Elevations in ostensibly anabolic hormones with resistance exercise enhance neither training-induced muscle hypertrophy nor strength of the elbow flexors

Daniel W. D. West, Nicholas A. Burd, Jason E. Tang, Daniel R. Moore, Aaron W. Staples, Andrew M. Holwerda, Steven K. Baker, and Stuart M. Phillips

1Exercise Metabolism Research Group, Department of Kinesiology; and 2Department of Medicine, McMaster University, Hamilton, Ontario, Canada

Submitted 7 October 2009; accepted in final form 6 November 2009

Hormones such as testosterone, growth hormone (GH), and insulin-like growth factor-1 (IGF-1) are important for skeletal muscle anabolism during growth and development (e.g., 23). It has been suggested that acute elevation of this triad of hormones that occurs after performance of intense resistance exercise is a significant contributor to the gains in strength and hypertrophy that are observed with resistance training (14, 19, 24, 30). The magnitude of the increase in concentration of these hormones is largely dependent on parameters of the resistance exercise. Specifically, large elevations of systemic hormones are observed acutely postexercise after resistance exercise bouts consisting of a high amount of work (8, 9, 28). Additionally, higher intensity exercise (16, 19) with short rest intervals (4, 17) that is performed with large muscle groups (10, 19) are also associated with large rises in these hormones. In fact, training principles have been constructed to maximize the postexercise rise in these hormones based on the interpretation that exercise-induced increases in systemic hormones like testosterone and GH will enhance muscle hypertrophy (15, 27, 31). Specifically, it has been suggested that small exercising muscle groups (e.g., biceps brachii), which are incapable of inducing large increases in systemic anabolic hormones when used in isolation, should be trained concurrently with large exercising muscle masses that can elevate testosterone and GH (10, 19). However, the hypothesis that exercise-induced rises in testosterone and GH enhance increases in strength and muscle hypertrophy with training remains untested.

We have previously demonstrated that acute changes in muscle protein synthesis following resistance exercise and feeding (38) can qualitatively predict gains in muscle fiber cross-sectional area (CSA) and lean body mass (11). Recently, we have shown that exercise-induced hormones that are hypothesized to be anabolic and affect hypertrophy do not acutely enhance fed-state myofibrillar protein synthesis after acute resistance exercise (36). Despite the ability of the postexercise protein synthetic response (38) to predict phenotypic adaptations to training (11), it is unknown whether the synthetic responses that we observed after a single acute bout of exercise (36) would ultimately translate into similar increases in muscle size and strength when hormone availability is manipulated following repeated bouts of resistance exercise (i.e., training). Thus the aim of our present study was to determine if increases in muscle strength and hypertrophy are enhanced by exercise-induced increases in hormone availability with resistance training. Based on our previous acute finding that myofibrillar protein synthesis was not enhanced by acute elevations in circulating testosterone, GH, and IGF-1, we hypothesized that repeatedly elevating the acute postexercise availability of these hormones would not enhance the muscle hypertrophy and strength gains achieved from a progressive resistance training program. Previous work demonstrating an effect of elevated endogenous hormones on gains in isometric strength has used a similar design (10). Thus, as a reasonable proof of principle study, if acute rises in GH, IGF-1, and testosterone were to result in differential phenotypic changes, then an inherently superior design for testing this thesis is a unilateral within-person design, where interindividual differences in potential for hypertrophy and strength gains, which are substantial (12), are minimized.
HORMONES AND MUSCLE HYPERTROPHY IN HUMANS

METHODS

Subjects. Twelve healthy young men (21.8 ± 0.4 yr, 1.78 ± 0.02 m, 74.1 ± 3.3 kg; means ± SE) volunteered to participate in the study after being informed of the procedures and potential risks involved in the investigation. Subjects were recreationally active with no formal weightlifting experience. Participants provided consent to an agreement that was approved by the Research Ethics Board of Hamilton Health Sciences and that was written in accordance with standards set by the Declaration of Helsinki.

