decisions about proper training strategy.

Thus, the aim of the current study was the analysis of the oxidative stress biomarkers, as well as the urea-induced resistance of erythrocytes in mares and stallions of ponies involved in recreational horseback riding in Pomeranian regions.

**Materials and methods.** **Horses.** Ten healthy adult Hucul ponies (5 stallions and 5 mares), 5-11 years old, from Pomeranian regions in Poland (Ustka city, 54°34′43″N 16°52′09″E, Słupsk region, Pomeranian Voivodship, northern Poland, Fig. 1) were used in our study. All horses participated in recreational horseback riding and were subjected to the resembling type of management. Horses were housed in individual boxes under natural spring photoperiod (sunrise at 06:00 h, sunset at 19:00 h), with feeding (hay and oat) provided twice a day, at 08.00 and 18.00 h, and water available ad libitum. All horses were thoroughly examined clinically and screened for hematological, biochemical, and vital parameters, which were within reference ranges. The females were non-pregnant.

**Training.** Training started at 10:00 AM, lasted 1 hour, and included a ride of cross country by walking (10 min), trotting (15 min), walking (10 min), galloping (15 min), and walking (10 min). Environmental conditions during the ride had ranged from 18 to 23 °C.

**Blood samples.** Blood was drawn from jugular veins of the animals in the morning, 90 minutes after feeding, while the horses were in the stables (between 8:30 and 10 AM), and immediately after the exercise session (between 11:00 AM and 2:00 PM). Blood was stored into tubes with K3-EDTA and held on ice until centrifugation at 3,000g for 15 minutes. The plasma was removed. The erythrocyte suspension (one vol-
ume) was washed with five volumes of saline solution three times and centrifuged at 3,000g for 15 minutes. Plasma aliquots were frozen and stored at –25°C until analyzed. Erythrocyte suspensions were used for the determination of the level of 2-thiobarbituric acid reacting substances (TBARS), carbonyl derivatives of protein oxidative modification, total antioxidant capacity (TAC), as well as for acid-induced erythrocyte hemolysis assay.

**2-Thiobarbituric acid reactive substances (TBARS) assay.** The lipid peroxidation was determined by quantifying the concentration of 2-thiobarbituric acid reacting substances (TBARS) for determining the malondialdehyde (MDA) concentration by the Kamyshnikov (2004). This method is based on the reaction of the degradation of lipid peroxidation product, MDA, with 2-thiobarbituric acid (TBA) under high temperature and acidity to generate a colored adduct that is measured spectrophotometrically. The µmol of MDA per l was calculated by using 1.56·10^5 mM^1·cm^1 as the extinction coefficient.

**The carbonyl derivative contents of oxidative modification of proteins (OMP) assay.** The rate of protein oxidative destruction was estimated from the reaction of the resultant carbonyl derivatives of amino acid reaction with 2,4-dinitrophenyl hydrazine (DNPH) as described by Levine and co-workers (1990) and as modified by Dubinina and co-workers (1995). DNPH was used for determining carbonyl content in soluble and insoluble proteins. The carbonyl contents were calculated from the absorbance measurement at 370 nm and 430 nm, and an absorption coefficient 22,000 M^1·cm^1. Carbonyl groups were determined spectrophotometrically from the difference in absorbance at 370 nm (aldehydic derivatives, OMP_370) and 430 nm (ketonic derivatives, OMP_430) and defined as 1 nmol per mL of blood.

**Total antioxidant capacity (TAC) assay.** The TAC level in the erythrocytes was estimated by measuring the TBARS level following Tween-80 oxidation by Galaktionova et al. (1998). Plasma inhibits the Fe^{2+}/ascorbate-induced oxidation of Tween 80, resulting in a decrease in the TBARS level. The absorbance of the obtained solution was measured at 532 nm. The absorbance of the blank was defined as 100 %. The level of TAC in the sample (%) was calculated concerning the absorbance of the blank.
Assay of Osmotic-induced Hemolysis of Erythrocytes. The osmotic resistance of erythrocytes in solutions with different NaCl concentrations was measured spectrophotometrically at the wavelength of 540 nm as described by Mariańska et al. (2003). The method is based on the determination of differences between osmotic resistance of erythrocytes to a mixture containing different concentration of sodium chloride (0.1 %, 0.2 %, 0.3 %, 0.4 %, 0.5 %, 0.6 %, 0.7 %, 0.8 %, 0.9 %). The absorbance of the mixture contained erythrocytes and distilled water was determined as 100% hemolysis (blank). The degree of hemolysis in every test tube (%) was calculated by the absorbance of the blank. Hemolysis of erythrocytes (%) in every test tube with different sodium chloride concentrations was expressed as a curve [12].

