An Evidence-Based Narrative Review of Mechanisms of Resistance Exercise–Induced Human Skeletal Muscle Hypertrophy

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ABSTRACT

LIM, C., E. A. NUNES, B. S. CURRIER, J. C. MCLEOD, A. C. Q. THOMAS, and S. M. PHILLIPS. An Evidence-Based Narrative Review of Mechanisms of Resistance Exercise–Induced Human Skeletal Muscle Hypertrophy. Med. Sci. Sports Exerc., Vol. 54, No. 9, pp. 1546–1559, 2022. Skeletal muscle plays a critical role in physical function and metabolic health. Muscle is a highly adaptable tissue that responds to resistance exercise (RE; loading) by hypertrophying, or during muscle disuse, RE mitigates muscle loss. Resistance exercise training (RET)–induced skeletal muscle hypertrophy is a product of external (e.g., RE programming, diet, some supplements) and internal variables (e.g., mechanotransduction, ribosomes, gene expression, satellite cells activity). RE is undeniably the most potent nonpharmacological external variable to stimulate the activation/suppression of internal variables linked to muscular hypertrophy or countering disuse-induced muscle loss. Here, we posit that despite considerable research on the impact of external variables on RET and hypertrophy, internal variables (i.e., inherent skeletal muscle biology) are dominant in regulating the extent of hypertrophy in response to external stimuli. Thus, identifying the key internal skeletal muscle–derived variables that mediate the translation of external RE variables will be pivotal to determining the most effective strategies for skeletal muscle hypertrophy in healthy persons. Such work will aid in enhancing function in clinical populations, slowing functional decline, and promoting physical mobility. We provide up-to-date, evidence-based perspectives of the mechanisms regulating RET-induced skeletal muscle hypertrophy. Key Words: SKELETAL MUSCLE, HYPERTROPHY, RESISTANCE EXERCISE, PROTEIN SYNTHESIS, ANABOLIC MECHANISMS

Skeletal muscle plays a critical role in physical function, athletic performance, and metabolic health, and low muscle mass is associated with greater mortality in healthy adults and adults with comorbidities (1). The regulation of skeletal muscle mass is influenced by several variables that can broadly be categorized into external or internal system variables. Resistance exercise training (RET) is the most potent nonpharmacological external means of increasing skeletal muscle mass (2) and is an external system variable. In contrast, internal variables are inherent systemic or, more often, local (within the muscle) biological processes that mechanistically underpin hypertrophy in response to external stimuli like RET (Fig. 1). An important question is to what extent manipulation of external variables influences internal variable responses to affect the outcome—hypertrophy. In our view, identifying the key skeletal muscle molecular targets activated by resistance exercise (RE) that, with repetition, will underpin RET-induced skeletal muscle hypertrophy is critical.

At the molecular level, RET-induced skeletal muscle hypertrophy (defined here as an increase in axial cross-sectional area (CSA) of a muscle/muscle fiber) occurs in adult humans because of the accrual of cellular proteins (e.g., myofibrillar, sarcoplasmic, mitochondrial) within preexisting muscle fibers. Surprisingly, we remain largely unaware of the structural adaptations associated with RET-induced skeletal muscle hypertrophy. Nonetheless, an axiom is that hypertrophy requires, among other processes, net muscle protein accretion, which occurs when the rate of muscle protein synthesis (MPS) exceeds that of muscle protein breakdown (MPB)—the algebraic difference between which is commonly referred to as net
protein balance. In contrast, extended periods of negative net protein balance (MPB > MPS) manifest as skeletal muscle atrophy, which occurs under a variety of systemic scenarios, such as reduced physical activity and bed rest, or local, including limb immobilization (3). Importantly, many chronic diseases, including cancer, chronic obstructive pulmonary disorder (COPD), cardiovascular disease, sepsis, uremia, and burns, also have muscle-wasting components (4). Therefore, identifying the mechanisms that regulate muscle protein turnover to favor net anabolism is as pertinent a mission clinically as it is for athletes and possibly more so.

The molecular control of MPS and MPB is complex, and many protein signaling cascades dictate net muscle protein balance. Nonetheless, we are still deciphering what signals trigger rises in MPS and thus could potentially contribute to skeletal muscle hypertrophy. The purpose of this review is to provide an up-to-date synopsis of the main molecular mechanisms regulating RET-induced skeletal muscle hypertrophy in humans.

