Endothelial function in athletes in the process of adaptation to various training exercise modes

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Key words: endothelial function, endurance, strength, speed.

A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article.

Studies on endothelial dysfunction and its relationship with adaptive disorders in highly skilled athletes are few in number and mainly carried out in cyclic kinds of sports due to larger volumes and higher intensities of training loads associated with endurance performance gain. Stress and sex hormones and growth hormone play an important role in the regulation of endothelial function, but factors, which can stimulate negative vascular changes, remain a matter of debate. It is also unclear, whether changes in the vascular system depend on the type, mode or intensity of physical activity.

The aim of the work was to perform a comparative assessment of endothelial function and to study the role of some hormones in its regulation in highly qualified athletes in the process of adaptation to various training loads.

Material and methods. After signing a written informed consent, the study involved 104 athletes (80 men and 24 women) qualifying from the First-Class athletes to Masters of Sports of International Class (MSIC): 63 – athletes, who mainly trained endurance performance (triathletes, swimmers, long-distance runners, rowers), 31 – strength performance (weightlifters, powerlifters, kettlebell lifters), 10 – speed performance (sprinters). The mean age of the examined athletes was 21.75 ± 3.32 years. Among them, there were Masters of Sports of International Class (MSIC) – 2 athletes, Masters of Sports (MS) – 25, Candidates Master of Sports (CMS) – 48, First-Class athletes – 29.

Plasma levels of endothelin-1, 6-keto-PG, erythropoietin, growth hormone, testosterone, free testosterone were determined by enzyme-linked immunosassay on a Seac ELISA-Reader Sirio S (Seac Radim Company, Italy).

Results. There was no statistically significant difference in endothelin-1 levels between the endurance and strength athletes, but the strength-trained athletes showed a tendency towards endothelin-1 level predominance (0.77 ± 0.04 fmol/ml vs. 0.72 ± 0.06 fmol/ml; P = 0.176) and 2 times (P = 0.017) higher levels of 6-keto-PG. The endothelin-1 and 6-keto-PG levels did not differ significantly between the endurance and speed athletes. The strength-trained athletes exhibited 5.2 % (P = 0.016) higher endothelin-1 levels than speed-trained athletes. However, these athletes did not differ statistically in the 6-keto-PG (292.30 ± 70.38 pg/ml against 106.92 ± 74.44 pg/ml; P = 0.834) level. A positive correlation was found between the levels of erythropoietin and 6-keto-PG in the endurance-trained (r = 0.57; P = 0.00001) and strength-trained (r = 0.46; P = 0.013) athletes.

Analysis of testosterone and free testosterone levels did not reveal statistically significant differences between endurance-, strength- or speed-trained athletes. At the same time, there was a trend towards higher levels of testosterone and free testosterone in the strength-trained athletes as compared to those in the endurance- or speed-trained athletes. The strength athletes showed a positive correlation (r = 0.46; P = 0.013) between the levels of free testosterone and endothelin-1. The highest level of somatotropic hormone was in the speed-trained athletes (11.74 ± 3.13 mIU/l), 2 times less – in the endurance-trained athletes (5.69 ± 1.19 mIU/l), and the lowest one – in the strength-trained athletes (2.66 ± 1.32 mIU/l). A positive correlation between the growth hormone and erythropoietin levels (r = 0.29; P = 0.038) was revealed in the endurance athletes.

Conclusions. The endurance and speed athletes did not differ in the levels of endothelin-1, 6-keto-PG, erythropoietin, and testosterone. The strength-trained athletes showed signs of endothelial dysfunction: higher endothelin-1 levels with significantly reduced growth hormone and a tendency of increase in the serum testosterone level, as well as a compensatory increase in endurance performance gain. Stress and sex hormones and growth hormone play an important role in the regulation of endothelial function, but factors, which can stimulate negative vascular changes, remain a matter of debate. It is also unclear, whether changes in the vascular system depend on the type, mode or intensity of physical activity.