Statistical analysis. Results are expressed as mean ± S.E.M. All variables were tested for normal distribution using the Kolmogorov-Smirnov test (p>0.05). To find significant differences (significance level, p≤0.05) between at the rest and after training, the Wilcoxon signed-rank test was applied to the data. All statistical analyses were performed using STATISTICA 8.0 software (StatSoft, Krakow, Poland).

Results and discussion. Lipid peroxidation is a complex phenomenon involving the generation of many products [5]. Aldehydes, especially MDA, frequently used as markers of oxidative stress in response to exercise [19], can be measured by spectrophotometric or spectrofluorescentic assay. The most common method used to assess the MDA level during exercise training is the 2-thiobarbituric acid reactive substances (TBARS) assay. The marker of lipid peroxidation during the training session was measured through analysis of the TBARS level and shown in Fig. 2.

Fig. 2. The level of lipid peroxidation determined by quantifying of 2-thiobarbituric acid reactive substrates (TBARS) level (µmol MDA·L⁻¹) in the erythrocytes of ponies involved in recreational horseback ridings at the rest and after a training session. Values expressed as mean ± S.E.M.

TBARS level in the erythrocytes of mares exhibited a non-significant decrease (by 1.8 %, p>0.05) immediately after exercise as compared to the resting period. Also, the TBARS level in the erythrocytes of stallions was non-significantly decreased by 6.38 % (p>0.05) after the training sessions. There were no significant differences in the erythrocytes TBARS level between the resting period and after exercise both mares and stallions. Both at the rest and after the training session, the TBARS level in the
erythrocyte suspensions of mares was non-significantly higher by 4.5 % (p>0.05) and by 8.9 % (p>0.05) compared to stallions (Fig. 2).

The effect of a training session on carbonyl contents in the erythrocyte suspensions of mares and stallions of ponies are shown in Fig. 3. There were no statistically significant changes in the carbonyl derivatives of protein destruction in the erythrocytes of horses during training sessions. Decreased level of aldehydic (by 6.3 % and 9.3 %, p>0.05, respectively) and ketonic derivatives (by 25.5 % and 8.1 %, p>0.05, respectively) in the erythrocytes of mares and stallions after training sessions compared to the resting period was observed (Fig. 3).

Similarly to the TBARS level, at the rest period, significantly higher levels of aldehydic and ketonic derivatives were observed in the erythrocytes of mares (by 39.6 % and 3.4 %, p>0.05) compared to the stallions. In a like manner, differences of aldehydic and ketonic derivatives of oxidatively modified proteins between mares and stallions after training sessions were noted, i.e. levels of aldehydic derivatives were higher in the mares by 41.6 % (p>0.05) compared to the stallions, while ketonic derivatives were higher in the stallions by 16 % (p>0.05) compared to the mares (Fig. 3).

It is well known that total antioxidant capacity (TAC) includes enzymatic antioxidant such as superoxide dismutase, catalase, glutathione peroxidase, as well as some macromolecules (albumin, ceruloplasmin, urea, glutathione, ferritin, etc.), and its assessment may contain more information than a single review of its constituent parts [24]. The total antioxidant capacity (TAC) in erythrocyte suspensions is shown in Fig. 4.

TAC level in the erythrocytes of mares exhibited a non-significant increase (by 4.8 %, p>0.05) immediately after exercise as compared to the resting period. On the other hand, the TAC level in the erythrocytes of stallions was non-significantly decreased by 2.9 % (p>0.05) after the training sessions. There were no significant differ-
ences in the erythrocytes TAC level between the resting period and after exercise both mares and stallions. Both at the rest and after the training session, the TAC level in the erythrocyte suspensions of stallions was non-significantly higher by 12.8 % (p>0.05) and by 4.5 % (p>0.05) compared to mares (Fig. 4).

