Acute Resistance Exercise Program Variables and Subsequent Hormonal Response

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Abstract

Resistance Exercise (RE) is a widely practiced activity both in leisure time and in training periods for competitive athletes. Recent advances in molecular biology and muscle physiology have elucidated some of the mechanisms that regulate muscle growth. However, these molecular advances require application in acute program variables of RE. Therefore, we present an updated paradigm of resistance exercise variables and the effect manipulating these has on signalling pathways and hormonal response to optimise RE adaptations. We herein explore the effect of altering (i) choice of exercise; (ii) order of exercises performed; (iii) load (weight/resistance); (iv) volume (i.e. repetitions x sets x load) and (v) rest period between sets. Manipulating these variables has a consequential effect on signalling pathways, hormone response and neural adaptations that may influence protein synthesis and therefore gradual protein accretion leading to increased muscle size and strength.

Keywords: Resistance exercise; Weight training; Growth hormone; Testosterone; Cortisol

Introduction

Resistance Exercise (RE) is one of the most widely practiced forms of physical activity. This type of exercise is used to enhance athletic performance, augment musculo-skeletal health and alter body aesthetics. The health benefits of RE are primarily as a preventative or countermeasure to circumstances where muscle weakness compromises optimal function (injury or prolonged inactivity, sarcopenia or musculo-skeletal disorders) however, it also has a positive effect on skeletal and metabolic health as well as potential psychological benefits. This review addresses factors that can impact upon RE outcomes and attempts to outline recommendations for favourable physiological adaptations.

Training Design

Acute RE program variables, originally defined by Kraemer characterise the type and magnitude of the RE stimulus [1]. These variables are: (i) choice of exercise; (ii) order of exercises performed; (iii) load (weight/resistance); (iv) volume (i.e. repetitions x sets x load) and (v) rest period between sets.

Choice of exercise: Exercise selection determines the muscle group exercised and the speed of muscle contraction (eccentric, concentric and/or isometric). Multiple-joint exercises such as bench press and squat require complex neural responses and have generally been found most effective for overall strength increases because they allow greater weight to be lifted [2,3]. Exercising large muscle groups provides a larger hormone response than exercising small muscle groups [4,5]. An increased level of hypertrophy in response to RE has previously been reported in the upper body muscles compared to lower extremity muscles [6]. Welle et al. [6] reported Anatomical Cross-Sectional Area (ACSA) of elbow flexors to increase by 9% and 22% in old and young subjects, respectively; whereas, knee extension ACSA increased by only 6% and 4% respectively. Abe et al. [7] found a similar response when assessing muscle thickness by ultrasound. A logical explanation for this is that the lower body musculature is habitually activated and loaded to a higher level during diurnal activities than the upper body musculature [8]. Alternatively, Kadi et al. [9] suggested that intramuscular differences in androgen receptor concentrations may explain the increased potential for upper body hypertrophy.

Muscle contraction velocity is inversely related to the capacity for maximum load [10]. However, isokinetic training has been shown to increase muscular strength specific to the training velocity [11] yet, training at a moderate velocity (180-240°s-1) produces the greatest carryover of strength [12]. The velocity of muscular contraction also impacts signalling responses and adaptations to RE [13]. Eccentric contractions noticeably increase the 70 kDa ribosomal protein S6 kinase (p70 S6K) phosphorylation, while maximal concentric and isometric contractions provide smaller effects [14]. Additionally, eccentric muscle actions stimulate mitogen-activated protein kinase (MAPK) signalling [15]. When force is equated, concentric, isometric and eccentric actions produce similar signalling responses [16], however it is well reported that skeletal muscle can generate ~30% more tension during eccentric contractions than during concentric muscle actions [14]. This inequality in muscle tension has been proposed as an explanation for greater signalling response during eccentric contractions [14]. Furthermore, hypertrophy is attenuated after concentric only RE (matched with conventional concentric/eccentric for total work) [17]. Together, these studies suggest that signalling is similar for all muscle contraction velocities. However, force production capacity is greater during eccentric contractions and may be a more potent stimulus for muscle signalling. Although inclusion of eccentric contractions may be beneficial for RE adaptations, excessive eccentric muscle actions may produce excessive muscle damage. However, this myofibrillar disruption may be causally related to hypertrophy but this is still speculative.

Order of Exercises Performed: Exercise order significantly affects

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the acute expression of muscular strength [18]. Spreuwenberg et al. [18] reported that performing a squat last, as opposed to first during a RE session significantly reduced the number of repetitions performed. Strength and power during multiple joint exercises may be reduced when performed after several other exercises in a RE session [19]. A reduction in muscle activation (measured by electromyography) and metabolic fatigue (reductions in glycogen and/or phosphocreatine) are likely explanations for reduced strength and/or muscular endurance [13,20]. Interestingly, when exercises are sequenced based on agonist/antagonist muscle group relationships, muscle force and power may be potentiated [21]. Considering that multiple-joint exercises have been shown effective for increasing strength and hypertrophy, maximising performance of these exercises early in a workout may be necessary for optimal strength gains [18]. In fact, the American College of Sports Medicine Position Stand recommends that workouts include large muscle groups before small muscle groups, multiple-joint before single-joint exercises, high-intensity before lower-intensity exercises [22]. The same authors offer alternative recommendations; rotation of upper and lower body exercises or agonist-antagonist exercises (commonly known as “supersets”).