MECHANOTRANSDUCTION

As an external stimulus, RE results in mechanical loading of skeletal muscle; however, the mechanisms by which skeletal muscle ‘senses’ and then initiates responses culminating in hypertrophy are still being unraveled (5). Several protein complexes have been identified as candidate mechanosensors that act as molecular transducers during myofiber contraction. The extracellular matrix is thought to play a critical role in mechanotransducing signals into biochemical signals that ultimately regulate the control of skeletal muscle mass (5). In the presence of mechanical stimuli, phospholipase Cγ1 colocalizes around focal adhesion kinase (FAK)—a densely localized protein within the costamere—and catalyzes the conversion of phosphatidylinositol 4,5-biphosphate to phosphatic acid (PA) in HEK293T cells (6,7). The synthesized PA activates the HIPPO pathway effectors Yes-associated protein 1 (YAP) and transcriptional coactivator with PDZ-binding motif (TAZ) through signaling cascade (7). YAP and TAZ not only control cell growth in Drosophila melanogaster and some mammalian tissues (8,9) but also regulate myoblast proliferation and differentiation (10,11). Furthermore, although no mechanistic link has been elucidated regarding YAP and mechanistic target of rapamycin complex 1 (mTORC1) signaling, animal studies demonstrated that elevation of YAP expression is sufficient to augment skeletal muscle mass during inhibition of mTORC1 via rapamycin (12,13). Also, YAP and TAZ may play a role in mechanically induced anabolic signaling through the elevated expression of the genes Slc7a5 and Slc3a2 (14). These genes encode for proteins that are the leucine amino acid transporters, which could sensitize mechanically loaded muscle to leucine stimulated MPS (14). In addition, PA may indirectly modulate skeletal muscle hypertrophy via the HIPPO signaling pathways and activate mTORC1 (15). Researchers demonstrated that RE was sufficient for elevating local PA concentration and inhibiting the production of PA ablated markers of mTORC1 activity after mechanical overload of skeletal muscle (15). In sum, costamere-based protein sensors may be necessary for hypertrophic signaling in the immediate postexercise period.

Titin is a large elastic protein structure that spans half the length of each sarcomere from the Z-disk to the M-band (16) and is a primary contributor to the passive force generated during eccentric contraction (17). Titin contains a stretch-activated kinase domain, and the stretch of the sarcomere during myofibrillar
contraction exposes several amino acids in the ATP-binding pocket of titin, thereby activating the protein kinase (18). On the other hand, the role of titin during concentric contraction remains undefined. Because of its mechanosensory properties, titin has been proposed to serve as a mechanosensors and regulator of anabolic stimuli (19,20). However, no existing mechanisms are known connecting titin and mTORC1 signaling. In contrast, titin activation is related to autophagy signaling via muscle ring-finger protein (Murf)1/2–proteasome and therefore may be involved in protein turnover and regulation of muscle mass (21,22).

Filamin-C Bag3 is another Z-disk localized protein structure that has been linked with the regulation of muscle size in response to mechanical stimuli (13). Filamin-C is a V-shaped homodimer protein and, in response to mechanical loading, is proposed to interact with Bag3 and, together, regulate two known hypertrophic mechanisms. First, Bag3 increases mechanical-induced activation of YAP through binding-inhibition HIPPO hypertrophic mechanisms. It appears titin, and Filamin-C Bag3 may play a significant role in regulating muscle mass; however, the mechanisms are far from being understood.

**MTOR SIGNALING PATHWAY**

The mTOR complex includes a serine/threonine kinase that centers two protein complexes in mammals—mTORC1 and mTOR complex 2 (mTORC2). Both complexes contain the subunits DEP-domain–containing mTOR-interacting protein (DEPTOR) and mammalian lethal with SEC13 protein 8 (mLST8) (25). However, mTORC1 and mTORC2 differ in rapamycin sensitivity, functions, and additional subunits. Broadly, mTORC1 is characterized as a rapamycin-sensitive regulator of cell size with the subunits regulatory-associated protein of mTOR (Raptor) and proline-rich AKT substrate 40 kDa (PRAS40) (25). Upstream stimuli, such as nutrients (i.e., leucine), growth factors (i.e., insulin-like growth factor-1 (IGF-1)) and mechanical stimuli (i.e., RE), are converted to intracellular signals and subsequently detected by mTORC1 subunits, such as Raptor (25). Conversely, mTORC2 is characterized as a rapamycin-insensitive regulator of cytoskeletal structure and cell survival with the subunits: rapamycin-insensitive component of mTOR (Rictor), mitogen-activated protein kinase (MAPK)–interacting protein (mSin1), and protein associated with Rictor 1 or 2 (PROTOR 1/2) (26).

