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New fundamental resistance exercise determinants of molecular and cellular muscle adaptations

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Abstract Physical activity relies on muscular force. In adult skeletal muscle, force results from the contraction of postmitotic, multinucleated myofibres of different contractile and metabolic properties. Myofibres can adapt to (patho-)physiological conditions of altered functional demand by radial growth, longitudinal growth, and regulation of fibre type functional gene modules. The adaptation's specificity depends on the distinct molecular and cellular events triggered by unique combinations of conditional cues. In order to derive effective and tailored exercise prescriptions, it must be determined (1) which mechano-biological condition leads to what molecular/cellular response, and (2) how this molecular/cellular response relates to the structural, contractile, and metabolic adaptation. It follows that a thorough mechano-biological description of the loading condition is imperative. Unfortunately, the definition of (resistance) exercise conditions in the past and present literature is insufficient. It is classically limited to load magnitude, number of repetitions and sets, rest in-between sets, number of interventions/week, and training period. In this review, we show why the current description is insufficient, and identify new determinants of quantitative and/or qualitative effects on skeletal muscle with respect to resistance exercise in healthy, adult humans. These new mandatory determinants comprise the fractional and temporal distribution of the contraction modes per repetition, duration of one repetition, rest in-between repetitions, time under tension, muscular failure, range of motion, recovery time, and anatomical definition. We strongly recommend to standardise the design and description of all future resistance exercise investigations by using the herein proposed set of 13 mechano-biological determinants (classical and new ones).

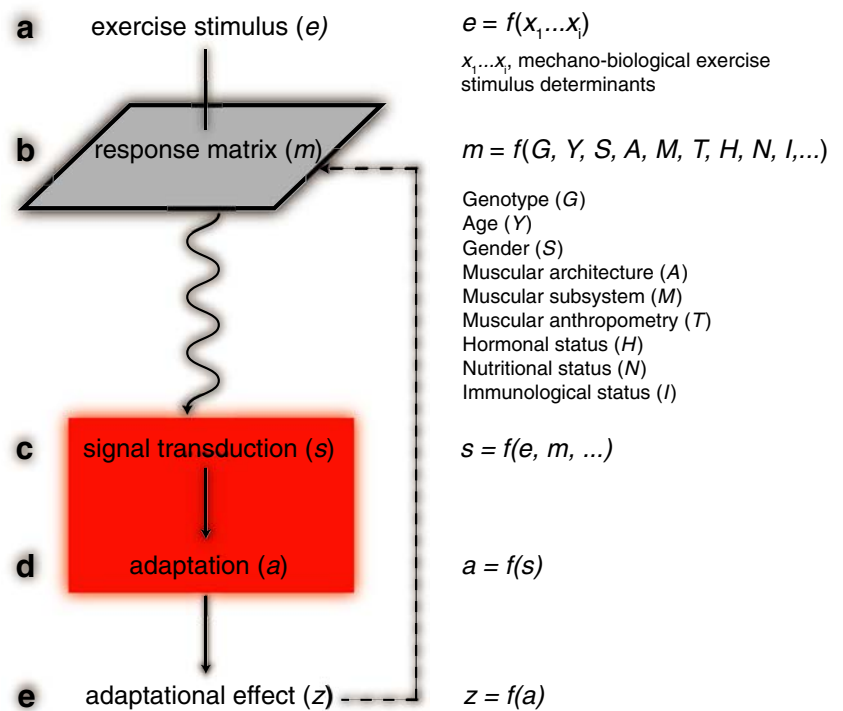
Keywords Exercise · Skeletal muscle · Cellular mechanotransduction · Skeletal muscle satellite cells · Hypertrophy

From stimulus to adaptational effect

Physiological conditions such as resistance exercise perturb the skeletal muscle's tensional integrity (Ingber 2003a, b). These perturbations are mechano-chemically transduced into a molecular and cellular response within and between myofibres and satellite cells (i.e. muscle stem cells) (Tidball 2005). The mechano-chemical transduction is based on the genetic background, age, gender, and several other factors. Finally, this molecular and cellular response leads to specific structural adaptations that result in task-specific functional enhancement (i.e. the adaptational effect) (Fig. 1). However, a causal connection only exists between the molecular/cellular response (i.e. signal transduction) and the adaptation (Fig. 1). "More strength", i.e. the adaptational effect, is not necessarily the result of more muscle mass, since several distinct adaptations can lead to the same effect (at least in the short term). Conversely, training in the "6–12-repetition-maximum zone with multiple sets for 2–3 days week⁻¹", the so-called "hypertrophy training" (Kraemer and Ratamess 2004), doesn't necessarily mean that muscle hypertrophy (i.e. an increase in muscle mass due to the increase in the size, as opposed to the number, of preexisting skeletal muscle fibres) will result. This is due to the fact that the ability to exercise is distinct from the ability to adapt. It has recently been shown for a large cohort of men and women that significant variability in muscle size and strength gain exists after unilateral resistance exercise of the elbow flexors (Hubal et al. 2005). While some subjects showed little to no gain, other showed profound changes. Also, sex differences were apparent. Men had a slight advantage in relative size gains compared to women, whereas women outpaced men considerably in relative gains in strength. Age differences exhibit a profound

