# Adaption of Maximal Glycolysis Rate after Resistance Exercise with Different Volume Load





## Authors Nico Nitzsche, Julian Christian Lenz, Pjotr Voronoi, Henry Schulz

#### Affiliation

Institute of Human Movement Science and Health, Technische Universitat Chemnitz, Chemnitz, Germany

#### Key words

resistance exercise, lactate, anaerobic power, anaerobic metabolism, muscle adaption

received 22.02.2020 revised 24.03.2020 accepted 25.03.2020

#### Bibliography

DOI https://doi.org/10.1055/a-1146-4236 Sports Medicine International Open 2020; 4: E39–E44 © Georg Thieme Verlag KG Stuttgart - New York ISSN 2367-1890

### Correspondence:

Dr. Dr. Nico Nitzsche Technische Universitat Chemnitz, Institute of Human Movement Science and Health Thüringer Weg 11 09126 Chemnitz Germany Tel.:+4937153137326, Fax :+49 371 531-834942 nico.nitzsche@hsw.tu-chemnitz.de

#### ABSTRACT

The aim of this study was to investigate the effect of six-weeks of resistance training with different volume load on the maximum glycolysis rate. 24 male strength-trained volunteers were assigned in a high volume low load (50% of their 1RM with 5 sets and reps up to muscle failure) and a low volume high load (70% of their 1RM with 5 sets of ten reps) resistance exercise group. The resistance training performed 3 days per week over 6 weeks. The maximum glycolysis rate was determined using isokinetic force testing before and after the intervention. There was a significant increase in glycolysis rate over the training period across all subjects (p = 0.032). High volume low load exercise increased significantly from 0.271 ± 0.067 mmol·l<sup>-1</sup>·s<sup>-1</sup> to  $0.298 \pm 0.067$  mmol·l<sup>-1</sup>·s<sup>-1</sup> (p = 0.022) and low volume high load exercise showed no significant changes from 0.249±0.122  $mmol \cdot l^{-1} \cdot s^{-1}$  to 0.291 ± 0.089 mmol  $\cdot l^{-1} \cdot s^{-1}$  (p = 0.233). No significant effect on glycolysis rate was observed between the training groups (p = 0.650). Resistance training increases glycolysis rate regardless of volume load.

# Introduction

Resistance training represents a great strain for anaerobic energy metabolism. This metabolic strain is an important prerequisite for muscle growth and adaption of energy metabolism [1]. In various studies, high-energy substrates, such as creatine phosphate (PCr) and adenylic acids (ATP, ADP), were examined during strength exertion or strength training and significant changes in concentration were observed during and immediately after exertion [2, 3]. Other studies show the high burden on anaerobic energy metabolism using lactate concentration [La + ], which is mostly considered post-load [4–7].

In the long term, regular strength training leads to the adaption of enzymes and substrates of energy supply and muscle volume [7–11]. It has been shown, for example, that the activity of anaer-

obic metabolism enzymes (lactate dehydrogenase (LDH), myokinase) in glycolytic muscle fibers is increased compared to oxidative fibers as a result of chronic strength training. Takada et al. [2] showed that the increases in inorganic phosphate (P<sub>i</sub>) and adenosine diphosphate (ADP) are associated with an increase in muscle volume following four weeks of resistance training. Haun et al. [11] determined increases in anaerobic energy metabolism enzymes (LDH, phosphofructokinase (PFK)) through resistance training lasting several weeks. Consequently, the energetic flow rate increases due to ATPs and an increased lactate accumulation occurs with unchanged lactate elimination. Based on assumptions and findings, metabolic stress is an important factor in muscle hypertrophy [1,7,11]. The studies suggest that increases in muscle volume are associated with an adaption of the anaerobic energy metabolism.