The mTORC1–induced increases in MPS have been a focal point in the context of skeletal muscle hypertrophy, and eukaryotic translation initiation factor 4E-binding protein 1 (4EBP-1) and p70S6 kinase 1 (S6K1) are two frequently investigated downstream targets. Skeletal muscle hypertrophy has been correlated with basal MPS (27) and RE-induced S6K1 (28) and 4EBP-1 phosphorylation (29). Notably, mTORC1 activation contributes to increases in MPS, but acute short-term (i.e., within hours) RE-induced increases in MPS are not always correlated with chronic RET-induced skeletal muscle hypertrophy (29). Mitchell et al. (29) observed no correlation between untrained men’s acute post-RE MPS rates and muscle hypertrophy after 16 wk of RET. Damas et al. (30) further demonstrated that the MPS after a single bout of RE at baseline were not correlated with the percent change in *vastus lateralis* CSA (%Δ VL CSA) after 10 wk of RET in young men; however, the MPS after a single bout of RE at weeks 3 and 10 was positively correlated with the %Δ VL CSA after 10 wk of RET. The authors suggested that RE-induced increases in MPS rates largely attenuate muscle damage in the early stages (~3 wk) of RET but, after that, support hypertrophy (30). Overall, mTORC1 seems to play a role in RET-induced skeletal muscle hypertrophy, but the mechanisms underpinning this complex process likely extend beyond merely stimulating mTORC1.

Translocation to the lysosome is critical for mTORC1 activation (31), and the intracellular positioning of mTORC1 after anabolic stimuli has been increasingly studied (32). RE induces mTORC1 translocation to the lysosome, and the mTORC1–lysosome complex subsequently translocates to the cell membrane with a proclivity for capillaries (33). mTORC1 relocation to the cell periphery may promote MPS because of increased proximity to upstream activators, translation initiation factors, and microvasculature (i.e., nutrients) (32). Albeit after endurance exercise, work in trained young men suggests that colocalization of mTORC1 with upstream activators could specifically regulate myofibrillar protein synthesis (34), which would certainly contribute to RET adaptations. The potential impact of nutrient provision (33,35,36) and anabolic properties of the lysosome (reviewed elsewhere [37]) remain avenues for future research on spatial regulation of mTORC1 after exercise.

Rapamycin-insensitive, in addition to rapamycin–sensitive components of mTOR signaling, may also contribute to RE-induced increases in MPS and hypertrophy (38), and this notion is supported by evidence from preclinical models. In rats, rapamycin administration ablated RE-induced increases in MPS completely at 6 h post-RE but only partially 18 h post-RE (39); furthermore, rapamycin administration did not completely ablate RET-induced skeletal muscle hypertrophy (40). In a cornerstone article, Drummond et al. (41) observed that rapamycin blunted MPS in humans after acute RE, particularly ~1 h post-RE. Although chronic rapamycin administration is not feasible in humans, these data cumulatively suggest that rapamycin–insensitive processes impact MPS several hours post-RE and chronic RET adaptations. Interestingly, AZD8055, an inhibitor of both mTOR complexes, completely inhibited RE-induced increases of MPS in rats (42), and tripartite motif-containing 28 (TRIM28) has recently been identified as a rapamycin–insensitive regulator of mechanical load-induced hypertrophy (43). Mechanical stimuli activate Ras kinases that subsequently activate extracellular signal-regulated...
kinases (ERK), which phosphorylate TSC2; thus, MAPK-ERK1/2 signaling may be an additional mechanism by which mTORC1 senses mechanical stimuli (26). ERK1/2 also phosphorylates kinases involved in protein translation (Fig. 2), such as p90 ribosomal protein S6 kinase (p90RSK) and MAP kinase-interacting kinase 1 (MNK1), although these may be mTORC1-dependent processes (26). Both ERK1/2 and mTORC1 may be required to stimulate MPS maximally after RE (41). In sum, the mTOR-related signaling pathways play a major role in skeletal muscle anabolism, and the precise contributions of mTORC1 and mTORC2 continue to be refined.

RIBOSOMAL BIOGENESIS

A ribosome is a protein- and RNA-containing (ribosomal RNA–rRNA) molecular machine that plays an indispensable role in protein translation. A translationally competent ribosome (80S) contains two subunits (one large (60S) and one small (40S)), formed by the intricate association of over 80 ribosomal-associated proteins and 4 rRNAs (44). Ribosome biogenesis consumes a large proportion of cellular energy and is the only molecular process requiring coordinated activation of all three RNA polymerases (45). The RE-induced increase in MPS occurs via two mechanisms (46). Increased translational efficiency is an increased rate of messenger RNA (mRNA) translation with a fixed pool of ribosomes, whereas increased translational capacity occurs when increased numbers of ribosomes are available to translate mRNA. Importantly, mTOR-dependent and mTOR-independent mechanisms regulate translational efficiency and capacity (39).

Ribosomal biogenesis has emerged as a regulatory of RE-induced skeletal muscle hypertrophy (47). Given that rRNA makes up ~85% of total RNA (48), any change in total RNA concentration reflects changes in ribosomal biogenesis. Work in preclinical synergist ablation (SA) models of skeletal muscle overload demonstrate that increased total RNA concentration is associated with skeletal muscle hypertrophy (47). Acknowledging that SA models are pertinent for identifying potential cellular and molecular mechanisms regulating rapid skeletal muscle hypertrophy, the extent of hypertrophy with SA is extreme and in the range of 40%–70% hypertrophy within days (49) to weeks (47). Such rates of hypertrophy do not reflect magnitudes of muscle growth observed in human models of RE. Work from animal models (39) suggests that both translational efficiency and capacity are important in sustaining increases in MPS. However, recent work from Kotani et al. (50) showed no increase in MPS, although three bouts of RE was sufficient to increase ribosomal content.