Experimental protocol. Using a within-person design, participants trained each arm on separate days under two different hormonal environments for 15 wk. In the low hormone condition (LH), one arm performed arm curl exercise only, while in the high hormone condition (HH) the contralateral arm performed the same arm curl exercise followed immediately by a bout of leg resistance exercises designed to elicit large increases in circulating hormones. In weeks 1–6, participants trained each arm three times over 2 wk; they trained in a manner that allowed 72 h between LH and HH training days (e.g., week 1: Monday*, Tuesday, Friday; week 2: Monday, Thursday*, Friday; HH session days shown with asterisk). This approach was taken to ensure that the enhanced muscle protein synthetic response, which can be elevated in the untrained state for ~48 h (22, 26), occurred exclusively on the background of basal hormone concentrations during LH before the hormonal spike that was associated with exercise on the HH day 72 h later. In weeks 7–15, an extra training session was added to enhance the training stimulus so that each arm trained twice per week with at least 48 h following each arm-only trial (e.g., week 7: Monday*, Tuesday, Thursday*, Friday; HH session days shown with asterisk). We have previously shown that resistance training shortens the duration (i.e., <28 h) for which muscle protein synthesis is elevated after exercise (33). Therefore, despite the inclusion of an additional training session per week, the attenuated time course of muscle protein synthesis after exercise in a more trained state would still have provided enough recovery to ensure that the LH arm was not exposed to the hormonal milieu of HH during the skeletal muscle remodeling process. Participants consumed 18 g of whey protein immediately before exercise and 18 g at 90 min after arm exercise in each condition to support maximal rates of muscle protein synthesis both in the presence and absence of elevated hormone concentrations as well as to reduce variability in nutrition surrounding the exercise bout.

Training. Subjects were familiarized with each exercise before strength testing and beginning the training program. Exercise in LH consisted of three to four sets of 8–12 repetitions at a load that was ~95% of their 10-repetition maximum (RM) such that voluntary failure occurred during the final set. Exercise in HH was performed in the contralateral arm and consisted of identical arm exercise to LH but added to enhance the training stimulus so that each arm trained twice per week with at least 48 h following each arm-only trial (e.g., week 1: Monday*, Tuesday, Friday; week 2: Monday, Thursday*, Friday; HH session days shown with asterisk). This approach was taken to ensure that the enhanced muscle protein synthetic response, which can be elevated in the untrained state for ~48 h (22, 26), occurred exclusively on the background of basal hormone concentrations during LH before the hormonal spike that was associated with exercise on the HH day 72 h later. In weeks 7–15, an extra training session was added to enhance the training stimulus so that each arm trained twice per week with at least 48 h following each arm-only trial (e.g., week 7: Monday*, Tuesday, Thursday*, Friday; HH session days shown with asterisk). We have previously shown that resistance training shortens the duration (i.e., <28 h) for which muscle protein synthesis is elevated after exercise (33). Therefore, despite the inclusion of an additional training session per week, the attenuated time course of muscle protein synthesis after exercise in a more trained state would still have provided enough recovery to ensure that the LH arm was not exposed to the hormonal milieu of HH during the skeletal muscle remodeling process. Participants consumed 18 g of whey protein immediately before exercise and 18 g at 90 min after arm exercise in each condition to support maximal rates of muscle protein synthesis both in the presence and absence of elevated hormone concentrations as well as to reduce variability in nutrition surrounding the exercise bout.

Magnetic Resonance Imaging (MRI). MRI analyses were performed on a 1.5-T extremity scanner (OrthoOne, ON Medical Systems, Wilmington, MA) at the MRI Research Facility of the National Institute of Arthritis and Musculoskeletal and Skin Diseases (Bethesda, MD) to quantify CSA. Each scan was acquired using a surface receive coil and a T2-weighted turbo spin-echo sequence (TR = 2000 ms; TE = 50 ms; matrix = 256 × 256; field of view = 200 × 200 mm; slice thickness = 2 mm; number of excitations = 1). The volume of muscle CSA was determined from the images by using software specific for MRI analysis (OCT module, Inovance, Leuven, Belgium).

Blood samples were taken following the third as well as the final training session to characterize the hormonal response elicited by each condition (LH and HH) both early and late in the training period. Blood samples were analyzed for serum cortisol, testosterone, GH, dehydroepiandrosterone sulfate (DHEA-S) and IGF-1 using solid-phase, two-site chemiluminescence immunometric assays (Innolite; Intermedico, Hollliston, MA). All intra-assay coefficients of variation for these hormones were below 5%, and all assays included standards and daily quality assurance procedures. Free testosterone was calculated from total testosterone and sex hormone binding globulin (21). Lactate was measured on neutralized deproteinized whole blood using an enzymatic-colorimetric assay kit (Pointe Scientific, Canton, MI). Plasma insulin concentration was determined using Coat-a-Count insulin kits (Diagnostic Products, Los Angeles, CA). Blood amino acids were analyzed as previously described (25). Hematocrit was measured in triplicate using standard microcapillary methods. Change in plasma volume was calculated from hemoglobin concentration (cyanmethemoglobin technique, Pointe Scientific) and hematocrit using equations by Dill and Costill (7).