![Total antioxidant capacity](image)

**Fig. 4.** The total antioxidant capacity (TAC) in the erythrocytes of mares and stallions of ponies involved in recreational horseback ridings at the rest and after a training session. Values expressed as mean ± S.E.M.

Erythrocytes are the best indicators of increased generation of ROS and activation of oxidative stress. Therefore, the next goal of our study was the measurement of erythrocyte resistance to the hemolytic reagent (solutions with different concentrations of sodium chloride and urea) of the mares and stallions at the resting period and after exercise (Fig. 5). There were no significant differences in the percentage of hemolyzed erythrocytes between resting period and after training (incubation with 0.1 %-0.9 % NaCl and 0.12-0.15 mol/l of urea) in the mares and stallions of ponies (Fig. 5). Among mares, decreased of erythrocyte hemolysis was observed after training session compared to the rest period (by 5.18 %, 3.26 %, 8.67 %, 5.58 %, 5.89 %, 29.9 %, and 16.5 %, p>0.05 after incubation with 0.3%-0.9% NaCl, respectively). Among stallions, similar to the data in mares, decreased of erythrocyte hemolysis was observed after training session compared to the rest period (by 3.9 %, 2.1 %, 1.49 %, 10.7 %, 15.74 %, 5.1 %, 3.1 %, 42.7 %, and 28.8 %, p>0.05 after incubation with 0.1 %–0.9 % NaCl, respectively) (Fig. 5).

Comparison of erythrocyte hemolysis in mares and stallions at the rest period showed increased values of hemolysis in the stallions (by 5.2 %, 3.6 %, 13.3 %, and 19.5 % at incubation with 0.4 %, 0.5 %, 0.8 %, and 0.9 %, respectively). After the training session, decreased hemolysis was observed in the stallions compared to mares (by 5.81 %, 5 %, 2.87 %, 4.39 %, 3.65 %, 6.1 %, and 7.3 % at incubation with 0.1 %, 0.2 %, 0.4 %, 0.5 %, 0.6 %, 0.7 %, and 0.8 %, respectively) (Fig. 5).
Fig. 5. Osmotic resistance of erythrocytes, measured by hemolysis of erythrocytes incubating in the solution with different concentrations of saline, in mares and stallions of ponies involved in recreational horseback ridings at the rest and after a training session. Values expressed as mean ± S.E.M.

Conclusions:

1. Marker of lipid peroxidation (TBARS level) in the erythrocytes of both mares and stallions exhibited a non-significant decrease immediately after exercise as compared to the resting period. There were no significant differences in the erythrocytes TBARS level between the resting period and after exercise both mares and stallions. Both at the rest and after the training session, the TBARS level in the erythrocyte suspensions of mares was non-significantly higher compared to stallions.

2. Non-statistically decrease in levels of aldehydic and ketonic derivatives in the erythrocytes of mares and stallions after training sessions compared to the resting period was observed. Similarly to the TBARS level, at the rest period, significantly higher levels of aldehydic and ketonic derivatives were observed in the erythrocytes of mares compared to the stallions. In a like manner, differences of aldehydic and ketonic derivatives of oxidatively modified proteins between mares and stallions after training sessions were noted, i.e. levels of aldehydic derivatives were higher in the mares by 41.6 % (p>0.05) compared to the stallions, while ketonic derivatives were higher in the stallions by 16% (p>0.05) compared to the mares.

3. Total antioxidant capacity (TAC) level in the erythrocytes of mares exhibited a non-significant increase immediately after exercise as compared to the resting period. On the other hand, the TAC level in the erythrocytes of stallions was non-significantly decreased after the training sessions. There were no significant differences in the erythrocytes TAC level between the resting period and after exercise both mares and stallions. Both at the rest and after the training session, the TAC level in the erythrocyte suspensions of stallions was non-significantly higher compared to mares.