UDC 616.1-018.74:796.015.5/.6:796.071.2

DOI: 10.14739/2310-1210.2021.1.224881
Результати.
У спортсменів, які удосконалювали якості витривалості або сили, відсутня статистично вірогідна різниця за вмістом ендотеліну-1, проте у спортсменів, які розвивали якість сили, встановлена тенденція до переважання вмісту ендотеліну-1 (0,77 ± 0,04 фмоль/мл проти 0,72 ± 0,06 фмоль/мл; p = 0,176) та зв'язок (r = 0,46; p = 0,017) більший вміст 6-кето-ПГ. За вмістом ендотеліну-1 і 6-кето-ПГ у спортсменів, які розвивали якості витривалості або сили, проявлялося вірогідно не відрізнялася. У спортсменів, які розвивали якість сили, ендотелину-1 більший, ніж у спортсменів, які розвивали якість швидкості, на 5,2 % (p = 0,016). Але цей спортивний статистично не відрізнявся за вмістом ендотеліну-1 (292,30 ± 70,38 пг/мл проти 106,92 ± 74,44 пг/мл; p = 0,834). Між вмістом ендотеліну та 6-кето-ПГ встановлено позитивний кореляційний зв'язок у спортсменів, які удосконалювали якість витривалості (r = 0,57; p = 0,00001), та спортсменів, які розвивали якість сили (r = 0,46; p = 0,013).

Аналіз вмісту тестостерону та вільного тестостерону не показав статистично вірогідну різницю між групами спортсменів, які розвивали якості витривалості або сили, або вільного тестостерону. Але визначено тенденцію до переважання вмісту тестостерону та вільного тестостерону у спортсменів, які удосконалювали якості сили, навіть аналогічними показниками у спортсменів, які розвивали якості витривалості або швидкості. У спортсменів, які розвивали якість сили, вміст вільного тестостерону та ендотеліну-1 виявили позитивний кореляційний зв'язок (r = 0,46; p = 0,013). Найбільший вміст соматотропного гормона відзначено у спортсменів, які розвивали якість швидкості (11,74 ± 3,13 мкМОд/мл), удвічі менший – у спортсмені, які удосконалювали якість витривалості (5,69 ± 1,19 мкМОд/мл), найменший – у силових атлетів (2,66 ± 1,32 мкМОд/мл). У спортсмені, які удосконалювали якість витривалості, виявили позитивний кореляційний зв’язок між вмістом гормона росту та еритропоетину (r = 0,29; p = 0,038).

Функція эндотелия у спортсменов в процессе адаптации к тренировочным нагрузкам различной направленности

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Исследовано эндотелиальную дисфункцию и ее связи с адаптационными нарушениями у высокоактивированных спортсменов немолодогенерации. В основном их проявляют в циклических видах спорта, что обусловлено более интенсивными и объемными тренировочными нагрузками, направленными на развитие выносливости. Важную роль в регуляции функции эндотелия играют гормоны стресса, половые гормоны и гормон роста, но вопрос о том, какие факторы могут стимулировать негативные сосудистые изменения, остается дискуссионным. Также неясно, зависят ли изменения сосудистой функции эндотелия от вида, направленности и интенсивности физических нагрузок.

Цель работы – проведение сравнительной оценки показателей функции эндотелия и изучение роли некоторых гормонов в ее регуляции у высокоактивированных спортсменов в процессе адаптации к тренировочным нагрузкам различной направленности.

Материалы и методы. После подписания письменного информированного согласия в исследование включены 104 спортсмена (80 мужчин и 24 женщины) уровня мастерства от 1 разряда до МСМК – 2 спортсмена, майстрів спорту (МС) – 25, кандидатів у майстри спорту – 48, першорозрядників – 29. Методом ИФА определили уровни эндотелина-1, 6-кето-ПГ, эритропоэтина, гормона роста, тестостерона, свободного тестостерона.