Statistics. This study was a within-subject repeated-measures design. Blood lactate and serum hormone concentrations were analyzed isometric maximal voluntary contraction (MVC) strength was tested using a custom dynamometer. To isolate the elbow flexors as the sole producers of force, participants were seated with their upper arm abducted in the horizontal plane and with a pad clamped firmly down on the top of their shoulder. The forearm was supinated and tightly fastened using straps on an aluminum plate that was attached to a steel shaft. The participant’s arm was positioned at 120° (180° = full extension) with the elbow visually aligned with the axis of rotation. A strain gauge was used to detect torque produced by
using three-factor repeated-measures ANOVA statistics with training (pre and post), condition (LH and HH), and time (preexercise and 0, 15, 30, and 60 min postexercise) as within-subject factors. Two-factor (training × condition) repeated-measures ANOVA was used to analyze area under the curve data; two-factor (time × condition) repeated-measures ANOVA was used to analyze hematocrit and plasma volume change early in the training period. One-factor (time) repeated-measures ANOVA was used to analyze mean insulin and amino acid concentrations across training in each condition. Hypertrophy and strength data were analyzed using two-factor (training × condition)

Fig. 1. Whole blood lactate (A) and serum growth hormone (GH; B), IGF-1 (C), total testosterone (D), free testosterone (E) and cortisol (F) concentrations at rest and after low hormone (LH) and high hormone (HH) exercise protocols. Insets: net area under the curve (AUC; rest = 0); closed bars, HH; open bars, LH. Significantly greater than LH for corresponding time points and for AUC, *P < 0.01, †P < 0.001. Values are means ± SE.
Hematocrit increased, and plasma volume decreased, to a basal levels or in the pattern of response (data not shown).

Compared with LH and with no training-induced changes in other hormones with a significant rise at 0 and 15 min in HH training period. DHEA-S exhibited a pattern broadly similar to attenuation of the cortisol response after HH at the end of the after LH and were markedly elevated after HH; there was an availability was present both at the beginning and end of 15 wk /H11021

Table 1. Hematocrit and change in plasma volume

<table>
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<tr>
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<th>0</th>
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<tr>
<td>Hematocrit, %</td>
<td>HH</td>
<td>43.7 ± 1.2</td>
<td>49.5 ± 1.0†</td>
<td>46.2 ± 1.6†</td>
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<td>LH</td>
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<td>46.0 ± 0.9</td>
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<td>Plasma volume, %Δ</td>
<td>HH</td>
<td>−19.8 ± 1.0†</td>
<td>−9.1 ± 2.2</td>
<td>−5.6 ± 1.4</td>
<td>−5.8 ± 1.9</td>
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<tr>
<td></td>
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<td>−4.1 ± 1.2</td>
<td>−3.2 ± 2.2</td>
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Values are means ± SE. Hematocrit: there was a greater increase in hematocrit after the high hormone (HH) protocol (condition × time interaction, P < 0.001). Different from low hormone (LH) at same time: †P < 0.05; ‡P < 0.01; ‡P < 0.001. Plasma volume: there was a greater decrease in plasma volume after the HH protocol (condition × time interaction, P < 0.001). †Different from LH at same time, P < 0.01.

Results

Blood analyses. LH elicited a minimal rise in lactate concentration (~2 mmol/l at peak), whereas HH elicited a marked increase (~10.5 mmol/l at peak); nearly identical responses were elicited before and after training within each condition (Fig. 1A). Figure 1, B–E, shows the hormone responses of GH, IGF-1, and total and free testosterone, respectively, after each condition during the first and last weeks of training. There was no change in the concentration of any of these hormones after exercise in LH. In contrast, there was a marked increase in GH, IGF-1, and total and free testosterone that peaked ~15 min after HH (P < 0.001); this divergent pattern in hormone availability was present both at the beginning and end of 15 wk of training. Cortisol concentrations were similar to basal levels after LH and were markedly elevated after HH; there was an attenuation of the cortisol response after HH at the end of the training period. DHEA-S exhibited a pattern broadly similar to other hormones with a significant rise at 0 and 15 min in HH compared with LH and with no training-induced changes in basal levels or in the pattern of response (data not shown). Hematocrit increased, and plasma volume decreased, to a greater extent after HH compared with after LH (Table 1).