4. Among both mares and stallions, a non-significantly decreased erythrocyte hemolysis was observed after training sessions compared to the rest period. A comparison of erythrocyte hemolysis in mares and stallions at the rest period showed increased values of hemolysis in the stallions. After the training session, decreased hemolysis was observed in the stallions compared to mares.
Efforts should be directed toward a thorough characterization of antioxidant defenses, as well as the correlation links between oxidative stress biomarkers and antioxidant defenses including age- and gender-related differences in the training programs of horses involved in the recreational horseback ridings.

References
1. Andrichuk, A., & Tkachenko, H. (2017). Effect of gender and exercise on haematological and biochemical parameters in Holsteiner horses. *J. Anim. Physiol. Anim. Nutr. (Berl).* 101(5), e404–e413.
2. Andrichuk, A., & Tkachenko, H. (2015). Seasonal variations of hematological indices in equines involved in recreational horse riding. *Journal of Ecology and Protection of the Coastline (Baltic Coastal Zone)*, 19, 11–22.
3. Andrichuk, A., & Tkachenko, H., Kurhaluk, N. (2014). Gender differences of oxidative stress biomarkers and erythrocyte damage in well-trained horses during exercises. *Journal of Equine Veterinary Science*, 34, 978–985.
4. Andrichuk, A., Tkachenko, H., Kurhaluk, N., & Tkachova, I. (2013). Oxidative stress biomarkers and erythrocyte hemolysis in trained Ukrainian warmblood horses under the influence of exercises. *The Animal Biology*, 15(4), 9–24.
5. Chiaradia, E., Avellini, L., Rueca, F., Spaterna, A., Porciello, F., Antonioni, M.T., & Gaiti, A. (1998). Physical exercise, oxidative stress and muscle damage in racehorses. *Comp. Biochem Physiol. B Biochem. Mol. Biol.*, 119(4), 833–836.
6. ÇimenBurak, M. Y. (2008). Free radical metabolism in human erythrocytes. *Clinica Chemica Acta*, 390, 1–11.
7. Deaton, C. M., & Marlin, D. J. (2003). Exercise-associated oxidative stress. *Clinical Techniques in Equine Practice*, 2, 278–291.
8. Fazio, F., Assenza, A., Tosto, F., Casella, S., Piccione, G., & Caola, G. (2011). Training and haematochemical profile in Thoroughbreds and Standardbreds: A longitudinal study. *Livestock Science*, 141, 221–226.
9. Hinchcliff, K. W., & Geor, R. J. (2008). The horse as an athlete: a physiological overview. In: *Equine Exercise Physiology. The Science of Exercise in the Athletic Horse*. Eds. K. W. Hinchcliff, R. J. Geor, A. J. Kaneps. Elsevier, 2–11.
10. Horohov, D. W., Sinatra, S. T., Chopra, R. K., Jankowitz, S., Betancourt, A., & Bloomer, R. J. (2012). The effect of exercise and nutritional supplementation on pro-inflammatory cytokine expression in young racehorses during training. *Journal of Equine Veterinary Science*, 32, 805–815.
11. Kirschvink, N., de Moffarts, B., & Lekeux, P. (2008). The oxidant/antioxidant equilibrium in horses. *Vet. J.*, 177(2), 178–191.
12. Mariańska, B., Fabijańska-Mitek, J., & Windyga, J. (2003). *Laboratory studies in hematology. Textbook for Medical Students*, PZWL, Warsaw [in Polish].
13. Nikolaidis, M. G., & Jamurtas, A. Z. (2009). Blood as a reactive species generator and redox status regulator during exercise. *Archives of Biochemistry and Biophysics*, 490, 77–84.
14. Petibois, C., & Déléris, G. (2005). Erythrocyte adaptation to oxidative stress in endurance training. *Archives of Medical Research*, 36, 524–531.
15. Piccione, G., Casella, S., Gianetto, C., Messina, V., Monteverde, V., Caola, G., & Gutta-dauro, S. (2010). Haematological and haematochemical responses to training and competition in Standardbred horses. *Comp. Clin. Pathol.*, 19, 95–101.
16. Radák, Z., Sasvári, M., Nyakas, C., Taylor, A.W., Ohno, H., Nakamotou, H., & Goto, S. (2000). Regular training modulates the accumulation of reactive carbonyl.
derivatives in mitochondrial and cytosolic fractions of rat skeletal muscle. Arch. Bioch. Biophys., 383, 114–118.