In RE-trained humans, one bout of RE resulted in no measurable increase in total RNA content, despite elevated rates of MPS (46). Indeed, markers of ribosomal gene expression and transcription plateau after ~2 wk of RET (51) and, in some instances, return to baseline after 12 wk of RET (52). It raises the possibility that the rise in ribosomal content in response to
unaccustomed exercise is initially rapid and is nonspecific to the exercise stimulus. Nonetheless, as RET progresses, ribosomal content declines, as does non–stimulus-specific mRNA content, the protein synthetic response becomes more efficient and specific to the exercise stimulus, and translational efficiency is elevated (Fig. 3) (53). Brook et al. (54) recently observed integrated (i.e., days to weeks) RNA synthesis, which would predominantly reflect ribosomal biogenesis, was increased above basal rates over the 0- to 6-wk period with RET, whereas MPS was not significantly increased above basal level during this period; however, this observation does not necessarily mean that ribosomal biogenesis is not relevant for RET-induced skeletal muscle hypertrophy (55). In sum, protein synthetic responses and transcriptional programs rapidly adapt to the RET stimulus, and further increasing translational capacity would not be required and would likely decline. The pertinent question is, do underlying changes in translational capacity with RET limit skeletal muscle hypertrophy?

RET may lead to a heterogenous hypertrophic response across individuals. Phillips et al. (56) clustered individuals who had completed 20 wk of RET into quartiles of RET-induced changes in lean body mass and demonstrated that individuals with the greatest hypertrophy had a downregulation of genes encoding ribosomal proteins. In addition, several studies demonstrated similar increases in RNA content between individuals who show no change, or a profound increase in skeletal muscle hypertrophy, after RET (57,58). Furthermore, the genes encoding rRNAs (45S and 5S) are tandemly repeated, meaning individuals have numerous copies of rRNA genes; interestingly, there seems to be significant individual heterogeneity in rDNA copy number (59). Figueiredo et al. (60) demonstrated that rDNA copy number was positively correlated with 45S pre-rRNA expression 24 h after a bout of RE. Furthermore, after a bout of RE, hypomethylation was observed at rDNA enhancer(s) sites and binding domains for the transcription factor MYC (60), which is implicated in RNA polymerase I activity and ribosome biogenesis. A limitation with the studies mentioned previously is that the lack of a control group makes it difficult to discern how much change is due to the intervention (i.e., RET), and how much change is simply due to random error (61,62). Nonetheless, although the work is suggestive that differential responders to RET exist and that ribosomal biogenesis may be an important determinant for explaining differential responders to RET, the current data are, in our view, inconclusive, and future work is required.

GENE EXPRESSION

With the advent of “omics” technologies providing a global and unbiased perspective on understanding molecular transducers of skeletal muscle adaptations, we know that exercise results in changes in the abundance of more than 2000 gene transcripts (of a possible 45,000 known genes) (63). Also, the changes in abundance (64) and the ratio of posttranslational modifications of proteins in skeletal muscle can be detected (e.g., ~10,000 phosphorylation sites) (43). The incorporation of next-generation sequencing (i.e., RNA sequencing) to correctly interrogate the breadth and the complexity of the mammalian transcription is limited (65); for example, the top 1% of most highly expressed protein-coding genes commonly encompass up to 40% of sequencing reads (66). Furthermore, grouping differentially expressed genes into functional categories

![Figure 3](http://www.acsm-msse.org)
Pillon et al. (69) assessed changes in gene expression in response to a RET regime without incorporating the large sample size. However, despite characterizing that RET mainly upregulated the mRNA genes of 2000 genes affected by RET. Also, Gene Ontology analysis identified that over 600 genes correlated with muscle growth and a role in hypertrophy is unknown. Indeed, Raue et al. (70) identified that over 600 genes correlated with muscle growth and strength changes after 12 wk of RET. However, many of the growth-related genes were generic features of exercise adaptation(s) (70) and not specific to RET-induced skeletal muscle hypertrophy, per se (56). Rather than averaging transcriptional responses across a cohort of individuals, we propose that if we leverage individual responses (i.e., skeletal muscle hypertrophy) to a RET regime, one can determine the transcriptional signature specific to skeletal muscle hypertrophy.

