Low-Load Bench Press Training to Fatigue Results in Muscle Hypertrophy Similar to High-Load Bench Press Training

Riki Ogasawara¹,², Jeremy P. Loenneke³, Robert S. Thiebaud³, Takashi Abe¹,⁴
¹Graduate School of Frontier Sciences, University of Tokyo, Kashiwa, Japan; ²College of Sport and Health Science, Ritsumeikan University, Kusatsu, Japan; ³Department of Health and Exercise Science, University of Oklahoma, Norman, USA; ⁴Department of Health, Exercise Science, & Recreation Management, University of Mississippi, Oxford, USA.

Email: t12abe@gmail.com

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ABSTRACT

The purpose of this study was to determine whether the training responses observed with low-load resistance exercise to volitional fatigue translates into significant muscle hypertrophy, and compare that response to high-load resistance training. Nine previously untrained men (aged 25 [SD 3] years at the beginning of the study, standing height 1.73 [SD 0.07] m, body mass 68.9 [SD 8.1] kg) completed 6 weeks of high load-resistance training (HL-RT) (75% of one repetition maximal [1RM], 3-sets, 3x/wk) followed by 12 months of detraining. Following this, subjects completed 6 weeks of low load-resistance training (LL-RT) to volitional fatigue (30% 1 RM, 4 sets, 3x/wk). Increases (p < 0.05) in magnetic resonance imaging-measured triceps brachii and pectoralis major muscle cross-sectional areas were similar for both HL-RT (11.9% and 17.6%, respectively) and LL-RT (9.8% and 21.1%, respectively). In addition, both groups increased (p < 0.05) 1RM and maximal elbow extension strength following training; however, the percent increases in 1RM (8.6% vs. 21.0%) and elbow extension strength (6.5% vs. 13.9%) were significantly (p < 0.05) lower with LL-RT.

Both protocols elicited similar increases in muscle cross-sectional area, however differences were observed in strength. An explanation of the smaller relative increases in strength may be due to the fact that detraining after HL-RT did not cause strength values to return to baseline levels thereby producing smaller changes in strength. In addition, the results may also suggest that the consistent practice of lifting a heavy load is necessary to maximize gains in muscular strength of the trained movement. These results demonstrate that significant muscle hypertrophy can occur without high-load resistance training and suggests that the focus on percentage of external load as the important deciding factor on muscle hypertrophy is too simplistic and inappropriate.

Keywords: Bench Press; Training Intensity; Muscle CSA; MRI; Strength

1. Introduction

As a muscle is overloaded from increased mechanical work, the added stress increases skeletal muscle amino acid transporter expression [1], which in turn enhances the synthesis of the contractile proteins, actin and myosin [2]. These acute positive balances between muscle protein synthesis (MPS) and muscle protein breakdown (MPB) lead to skeletal muscle hypertrophy over time which occurs from both an increase in the thickness and number of myofibrils [see molecular pathway review by Adams [3]. Although skeletal muscle hypertrophy occurs in both slow twitch (ST) and fast twitch (FT) fibers, the latter has the greatest potential for growth [4]. Therefore it is been hypothesized that skeletal muscle hypertrophy can occur independent of exercise load, as long as FT fibers are activated [5,6].

Conventional thought is that at least 70% of one’s repetition maximum (1 RM) must be lifted repeatedly to observe a meaningful increase in muscular size [7]. However, acute molecular research indicates that external exercise load may be of less importance when adequate volume of resistance exercise is completed. To illustrate, when four sets of resistance exercise was performed at 30% 1 RM to volitional fatigue, myofibril MPS was elevated to the same level as 90% 1 RM to volitional fatigue (not work matched) [8]. This is contrary to what has commonly been reported in the literature which states that training to volitional fatigue is not an effective stimulus unless a sufficient external load as defined by percentage of 1 RM (~80% 1 RM) is lifted. The common thought has always been that higher repetition training...
cannot produce a stress that is adequate enough to recruit and fatigue the highest threshold motor units [9].

Interestingly, Campos et al. [10] provide the only evidence to date that resistance exercise to volitional fatigue at higher loads is more effective than training at lower loads for skeletal muscle hypertrophy (4 sets 3 -4 RM vs. 2 sets 20-28 RM). However, using the identical methods of Campos et al. [10], Leger et al. [11] observed significant increases in muscle hypertrophy, muscular strength, and endurance independent of the external load lifted. One possible reason for the difference could be due to the older less active subjects used in latter study (36 vs 22 yrs). In addition, the volume of exercise (2 sets) may have been inadequate to recruit the higher threshold motor units in the younger more active subjects used in the Campos et al. [10] paper.