17. Radák, Z., Zhao, Z., Koltai, E., Ohno, H., & Atalay, M. (2013). Oxygen consumption and usage during physical exercise: the balance between oxidative stress and ROS-depend adaptive signaling. Antioxid. Redox Signal., 18(10), 1208–1246.

18. Tkachenko, H., Pažontka-Lipiński, P., & Witaszek, P. (2016). Seasonal alterations in exercise-induced oxidative stress of horses involved in recreational horse-back ride. Globalisation and regional environment protection. Technique, technology, ecology. Eds T. Noch, W.Mikolajczewska, A.Wesolowska. Gdańsk, Gdańsk High School Publ.,193–212.

19. Urso, M. L., & Clarkson, P. M. (2003). Oxidative stress, exercise, and anti-oxidant supplementation. Toxicology, 189, 41–54.

20. Watson, T. A., MacDonald-Wicks, L. K., & Garg, M. L. (2005). Oxidative stress and antioxidant in athletes undertaking regular exercise training. International Journal of Sport Nutrition and Exercise Metabolism, 15, 131–146.

ВЫЗВАННЫЕ ФИЗИЧЕСКИМ ТРЕНИНГОМ ИЗМЕНЕНИЯ В СОДЕРЖАНИИ БИОМАРКЕРОВ ОКИСЛИТЕЛЬНОГО СТРЕССА ЭРИТРОЦИТОВ У ПОНИ, УЧАСТВУЮЩИХ В РЕКРЕАЦИОННОЙ ВЕРХОВОЙ ЕЗДЕ

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Целью исследования был анализ биомаркеров окислительного стресса, а также устойчивости эритроцитов у кобыл и жеребцов пони, участвующих в рекреационной верховой езде. В нашем исследовании были использованы десять здоровых взрослых пони (5 жеребцов и 5 кобыл), в возрасте 5-11 лет, из Поморских регионов Польши (Устка, Поморское воеводство, Польша). Все лошади участвовали в рекреационной езде. Тренировка продолжалась 1 час и включала выезду по пересеченной местности с ходьбой (10 минут), рысь (15 минут), ходьбу (10 минут), галоп (15 минут) и ходьбу (10 минут). Кровь отбирали из яремных вен животных утром, через 90 минут после кормления и сразу после тренировки. Маркер перекисного окисления липидов (вещества, реагирующие с 2-тиобарбитуровой кислотой, TBARS), альдегидные и кетоновые производные окислительно модифицированных белков, уровень общей антиоксидантной активности (TAC) в эритроцитах как кобыл, так и жеребцов, показали незначительные изменения сразу после физической нагрузки по сравнению с периодом отдыха. Как в покое, так и после тренировки уровни TBARS, альдегидных и кетоновых производных окислительно модифицированных белков между кобылами и жеребцами после тренировки. Как в покое, так и после тренировки уровень TAC в эритроцитарных супсепшнях жеребцов был незначительно выше по сравнению с жеребцами. Аналогичным образом были отмечены различия в уровнях альдегидных и кетоновых производных окислительно модифицированных белков между кобылями и жеребцами после тренировок. Как в покое, так и после тренировки уровень TAC в эритроцитарных супсепшнях жеребцов был незначительно выше по сравнению с кобылами. Как у кобыл, так и у жеребцов, после тренировок наблюдалось незначительное снижение гемолиза эритроцитов по сравнению с периодом отдыха. Сравнение гемолиза эритроцитов в период отдыха показало увеличение значений гемолиза у жеребцов. После тренировки наблюдалось снижение гемолиза у жеребцов по сравнению с кобылами. Усилия должны быть направлены на тщательную характеристику антиоксидантной защиты, а
також кореляційних связей між біомаркерами окислительного стресу і антиоксидантної захисту, включаючи вікові і гендерні відмінності в програмах тренувань коней, які беруть участь у рекреаційній верховій їзді.

Ключові слова: речовини, що реагують з 2-тіобарбітурової кислотою (ТБК-активні продукти), альдегідні і кетонові похідні, окиснювально-модифіковані білки, загальна антиоксидантна активність, гемоліз еритроцитів, тренінг.