🖗 Thieme

The rate of lactate accumulation (vLa<sub>max</sub>) (also known as the glycolysis rate) can thus be regarded as an indirect measure of glycolysis activity [12, 13]. This rate can be determined based on the acute increase in [La + ] as a result of the load as a function of the load time ( $t_{load}$ ). Thus, the activity of glycolysis can be estimated quite simply on the basis of the change in [La + ] as a function of the t<sub>load</sub> and show the metabolic stress. This measure describes the efficiency of the anaerobic energy metabolism, which increases significantly during strength training and is dependent on movement speed [14]. Good reproducibility of vLa<sub>max</sub> was shown during isokinetic force loading as well as sprint loading [15, 16]. The adaptation of the vLa<sub>max</sub> has been examined in endurance studies. So far, only one study is available that shows a reduction of vLa<sub>max</sub> following endurance training that lasted several weeks [17]. This study compared sprint interval (SIT) training with continuous endurance training (CET) and showed that after 6 weeks SIT, there was a significant reduction in the  $vLa_{max}$ . This remained unchanged at CET. The adaptation of the  $\nu La_{max}$  through resistance training lasting several weeks has not yet been investigated.

In resistance training, protocols are generally used that contain different exercise volume load (volume load = set × reps × load; e. g.,  $2500 = 5 \text{ sets} \times 10 \text{ reps} \times 50 \text{ kg}$ ; load based in % of 1RM) [18, 19]. The comparison of training protocols showed that high-volume training causes significantly higher lactate concentrations compared to low volume with high load after the exertion [20, 21]. After an 8-week resistance exercise period, Mangine et al. [20] did not observe a change in [La + ] immediately after the last exercise session in a high-volume and a strength protocol. This [La + ] is the result of the acute load and does not show the efficiency of the anaerobic glycolysis.

With a focus on muscle volume, a few studies examined volume following resistance training with different volume loads. For example, the impact of volume load in resistance exercise was examined [22]. A high-load protocol of 3 sets of 10 reps at 75% of 1 RM was compared with a low load of 4 sets of reps until failure at 30% 1 RM on pectoralis major and triceps brachii hypertrophy. The resistance training lasted 6 weeks with 3 training days per week. The results showed that both resistance training protocols had comparable muscle cross section increases. An another study by Schoenfeld et al. [23] examined different training volumes with a strength training protocol (7 sets, 3 reps at 3 RM) and a hypertrophy training program (3 sets, 10 reps at 10 RM) over 8 weeks (3 × weekly). The results showed comparable increases in muscle thickness in the biceps brachii in both groups. Both studies showed no significant impact on muscle hypertrophy at different volume loads. However, adaptations of the energy metabolism based on vLa<sub>max</sub> have hardly been considered in this context. If metabolic stress is important for muscle hypertrophy, it would be helpful to know how the metabolic measure vLa<sub>max</sub> adapts through resistance training at different volume loads.

Based on the available studies, it is assumed that resistance training over several weeks leads to a change in glycolytic enzymes and [La + ] in the afterload. No statements can be made about the changes in the vLa<sub>max</sub> caused by resistance training. The aim of this study was to investigate the effect of 6 weeks of resistance training with different training volumes load on vLa<sub>max</sub>. If changes in vLa<sub>max</sub> are associated with changes in performance from strength training over several weeks, the  $vLa_{max}$  could be an important parameter of anaerobic performance in resistance training. It may be possible to assess metabolic adaptations with varying training volumes in strength training using  $vLa_{max}$ .

# Materials and Methods

## Experimental design and subjects

After receiving information and giving written consent to participate in the study, 24 healthy male strength-trained subjects were assigned to one of two groups (high-volume, low-load = HVLL, lowvolume, high-load = LVHL) with different training volume loads. HVLL (n = 14; age 25.0 ± 4.3 years; height 179.7 ± 7.1 cm; body mass 83.6 ± 11.0 kg; body mass index 25.9 ± 3.1 kg m<sup>-2</sup>) trained at 50 % of the 1RM with 5 sets and reps up to muscle failure. The subjects in LVHL (n = 10; age 24.6 ± 2.8 years; height 178.1 ± 6.0 cm; body mass  $80.5 \pm 11.2$  kg; body mass index  $25.4 \pm 3.3$  kg m<sup>-2</sup>) trained at 70% of the 1RM with 5 sets of 10 reps. All test subjects were free of injuries and chronic diseases. Furthermore, all subjects had more than 2 years of training experience and had a training scope of 1 to 4 training units per week at the beginning of the study. The training scope ranged from 1.5 up to 5.0 hours per week. The study meets the ethical standards in sports and exercise science and was approved by the local ethical committee (V-361-17-HSchvla<sub>max</sub>-12122019) [24].