Результаты. У спортсменов, которые совершенствовали качество выносливости или сили, не установлена статистически достоверная разница по содержанию эндотелина-1, однако у спортсменов, совершенствовавших качество сили, отмечена тенденция к преобладанию содержания эндотелина-1 (0,77 ± 0,04 фмоль/мл против 0,72 ± 0,06 фмоль/мл; p = 0,176) и в 2 раза (p = 0,017) большее содержание 6-кето-ПГ. По содержанию эндотелина-1 и 6-кето-ПГ спортсмены, совершенствовавшие качество выносливости или сили, достоверно не отличались. У спортсменов, которые совершенствовали качество сили, содержание эндотелина-1 больше, чем у спортсменов, которые совершенствовали качество скорости, на 5,2 % (p = 0,016). Однако эти спортсмены статистически не отличались по содержанию 6-кето-ПГ (292,30 ± 70,38 пг/мл против 106,92 ± 74,44 пг/мл, p = 0,834). Между содержанием эритропоэтина и 6-кето-ПГ установлена положительная корреляционная связь у спортсменов, совершенствовавших качество выносливости (r = 0,57; p = 0,00001), и спортсменов, которые совершенствовали качество сили (r = 0,46; p = 0,013). Анализ содержания эндотелина и эритропоэтина у спортсменов без тренировочного стимула не показал статистически достоверную разницу между группами спортсменов, которые совершенствовали качество выносливости, сили или скорости. Тем не менее отмечалась тенденция к преобладанию содержания эндотелина и эритропоэтина у спортсменов, совершенствовавших качество сили, над аналогичными показателями у спортсменов, совершенствовавших качество выносливости или скорости. У спортсменов, который развивали качество сили, между содержанием свободного тестостерона и эндотелина-1 установлена положительная корреляционная связь (r = 0,46; p = 0,013). Наибольшее содержание соматотропного гормона отмечено у спортсменов, совершенствовавших качество скорости (11,74 ± 3,13 мкМОд/мл), в 2 раза меньше – у спортсменов, совершенствовавших качество выносливости (106,92 ± 74,44 пг/мл; p = 0,834). Між вмістом ендотеліну та 6-кето-ПГ встановлено позитивний кореляційний зв’язок у спортсменів, які розвивали якість витривалості (r = 0,57; p = 0,00001), та спортсменів, які розвивали якість сили (r = 0,46; p = 0,013).
Special attention is now being paid to studying vascular endothelial dysfunction as its biological role in the development of pre-pathological and pathological conditions remains poorly known, and the prevention of endothelial dysfunction can be important for achieving performance benefits.

Higher volumes of exercise can accelerate the development of endothelial dysfunction, and high-intensity exercise can increase the risk of cardiovascular disease (CVD). The severity of vascular damage depends on training volumes, undertaken by an athlete throughout life. CVD risk is likely to be associated with the type of sports discipline practiced [9].

However, there was evidence of no differences in endothelial function in athletes, who performed different intensity and duration of exercise [15]. In modern high-achievement sports, intense physical activity leads to multidirectional changes in the body; in particular, contributes to the development of universal biochemical and physiological phenomena — chronic oxidative stress and associated tissue hypoxia [4], which is one of the components of vascular endothelial dysfunction [8].

Endothelial cells are less sensitive to hypoxia and are damaged by ischemia much less than other cells due to their ability to switch to anaerobic energy metabolism, as well as to synthesize heat shock proteins, glucose-regulated proteins, enzymes involved in glycolysis (glyceraldehyde 3-phosphate dehydrogenase and non-neuronal enolase), that increases the resistance of cells to damage [23]. Stress hormones, sex hormones and growth hormone (GH) play an important role in the regulation of endothelial function, but what kind of factors can stimulate negative vascular changes, remains a matter of debate. It is also unclear, whether changes in the vascular system depend on the type, mode or intensity of physical activity [7].