**Load:** Load represents the amount of weight lifted or the resistance used during an exercise. Altering this load affects the acute metabolic hormonal neural and cardiovascular responses to RE [23-26]. The maximal load that can be used is heavily dependent on other acute program variables such as exercise order, exercise selection (specifically the muscle action), volume and rest interval length. Load is typically prescribed as a percentage of one repetition maximum (1RM) (e.g. 80% 1RM) or as a weight that allows a certain number of repetitions (e.g. 6RM). Typically in RE, there is an inverse relationship between the load and the volume (i.e. as the load increases the number of repetitions decreases). Loads of below 50% 1RM have been shown to increase dynamic muscular strength in untrained individuals [27,28]. Hakkinen et al. [26] reported that at least 80% 1RM was required to stimulate further neural adaptations and strength during RE in experienced weight lifters. In agreement with these findings, several authors indicated that loads of approximately 5-6RM were optimal for increasing maximal dynamic strength [29-31]. Although significant increases in strength have been reported using loads of 8-12RM [32,33], these loads may be insufficient to stimulate adaptations in experienced weight lifters [26,32,33].

In RE research, rarely is the effect of load on muscular adaptations examined without being affected by confounding variables such as volume. However, Low-Frequency Electrical Stimulation (LFES, simulating low-force contractions) and High-Frequency Electrical Stimulation (HFES, simulating high-force contractions) allow the effect of load on RE adaptations to be analysed [34,35]. LFES increases Adenosine Monophosphate-Activated Protein Kinase (AMPK) activity which promotes glucose and fatty acid oxidation and inhibits protein synthesis (Figure 1). Simply, AMPK promotes energy release and inhibits energy consumption when cellular energy is low (marked by high Adenosine Monophosphate (AMP) and low glycogen) [13]. HFES stimulates protein kinase B (Akt)-mammalian target of rapamycin (mTOR) signalling. This response is critical for increasing muscle protein synthesis and subsequent hypertrophy [36,37]. mTOR signalling increases protein synthesis by increasing the number of messenger RNA translated per ribosome [38]. According to Henneman’s size principal smaller motor units (primarily type I) are recruited prior to larger motor units (primarily type II) until force production is equal to force requirement [39]. Low-force activities recruit primarily type I, slow fatiguing motor units. As the load increases, higher threshold motor units are recruited, encompassing more type II fast-fatiguing fibres until near maximal-loads where the entire spectrum of motor units is activated. As the size principal is related to force, and force=mass×acceleration, rapidly accelerating a load can recruit higher-threshold motor units [40]. Linnamo et al. [41] reported that performing ‘explosive’ concentric contractions using ~40% of maximal isometric force increased Electromyographic (EMG) activity when compared with the same exercise at ~67% of maximal isometric force at a slow velocity. EMG activity has also been found to increase with the onset of fatigue [42]. This indicates increased contribution of higher threshold motor units to maintain force output as lower threshold units begin to fatigue. Therefore, a heavy weight, ‘explosive’ contractions, or fatigue may all increase the number of type II motor units recruited. This may affect the adaptations to RE as only motor units recruited respond to RE type II fibres have a greater capacity for hypertrophy and different muscle groups contain varied percentages of type I and type II fibres [43]. RE to maximize hypertrophy typically include moderate to high loads [13,43-45]. These loads allow the volume to be high and the rest periods to be short to maximize hormonal response (Figure 1) [44].

Traditionally, 1-6RM, 8-12RM and 15+RM loads are recommended to maximize strength, hypertrophy and local muscular endurance, respectively. Campos et al. confirmed these preconceptions by investigating the adaptations following an 8 week RE program in groups using 3-5RM loads (strength protocol), 9-11RM loads (hypertrophy protocol) and 20-28RM loads (muscular endurance protocol) [28]. The investigation revealed a step-wise increase in strength (strength protocol > hypertrophy protocol> muscular endurance protocol), local muscular endurance (muscular endurance protocol-hypertrophy protocol >strength protocol). Increased muscle fiber Cross-Sectional Area (CSA) occurred only in the hypertrophy and strength protocol groups with no significant difference observed between the two. Maximal aerobic power increased solely in the muscular endurance protocol group. Campos et al. however, did not control for volume of training, which has been suggested as a strong indicator for muscular adaptations [28,46].