#### Measurements

Before the intervention, anthropometric data and the one repetition maximum (1RM) were recorded. A maximum isokinetic strength test (Con-Trex<sup>®</sup> Multi Joint System, Physiomed, Schnaittach, Germany) was performed before and after the intervention to determine anaerobic performance and capacity. A concentric isokinetic strength test was carried out with an ankle velocity of 180 °s<sup>-1</sup> and 10 reps (15 s load time). The maximum (P<sub>max</sub>) and mean maximum power (meanP<sub>max</sub>, mean of ten reps) of the thigh extensors and flexors were evaluated. P<sub>max</sub> was standardized to body mass.

To determine the lactate concentration [La +], capillary blood samples (20µl) were taken from the earlobe before  $[La +]_{pre}$ , immediately after exercise, and up to the ninth minute post-exercise (up to the third minute at 30-second intervals, from the third to the ninth minute at 60-second intervals). The calculation of the maximum glycolysis rate (vLa<sub>max</sub>) was based on the pre-load lactate concentration  $[La +]_{pre}$ , maximum lactate concentration in the postload  $[La +]_{max}$ , the loading time (t<sub>load</sub>), and the alactic time interval of 3 seconds [13, 25]. The reproducibility of the maximum glycolysis rate by an isokinetic strength test showed a high correlation of r > 0.67 [15].

## Intervention

Two to five days before the training intervention, the 1RM was recorded to calculate the training load for each exercise [26]. Both groups trained their lower extremities 3 times a week for 6 weeks. The strength training program was carried out on sequential machines (Gym80, Gelsenkirchen, Germany). The exercises consisted of leg press (LP), leg extension (LE) and leg flexor (LC, in the prone position) sets and were performed bilaterally in random order. Both training groups performed 5 sets each with a break of 90 seconds between series. HVLL completed the maximum possible number of repetitions until local muscle failure at 50% of the 1RM. LVHL completed 5 sets of 10 repetitions each at 70% of the 1RM. There was a regeneration period of 24 to 48 hours between each training day. The exercise sessions were observed by a practiced coach.

The absolute exercise volume load (EV) was calculated from the product of the training weight (load) and the number of repetitions (rep), which was then summed up over all sets and training sessions (TS) [19]. At relative EV (per TS), the total EV was relativized to TS.

#### Statistical analysis

The arithmetic mean (mean), standard deviation (±SD), minimum (MIN), and maximum (MAX) were calculated for all data (Microsoft Excel Version 16.0; Microsoft, Redmond, WA, USA). The inferential statistical analysis was performed using IBM SPSS Statistics Version 22 (IBM Corp., Armonk, NY, USA). The test for normal distribution was performed using the Shapiro Wilk test. Homoscedasticity was checked using Levene's test. If the test requirements were met, the training group and training time were checked for significant effects on the dependent variables using two-way variance analysis. If the requirements for a parametric test were not met, a Friedman test was used to check for significant main effects. Comparisons between the two groups were then made using the Mann-Whitney U test. Pre-post comparisons within the group were performed using a dependent t-test and Wilcoxon's test. The effect sizes were determined for pre-post comparisons using Cohen's d (d).  $\eta^2$  was used as an effect measure in variance-analytical comparisons. The interpretation of effect size's based on [27]. The test power for vLa<sub>max</sub> was determined post-hoc (G \* Power, Version 3.1.9.2; Düsseldorf, Germany). Correlation analyses using Spearman (no normal distribution) were used to check for changes in performance associated with changes in vLa<sub>max</sub>. The level of significance was set at  $p \leq 0.05$ .

# Results

There were no significant differences in anthropometric data or performance between the two training groups before the training intervention (p>0.05). The training frequency (TF) after six weeks was 17.07  $\pm$  1.27 (94.9%) in HVLL and 16.8  $\pm$  2.35 (93.3%) in LVHL (p = 0.841). The relative EV was 10868  $\pm$  2960 kg for HVLL and 4908  $\pm$  1989 kg for LVHL (p = 0.000; d = 2.286). HVLL had an approximately 2.2 times higher EV compared to LVHL. In HVLL, this resulted in 18.43  $\pm$  2.93 (13.91–25.92) reps per set and exercise. The LVHL group always completed 10 reps per set and exercise.