The aforementioned evidence has led to the formation of the metabolite/volume threshold theory [5]. This theory states that, assuming an adequate exercise volume is achieved, the recruitment of FT fibers appears to be the large driving force of skeletal muscle hypertrophy whereas the external load lifted and systemic endogenous hormone elevations may not be as important as previously thought [12,13]. Much of this theory was based on acute myofibril MPS and it is acknowledged that although these acute studies are hypothesized to be predictive of chronic adaptations, they are not definitive as incongruences may exist between the acute and chronic changes following resistance training [14,15]. Therefore, the purpose of this study was to determine whether the training responses observed with low-load resistance exercise to volitional fatigue translates into significant muscle hypertrophy, and compare that response to high-load resistance training. Low load knee extensor exercise to fatigue has shown that muscle hypertrophy (whole muscle and fiber level) occurs at levels similar to higher loads [16], however it is currently unknown whether this is also true for upper body resistance exercise. Bench press is one of the major exercises for developing the upper body, however, very few studies report muscle size changes in the chest and upper arm following a single mode of high-load bench press training [17,18]. In the present study, within subject experimental design was chosen to reduce biological variability. Further, due to possible differences in systemic endogenous hormones with each loading scheme and the cross-training neural adaptations associated with a unilateral training model [19], each subject completed both exercise protocols separated by over a year (12 months). All subjects began with high-load resistance training as this design also allowed us to investigate the muscle size and strength changes to one year of detraining with traditional high load exercise. Although the order of training was not randomized, it increased our statistical power to investi-

### Table 1. Physical characteristics of the subjects.

|                | Height (m) | Body mass (kg) | Body mass index (kg/m²) |
|----------------|------------|----------------|------------------------|
| HL-RT pre      | 1.73       | 68.9 (8.1)     | 23.0 (2.8)             |
| HL-RT post     | 69.5 (8.5) | 23.2 (2.8)     |
| LL-RT pre      | 1.74       | 68.8 (8.0)     | 22.9 (2.8)             |
| LL-RT post     | 69.4 (7.9) | 23.1 (2.5)     |

HL-RT, high-load resistance training; LL-RT, low-load resistance training; *p < 0.05, pre vs. post.
sec per muscle action. Training load was adjusted to the new 1RM determined at 3 weeks in both training protocols. For the HL-RT, if subjects were able to perform 12 repetitions or more during a training session, the training load was increased ~5% for the next training session. To ensure adequate training load, all training sessions were surveyed and supervised by trained personnel. All subjects successfully completed every training session.

2.3. Measurements Schedule

Subjects testing took place before the start of the study (pre) and 3 - 4 days after (post) the 6-week training period. The magnetic resonance imaging (MRI) measurement was obtained between 16:00 and 19:00 hours. The strength measurement was determined on the same day or the following day after the MRI measurement. All measurements were balanced for the time of day.

2.4. Strength Measurement

All subjects completed 2 - 3 familiarization sessions to receive instruction on proper technique and to practice the 1 RM and maximal voluntary isometric strength (MVC) tests. The 1RM was assessed with the free-weight bench press exercise. The 1 RM was determined by progressively increasing the weight lifted until the subject failed to lift the weight through a complete range of motion. Usually 5 trials were required to complete a 1 RM test. Adequate amount of recovery time was permitted between 1RM trials (3 - 5 min) [20]. MVC of the elbow extensors (right arm) was measured by using an isokinetic dynamometer (Biodex System 3, Biodex Medical Systems Inc., Shirley, NY, USA). The subjects were comfortably seated on a chair and the arm was positioned on a firm and stable table at chest level with an elbow joint angle of 90˚ (0˚ at full extension). The upper arm was maintained in the horizontal plane while the subject’s wrist was fixed at the end of the lever arm in a position halfway between supination and pronation. The elbow extensor force was measured with a transducer, while a diagonal strap was secured over the elbow to maintain a stationary position during the MVC. Subjects were instructed to contract as fast and forcefully as possible. MVC was measured twice. If MVC torque for the first two MVCs varied by >5%, up to two additional MVCs were performed. Each effort was held for ~5 s. The coefficient of variation (CV) for this measurement from test to retest was 3.1% [21]. Both MVC and 1RM tests (same day and about 20 min apart between two tests) were performed before training and after 3 and 6 weeks of training.

2.5. Muscle Size Measurements

Multi-slice MRI images of the upper arm and chest were obtained using a MRI scanner (General Electric Yokogawa Signa 0.2-T, Milwaukee, WI, USA). A T1-weighted, spin-echo, axial plane sequence was performed with a 520 ms repetition time and a 20 ms echo time. Subjects rested quietly in the magnet bore in a supine position with their arms extended. The lateral epicondyte of the humerus was used as the origin point, and continuous transverse images with 1.0 cm slice thickness (0.2 cm interslice gap) were obtained from the lateral epicondyle of the humerus to the acromial process of the scapula for each subject (Figure 1). All MRI data were transferred to a personal computer for analysis using specially designed image analysis software (TomoVision Inc., Montreal, Canada). For each slice, skeletal muscle tissue cross-sectional area (CSA) was digitized. Triceps brachii (TB) and pectoralis major (PM) muscle CSA of 3 continuous slices for the muscle belly were averaged to represent a single data point for statistical analysis, respectively. We have previously determined that the CV of this measurement was less than 1% [21].