Research examining periodization has demonstrated a need for variable-intensity loading schemes in RE [45,47]. Rhea et al. reported that daily undulating periodization elicited greater strength gains than the classic linear schedule [48]. An example of daily undulating periodization would be training at 70, 85, and 95% of 1RM on Monday, Wednesday, and Friday, respectively. The mechanisms behind increased strength due solely to periodization are unclear. Selye’s General Adaptation Syndrome suggests that a system will adapt to an unaccustomed stress (i.e. overload). Greater volumes and intensities of RE result in strength adaptations to a certain level presumably, due to an overload of the neuromuscular system [46,49]. Therefore, variations...
in training may increase the neuromuscular overload by continually applying an unaccustomed stress. The American College of Sports Medicine Position Stand recommends that workouts cycle training loads of 80-100% 1RM to maximize muscular strength and 70-100% 1RM to maximize hypertrophy [22].

**Volume:** Exercise volume is typically described as the total number of repetitions performed during a training session multiplied by the resistance used and is reflective of the duration which muscles are being stressed [50]. Therefore, manipulation of training volume can be achieved by altering the number of exercises per session, the number of sets per exercise, the number of repetitions per set, or the resistance used. Volume has been shown to affect neural hypertrophic metabolic and hormonal responses to RE [22,24,50,51]. For example, multiple-set workouts invoke significantly greater Growth Hormone (GH) and Testosterone (T) responses than single-set programs [52]. Long-term training that employs multiple sets, compared to single sets appears to be superior for strength development and hypertrophy [51]. It is unclear however, whether a dose response exists between volume and RE adaptations. Studies incorporating two three four to five and six or more sets per exercise have all produced significant increases in strength in trained and untrained individuals. Interestingly, similar strength increases have been reported when comparing two and four sets and two and three sets [32,33,53-57]. In contrast to Ostrowski et al. [56], Berger [29] reported that three sets were indeed superior to two sets for the development of strength. The optimal number of sets per exercise still remains an equivocal matter with varied opinions even between review and meta-analytical papers. A recent meta-regression reported that multiple sets were associated with a larger effect size than a single set [58]. In a dose-response manner, two to three sets per exercise were associated with a significantly greater effect size than one set. However, no significant difference between one set per exercise and four to six sets per exercise, or between two to three sets per exercise and four to six sets per exercise was observed. Despite no significant difference observed between one set and four to six sets, a trend for four to six sets to result in greater strength increases existed. This was not significant due to the large standard deviation observed. The authors concluded by advising that two to three sets per exercise are associated with 46% greater strength gains than one set, in both trained and untrained subjects. An alternative view has been proposed by Peterson et al. [46], who reported that approximately eight sets per muscle group produced the largest effect size in athletes. Peterson et al. [46] investigated 177 studies and 1,803 effect sizes and advised the following RE prescription: For athletes, maximal strength gains are elicited at a mean training intensity of 80% of 1RM (agreeing with the American College of Sports Medicine Position Stand) two days per week, with a training volume of eight sets per muscle group [22]. RE trained non-athletes exhibit maximal strength gains with a training intensity of 80% 1RM, 3 days per week, with a training volume of 4 sets per muscle group. Untrained individuals experienced maximal strength gains at 60% 1RM, three days per week and four sets per muscle group. These authors demonstrated that exercise prescription for maximum strength gains should vary between populations. However, trained non athletes commonly train for hypertrophy rather than strength and therefore, the above prescriptions may not be relevant to this population.

Hypertrophy involves a proportionate increase in the net accretion of actin and myosin as well as other structural proteins. Mechanical loading leads to a series of intracellular events that ultimately regulate gene expression and protein synthesis [22]. Protein synthesis in human muscle increases after only one bout of RE and peaks approximately 24 h post exercise [59]. This anabolic environment remains elevated from ~2 h post exercise up through 36-48 h post exercise [60] (depending on training status). For previously untrained individuals neural adaptations dominate early stages of training with muscle hypertrophy occurring within six weeks [61]. Similarly to strength training, hypertrophy has shown to be enhanced with multiple-set over single-set training [23]. In untrained individuals however, as with strength, general nonspecific program design appears to be an adequate overload to stimulate hypertrophy [62]. For more experienced individuals, periodized training schemes appear to be the most effective way to maximize hypertrophy. RE targeting hypertrophy commonly include moderate to very high loading, relatively high volume, and short rest intervals as this design produces a greater elevation in T and GH than a strength program with long rest, high-load and low-volume [22]. Total work, in combination with loading has been implicated for gains in hypertrophy [63]. This result has been supported by recent meta-regression analysis of multiple-set, high-volume programs compared to single-set programs [33,64]. Goto et al. reported that the addition of a high-repetition low-load set to a traditional strength training protocol increased muscle CSA [65]. However, this combination protocol was compared with a strength only protocol and a typical hypertrophy regimen was not analysed. Campos et al. examined eight weeks of a high-load RE protocol and reported that neither type I nor type II muscle fibres exhibited hypertrophy [28]. Therefore, the American College of Sports Medicine Position Stand advised that a combination of strength training and typical hypertrophy training is most effective for advanced hypertrophy training as it compromises a load sufficient to overload the musculature and a volume high enough to elicit an elevated endocrine response [22]. As with many RE variables, volume is difficult to isolate as and therefore most experiments manipulate load and volume concomitantly. It seems logical however, that increased load would affect muscle signalling. High-volume RE would deplete muscle glycogen stores and therefore stimulate AMPK activity [13]. As previously reported AMPK promotes muscular endurance adaptations and inhibits protein synthesis (via inhibition of Akt-mTOR signalling). This relationship has been supported by research that demonstrates that low muscle glycogen potentiates exercise-induced AMPK activity and attenuates Akt signalling [64-66]. Comparatively, adequate exercise volume appears necessary for optimal gains in muscular strength and size [51]. Therefore, making conclusions and recommendations about RE volume and muscle signalling is difficult and a further research to establish the threshold RE volume at which AMPK begins to inhibit Akt-mTOR signalling is required.