A significant time effect of vLa<sub>max</sub> was observed after 6 weeks of strength training (p = 0.032; d = 0.974;  $\eta_{part}^2 = 0.192$ ). vLa<sub>max</sub> showed an increase from 0.262 ± 0.092 to 0.295 ± 0.075 mmol·l<sup>-1</sup>·s<sup>-1</sup> (**▶ Fig. 1**). In HVLL, vLa<sub>max</sub> increased significantly (p = 0.022; d = 0.406), but in LVHL the increase of vLa<sub>max</sub> was not significant (p = 0.233; d = 0.384). A significant group effect on vLa<sub>max</sub> could not be determined (p=0.650; d=0.201;  $\eta_{part}^2$ =0.010).

The [La + ]<sub>pre</sub> showed no significant differences between pre-test and post-test in either group (p>0.05). [La + ]<sub>max</sub> showed a significant time effect (p = 0.001; d = 0.685;  $\eta_{part}^2$  = 0.376). The maximum lactate concentration increased from 3.85 ± 0.965 mmol l<sup>-1</sup> to 4.45 ± 0.771 mmol l<sup>-1</sup>. [La + ]<sub>max</sub> increased significantly in HVLL (p=0.012; d=0.410;  $\eta_{part}^2$ =0.042) and in LVHL (p=0.039; d=1.03;  $\eta_{part}^2$ = 0.212). A significant group effect was not observed for [La + ]<sub>max</sub> (p=0.130; d=0.652;  $\eta_{part}^2$ = 0.096).

The isokinetic strength test showed significant time effects of mean  $P_{max}$  (p = 0.000; d = 2.268), relative mean  $P_{max}$  (p = 0.004; d = 1.436),  $P_{max}$  (p = 0.000; d = 2.016), and rel  $P_{max}$  (p = 0.004; d = 1.436). There was no significant group effect (p > 0.05; d < 0.4). Significant increases in the parameters of the isokinetic strength test were found within groups (p < 0.05) ( $\blacktriangleright$  **Table 1**). HVLL showed a significant increase in mean  $P_{max}$  (p = 0.004; d = 0.370), relative mean  $P_{max}$  (p = 0.008; d = 0.445),  $P_{max}$  (p = 0.004; d = 0.314) and relative  $P_{max}$  (p = 0.015; d = 0.335). LVHL also showed a significant increase in mean  $P_{max}$  (p = 0.010; d = 0.357), relative mean  $P_{max}$ 



▶ Figure 1 Pre-Post comparison of the mean values for vLa<sub>max</sub>. left means of both groups, right means of group HVLL and LVHL (error bars show the standard deviation)

	HVLL		LVHL	
	Pre	Post	Pre	Post
meanP <sub>max</sub> (W)	385.1±74.0	416.5±94.6*	378.8±169.5	434.5±141.5+#
	(298.8-552.1)	(298.1-641.2)	(156.2–755.9)	(304.4–764.9)
relative mean P <sub>max</sub> (W·kg <sup>-1</sup> )	4.60±0.59	4.90±0.75*	4.68±1.77	5.35 ± 1.39 * #
	(3.73–5.54)	(3.86-6.00)	(3.73–5.54)	(1.80-3.71)
P <sub>max</sub> (W)	424.3±80.2	452.3±97.5*	405.6±173.2	462.2±138.7*~
	(342.3–584.8)	(329.2-686.6)	(190.7–797.2)	(330.7–780.0)
relative P <sub>max</sub> (W·kg <sup>-1</sup> )	5.08±0.69	5.33±0.80*	5.01±1.77	5.70±1.34 * #
	(4.04–6.26)	(4.26-6.43)	(2.20-8.17)	(4.02-8.03)
[La+] <sub>pre</sub> (mmol·l <sup>-1</sup> )	0.59±0.17	0.68±0.18	0.96±0.22	0.94±0.34
	(0.50–1.90)	(0.55–1.53)	(0.60–1.33)	(0.50-1.51)
vLa <sub>max</sub> (mmol·l <sup>-1</sup> ·s <sup>-1</sup> )	0.271±0.067	0.298±0.067*	0.249±0.122	0.291 ± 0.089~
	(0.133-0.371)	(0.196-0.421)	(0.034-0.465)	(0.104-0.402)
[La + ] <sub>max</sub> (mmol·l <sup>-1</sup> )	4.03±0.859	4.4±0.887*	3.61±1.09	4.53±0.611*~
	(2.21-5.16)	(2.95-6.05)	(1.54–5.33)	(3.7-6.09)