We recently discovered a set of 141 genes correlated with the muscle growth response to chronic muscle loading in humans (n = 100) (63). The signature showed that muscle loading regulated the untranslated regions (UTR) of mRNA (length of their 3’ or 5’UTR), and this regulated-UTR length was closely correlated with muscle growth, despite levels of mRNA remaining unchanged (>1000 genes) (63). For example, the increase in length of BCAT2 3’UTR or EXT1 5’UTR was strongly related to gain in muscle mass after RET. Ours was the first study linking UTR regulatory events to skeletal muscle hypertrophy via RET; thus, it provided potential clues to the reported discordance between mRNA and corresponding protein levels (64). Also, performing within-individual paired muscle tissue analysis in this study strengthened the reliability of the obtained results by reducing the response heterogeneity by ~40% (71). Our study identified that RET activated the genes associated with extracellular matrix remodeling, angiogenesis, and mitochondria (e.g., FKBP1A, BCAT2, NID2) as central pathways for muscle growth (63). Collectively, utilizing transcriptome technology and leveraging individual heterogeneity in response to RET may help determine molecular regulators for RET-induced skeletal muscle hypertrophy. Nonetheless, the best approach to determine the molecular responses to RET will be to perform reliable gene expression profiling that is complementary to reliable high-throughput protein expression methods.

**RE-INDUCED ACUTE CHANGES IN SYSTEMIC ANABOLIC HORMONES**

Canonical ostensibly anabolic hormones (e.g., testosterone, growth hormone (GH) and its various isoforms, and IGF-1), the concentrations of which are moderately (usually well within the diurnal variation of the hormone) and transiently increased for 15–30 min after RE, have been proposed to be internal stimuli having causative roles in RET-induced skeletal muscle hypertrophy (72). However, despite numerous studies designed to probe this question directly, our group and others have found no support for the thesis that acute changes in serum anabolic hormones induced by RE are mechanistically responsible for skeletal muscle hypertrophy (28,73) or increments in MPS (74). Notably, serum cortisol (i.e., a catabolic hormone for skeletal muscle) was the only hormone shown to be associated with the change in type II fiber CSA resulting from RET (75).

Hypotheses for the potential role of acute changes in anabolic hormones mediating skeletal muscle hypertrophy originate from the observation that RE is an effective physiological stimulus for GH release (76). During an RE session, serum GH levels increase 10–20 min after initiation and peak at the end of the RE, returning to baseline values about 30 min post-RE (76). A relevant question is whether GH is a mediator of muscular growth at all? For example, GH infusion studies mimicking the response to RE show stimulation of MPS and decrement in MPB (77). However, RE-induced increase in serum GH was not associated with MPS (78). Furthermore, there was no additional effect in quadriceps protein synthesis and circumference during a RET program even when young men administered 40 μg·kg⁻¹ of GH for 12 wk (79). Similarly, one bout of RE increased serum IGF-1 levels from 40–50 (resting levels) to 60–70 nM (74). However, the increased IGF-1 levels returned to the basal level within 30 min of post-RE (76), and there was no correlation between systemic changes in the IGF-1 level and MPS (74) or skeletal muscle hypertrophy over time (29,74). In addition, IGF-1 (15 μg·kg⁻¹·d⁻¹) administered for 1 yr in older women, who usually have lower basal GH and IGF-1 than young adults, did not change body composition (or any other measured outcome variable) compared with a placebo group (80). Therefore, changes in serum GH or IGF-1 in response to RE or exogenous administration do not seem to influence skeletal muscle hypertrophy.

Sex steroids are anabolic hormones that have been repeatedly investigated in skeletal muscle hypertrophy studies. Testosterone is an androgenic hormone, and previous studies have been shown that exogenous administrations of testosterone in supraphysiological doses to healthy eugonadal men (81,82) and replacement doses to hypogonadal men (83,84) significantly increase muscle mass and lean body mass. Also, administration of testosterone adjuvant to RET promoted muscle mass increase in older adults who have lower baseline levels of endogenous testosterone (85). RE endogenously increases the systemic concentration of testosterone by 2–4 times above baseline for ~15–30 min in healthy young men (86). Contrary to exogenous testosterone administration, this transient ~30 min spike in serum testosterone has a minimum impact on daily testosterone physiological fluctuation, and it is far lower (4- to 6-fold) than the concentrations reached by exogenous administrations of testosterone (Fig. 4) (86). Furthermore, we have repeatedly shown
no association between changes in systemic testosterone concentrations and skeletal muscle hypertrophy response to RET (73,75). Instead, we have shown that androgen receptor (AR) content in skeletal muscle seems more relevant as a variable to predict the hypertrophic response to RET (73). In addition, it has been shown that RE increases AR binding to DNA, improving anabolic signaling (89). It is also worth noting that healthy eugonadal women, with ~10-fold lower circulating testosterone than men, show similar relative hypertrophy and strength gains in response to RET (90), which is an observation that is difficult to reconcile with testosterone being a mechanistically important, rather than a possibly permissive hormone in RET-induced skeletal muscle hypertrophy.