**Rest intervals:** Rest periods between sets significantly influence the adaptations and responses to RE. Short rest periods are recommended for hypertrophy as short rest periods augment the GH response [67]. However, short rest intervals have a detrimental effect of subsequent sets and over several weeks, attenuate strength increases [42,68]. When training to optimise strength gains, longer rest periods are advised as performance may be compromised with one-min compared to three min rest periods [69]. Strength recovery may not even be complete after 2 mins with several studies suggesting that 3 ~ 5 min rest interval produce less performance decrements than 30 s - 2 min [23,71,70]. The American College of Sports Medicine Position Stand advised rest periods of at least 2 – 3 min for core exercises using heavy loads (e.g. Olympic lifts) and one ~ two min for assistance exercises for the development of strength [22]. The authors advised identical rest periods to maximize hypertrophy in advanced RE training. The aim of hypertrophy training is to produce an anabolic environment, which appears to be best done with short rest periods. However, Robinzon et al. [72] reported no difference in muscle girths, body mass or skin folds in recreationally trained participants when utilising 30, 90 and 180 s
rest intervals. In agreement, Ahtiainen et al. [73] found no difference in muscle CSA after three months of training with 5 min or 2 min rest periods. Intuitively, short rest periods would increase AMPK activity due to the metabolic stress associated with little rest and therefore inhibit protein synthesis following RE, explicating why short rest periods attenuate strength gains [42,69]. However, there is some debate as to whether short rest periods inhibit protein synthesis via AMPK activity as GH is increased with short rest periods, therefore promoting hypertrophy. Also, the importance of increased AMPK activity – 1 – 2 h post-exercise remains unclear [13]. Regardless of these indefinities, short rest periods are commonly prescribed when hypertrophy is the aim [13,22].

Resistance training and the endocrine system

Dependent upon RE variables, RE incites an anabolic hormonal response, including T, GH and Insulin-Like Growth Factor-1 (IGF-1), but also Cortisol (C, a catabolic hormone) [22]. The acute program variables discussed in section 1 dictate the magnitude of this hormonal response. Typical hypertrophy protocols (moderate to high load, short rest periods and high volume) recruiting large muscle groups appear to produce the greatest endocrine response (Figure 2) [3].

Testosterone

T is considered to be of high importance in resistance training adaptations in men [74]. T is a steroid hormone secreted from the Leydig cells of the testes under hypothalamic and pituitary control. T has both anabolic and anticatabolic effects on muscle tissue [75]. The direct action of T on muscle is supported by the presence of cytoplasmic receptors for T in skeletal muscle homogenates [76]. T promotes protein synthesis in muscle tissue with the result being increased muscle mass and strength [77]. Kvorning et al. [78] suppressed endogenous production of T with goserelin in healthy young males and reported serum T below 10% of normal levels strongly attenuated the increase in lean mass and muscle strength and actually increased fat mass during a resistance training protocol. The placebo group in the study adapted to the resistance training protocol by significantly increasing isometric strength and lean leg mass. These results demonstrate a direct link between endogenous T and adaptations to resistance training. Hansen et al. [79] subjected 16 untrained men to nine weeks of training either unilateral arm alone, or unilateral arm and legs. It was reported that the group who training both legs and arm in the same training session exhibited a mean isometric elbow flexion strength gain of 37% compared to 10% in the arm only group after nine weeks. In fact, even the untrained arm showed a significant increase in strength for the legs and arm group over the nine week training program. In contrast, West et al. [5] conducted a similar study but utilised a within-participants design and noted that despite a higher hormonal response to training, subjects achieved significantly different hypertrophy and strength gains when training one arm and legs compared to one arm alone. These authors summarised by suggesting that acute systemic hormone responses are not responsible for muscle hypertrophy and that local mechanisms are primary in producing gains in strength and hypertrophy. However, West et al. [5] tested strength dynamically, whereas Hansen et al. [79] tested isometric strength and this may explain some difference.