Endothelial dysfunction is recognized as a universal mechanism modulated by all risk factors associated with cardiovascular disorders. To assess the adaptation level to training loads in athletes, the dynamics of changes in the vascular endothelium regulators is used [1]. However, studies on endothelial dysfunction and its relationship with adaptive disorders in highly skilled athletes are few in number [13] and mainly carried out in cyclic kinds of sports due to larger volumes and higher intensities of training loads in athletes, the dynamics of changes in the vascular system depend on the type, mode or intensity of physical activity [7].

Therefore, the aim of the work was to perform a comparative assessment of endothelial function and to study the role of some hormones in its regulation in highly qualified athletes in the process of adaptation to various training loads.

Material and methods

After signing a written informed consent, the study involved 104 athletes (80 men and 24 women) qualifying from the First-Class athletes to Masters of Sports of International Class (MSIC): 63 – athletes, who mainly trained endurance performance (triathletes, swimmers, long-distance runners, rowers), 31 – strength performance (weightlifters, powerlifters, kettlebell lifters), 10 – speed performance (sprinters). The mean age of the examined athletes was 21.75 ± 3.32 years. The mean age of the examined athletes was 21.75 ± 3.32 years. Among them, there were Masters of Sports of International Class (MSIC) – 2 athletes, masters of sports (MS) – 25, candidates masters of sports – 48, athletes of the 1 category – 29.

Plasma levels of endothelin-1, erythropoietin, 6-keto-prostaglandin, testosterone, free testosterone and GH were determined by enzyme-linked immunoassay on a Seac ELISA-Reader Sirio S (Seac Radim Company, Italy).

The results were processed by methods of variation statistics using the software package Statistica 13.0 (StatSoft, USA), license number JPZ8041382130ARCN10-J). The Shapiro–Wilks test was used to test the hypothesis of normal distribution of quantitative indicators. The quantitative indicators were presented in the form of arithmetic mean and standard error (M ± m), Me (Q25; Q75) taking into account the normality of the data distribution; qualitative indicators – in the form of absolute and relative frequency. The quantitative indicators of independent groups were compared by the method of parametric statistics using the two-sample Student’s t-test with a two-sided test index for a statistical significance value, and the non-parametric Mann–Whitney U test. The differences were considered statistically significant at a value of P < 0.05.

Results

Endothelial function in athletes was assessed by the levels of endothelin-1 and 6-keto-prostaglandin (6-keto-PG). There was no statistically significant difference in endothelin-1 levels between the endurance and strength athletes (Table 1, Fig. 1), but the strength-trained athletes showed a tendency towards endothelin-1 level predominance (0.77 ± 0.04 fmol/ml vs. 0.72 ± 0.06 fmol/ml; P = 0.176) and 2 times (P = 0.017) higher levels of 6-keto-PG (Table 1, Fig. 2). The levels of endothelin-1 and 6-keto-PG in the endurance- and speed-trained athletes did not differ significantly (0.72 ± 0.06 fmol/ml vs. 0.73 ± 0.05 fmol/ml; P = 0.176) and (138.24 ± 26.12 pg/ml vs. 106.92 ± 74.44 pg/ml; P = 0.834), respectively.

A comparison of endothelin-1 level between the strength and speed athletes showed significantly 5.2 % higher its concentrations in strength-trained athletes (0.77 ± 0.04 fmol/ml

**Table 1**

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Endurance (n = 31)</th>
<th>Strength (n = 31)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endothelin-1 (fmol/ml)</td>
<td>0.77 ± 0.04</td>
<td>0.72 ± 0.06</td>
<td>0.176</td>
</tr>
<tr>
<td>6-keto-Prostaglandin (pg/ml)</td>
<td>138.24 ± 26.12</td>
<td>106.92 ± 74.44</td>
<td>0.834</td>
</tr>
</tbody>
</table>

**Fig. 1**

**Fig. 2**
A positive correlation was found between the levels of 6-keto-PG did not differ statistically between these athletes, there was a tendency to a predominance of 6-keto-PG in the strength-trained athletes compared with the speed-trained athletes (292.30 ± 70.38 pg/ml against 106.92 ± 74.44 pg/ml, P = 0.834).