Estrogen may also be a relevant hormone acting to augment hypertrophy by decreasing muscle damage caused by exercise and upregulating anabolic signaling pathways relevant to muscle anabolism (e.g., insulin/IGF-1 and PI3K/Akt signaling) (72). However, there is no consensus on the role of estrogen in RET-induced skeletal muscle hypertrophy. Variables like the menstrual cycle phase and the testing of subjects before or after menopause are certain variables to control and investigate in future studies (91). We speculate, however, that the role of estrogen in skeletal muscle hypertrophy will follow the pattern of other androgenic hormones and be related to intrinsic muscle variables (receptor density and postreceptor signaling) as we and others have observed with testosterone and AR content (28,92).

**RE-INDUCED MUSCLE DAMAGE**

Muscle damage can significantly increase inflammatory mediators in skeletal muscle and induce satellite cell (SC) activation (93), affecting muscle regenerative processes. The gold standard method for assessing RE-induced muscle damage is via examination of ultrastructural changes, including z-band streaming or muscle swelling (edema) (94). Many, however, rely on indirect measures of proxy markers such as elevation in muscle soreness and creatine kinase (CK) activity in the blood, which is not a measure of damage per se. There is controversy regarding the validity of raised serum CK levels after RET as a relevant marker of myofiber damage and its relationship to MPS and hypertrophy (95). Damas et al. (95) demonstrated that MPS, in addition to markers of muscle damage (serum CK activity, indirect) and Z-disk streaming (direct), was highest after RE in untrained persons early in a RET program; however, neither measure was well correlated with MPS or RET-induced skeletal muscle hypertrophy. Nevertheless, after 10 wk of RET, the acute MPS was correlated with the degree of muscular hypertrophy observed, despite significantly lower muscle damage (95). Thus, muscle damage, which is progressively mitigated with chronic RE, is a poor proxy for MPS and skeletal muscle hypertrophy (95).

After RE, SC responds by activating the myogenic program to proliferate and either return to quiescence or differentiate, donating their nuclei to the existing myofibers (96). Damas et al. (97) reported increased SC content during the first week of RET, showing the more significant RET-induced muscle damage. However, there was no correlation between the SC content and MPS throughout RET (97), suggesting that SC may serve a more prominent role in myofiber repair during the initial stages of RET than the latter stages of RET showing muscle hypertrophy, which is contrary to previous dogma that muscle damage is concomitant and a prerequisite for muscle hypertrophy (98). Recent work from Roman et al. (99) demonstrated that local muscle damage can be repaired independent of SC through a mechanism related to nuclear migration. This alternative myofiber-autonomous repair mechanism challenges...
the role that SC may play in acute local muscle damage as myonuclear migration was sufficient for the local delivery of mRNAs necessary for efficient repair of the damaged sarcomeres (99).

Muscle damage induced by RE triggers an inflammatory response characterized by the release of several mediators (93) and proinflammatory cytokines (e.g., tumor necrosis factor alpha (TNF-α) and interleukin (IL) 1β)) that are known regulators of prooproteolytic activity in skeletal muscle. In comparison, preclinical studies have shown that IL-1β and TNF-α have proinflammatory effects in myoblast cells through mechanisms involving IL-6 and prostataglandins (100), and myotubes treated with IL-6 upregulate mTORC1 signaling and myotube protein synthesis (101). Nevertheless, when tested in humans, daily ingestion of anti-inflammatory medication during RE was reported to have no effects on muscle thickness in young (102) and hypertrophy in older adults (103). Our group found a correlation between the concentration of the IL-6 post-RE and changes in myofiber CSA in subjects submitted to RE (28). Therefore, inflammatory mediators might play a role in skeletal muscle hypertrophy, but this field demands further research exploring intrinsic and local mechanisms.

**METABOLITES**

Metabolites produced during muscular contractions have been posited to be potential internal determinants of RET-induced skeletal muscle hypertrophy (104). Because marked changes in metabolite concentrations always accompany RE (or any other form of muscle contraction), several different molecules are proposed to be involved in gene expression (105) and distinct protein signaling pathways (104). However, no causative research shows that any metabolite is a viable signaling candidate for triggering skeletal muscle anabolism in humans.

Elevated lactate, hydrogen ion, inorganic phosphate and reduced phosphocreatine are all elevated with muscle contraction (106). Based on this mechanism, it has been suggested that the reduced blood pH may promote muscle growth by potentiating GH release and increasing motor unit (MU) recruitment to maintain force output (107). Nonetheless, as pointed out in a preceding section of this review (see RE-Induced Acute Changes in Systemic Anabolic Hormones), changes in serum levels of GH (and its various isoforms) after RE are not correlated, mechanistically incongruent, and with stimulation of MPS (we note that collagen-predominant tissues like bone are markedly sensitive to GH) or hypertrophy (73,74). Furthermore, no additional muscle hypertrophy was observed after RET with blood flow restriction—the RET model used to elevate metabolites production by limiting blood flow—compared with traditional RET (108). Rather, RET with blood flow restriction showed weaker higher threshold MU recruitment (109). In addition, increased lactate concentration in plasma does not induce an additional increase in MPS (74) or CSA by MRI (76) after RE and RET, respectively. Overall, little-to-no evidence exists to suggest that any single metabolite, or even a plausible combination, influences RET-induced anabolic signaling or hypertrophy.