The correlation reported by Ahtiainen et al. [80] between changes in isometric strength and T suggest that T may be an important factor for strength development. Interestingly, participants who exhibited increased acute T response after the training period were able to increase CSA of muscle more than those with a lowered response. However, caution should be exerted when examining T concentrations correlated to strength or hypertrophy as T response to resistance exercise is transient and the accretion of protein leading to hypertrophy and strength increases are accumulative. Although T causes up regulation of androgen receptors, the increase in T is transitory and muscle protein synthetic response can be elevated for ~48 h although this time course is reduced with training [13,81]. The hormone response to resistance training has been repeatedly investigated with the current theory of interaction between endogenous T and androgen receptors in the recovery phase stimulating muscle hypertrophy, protein synthesis and strength [4,74,80]. This is furthermore supported by the fact that exogenous T (via supplementation) significantly increases muscle mass and strength, especially when combined with resistance training [82]. A reduction in T may cause an increased storage of fat via decreased fat oxidation, decreased resting energy expenditure, and increased adiposity [83,84].

Despite many authors suggesting that resistance exercise-induced hypertrophy is attributable to acute hormonal response recent findings have suggested otherwise [4,25,79,85,86]. West et al. [86] found that myofibrillar protein synthesis and phosphorylation of the 70 kDa S6 protein kinase (both thought to be predictors of hypertrophy) exhibited no difference between arm only and arm and legs training. The latter protocol elicited a marked elevation in serum T, GH and insulin-like growth factor-1 (IGF-1) from resting values, whereas the former did not. However, participants were supplemented with 25 g whey protein after both protocols which led West et al. [5] to suggest that high quality protein availability, rather than training designed to elevate hormone levels is crucial for muscle hypertrophy.

T indirectly stimulates secretion of IGF-1 and GH, known to be correlated with the magnitude of type I and type II muscle fiber hypertrophy [87]. Numerous studies have demonstrated that resistance exercise increase circulating T concentrations [5,9,25,85,86]. T exerts its influence on protein synthesis via androgen receptors in the muscle. T binds to and converts androgen receptors to a sequence-specific DNA binding factor capable of transferring to the nucleus and associating with DNA to regulate androgen-specific gene expression. Blocking of androgen receptors reduces muscle protein accretion, indicating the physiological importance of T-androgen receptor interactions for hypertrophy [88]. Androgen receptors are up regulated by T therefore the higher T concentrations in men likely explains the higher androgen receptor content found in men compared to women [38,89]. Kraemer et al. [90] reported that a resistance training protocol significantly increased resting testosterone concentrations and that this was augmented by essential amino acid supplementation. This would suggest that androgen receptors were up regulated as a result of this and indeed, hypertrophy (as a result of protein accretion) did occur

![Figure 2: Schematic map of the various hormone, physiological and food-derived factors that influence release of Growth Hormone (GH), and the main effects of GH, Insulin-like Growth Factor-1 (IGF-1), Insulin (IN) and GH on some of the key the metabolic processes that occur in various tissue cells.](image)
as a result of the 12-week training protocol. While endurance exercise may transiently increase T values the number of muscle fibres recruited is less than during resistance training and androgen receptors are only impacted if the motor unit is activated [91]. Therefore, endurance training is less effective at promoting hypertrophy than resistance exercise as resistance exercise (particularly when working to failure) recruits more muscle fibres [13].