► Table 1 Result of pre-test and post-test for all estimated parameters. Data are presented as mean ± standard deviation (min-max).

(p = 0.008; d = 0.421),  $P_{max}$  (p = 0.027; d = 0.361) and relative  $P_{max}$  (p = 0.009; d = 0.440).

A correlation analysis showed that there was a significant correlation between  $P_{max}$  and  $vLa_{max}$  prior to the training intervention in LVHL (r = 0.716; p = 0.02). This was not found in HVLL (r = 0.189; p = 0.521). In addition, a significant correlation between  $\Delta vLa_{max}$ and  $\Delta P_{max}$  was found across all subjects (r = 0.502; p = 0.012).

## Discussion

The aim of this study was to investigate the effect of 6 weeks of resistance training with different training volume loads on  $vLa_{max}$ . The  $vLa_{max}$  showed a significant time effect, but a significant group effect was not found. The increases in performance were in a mean linear relationship to the adaptation of the anaerobic energy metabolism using the  $vLa_{max}$ . Thus,  $vLa_{max}$  as a physiological measure of anaerobic energy metabolism seems to be a significant performance parameter.

We presume that the significant performance increases in the isokinetic strength test are the result of more activated muscle fibers [28]. The higher number of active muscle fibers results in greater anaerobic activity, as shown by the post-test increases in [La+]. Hommel et al. [17] found a significant reduction of vLa<sub>max</sub> after 2 weeks in sprint interval training, which was stabilized up to sixth week with increased maximal performance; oxygen uptake (VO<sub>2max</sub>) was unchanged. The extent to which the resistance training changed the VO<sub>2max</sub> cannot be answered, because the maximal oxygen uptake was not measured here. Ozaki et al. showed in an overview that changes in VO<sub>2max</sub> due to resistance exercise lasting several weeks were only available for untrained subjects [29]. There were inconsistent results in relation to the training volume. In part, no changes but also slight increases in VO<sub>2max</sub> were found. Furthermore, no significant change in mitochondria enzyme activity (e.g., citrate synthase and succinate dehydrogenase) was reported following RT, which has the potential to increase  $VO_{2max}$  [30, 31]. Due to the design of our study, it is not clear which temporal dynamics the adjustment of the maximum glycolysis rate has during the training period. This would have required measurements of anaerobic performance at intervals of one or two weeks.

In a six-week study with resistance training, muscle biopsy analyses showed adaptations of the anaerobic energy metabolism and hypertrophy of the muscle fibers. Numerous enzymes of the lactacid energy metabolism (e.g., PFK, LDH) increased their activities [11]. It is assumed that the higher activity of anaerobic enzymes causes an increase in glycolysis activity. This then results in a timedependent increase in the maximum glycolysis rate. For muscular work, this means more available ATP over the loading period. An increase in glycolytic enzymes is a possible explanation for the increased maximum glycolysis rate after the training intervention in both groups. However, the increases in glycolytic enzymes, such as PFK, appear to be only marginally caused by chronic strength training [10, 32]. Oxidative and therefore also metabolic stress is considered relevant for hypertrophic muscle adaptation [33]. However, it is important to point out that other factors are also of high importance for muscle hypertrophy. These include the influence of amino acids intake, growth hormone concentrations, and mechanical stress [34].