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Fig. 1 Simplified model for the transduction of exercise-related skeletal muscle perturbations into structural adaptations with associated adaptational effect. An exercise stimulus with defined mechano-biological characteristics (described in this paper) (a) is mechano-chemically transduced (c) into a quantitative and/or qualitative adaptation (d) of the muscle phenotype, based on the respective response matrix (b). The adaptation (d) is associated with the corresponding adaptational effect (e). A causal connection exists between (c) and (d) (red shading)



impact on the molecular response to resistance exercise, too (Hameed et al. 2003). Following a single bout of high load knee extensor resistance exercise, mechanogrowth factor (MGF) response was attenuated in older subjects. This is indicative of age-related desensitiveness to mechanical loading. Also, various human polymorphisms at genetic loci have a quantitative effect on muscle phenotype (Thompson et al. 2004). Such polymorphisms, also known as muscle quantitative trait loci, drive the genetic variation that underlies muscle size and strength.

Therefore, several prerequisites for the identification of effective and specific exercise perturbations with desired functional enhancement are to be met. First, causal relationships between changes at the molecular and cellular level and the resulting adaptation need to be identified. These relationships need to be established on the basis of several factors such as genetics, age, sex, etc. Further, it must be ascertained whether the adaptation leads to functional enhancement. Finally, the various causal connections between signal transduction and adaptation can be interrogated by multiple exercise perturbations in order to identify the most effective and specific exercise perturbations. However, for the identification of effective and specific exercise perturbations it is imperative to unequivocally define and describe the exercise stimulus in mechano-biological relevant terms. These mechano-biological terms should be directly associated with the molecular response. Unfortunately, in current investigations into the molecular response to exercise, perturbation design and definition remain largely undefined or ignored. This review aims at identifying mechano-biological determinants of exercise

conditions that have quantitative and/or qualitative effects on skeletal muscle phenotype. These mechano-biological determinants are proposed for standardised description of resistance exercise stimuli in both academic and recreational settings.

Fundamental mechanical stimuli decoded by skeletal muscle

Basically, muscles can adopt three strategies of quantitative or qualitative effect on muscular phenotype to adjust for altered functional demands (Goldspink 1985): (1) positive or negative longitudinal growth; (2) positive or negative radial growth and (3) contractile [myosin heavy chain (MyHC)] and metabolic tuning. These adaptational strategies can be adopted concurrently or separately, depending on the specificity of the (patho-)physiological condition. Generally, exercise-induced physiological conditions can be viewed as the perturbations of the muscle cells' tensional integrity. Perturbation of tensional integrity occurs by increasing or decreasing myocellular active and/or passive tension, as well as energy production or absorption. Additionally, these tensional and energetic alterations can be sustained for different duration. Thus, every exercise condition is coded by a specific combination of changes in constant or intermittent active and/or passive tension of different duration. These temporal changes in active and/or passive tension, together with the inferred structural insults, are then decoded at the molecular level and transduced into an appropriate

adaptational response. In the following sections, we provide a detailed description on how the three fundamental adaptations are regulated at the molecular and cellular levels. From this, we derive the relevant mechano-biological determinants for resistance exercise-induced muscular conditioning.

Molecular and cellular determinants of longitudinal muscle growth

Mechanical measurements of rabbit muscle strips before and after skinning indicate that total passive tension increases with increasing sarcomere length (second-order polynomial) (Prado et al. 2005), i.e. with increased strain. Total passive tension is believed to develop due to the lengthening of extramyofibrillar elements (especially the collagen content in the extracellular matrix) as well as to the lengthening of titin. Titin is a giant (~3–3.7 MDa) sarcomeric protein that contains a series of spring elements within its I-band region, which contribute to the elastic properties of myofibrils (Prado et al. 2005). External or internal forces applied to the myofibril lengthen or shorten the myofibril to above or below the slack length, respectively. The lengthening or shortening of the myofibril creates a titin force, which is directed to restore resting length (Miller et al. 2003). Myofibrils can be lengthened actively or passively. This means that myofibrils can lengthen while contracting (“lengthening contraction”, “eccentric contraction”, “active stretch”) or without concurrent contraction. In contrast, myofibrils can only actively shorten, i.e. while contracting (“shortening contraction”, “concentric contraction”). Consequently, passive tension in myofibrils can develop with or without concurrent lengthening contraction or with shortening contraction.