Reactive oxygen species and nitric oxide have been mentioned as potential mediators of skeletal muscle hypertrophy by activating MAPK signaling pathways and SC, respectively (110). In addition to recognizing that evidence supporting this claim is scarce, it is critical to consider the vast regulatory networks involved with RE-induced activation of MAPK (111) and SC (112) activated by mechanotransduction. Based on existing evidence, MAPK and SC activation should be recognized for their anabolic effect rather than reactive oxygen species and nitric oxide production per se.

Given that there are over 4200 metabolites in human serum, any metabolite may be directed/indirectly associated with anabolic signaling for muscle growth. However, the exercises that result in a lesser degree of skeletal muscle hypertrophy relative to RE (e.g., endurance or higher-intensity interval or sprint exercise) also result in significant increments in several metabolite concentrations similar to, or greater than, RE (113,114), further suggesting that metabolites are not the primary drivers of muscle hypertrophy.

**CLINICAL ILLNESS AND AGING AND THE MECHANISMS INVOLVED IN MUSCLE HYPERTROPHY**

As opposed to being the exclusive domain of athletes and bodybuilders, it is abundantly clear that RET is a useful therapeutic modality in clinical care. Importantly, we are beginning to gain critical mechanistic insight into how RET can affect diseased muscle to impart a less catabolic phenotype and greatly improve clinical outcomes. We highlight here some relevant and newer advances in these exciting areas.

Muscle loss in clinical illness(es) (e.g., cancer, COPD, cardiovascular disease, sepsis, and burns) and aging is, in part, a result of rates of MPB chronically exceeding rates of MPS. Specifically, proteolysis through the ubiquitin–proteasome system (UPS) has been considered a primary mechanism of muscle loss during clinical illness (115). Concomitantly, reduced PI3K–AKT/mTORC1 pathway activity has been considered the main mechanism underpinning an attenuated MPS response (115). Our understanding of mechanistic processes underlying muscle loss during illness is mostly derived from animal studies. Thus, much remains to be discovered about these complex mechanisms, particularly in humans, and there are currently no successful pharmacological treatments to prevent muscle wasting. However, previous studies have reported that RET can increase lean body mass, or prevent further losses, in several clinical populations, including cancer patients and survivors (1), patients with renal disease (116), and patients with Parkinson’s disease (117). Notably, RET counteracts skeletal muscle wasting and thus may be characterized as mitigation of muscle loss rather than true hypertrophy in several clinical populations. Also, previous studies highlighted the association between low muscle mass and poor clinical outcomes, such as treatment tolerability and survival.
in cancer patients (118,119). Understanding the mechanisms driving RET-induced skeletal muscle hypertrophy could improve therapeutics to improve clinical outcomes during clinical illness.

In many diseases (cancer, sepsis, diabetes, COPD, heart failure, and burns), increased systemic concentrations of inflammatory markers (e.g., TNF-α, IL-1β, and IL-6) have been shown to coordinate the changes in different mechanisms regulating muscle protein turnover and muscle regeneration (4). The increase in proinflammatory cytokines may promote proteolysis by stimulating the UPS and decreasing MPS response (122). Thus, reducing the resting concentration of proinflammatory agents and elevating circulating levels of anti-inflammatory cytokines (e.g., IL-1 receptor antagonist, soluble TNF-receptor, IL-10), such as with regular RET, may attenuate muscle loss in clinical illness.

Elevated systemic levels of glucocorticoids (e.g., cortisol) have been observed in many diseases (cancer, sepsis, diabetes, renal disease, COPD, and heart failure), and this increment can happen because of exogenous therapeutic administration or endogenous cortisol secretion as part of the stress response to the disease state (4). The excessive glucocorticoid level in systemic circulation activates protein breakdown signaling, including FOXO1, FOXO3, NF-κB, and reduces PI3K–AKT/mTORC1 signaling pathway activity, thereby inducing muscle atrophy (123). However, 7-wk RET increased thigh CSA measured by computer tomography in renal transplants patients receiving prednisone therapy (124). Also, because a diminished capillary number was shown in such patients, the previous study suggested the reduced muscle perfusion (i.e., delivery of amino acids and oxygen) as a reason for atrophy during clinical illness (124). Our laboratory also found the lower capillary number and reduced angiogenesis-related markers protein expression in coronary artery disease patients with reduced SC number and abnormal muscle fiber–type shifting (125). However, 4 and 12 wk of stair climbing-based high-intensity interval training improved the compromised muscle characteristics (125).