2.6. Statistical Analysis

All values are expressed as mean [SD]. TB and PM muscle CSA, 1RM, MVC data were analyzed using two-way ANOVA with repeated measures (group × time). Post hoc testing was performed using Tukey-Kramer when appropriate. Pre-training values of each training protocol were compared using a paired t-test. Pearson product-moment correlation coefficients determined the association between high-load and low-load hypertrophy changes in TB and PM muscle CSA. Significance was set at p < 0.05. All analyses were performed using JMP statistical software version 8.0 (SAS Institute, Cary, NC, USA).

Figure 1. Typical magnetic resonance imaging image showing transverse scan of the chest.
3. Results

There was no difference in body weight at pre-training between HL-RT (68.9 [8.1] kg) and LL-RT (68.8 [8.0] kg). After 6-week of training, body weight increased ($p < 0.05$) by 0.6 kg in the HL-RT and 0.6 kg in the LL-RT. During the LL-RT protocol, the average total number of repetitions for each exercise session was 141 [14].

Following 6 weeks of training, 1 RM and MVC strength increased ($p < 0.05$) significantly in both HL-RT and LL-RT protocols. However, the percent increases in strength were lower ($p < 0.05$) in the LL-RT (1 RM 8.6 [2.9]%, MVC 6.5 [4.9]%) than in the HL-RT (1 RM 21.0 [5.9]%, MVC 13.9 [7.5]%) (Figure 2). Before the start of the LL-RT, 1-RM and MVC strength had not returned to pre-training HL-RT 1-RM and MVC strength levels (Figure 2).

At the start of training, muscle CSA in the PM was the same between the HL-RT and LL-RT protocols, whereas muscle CSA in the TB was 2.2% higher ($p = 0.03$) in LL-RT than in HL-RT. The TB muscle CSA increased ($p < 0.01$) following LL-RT and HL-RT and the percent increase in muscle CSA was similar between the two training protocols (LL-RT 9.8 [4.6]%, HL-RT 11.9 [2.6]%) (Figure 3(a)). Similarly, absolute and relative increases ($p < 0.01$) in PM muscle CSA were similar between HL-RT and LL-RT (Figure 3B). A significant correlation was observed between percent increase in muscle CSA following HL-RT and LL-RT in the TB and PM muscles (Figure 4).

Figure 2. Changes in maximum dynamic (bench press one repetition maximum) and isometric (elbow extension) strength following 6 weeks of high-load (HL-RT) and low-load (LL-RT) resistance training. Pre, before training; wk3, after 3 weeks; Post, after 6 weeks. * $p < 0.05$ vs. pre-training, † $p < 0.05$ vs. HL-RT.

Figure 3. Changes in muscle cross-sectional area (CSA) in the triceps brachii (TB) and pectoralis major (PM) muscles following 6 weeks of high-load (HL-RT) and low-load (LL-RT) resistance training. Pre, before training; Post, after 6 weeks. * $p < 0.05$ vs. pre-training, † $p < 0.05$ vs. HL-RT.
4. Discussion

This study found that 1) LL-RT to volitional fatigue and HL-RT results in similar levels of skeletal muscle hypertrophy in the upper body and 2) significant correlations in the degree of muscle hypertrophy between LL-RT to volitional fatigue and HL-RT. This data suggests that skeletal muscle hypertrophy can occur independent of a higher load in the upper body as long as there is adequate exercise volume. In addition, one year of detraining from HL-RT results in a complete loss of muscle size, however muscle strength was decreased but still elevated above the pre-training level.

4.1. Muscle Hypertrophy

Six weeks of high-load (75% 1 RM) resistance training resulted in significant skeletal muscle hypertrophy. Interestingly, after 12 months of detraining the same subjects then performed low-load resistance training to volitional fatigue and found similar increases in skeletal muscle hypertrophy compared to that observed with high-load training. This is contrary to previous research [9,10] and recommendations [7] that report higher-loads to be superior. However, the research in which those recommendations were largely based were matched for work and it appears that in order for low-loads to increase muscle hypertrophy to levels similar to high-loads, exercise must be taken to volitional fatigue [5].