There was a high effect size (d = 0.97) of resistance training on  $vLa_{max}$  for both groups (all subjects). Within both groups, this was considered a medium effect (d = 0.38 – 0.40). Statements about which training protocol is more effective in increasing anaerobic performance cannot be made at present. No significant differences between groups were found for  $vLa_{max}$ . The post hoc power analysis showed that a variance-analytical comparison between the two groups would have required more than 500 subjects (for p-value of 0.05; d = 0.2) to show significant differences. The data show that there was a high dispersion of  $vLa_{max}$  within LVHL, which indicates a heterogeneous study cohort. A more homogeneous subject co-

hort may also have resulted in a significant increase in group LVHL. Scott et al. [19] pointed out that heterogeneous subjects in a resistance exercise protocol are problematic. Subjects can certainly be selected based on their training status; an assignment based on glycolysis activity currently seems difficult to us.

There are currently no studies that directly examine the effects of resistance training on  $\nu La_{max}$ . However, acute studies show that lower loads with exhaustive reps lead to significantly higher metabolic stress (lactate concentration) than higher loads with few reps [35]. It was also shown that higher reps (5 sets of 10 reps) compared to lower reps (10 sets of 5 reps) at the same load also lead to increased metabolic stress. This was reflected in a stronger reduction of ATP and PCr. The significantly higher [La + ] at 5 sets of 10 reps indicates the increased stress on glycolytic enzymes [36]. A training experiment at 70% of 1RM (with high volume) and 90% of 1RM (with low volume) led to similar results [21]. In this training study, the strong [La + ] increases found in acute studies at high EV did not lead to significantly different adjustments of vLa<sub>max</sub> between EV. The variations in load and volume used here are not directly comparable with existing studies. In order to clearly show the influence of the volume load, loads above 70% of the 1RM and below 50% of the 1RM in the other group should have been chosen.

The increases in  ${\rm La}_{\rm max}$  due to training interventions are possibly the result of increased glycolysis and have already been noted in previous studies [37]. If we presume that training-related changes in anaerobic enzymes occur only to a small extent [10], it cannot be ruled out that untrained volunteers would have shown more marked adjustments to vLa<sub>max</sub>. Lactate as an intermediate product of glycolysis and its change in [La + ] over the t<sub>load</sub> is considered here in this highly intensive anaerobic test (maximum one-legged strength test) as a measure of glycolysis activity. Adjustments of the anaerobic enzyme activities that regulate [La + ] can only be speculated in this study. In the future, a <sup>31</sup>phosphorus -magnetic resonance spectroscopy (31P-MRS) may help show concentrations of energetic substrates before and after several weeks of training. It should also be mentioned that lactate concentrations determined from capillary blood are dependent on the time constant of elimination. Furthermore, the lactate formed in the muscle is transported through different compartments [38]. Thus, the lactate concentrations determined from capillary blood will be lower than the acute reactions produced by the test in the muscle under stress [39]. Furthermore, there was no control of food intake in this study, there may be influences of increased or reduced glucose intake on [La+] [40].

# Conclusions

Based on the available data, six weeks of resistance training of the lower extremities can increase anaerobic performance using the glycolysis rate. Effects of the volume load could not be determined. Thus, the effectiveness of training protocols with high or low training-volume loads on  $vLa_{max}$  cannot yet be assessed.

# Funding

This research received no external funding.

# Acknowledgements

The authors thank Andrea Ludwig for support recording data.

## Conflict of Interest

The authors declare that they have no conflict of interest.