Thus, the levels of endothelin-1 and 6-keto-PG did not differ between the endurance and speed athletes, who shared the same pattern of endothelial function. At the same time, the strength athletes showed signs of endothelial dysfunction because they had higher levels of endothelin-1. However, this increase in vasoconstrictor was compensated by a 2-fold increase in the level of 6-keto-PG – a biologically active substance with vasodilating properties.

An analysis of erythropoietin level (Table 1, Fig. 3) revealed an upward trend in its concentration in the speed-trained athletes. The lowest level of erythropoietin was observed in the endurance athletes. However, the difference in erythropoietin levels between the groups of endurance-, strength- and speed-trained athletes was not statistically significant.

A positive correlation was found between the levels of erythropoietin and 6-keto-PG (Fig. 4) in the endurance-trained athletes (r = 0.57; P = 0.00001) and in the strength-trained athletes (r = 0.46; P = 0.013).

An analysis of testosterone and free testosterone (Table 1, Fig. 5, 6), despite the expectations, did not reveal a statistically significant difference between the groups of endurance-, strength- and speed-trained athletes. But even so, there was a trend towards higher levels of testosterone and free testosterone in the strength-trained athletes as compared to those in the endurance- or speed-trained athletes.

In the strength-trained athletes (Fig. 7), a positive correlation was found between the levels of free testosterone and endothelin-1 (r = 0.46; P = 0.013), which may indicate the ability of the active fraction of testosterone to induce endothelin-1 synthesis in the vascular endothelium.

The athletes of all three studied groups differed significantly in GH level (Table 1, Fig. 8). The highest level of somatotropic hormone was in the speed-trained athletes (11.74 ± 3.13 mUI/l), 2 times less – in the endurance-trained athletes (5.69 ± 1.19 mUI/l), and the lowest one – in the strength-trained athletes (2.66 ± 1.32 mUI/l).

A positive correlation between the GH and erythropoietin levels (r = 0.29; P = 0.038) was revealed in the endurance-trained athletes (r = 0.29; P = 0.038).

Thus, the endurance and speed training in the athletes was associated with a significant increase in the somatotropic hormone level, due to the need to ensure an adequate level of aerobic energy metabolism during muscular exercise.

### Table 1. The levels of endothelin-1, 6-keto-PG, erythropoietin, growth hormone, testosterone, free testosterone in endurance-, strength- and speed-trained athletes, M ± m, Me (Q25; Q75)

<table>
<thead>
<tr>
<th>Value, units of measure</th>
<th>Performance</th>
<th>Strength</th>
<th>Speed</th>
<th>P&lt;sub&gt;1-2&lt;/sub&gt; Endurance – Strength Mann–Whitney U Test</th>
<th>P&lt;sub&gt;1-3&lt;/sub&gt; Endurance – Speed Mann–Whitney U Test</th>
<th>P&lt;sub&gt;2-3&lt;/sub&gt; Speed – Strength Mann–Whitney U Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythropoietin, mlU/ml</td>
<td>7.66 ± 0.63</td>
<td>8.52 ± 1.47</td>
<td>10.57 ± 4.95</td>
<td>0.803</td>
<td>0.460</td>
<td>0.304</td>
</tr>
<tr>
<td>Endothelin-1, fmol/ml</td>
<td>0.72 ± 0.06</td>
<td>0.77 ± 0.04</td>
<td>0.73 ± 0.05</td>
<td>0.178</td>
<td>0.160</td>
<td>0.016</td>
</tr>
<tr>
<td>Testosterone, ng/ml</td>
<td>7.78 ± 0.59</td>
<td>8.14 ± 0.68</td>
<td>5.83 ± 1.55</td>
<td>0.792</td>
<td>0.126</td>
<td>0.873</td>
</tr>
<tr>
<td>Testosterone free, pg/ml</td>
<td>11.37 ± 1.18</td>
<td>13.00 ± 1.59</td>
<td>6.78 ± 2.05</td>
<td>0.459</td>
<td>0.071</td>
<td>0.873</td>
</tr>
<tr>
<td>Growth hormone, mUI</td>
<td>5.69 ± 1.19</td>
<td>2.66 ± 1.32</td>
<td>11.74 ± 3.13</td>
<td>0.014</td>
<td>0.021</td>
<td>0.011</td>
</tr>
<tr>
<td>6-keto-PG, pg/ml</td>
<td>138.24 ± 26.12</td>
<td>292.30 ± 70.38</td>
<td>106.92 ± 74.44</td>
<td>0.017</td>
<td>0.085</td>
<td>0.834</td>
</tr>
</tbody>
</table>