It has previously been reported that T responses to training may affect individual adaptation and the ability to exhibit hypertrophy [25,78,85]. For example, Kraemer et al. [92] reported that protocol-specific acute serum T increases were observed after strength and hypertrophy protocols. However, the direction and amplitude of hormonal responses has been shown to vary even in response to similar resistance training protocols [25,68,85]. Beaven et al. [25] investigated the effect of four different resistance exercise protocols on hormonal response in elite rugby players and reported large inter-individual variability for serum T. None of the four protocols: strength (3 X 5 reps at 85% 1RM), hypertrophy (4 X 10 reps at 70% 1RM), power (3 X 5 reps at 40% 1RM) and strength endurance (5 X 15 reps at 55% 1RM) induced a statistically significant increase in T. This was probably due to the large inter-individual variability within the study. The strength and endurance power protocols in the Beaven et al. [25] study elicited lower T responses than the strength and hypertrophy protocols. However, significance was not reached as there was large variability in hormonal responses to resistance exercise as indicated by large standard errors. The authors noted, however, that individual's hormonal response was pronounced in one or occasionally two of the protocols implemented. Therefore, the protocol considered to be optimal in terms on anabolism (defined by an absolute increase in free T concentration) varied amongst individuals. These investigators were the first to observe this phenomenon and offered it as a potential reason why studies have varied in terms of observed hormonal response to resistance exercise. Beaven et al. [25] reported four groups of responders (hypertrophy: n = 4; strength: n = 5; strength endurance: n = 4; power: n = 2). If these findings were representative across a wider population, pooling data would likely elicited skewed results depending on subject homogeneity. As suggested by Beaven et al. [25] different resistance exercise protocols may result in individualised T responses and as already suggested, a relationship between T response and muscular adaptations has been suggested [78]. Therefore, Beaven et al. [85] assigned a group of amateur rugby players to three weeks of the protocol which elicited the lowest T response (Tmin), and three weeks to the protocol which elicited the highest T response (Tmax) in a cross-over design. The protocols were the same as in the earlier investigation [25]. It was reported that body mass, 1RM bench press and leg press all showed significant increases after the period of Tmax training. When performing the Tmin period of training, 75% of athletes showed either no change or a significant decline in 1RM performance. The authors concluded by suggesting the ability of a resistance exercise protocol to induce an increase in free T is causatively linked to strength and bodyweight gains. The observation by Beaven et al. [25] that hormone response to certain resistance exercise protocols shows inter-individual variability has many practical implications. Athletes may be able to make further musculoskeletal gains by individually prescribed resistance exercise protocols based on acute T response.

Cortisol: C is a steroid hormone released by the adrenocortical glands under hypothalamic and pituitary control. C has catabolic effects on muscle tissue and has important metabolic functions such as influencing the metabolism of lipids, proteins and glucose [76]. It increases the mobilisation of fatty acids from fat reserves to active tissue and raises blood glucose [93]. Intense physical exercise causes an increase in C which may inhibit protein synthesis and consequently increases in muscle mass by its catabolic effect [93,94]. All protocols investigated by Beaven et al. [25] resulted in a decrease in C concentration with strength, hypertrophy, power and strength endurance protocols decreasing C by 38.2 ± 20.6%, 33.6 ± 20.6%, 44.3 ± 20.6% and 22.2 ± 20.6% respectively. These findings conflict with a number of studies who report acute increases in C in response to resistance exercise [25,68,95]. Beaven et al. [25] attributed these differences to the anticipatory anxious response to stressful events that cause an increase in C and suggested that by investigating individuals with experience of resistance training and using non-invasive saliva-obtained samples, the stress to individuals in the study was reduced. In exercising humans, C increases the availability of metabolic substrates for exercising musculature, protects from an over-reaction of the immune system to the stress of exercise and maintains normal vascular integrity [96]. C also prepares the body for the next bout of exercise explaining why post-exercise levels of C are raised and the return to basal concentrations is prolonged. An important role of the acute C response is to meet the greater metabolic demand of resistance training [97,98]. In previous reports, acute C response has been highest when the overall stress of the training period has been very high and the response has been linked to the volume and/or intensity of total work [68,95]. Long-term exercise may lead to an overall reduction of acute C response during exercise in men [32,91]. However, Cadore et al. [24] opposed this by suggesting that untrained and resistance trained men exhibited a similar C response to a resistance exercise stimulus. Nevertheless, an increased T response to resistance exercise in trained individuals was reported. These findings may suggest that the participants were in an increased state of anabolism due to an increase in protein synthesis rather than a reduction in protein degradation. However, this opposes the understanding that untrained individuals embarking on resistance training experience more drastic hypertrophy than trained individuals. This may be due to long-term resistance training attenuating the protein synthetic response to acute exercise resistance, by shortening the duration for which protein synthesis is elevated [59].