Insulin concentration peaked 30 and 45 min after the pre-exercise protein drink in LH and HH, respectively (Table 2). Amino acids appeared in the blood at a similar rate. In LH, amino acids reached a plateau and concentrations remained elevated at 75 min postdrink (P < 0.001). In HH, amino acids peaked 30 min postdrink, and then declined, before appearing to reach a steady-state above baseline 90 min postdrink (P < 0.05; Table 2). Note that the values in Table 2 do not reflect the second 18-g protein drink that was given 90 min after arm exercise in each condition.

Strength. Isometric strength increased 20 ± 4% (range: 3–49%) in LH and 19 ± 3% (range: 2–34%) in HH with training (P < 0.001, Fig. 2A), but there were no differences between conditions (condition × training interaction, P = 0.65). Similarly, 1 RM increased 23 ± 6% (2–56%) in LH and 25 ± 5% (2–53%) in HH (P < 0.001, Fig. 2B) with no effect of condition (P = 0.43). The 10 RM increased 46 ± 3% (33–62%) in LH and 47 ± 6% (20–100%) in HH (P < 0.001, Fig. 2C), but there were no differences between conditions (P = 0.63).

Muscle fiber and elbow flexor CSA. Type I muscle fiber CSA increased 9 ± 3% (0–16%) in LH and 11 ± 4% (0–26%) in HH with training (P < 0.01, Fig. 3A) while type II muscle fiber CSA increased 21 ± 4% (8–42%) in LH and 24 ± 6% (9–53%) in HH (P < 0.001, Fig. 3B); there were no differences between conditions for either fiber type (condition × training interaction, P = 0.2).
we found no differences in the increases in strength or hyper-trophy in muscle exercised under low or high hormone condi-
tions after 15 wk of resistance training. These findings are in
agreement with our hypothesis and previous work showing that
exercise-induced hormone elevations do not stimulate myofi-
brilla protein synthesis (36) and are not necessary for hyper-
trophy (37). Thus our data (36 and present observations), when
viewed collectively, lead us to conclude that local mechanisms
are of far greater relevance in regulating muscle protein accre-
tion occurring with resistance training and that acute changes
in hormones, such as GH, IGF-1, and testosterone, do not
predict or in any way reflect a capacity for hypertrophy.

We found no additional strength improvements in the arm
trained under the HH condition, which is in contrast to a study
(10) that is commonly cited (e.g., 14, 18–20, 30, 31) to support
the thesis that exercise-induced increases in hormone availabil-
ity enhance training adaptations. However, the finding of
greater isometric strength gains due to physiological hormone
elevations in the previous study (10) may have been related to
differences in baseline strength between different groups train-
ing with either low or high hormone concentrations. In the
present study there was no difference between arms in any of
our measures of strength before or after training. Moreover, in
agreement with our data, Hansen and co-workers (10) reported
no effect of exercise-induced rises in hormone levels on dy-
namic strength gains with training. Since our study was a
within-subject design, we can expect that a portion of the
increase in strength was due to cross-education effects, i.e.,
increased strength in the limb that is contralateral to the
training limb, which is ~7% of initial strength in the elbow
flexors (5). Additionally, any contribution to increased strength
due to cross-education would have likely benefited both arms
similarly and is a reflection of increased motorneuron output
rather than muscular adaptations (6).

In addition to measuring changes in strength, we quantified
whole muscle and fiber CSA to determine if exercise-induced
elevations in circulating hormones enhances muscle hypertro-
phy with resistance training. We found no effect of elevated
hormones on the degree of muscle hypertrophy measured by
either MRI or histochemical staining with 15 wk of training.
Instead, our data showed virtually identical increases in both
muscle fiber and whole muscle CSA regardless of hormonal
condition. These findings agree with the notion that local and
not systemic factors are responsible for initiating signaling
responses in type I and II fibers in response to resistance
exercise (34).