#### Fig. 1. The level of endothelin-1 in the endurance-, strength- and speed-trained athletes.

#### Fig. 2. The level of 6-keto-PG in the endurance-, strength- and speed-trained athletes.
Endothelial dysfunction is characterized by a shift in the endothelium towards a pro-inflammatory and prothrombotic state with a decrease in vasodilation or paradoxical vasoconstriction. Paradoxical vasoconstriction and impaired NO-dependent vasodilation are of particular clinical significance in conditions of mental and physical stress. Some studies have proven a decreased endothelial response and endothelial dysfunction presence in athletes [7,13].

In the work of Smirnov I.E. it has been shown that young swimmers of both sexes had the most significant increase in plasma endothelin-1 concentrations: 3 times – in female swimmers, 3.4 times – in male swimmers compared with...
the reference group levels. However, the serum nitric oxide content in all the examined young swimmers was not different from the control group. Derangements in endogenous production of nitric oxide and endothelin-1 in the examined swimmers was accompanied by a significant decrease (1.4 times) in the plasma concentration ratio of these bioregulators in both boys and girls, indicating a shifted balance towards higher endothelin-1 production [13].

In the study of V. V. Kolotyrkova et al. among weightlifters during the reactive hyperemia test before exercise, there was no change in the brachial artery diameter, and after exercise it decreased by 5.3 %. In the group of athletes before exercise, the brachial artery diameter increased by 2.9 %, after exercise there was the vessel dilation by 4.1 %. That is, there was an increase in endothelial function in response to exercise training. According to the researchers, the endothelial functional state in strength sports athletes should be considered as an adaptive factor to regular exercise, as there was a temporary cessation of blood flow due to compression of blood vessels by muscle tensing. Significant endothelium-dependent vasodilation in such conditions may result in steal syndrome [7].

Athletes trained in cyclic sports also showed a decrease in endothelial activity [7]. According to D. J. Green and co-authors, signs of early atherosclerosis and a tendency to thrombosis were revealed in 60.5 % of athletes in cyclic kinds of sports (rowing, skiing and cycling). The researchers believe, that constant exposure of endothelium to shear stress, increased blood pressure and production of endothelium-dependent factors may stimulate the development of atherosclerosis [19].

The researchers attributed different responses to the reactive hyperemia test before and after exercise among athletes in cyclic and strength kinds of sports to variations in the endothelial nitric oxide synthase (eNOS) expression. Strength athletes demonstrated a sharp decrease in plasma eNOS after exercise training, while its contrasting increase was found in track-and-field athletes. In addition, the studies showed that the reactive hyperemia test induced NO-mediated brachial artery dilation [19].

Our study has shown no statistically significant differences in endothelin-1 and 6-keto-PG levels in the endurance- and speed-trained athletes. Meanwhile, the strength-trained athletes showed signs of endothelial dysfunction. They exhibited higher levels of endothelin-1, but the increased vasoconstrictor level was compensated by significantly increased level of 6-keto-PG.

When analyzing the levels of hormones with a potential effect on endothelial function, the significant difference only in somatropic hormone has been found.