## References

- Freitas de MC, Gerosa-Neto J, Zanchi NE et al. Role of metabolic stress for enhancing muscle adaptations: Practical applications. World J Methodol 2017; 7: 46–54. doi:10.5662/wjm.v7.i2.46
- [2] Takada S, Okita K, Suga T et al. Low-intensity exercise can increase muscle mass and strength proportionally to enhanced metabolic stress under ischemic conditions. J Appl Physiol 2012; 113: 199–205. doi:10.1152/japplphysiol.00149.2012
- [3] Yanagisawa O, Sanomura M. Effects of low-load resistance exercise with blood flow restriction on high-energy phosphate metabolism and oxygenation level in skeletal muscle. Interv Med Appl Sci 2017; 9: 67–75
- [4] Arazi H, Mirzaei B, Heidari N. Neuromuscular and metabolic responses to three different resistance exercise methods. Asian J Sports Med 2014; 5: 30–38
- [5] Rogatzki MJ, Wright GA, Mikat RP et al. Blood ammonium and lactate accumulation response to different training protocols using the parallel squat exercise. J Strength Cond Res 2014; 28: 1113–1118
- [6] Wirtz N, Wahl P, Kleinöder H et al. Lactate kinetics during multiple set resistance exercise. J Sports Sci Med 2014; 13: 73–77
- [7] Crewther B, Cronin J, Keogh J. Possible stimuli for strength and power adaptation: Acute metabolic responses. Sports Med 2006; 36: 65–78. doi:10.2165/00007256-200636010-00005
- [8] Fernandes T, Soci U, Melo S et al. Signaling pathways that mediate skeletal muscle hypertrophy: effects of exercise training. In :Cseri J, ed. Skeletal Muscle - From Myogenesis to Clinical Relations. London: InTech; 2012
- [9] Burgomaster KA, Howarth KR, Phillips SM et al. Similar metabolic adaptations during exercise after low volume sprint interval and traditional endurance training in humans. J Physiol 2008; 586: 151–160; doi:10.1113/jphysiol.2007.142109
- [10] Tesch PA. Skeletal muscle adaptations consequent to long-term heavy resistance exercise. Med Sci Sports Exerc 1988; 20: 132–134
- [11] Haun CT, Vann CG, Osburn SC et al. Muscle fiber hypertrophy in response to 6 weeks of high-volume resistance training in trained young men is largely attributed to sarcoplasmic hypertrophy. PLoS One 2019; 14: e0215267. doi:10.1371/journal.pone.0215267
- [12] Heck H, Schulz H. Diagnostics of anaerobic power and capacity. Dtsch Z Sportmed 2002; 53: 202–212
- [13] Mader A. Aussagekraft der Laktatleistungskurve in Kombination mit anaeroben Tests zur Bestimmung der Stoffwechselkapazität. In: Clasing D, Ed. Stellenwert der Laktatbestimmung in der Leistungsdiagnostik: 32 Tabellen. Stuttgart u. a.: G. Fischer. 1994: 133–152
- [14] Nitzsche N, Zschäbitz D, Baumgärtel L et al. The effect of isokinetic resistance load on glycolysis rate. In: Ferrauti A, Platen P, Grimminger-Seidensticker E et al., (Eds.) 22nd Annual Congress of the European College of Sport Science: 5–8th July 2017, Metropolis Ruhr - Germany: Book of Abstracts. Bochum: Bochumer Universitätsverlag Westdeutscher Universitätsverlag 2017; 558
- [15] Nitzsche N, Baumgärtel L, Maiwald C et al. Reproducibility of blood lactate concentration rate under isokinetic force loads. Sports 2018; 6: 1–8. doi:10.3390/sports6040150

- [16] Adam J, Ohmichen M, Ohmichen E et al. Reliability of the calculated maximal lactate steady state in amateur cyclists. Biol Sport 2015; 32: 97–102; doi:10.5604/20831862.1134311
- [17] Hommel J, Öhmichen S, Rudolph UM et al. Effects of six-week sprint interval or endurance training on calculated power in maximal lactate steady state. Biol Sport 2019; 36: 47–54. doi:10.5114/biolsport.2018.78906
- [18] ACSM Progression models in resistance training for healthy adults. Med Sci Sports Exerc 2009; 41: 687–708. doi:10.1249/ MSS.0b013e3181915670
- [19] Scott BR, Duthie GM, Thornton HR et al. Training monitoring for resistance exercise: Theory and applications. Sports Med 2016; 46: 687–698. doi:10.1007/s40279-015-0454-0
- [20] Mangine GT, Hoffman JR, Gonzalez AM et al. The effect of training volume and intensity on improvements in muscular strength and size in resistance-trained men. Physiol Rep 2015; 3: e12472. doi:10.14814/ phy2.12472
- [21] Gonzalez AM, Hoffman JR, Townsend JR et al. Intramuscular anabolic signaling and endocrine response following high volume and high intensity resistance exercise protocols in trained men. Physiol Rep 2015; 3: e12466. doi:10.14814/phy2.12466
- [22] Ogasawara R, Loenneke JP, Thiebaud RS et al. Low-load bench press training to fatigue results in muscle hypertrophy similar to high-load bench press training. Int J Clin Med 2013; 04: 114–121. doi:10.4236/ ijcm.2013.42022
- [23] Schoenfeld BJ, Ratamess NA, Peterson MD et al. Effects of different volume-equated resistance training loading strategies on muscular adaptations in well-trained men. J Strength Cond Res 2014; 28: 2909–2918. doi:10.1519/JSC.000000000000480
- [24] Harriss DJ, MacSween A, Atkinson G. Ethical standards in sport and exercise science research: 2020 update. Int J Sports Med 2019; 40: 813–817. doi:10.1055/a-1015-3123
- [25] Nitzsche N, Zschäbitz D, Baumgärtel L et al. Einfluss der Methode zur Bestimmung des alaktaziden Zeitintervalls bei isokinetischen Kraftbelastungen. In: Schwirtz A, Mess F, Demetriou Y et al., Eds. Innovation & Technologie im Sport. Hamburg: Feldhaus Edition Czwalina 2017; 215
- [26] Haff GG, Triplett NT., Eds. Essentials of strength training and conditioning. 4th ed. Champaign, IL, Windsor, ON, Leeds: Human Kinetics; 2016
- [27] Cohen J. Statistical Power Analysis for the Behavioral Sciences. 2nd ed., Hoboken: Taylor and Francis; 2013