By strictly supervising each session of a progressive training
program, we successfully maintained divergent hormone pro-
files in the acute recovery period after exercise. That is, LH and
HH elicited low and high hormone (and lactate) responses,
respectively, at both the beginning and through to the end of
training. The mechanisms that drive increases in exercise-
induced hormones, as well as the biological relevance of
exercise-induced hormones, are unclear but are more likely
related to metabolic stress and/or fuel mobilization rather than
muscle anabolism. It is important to note that the transient
physiological increase in hormone availability that can occur
after resistance exercise is in contrast to the continued marked
elevation that is observed with pharmacological administra-
tion, which can have an effect on overall muscle mass depend-
ing on the hormone. For example, supraphysiological doses of
testosterone are clearly potently anabolic to skeletal muscle (3)

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**DISCUSSION**

In the present study, we were able to effectively manipulate
endogenous hormone concentrations so that, during the acute
postexercise time period when amino acids were readily avail-
able for protein synthesis, one arm was repeatedly exposed to
marked increases in GH, IGF-1, as well as total and free
testosterone concentration. The contralateral arm was exposed
to only basal levels of these hormones. Despite vast differences
in hormone availability in the immediate postexercise period,
we found no differences in the increases in strength or hyper-

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**Fig. 2.** Maximal voluntary contraction (MVC; A), one-repetition maximum (1 RM; B), and 10-repetition maximum (10 RM; C) before and after training in LH and HH. *Main effect of training, $P < 0.001$; there were no interactions (training $\times$ condition) for any strength measure ($P = 0.65, 0.43, 0.63$ for MVC, 1 RM and 10 RM, respectively). Values are means $\pm$ SE.
whereas GH supplementation has little effect on the exercise-induced increase in muscle mass (39). Furthermore, there is evidence that a minimal basal level of testosterone is required to support strength and hypertrophy gains, which are otherwise attenuated (20). Therefore, the hormone-sensitive processes that underpin muscle anabolism at hypo- and supraphysiolog- ical hormone levels are not being activated appreciably by exercise-induced increases in hormone availability or at least do not result in any measurable enhancement of strength or hypertrophy.

In our view, resistance exercise provides an intrinsic stimulus to the working muscle, which drives hypertrophy, and whereas physiological systemic hormone concentrations may be permissive for the hypertrophic process, exercise-induced elevations do not enhance or in any way predict hypertrophy. Resistance exercise results in the phosphorylation of critically important signaling pathway proteins that are correlated with the extent of muscle hypertrophy in rodents (2) as well as humans (35). These data (2, 35) suggest that local mechanisms within the muscle are of paramount importance in determining muscle hypertrophy. Clearly further research is required to clarify how these mechanisms, possibly in concert with local growth factors (1), combine with other mechanical signals to stimulate muscle protein synthesis and facilitate muscle protein accretion. Ultimately, muscle hypertrophy is specific to the trained muscle group, and strength adaptations are specific to the characteristics of the training regime (29); our data suggest that these adaptations are dissociated from acute exercise-induced hormonal rises.

In summary, transient resistance exercise-induced increases in endogenous purportedly anabolic hormones do not enhance muscle strength or hypertrophy following 15 wk of resistance training. Instead, our data are consistent with the notion that local mechanisms are of primary relevance in producing gains.

Fig. 3. Type I (A) and II fiber (B) cross-sectional area (CSA) of the biceps brachii before (Pre) and after (Post) training in LH and HH; main effect of training, *P < 0.01, †P < 0.001. Elbow flexor CSA before and after training (C) in LH and HH; main effect of training, †P < 0.001. Elbow flexor CSA as a function of distance from the elbow joint line before and after training (D) in LH and HH; main effect of training, †P < 0.001. There were no interactions (training × condition) for either fiber or whole muscle CSA (type I, P = 66; type II, P = 0.55; CSA, P = 0.27). Values are means ± SE.
in muscle strength and hypertrophy with resistance training. These findings, combined with our previous work (36, 37), provide multiple lines of evidence that exercise-induced elevations of purportedly anabolic hormones are not necessary for, and do not enhance, muscle anabolism in young men. Our data indicate that exercise-induced changes in concentrations of systemic hormones do not reflect the underlying processes of muscle protein accretion and cannot be used as a proxy marker of muscle hypertrophy.

ACKNOWLEDGMENTS

We thank Todd Prior and Tracy Rerecich for technical assistance and our participants for their time and effort. We are grateful to Gianni Parise and Warren Foster for use of equipment for muscle fiber analysis and to Dean Inglis and Jonathan Adachi for assistance with MRI.

We thank Profet for the generous donation of whey protein isolate.

GRANTS

This work was supported by a Natural Science and Engineering Research Council (NSERC) of Canada grant to S. M. Phillips. D. W. D. West, J. E. Tang, A. W. Staples, and D. R. Moore are Canadian Institutes of Health Research (CIHR) Canada Graduate Scholarship Award recipients, and all authors acknowledge those sources of support during the conduct of this research.

DISCLOSURES

No conflicts of interest are declared by the authors.

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