It is known, that one of the most permanent biological effects of GH is the dilation of peripheral blood vessels. This effect is thought to be mediated by insulin-like growth factor-1 (IGF-1), for which there are high-affinity binding sites on the endothelial cells. IGF-1 directly stimulates the release of NO, which is a known mediator of vasodilation, inhibitor of platelet aggregation, leukocyte adhesion and smooth muscle cell growth. In healthy volunteers, infusions of IGF-1 caused a pronounced dilation of the forearm blood vessels, which was prevented by the administration of L-NG-monomethylarginine [2]. Recombinant growth hormone (rGR) replacement therapy had a positive effect on the vascular parameters: improvements of endothelial function, vascular elasticity, a reduction in total cholesterol, LDL cholesterol and an increase in HDL cholesterol. In GH-deficient adults, rGR treatment reversed early atherosclerotic changes in major arteries. In a small number of GH-deficient adults, rGR replacement therapy for 6-24 months reduced the intima-media complex thickness of the common carotid artery to normal [16]. Discontinuation of rGR therapy in adults (but not in adolescents) with severe GH deficiency led to an increase in the intima-media complex thickness and cardiovascular risk [17].

We have found the highest levels of GH in the speed-trained athletes, the endurance-trained athletes have shown half of this level, and the lowest levels of the hormone have been revealed in the strength-trained athletes.

Changes in GH level in athletes were also obtained in another study [12]. An increase in GH and a decrease in insulin levels after muscle work (30-minute ergometric cycling load with an intensity of 75 % of ischemic preconditioning (IPC)) were observed in endurance-trained athletes. These changes occurred due to the need to ensure proper muscle energy metabolism, as enforced by somatotropin – an insulin antagonist. In individuals with muscle adaptations to the load effect, there was a significant increase in insulin at constant levels of somatotropin. Athletes, who developed anaerobic capacity through graduated ergometric loads, were characterized by high stability of GH and insulin. According to the researchers, such changes in the plasma GH and insulin concentrations at rest and after graduated ergometric loads indicated the presence of specific shifts in hormonal and metabolic status due to the level and peculiarity of daily physical activity [12].

Although the erythropoietin levels showed no significant differences between the endurance-, strength- and speed-trained athletes, it exhibited a clear upward trend in the speed-trained athletes as compared to this hormone levels in the strength athletes, and the lowest levels of erythropoietin were determined in the endurance athletes. The most interesting fact was the positive correlation between the levels of 6-keto-PG and erythropoietin that we thought was indicative of the endothelioprotective properties of the latter. The association between the erythropoietin and 6-keto-PG levels was observed in both the endurance- and strength-trained athletes, i.e. regardless of the training mode. The data obtained suggest the positive effect of erythropoietin on endothelium-independent vasodilation. A possible pathophysiological mechanism of this erythropoietin action should be considered as its ability to increase circulating plasma titer of endothelial progenitor cells, which effectively help to maintain the vascular endothelium integrity. The process of endothelial progenitor cells homing or recruitment of circulating ones to endothelial injury or ischemic sites is regulated by key angiogenic chemokine stromal cell-derived factor-1 (SDF-1). In conditions of ischemia, inflammation, discrete regions of hypoxia in the bone marrow undergo an increase in the hypoxia inducible factor-1 (HIF-1) transcriptional activity, which is responsible for an expression of SDF-1. Nitric oxide, estrogens, high-density lipoproteins, vascular endothelial growth factor and erythropoietin also increase the plasma titer of endothelial progenitor cells and recruit them to the site of injury through activating the phosphotidylinositol-3-phosphate...
endothelial cells [5].

under the influence of the laminar shear stress of the blood, (PIP3)/Akt pathway. Cell adhesion molecules, such as P/E-selectin and ICAM-1, mediate the binding of endothelial progenitor cells to the injured endothelium. Subsequently, under the influence of the laminar shear stress of the blood, the attached endothelial progenitor cells differentiate into endothelial cells [5].

We have not obtained the statistically significant association between testosterone or free testosterone and exercise training mode in our study. However, there was a trend towards higher levels of testosterone and free testosterone in the strength-trained athletes as compared to those in the endurance- or speed-trained athletes.