The ratio of T/C is being intensively discussed and examined as a possible indicator of the anabolic/catabolic status of athletes. From a physiological view, the formation of such a quotient is problematic, as a number of other hormones also affect the anabolic/catabolic status of the body [99]. Addlercruetz et al. [100] however, suggested the T/C ratio as the most sensitive indicator of physical overload. Defining values of overload using this ratio have proven difficult due to high inter-individual variability [101]. Therefore, the ratio may be useful when investigating within-subjects effects however, caution should be used when applying this ratio to between subjects design. (Figure 3)
IGF-1: IGF-1 is secreted as it is produced in the liver in response to GH-stimulated DNA synthesis [102] and remains the most prescribed signal for activating the signal transduction necessary for the initiation of protein translation after mechanical loading of the muscle [103]. IGF-1 is a potent activator of the Akt/mTOR signalling pathway with inhibition shown to prevent protein synthesis after RE [104, 105]. RE has been shown to increase concentrations of circulating and muscle IGF-1 although a number of studies have reported no change [106-108]. The lack of change has been attributed to a delayed response of IGF-1 (3-9 h), following GH-stimulated mRNA synthesis with peak values not occurring until 16-28 h post-exercise [77,107]. RE alters concentrations of IGF binding proteins that influence the biological activity of IGF-1 [108,109]. IGF-1 stimulates muscle hypertrophy via phosphatidylinositol-3 kinase-(PI-3K) Akt-mTOR signalling. IGF-1 also increases the proliferation and differentiation of satellite cells to aid hypertrophy [110] and in response to mechanical overload, mechano-growth factor (MGF; a splice variant of IGF-1) stimulates satellite cell activation [76]. IGF-1 is known to stimulate myoblast proliferation and differentiation in vitro as well as muscle protein synthesis [17,111]. Local expression of IGF-1 in skeletal muscle appears to be load dependent and acts independently of changes in serum GH or IGF-1 [112]. However, Wilborn et al. [113] reported increased mRNA expression of IGF-1 for protocols of 60-65% 1RM 4 x 18-20 repetitions and 80-85% 1RM 4 x 8-10 repetitions. No significant effect for RE protocol was reported for IGF-1, IGF-1 receptor or MGF suggesting that these isoforms may not be load dependent. Interestingly, volume was not controlled for by Wilborn et al. [113] and may have contributed to this indifference as the 60-65% 1RM group experienced greater training volume. It would have been interesting if Wilborn et al. [113] had continued this study over a longer period to determine if hypertrophy, like muscle IGF-1, was equal for both protocols over time. Muscle IGF-1 is thought to induce myofiber hypertrophy by autocrine and/or paracrine action [111]. The effectiveness of muscle IGF-1 is dependent not only on its expression but also on its availability, which is controlled by a family of six IGF binding proteins and by the number of the IGF-1 receptor. As already suggested, expression of IGF-1 in muscle is independent of serum GH and IGF-1 and therefore, exogenous supplementation of serum GH or IGF-1 does not appear to stimulate muscular hypertrophy in the absence of applied load [114]. However, administration of IGF-1 directly into skeletal muscle does increase muscle mass suggesting that any increase in muscle IGF-1 availability may lead to hypertrophy [115]. Bailman et al. [108] hypothesised that muscle IGF-1 after a single bout of resistance exercise would be higher after eccentric contractions compared with concentric contractions. However, the results revealed that although MVC was decreased for longer and muscle soreness was perceived higher after eccentric exercise, serum IGF-1 expression showed no significant difference between the muscle actions. Nevertheless, local IGF-1 is independent of serum IGF-1 and therefore IGF-binding protein-4 (known to inhibit IGF-1) mRNA was reduced significantly after eccentric loading which lead to an increase in skeletal IGF-1 mRNA. This would lead to increased IGF-1 availability within the muscle after mechanical loading. Therefore, the authors suggested that IGF-1 was linked to mechanisms involved in tissue regeneration caused by muscle damage. However, muscle damage may not lead to hypertrophy, however as downhill running and other endurance exercises with large eccentric component cause significant muscle damage, yet endurance exercise is not a potent hypertrophic stimulus [91].

Insulin: Insulin causes cells in the liver, muscle, and adipose tissue to uptake glucose from the blood, storing it as glycogen in the liver and muscle, and stopping use of fat as an energy source. During exercise, insulin secretion is reduced to allow more glucose to the muscle cells. However, it is the aim during RE to minimise this reduction in insulin as it has been shown to have significant effect on muscle protein synthesis when adequate amino acid concentrations are available, by reducing protein catabolism [115]. Exercise-induced mRNA transcription and synthesis of proteins is only fully activated when plasma insulin levels are elevated [116]. Elevations in insulin are expected to inhibit AMPK activity, therefore promoting protein synthesis. Insulin likely exerts its effect on translation through the Akt-mTOR signal transduction pathway [117]. Serum insulin concentrations parallel changes in blood glucose, and the response is enhanced when protein and carbohydrates are ingestion prior to, during, or after a workout [107,115]. A potent anabolic hormone in its normal physiological range; insulin is mostly affected by blood glucose concentrations and dietary intake. Ingestion of carbohydrates, amino acids or a combination of both prior to, during, or after a workout is recommended for maximising insulin’s effects on tissue anabolism [91]. Supplementation before or during RE may maximize benefits as a consequence of increased muscular blood flow and subsequent amino acid delivery.