In addition to altered metabolism, clinical illness patients experience significantly reduced physical activity. Muscle disuse can negatively modulate skeletal muscle remodeling leading to muscle atrophy by decreasing the anabolic signals activated by mechanical stimuli (i.e., mechanotransduction), such as the mTOR signaling pathway (3). Disuse atrophy has been linked to anabolic resistance in response to hyperaminoacidemia (i.e., feeding) (3). Furthermore, in vitro and animal model data (126) indicate that ceramide accumulated during inactivity may inhibit factors downstream of PA (see the previous section on mechanosensors). However, regular exercise training improved MPS response and reduced ceramide in obese patients with higher muscle ceramide content (127). Reduced physical activity during clinically illness attenuates the activation of anabolic pathway downstream (e.g., HIPPO and mTOR) that can be promoted by mechanical stimuli (i.e., loss of mechanotransduction). Also, the increased level of inflammatory markers and glucocorticoids decrease the activation of anabolic signaling pathway (PI3K–AKT/mTOR signaling) and inactivate FOXO transcription factors, thereby promoting gene expression–associated protein degradation. Although more complex mechanisms are involved in the muscle loss during clinical illness, improving the deteriorated variables, explained previously, that may be able to be improved by RET could be an effective strategy to maintain muscle mass in disease patients.

Although aging is not an illness per se, there are undoubtedly factors common in aging and certain disease states that are likely playing a role in the age-related sarcopenic loss of muscle mass such as increased inflammation factors (128) and loss of proteostasis (129). Besides that, although multifactorial in origin, reduced number and regenerative capacity of SC (130,131), fiber denervation (132), and deregulated intracellular communications (e.g., GH/IGF-1, testosterone, and myostatin) (72,133) have been considered as a cause of muscle loss in aging. Although a deep exploration of the mechanisms that underpin sarcopenic muscle loss cannot be undertaken here, we refer the reader to a recent review on the topic (134). A common finding that occurs with aging and in many muscle wasting disease states is that the response of MPS to normally robust anabolic stimuli is attenuated. This so-called anabolic resistance of MPS, noted in response to RE and protein ingestion (i.e., hyperaminoacidemia), likely relates to cellular mechanisms and signaling responses being attenuated. In addition, a persistent but low-grade, sterile inflammatory state (inflammaging) is likely also playing a role in suppressing MPS and possibly increasing proteolysis by activating UPS (135). Previous studies have shown that these age-related negative adaptations in skeletal muscle could be alleviated by performing RET, resulting in an increase in SC number (130), innervating MU (136), and reduction of inflammation (137). Also, despite the presence of anabolic resistance and low-grade inflammation, older persons almost invariably experience gains in strength with RET but a lesser degree of hypertrophy relative to their younger-age counterparts (138). Nonetheless, RET is a powerful consistent stimulus that should be a primary form of exercise prescribed to counteract age- and diseaserelated muscle loss. Figure 5 outlines several factors that are likely to play a role in age-related sarcopenic muscle loss, including, in our view, a primary contributor, which is periodic disuse events (2).

**CONCLUSIONS AND FUTURE DIRECTIONS**

Skeletal muscle hypertrophy is a complex process resulting from an intricate interplay between external and internal variables (Fig. 1), and RET is the most potent external variable
that initiates a cascade of events that induce muscle hypertrophy (Fig. 6). Thus, understanding the internal variables activated by RE could provide valuable insight to induce skeletal muscle hypertrophy. However, compared with preclinical models, determining mechanisms from human studies is more challenging because of various factors (e.g., limitation of muscle tissue volume, experimental technical difficulty, ethics, and more). Nevertheless, mechanotransduction, translational capacity, and transcription seem to be very promising in identifying key mechanisms for RET-induced skeletal muscle hypertrophy in humans. Also, the anabolic mechanisms regulated by RET could be an important target to maintain muscle mass during disease and aging in general, although there are more complex mechanisms interfering with muscle homeostasis.

The search for molecular signatures identifying the transcripts involved in skeletal muscle hypertrophy, particularly in clinical populations, creates several avenues for future investigation. Activation of mTOR is clearly a key component of muscle anabolism, but additional factors also seem to contribute to skeletal muscle hypertrophy beyond this protein kinase. Ribosomal biogenesis (i.e., translational capacity) and translational efficiency seem relevant and associated with acute RET responses in training-naive subjects, although the role of such variables in the long-term skeletal muscle hypertrophic response requires further research. Studies applying new methods, using in vivo measurement of rRNA synthesis, might bring additional input to assess the role of translational capacity in skeletal muscle hypertrophy, and endocrine-related factors, such as AR.
content in skeletal muscle and female sex hormones, may yet help us understand the role of these variables in skeletal muscle physiology and hypertrophy. More work is still required, but with the rapid development of technology, the up-to-date techniques in skeletal muscle, such as single-cell isolation and single-cell RNA-seq (139,140), could be considered to accelerate to uncover the mechanisms underpinning RET-induced skeletal muscle hypertrophy in humans.

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