- [28] Tesch PA, Thorsson A, Essén-Gustavsson B. Enzyme activities of FT and ST muscle fibers in heavy-resistance trained athletes. J Appl Physiol 1989; 67: 83–87. doi:10.1152/jappl.1989.67.1.83
- [29] Ozaki H, Loenneke JP, Thiebaud RS et al. Resistance training induced increase in VO2max in young and older subjects. Eur Rev Aging Phys Act 2013; 10: 107–116. doi:10.1007/s11556-013-0120-1
- [30] Bell GJ, Syrotuik D, Martin TP et al. Effect of concurrent strength and endurance training on skeletal muscle properties and hormone concentrations in humans. Eur J Appl Physiol 2000; 81: 418–427. doi:10.1007/s004210050063
- [31] Nelson AG, Arnall DA, Loy SF et al. Consequences of combining strength and endurance training regimens. Phys Ther 1990; 70: 287–294. doi:10.1093/ptj/70.5.287
- [32] Tesch PA, Alkner BA. Acute and chronic muscle metabolic adaptations to strength training. In: Komi PV, Ed. Strength and Power in Sport. Oxford, UK: Blackwell Science Ltd; 2003: 265–280
- [33] Hoppeler H, Baum O, Lurman G et al. Molecular mechanisms of muscle plasticity with exercise. Compr Physiol 2011; 1: 1383–1412. doi:10.1002/cphy.c100042
- [34] Hoppeler H. Molecular networks in skeletal muscle plasticity. J Exp Biol 2016; 219: 205–213. doi:10.1242/jeb.128207
- [35] Buitrago S, Wirtz N, Yue Z et al. Effects of load and training modes on physiological and metabolic responses in resistance exercise. Eur J Appl Physiol 2012; 7: 2739–2748
- [36] Gorostiaga EM, Navarro-Amézqueta I, Calbet JAL et al. Energy metabolism during repeated sets of leg press exercise leading to failure or not. PLoS One 2012; 7: e40621. doi:10.1371/journal.pone.0040621
- [37] Nitzsche N, Jürgens S, Schulz H. Effekte eines achtwöchigen progressiven Rope-Trainings auf die Leistungsfähigkeit der oberen Extremitäten. Ger J Exerc Sport Res 2019; 49: 493–502. doi:10.1007/ s12662-019-00587-0
- [38] Zouloumian P, Freund H. Lactate after exercise in Man: II. Mathematical model. Eur J Appl Physiol Occup Physiol 1981; 46: 135–147
- [39] Bangsbo J, Johansen L, Graham T et al. Lactate and H + effluxes from human skeletal muscles during intense, dynamic exercise. J Physiol 1993; 462: 115–133
- [40] Mikulski T, Ziemba A, Nazar K. Influence of body carbohydrate store modification on catecholamine and lactate responses to graded exercise in sedentary and physically active subjects. J Physiol Pharmacol 2008; 59: 603–616