GH: The acidophilic cells of the anterior pituitary secrete molecules that make up the family of GH polypeptides. The most commonly studied GH isoform, the 22kD molecule, consists of 191 amino acids [118]. Other isoforms also exist and appear to function similarly to the 22kD molecule in promoting tissue anabolism. RE has been shown to acutely elevate many of the GH variants [119]. GH binding to its receptor initiates Janus kinase 2 (JAK2) signalling. JAK2 signalling activates PI-3K. As PI-3K is proximal to Akt-mTOR signalling, it appears that RE-induced elevations in GH promote translational efficiency and muscle anabolism [13]. Animal studies support this, as GH injection into pigs resulted in enhanced translational efficiency and protein synthesis [120,121]. This RE-induced elevation in GH has been shown through 30 min post-exercise [77]. The magnitude of this response appears dependent upon the amount of muscle mass recruited contraction volume intensity and rest intervals [3,21,67,68,80,95]. As with T, greater GH acute responses have been reported with workouts consisting of multiple-sets [68]. GH appears to be highly influenced by the volume of the RE protocol. GH response appears to be most substantial when a protocol elicits a high blood lactate concentration [67]. To evoke a substantial blood lactate response, a RE protocol should include moderate to high load, high volume, short rest intervals and activate large muscle groups. High correlations between serum GH and blood lactate have been reported and Kraemer et al. [77] proposed the accumulation of H+ produced by lactic acid acidosis as the primary factor influencing GH release [95]. Mulligan et al. [122] supported this hypothesis by suggesting that GH response was attenuated following induced alkalosis during high-intensity cycling. Although endurance exercise may transiently increases many of the anabolic hormones, prolonged endurance exercise is not a potent stimulus of hypertrophy as it does not recruit enough muscle fibres (androgen receptors are only activated if the motor unit is activated), specifically type II fibres (known to be most susceptible to hypertrophy) or stimulate protein synthesis via the Akt-mTOR pathway. Breath-holding, Hypoxia, protein catabolism and acid base shifts have also been reported to influence GH release [77]. Therefore, resistance exercise is a potent stimulus for increasing GH as long as exercise intensity and volume is sufficient [123]. A number of studies have reported a correlation between the total work of a RE protocol and the acute GH response [95,122]. Mostly, these studies compare typical strength protocols (high load, low repetitions, and long rest) with typical hypertrophy protocols (moderate load, moderate repetitions, short rest) or a typical strength endurance protocol (low load, high repetitions and short rest). Commonly the strength endurance protocol results in the greatest
maximize hypertrophy, load needs to be substantial enough to recruit fast twitch muscle fibres but light enough to enable short rest periods and a volume capable of causing metabolic stress. This type of protocol maximizes hormonal response. Approximately 5-6RM appears optimal for increased strength in untrained participants but as individuals become more experienced in RE, a greater load is required to stimulate overload. Commonly, compound exercises (Olympic lifts, dead lifts, squats, bench press) have been prescribed to maximize hypertrophy as they cause a high metabolic stress and stimulate a large hormonal response, thereby increasing protein synthesis. However, hypertrophy has been reported in smaller muscle groups without an anabolic hormone response to the protocol. This suggests that a combination of central and peripheral factors stimulate hypertrophy. Compound exercises are advised for athletes as in the majority of events, large muscle groups are the most utilised (often through locomotion) however, RE should be tailored to the individual athlete’s needs. A combination of eccentric and concentric muscle actions is advised for strength and hypertrophy training. Muscle adaptations appear specific to training and isotonic muscle actions are common in most weight lifting events. The exclusion of eccentric muscle actions in a RE protocol has been shown to attenuate muscle growth.

Superior results have been reported for all protocols when incorporating periodization into a training program. A daily-undulating-periodization; characterised by cycling training loads over a short period of time (E.g. 6RM Monday, 10RM Wednesday, 14RM Friday) appears superior to linear periodization and non-periodized training possibly due to more regular overload.

Availability of amino acids is important for protein synthesis associated with hypertrophy and inclusion of CHO may maximize the effect of insulin with creatine also increasing lean body mass. Therefore, a combination of CHO, protein and creatine would be most beneficial to consume pre-, during-, and post-exercise to maximize hypertrophy and recovery. However, creatine is not recommended for athletes who compete in events that require acceleration of one’s own body mass. A number of other factors affect the adaptations to RE including age, genetics, gender, time of day trained at, posture and living habits but these are outside the scope of this review.

With regards to previous and future research, it is clear that acute program variables are difficult to examine in isolation and this issue proves problematic in RE research. Therefore, it is difficult to make any certain conclusions about acute RE variables as altering one will normally impact on another. The issue of hypertrophy in the absence of elevated anabolic hormones is an interesting phenomenon that deserves further investigation. Although this phenomenon has been observed, hypertrophy programs have previously been prescribed based on anabolic hormone response and the findings of Cadore et al. [24] suggest that a hypertrophy is superior in increasing muscle growth compared to a strength or muscular endurance protocol. Therefore, another mechanism other than hormone response may determine why a program utilising repetitions of approximately 10 is superior in eliciting hypertrophy. Future investigations could aim to discover this mechanism. With regards to acute program variables, it is still unclear whether a dose response exists; i.e. would utilising 5, 4, 3, 2 and, 1 sets of an exercise result in a stepwise increase in strength or hypertrophy. Then, could this be extrapolated as far as 10 sets and beyond. This hole in the literature is apparent for all program variables, however may be most applicable to volume.
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