This study confirms acute research from Burd et al. [8] who found similar increases in myofibril MPS independent of exercise load when exercise was taken to volitional fatigue. This might be related to the significant increase in muscle time under tension when repetitions are taken to volitional fatigue as this has recently been found to be an important variable in the synthetic response [2]. In addition, MPS from resistance training occurs primarily from the activation of signaling proteins, primarily S6K1, which are approximately 3 to 4-fold higher in FT fibers compared to ST [22]. Furthermore, phosphorylation of this signaling protein has shown to be predictive of skeletal muscle hypertrophy [23]. This suggests that skeletal muscle hypertrophy occurs independent of a higher exercise load, as long as FT fibers are activated from sufficient exercise volume [5,6]. It is acknowledged that the protein degradation response to low-load resistance training to volitional fatigue is not known, as research is typically completed under the assumption that synthesis rates and not degradation rates are more responsive to resistance exercise in healthy humans [24]. The similar levels of muscle hypertrophy between protocols suggest that this assumption is likely true for the upper body. This also supports recent research completed in the lower body, which found significant muscle hypertrophy with low load (30% 1 RM) knee extensor exercise to fatigue [16].

Interestingly, it should be mentioned that rodent data suggest that the myonuclei gained from resistance training are not lost following 3 months of detraining [25]. This has led some to speculate that this retention of myonuclei is important in the “muscle memory” response to exercise. Therefore, if one is trained following the cessation of training, it might be possible that the rebound in muscle hypertrophy is due to the myonuclei that were added with training and maintained through muscle atrophy. It is currently unknown how this translates to humans or how long this effect lasts, but we cannot rule out the possibility that this may be playing some role in the equal response between variables.

The percentage increases in muscle hypertrophy for the TB and PM were larger than what has been previously reported for the lower body. Unfortunately, the molecular mechanisms for upper body muscle hypertrophy are currently under studied when compared with what is known for the lower body. However, the results of the present investigation suggest that heavy resistance exercise induced activation of muscle protein metabolism may be more responsive in the upper body compared to the lower body. To illustrate, Seynnes et al. [26] observed a 7% increase in quadriceps femoris CSA following 35 days of lower body bilateral knee extensions. In
addition, Abe et al. [20] observed after a 6 week total body workout (70% 1 RM), that the quadriceps muscle thickness increased 5%, however the PM and TB increased 13% and 9%, respectively. Furthermore, using a MRI, muscle CSA increased 16% in the PM and 10% in the TB following 18 days of bench press training (75% 1RM) [27]. Yasuda et al. [28] also observed that 18 days of bench press training (75% 1 RM) resulted in an 18% increase in PM and a 10% increase in the TB. The current findings are in agreement with the previous research in the upper body which suggests that the upper body may have a higher capacity for muscle hypertrophy than the lower body.

4.2. Muscular Strength

Changes in strength between the LL-RT and HL-RT are another interesting finding from this study. Both groups had significant increases in strength following training; however, the percent increases in strength were significantly lower in the LL-RT protocol. An explanation of the smaller relative increases in strength may be due to the fact that detraining after HL-RT did not cause strength values to return to baseline levels thereby producing smaller changes in strength. Although subjects were told to return back to their pre-training lifestyle, it is possible that subjects maintained a level of activity high enough to maintain strength but not muscle mass. Further, it is possible that the neural adaptation to resistance exercise is longer lasting than the hypertrophic response. Indeed, there is evidence to support the finding that strength does not return to baseline levels despite detraining. In young women who did 20 weeks of strength training and then detrained for 30 - 32 weeks, strength levels significantly decreased but did not return to pre-training levels [29]. In addition, Bickel et al. [30] found that after 16 weeks of lower body training and 32 weeks of detraining that strength significantly decreased by 7% but remained 23% above baseline. Another study in older adults found that 2 years of training followed by 3 years of detraining produced significant decreases in dynamic strength but levels remained slightly above baseline values and significantly higher than control subjects [31]. Although the reasons for this maintenance of strength are unknown from the present investigation, it is possible that following detraining there was a partial maintenance of the increased volitional drive from the supraspinal center which may have maintained part of the increased muscle activation likely gained from HL-RT [32]. Therefore, the lower amounts of strength observed in the LL-RT group compared to the HL-RT may be more a function of the training effect rather than the intervention itself. Also, all subjects began training with high-load resistance training and finished with low-load training, therefore it remains unknown if the same strength effects would be observed if the protocols were reversed. Lastly, an alternative explanation is that the specificity of training may dictate the overall maximal gains in strength. For example, the results may suggest that the consistent practice of lifting a heavy load is necessary to maximize gains in muscular strength of the trained movement.

5. Conclusion

This study verifies that similar degrees of muscle hypertrophy can occur in the upper body independent of a high external load, provided enough muscular work is completed. This data seems to support that the acute myofibril MPS responses previously observed with LL-RT to fatigue do translate to chronic training adaptation. These results demonstrate that significant muscle hypertrophy can occur without high-load resistance training and suggests that the focus on percentage of external load as the important deciding factor on muscle adaptation (i.e. muscle hypertrophy) is too simplistic and inappropriate.

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