The study [10] has proved a significant impact of sports specialization and sports experience, as well as the amounts and types of food intake and daily physical activity on the plasma levels of sex hormones, such as testosterone, progesterone, estradiol, follicle-stimulating and luteinizing hormone. Performing a 30-minute ergometric bicycle load with an intensity of 75 % of the IPC led to a decrease in the serum concentration of sex hormones in persons with different levels and specifics of daily physical activity compared to baseline values. The combined use of muscular loading and nutritional supplementation was accompanied by a pronounced gradual recovery of serum hormone levels in both athletes training in different exercise modes and non-athletes relative to the baseline values. The serum concentration of sex hormones in the athletes, who trained in different modes of energy expenditure, was highly resistant to muscular load in contrast to the control group [10].

The literature data on the effects of male sex hormones on the cardiovascular system and endothelial function are contradictory. The negative effect of androgens on the course of cardiovascular pathology is confirmed by a decrease in HDL cholesterol during puberty-related surge of androgens in boys and in the testosterone esters treatment for adolescents with delayed puberty [11], early onset CVD in athletes who used anabolic androgenic steroids. The monograph [6] presents 17 cases of anabolic androgenic steroid-related CVD (11 acute myocardial infarctions, 2 strokes and 4 cases of cardiomyopathy) in young athletes engaged in bodybuilding.

However, according to C. B. Severo (2013), anabolic androgenic steroid-users and non-users did not differ regarding endothelium-independent function, but the brachial artery flow-mediated dilation was significantly reduced in anabolic androgenic steroid-users athletes compared to non-users (P = 0.03). In addition, flow-mediated dilation was positively associated with high-density lipoprotein cholesterol levels (r = 0.49, P = 0.03) [22].

Despite the prevailing ideas about the negative effects of testosterone and the cardioprotective effect of estrogens, such a dichotomy is conditional. Callou de Sa E.Q. et al. found an association between estradiol (E2), rather than testosterone, and coronary heart disease, confirmed by coronaryography, in young and middle-aged men [14]. Approximately 30–40 μg of E2, the most active estrogen, is synthesized per day in men. About 20 % of this hormone is secreted directly from the testes by Leydig cells, the remaining 80 % is derived from peripheral aromatization of circulating androgen precursors, mainly testosterone, in different tissues (fat, prostate, endothelium and other tissues).

Meanwhile, in clinical observations, androgen therapy in men with coronary heart disease has been found to increase ST-segment depression [3]; conversely, other studies have shown improvements in coronary blood flow and the functional class of chronic heart failure, a decrease in blood pressure and left ventricular myocardial mass [20]. Testosterone replacement therapy stimulates stem cells-derived endothelial progenitor cells via androgen receptors, contributing to myocardial healing and reduction in necrotic lesion size [18].

The reason given by the vast majority of studies for the protective effect of endogenous androgens on the heart and blood vessels is the conversion of the hormones into estrogens, but in the work of D. Liu et al., the protective effect of dehydroepiandrosterone sulfate on the endothelium was not prevented by estrogen receptor inhibition [21].

Thus, in the strength-trained athletes, we have found the highest level of endothelin-1, which was associated with elevated testosterone levels, as evidenced by the direct correlation, and the lowest growth hormone concentration. However, the increase in endothelin-1 level in strength-trained athletes was accompanied by the significant increase in the level of 6-keto-PG. It is clear that shear stress, induced by strength exercise, on endothelial cells is caused by an increase in vasoconstrictors, which contributes to changes in testosterone and growth hormone levels, but the balance is restored by a compensatory increase in the vasodilator 6-keto-PG synthesis.

Conclusions

The endurance and speed athletes did not differ in the levels of endothelin-1, 6-keto-PG, erythropoietin, and testosterone. The strength-trained athletes showed signs of endothelial dysfunction: higher endothelin-1 levels with significantly reduced growth hormone and a tendency of increase in the serum testosterone level, as well as a compensatory increase in 6-keto-PG to maintain the balance between vasoconstrictors and vasodilators.

Conflict of interest: authors have no conflict of interest to declare.

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Сведения о статье:

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