Molecular sensing of myofibrillar strain

In line with titin’s structural and elastic properties, mounting evidence indicates that myofibrillar strain mediates signalling pathways that involve titin’s Z-line region (Miller et al. 2003). Signalling pathways comprise (1) the titin-muscle LIM protein (MLP) pathway, (2) the N2A-muscle ankyrin repeat protein (MARPs) pathway, and (3) the titin-muscle RING finger protein (MuRF) pathway:

- (1) is part of a stretch-dependent myocardial signalling pathway whose impairment contributes to the pathogenesis of a subset of dilated cardiomyopathies in humans (Knoll et al. 2002) and is also induced by skeletal muscle injury due to eccentric exercise (Barash et al. 2004, 2005; Chen et al. 2002; Hentzen et al. 2006).
- (2) involves 3 homologous MARPs: CARP/MARP1, Ankrd2/Arpp/MARP2, and DARP/MARP3. MARPs show cytokine-like induction following cardiac injury,

muscle denervation, and eccentric exercise in vivo (CARP/MARP1) (Aihara et al. 2000; Barash et al. 2004; Baumeister et al. 1997; Kuo et al. 1999); strain in culture, immobilisation at stretched muscle length, and eccentric exercise in vivo (Ankrd2/Arpp/MARP2) (Barash et al. 2004, 2005; Kemp et al. 2000; McKoy et al. 2005); or during recovery after metabolic challenge (DARP/MARP3) (Ikeda et al. 2003), respectively.

- (3) is likely to be involved in the regulation of ubiquitin-proteasome-mediated myofibrillar protein degradation (see “Atrophy signalling”). MuRF1 was found to bind the C-terminal immunoglobulin domains of titin (Centner et al. 2001), and in the nucleus MuRF1 can bind the transcription factor, glucocorticoid modulatory element binding protein-1 (McElhinny et al. 2002). Furthermore, mechanical tension and the titin catalytic domain have been shown to regulate the nuclear localisation of MuRF2 and serum response factor (SRF) (Lange et al. 2005).

Recently, it has also been demonstrated in an in vivo rat model that CARP and MLP are sensitive to both muscle tissue stress and contraction mode, while Ankrd2/Arpp is sensitive only to contraction mode (Barash et al. 2005). This raises the possibility that strain can be sensed independently of stress.

Further evidence for the molecular transduction of passive tension comes from the ~1.5- and ~2-fold increase in the serine/threonine kinase Akt [protein kinase B (PKB)] activity following 5 and 10–20 min of passive stretch, respectively, of the fast-twitch rat extensor digitorum longus muscle but not in the slow-twitch soleus muscle (Sakamoto et al. 2003). Once activated, Akt phosphorylates an array of substrates, including proteins that mediate protein synthesis, gene transcription, cell proliferation, and survival (Vivanco and Sawyers 2002). In mammals, there are three forms of Akt, (Akt1-3), encoded by distinct genes (Vivanco and Sawyers 2002). Expression of a constitutively active form of Akt1 in skeletal muscle cells, either in vitro (Takahashi et al. 2002) or in mice and rats (Bodine et al. 2001b; Lai et al. 2004; Pallafacchina et al. 2002), causes hypertrophy. Conversely, Akt1 inhibits atrophy in vitro and in mice (Sandri et al. 2004; Stitt et al. 2004). Based on their finding that passive stretch has no effect on Akt activity in rat slow-twitch soleus muscle (in contrast to rat extensor digitorum longus muscle), the authors suggested that susceptibility to mechanical stretch is fibre type-specific (Sakamoto et al. 2003). This notion is also supported by the recent finding that the relative importance of titin and the extracellular matrix for total passive tension can vary between different adult rabbit skeletal muscles (Prado et al. 2005). Slow-twitch rabbit muscles exhibit low titin-based passive tension but this tension is highly variable in fast-twitch muscles. Furthermore, titin-based passive tension, but not extramyofibrillar passive tension correlates with the muscle type (Prado et al. 2005).

Cellular sensing of mechanical stretch

Satellite cells are muscle precursor cells that lie between the basal lamina and sarcolemma of skeletal muscle fibres (Mauro 1961). In normal adult muscle, satellite cells are mitotically and metabolically quiescent (Schultz et al. 1978). With appropriate environmental signals, satellite cells enter into the cell cycle, i.e. are “activated”, to provide the precursors needed for new muscle formation in growth and repair (Charge and Rudnicki 2004; Hill et al. 2003; McKinnell et al. 2005). Results from *in vitro* stretch assays demonstrate that mechanical stretch can result in satellite cell activation (Anderson 2000; Anderson and Pilipowicz 2002; Tatsumi et al. 2001; Wozniak et al. 2003). This mechanical stretch induces hepatocyte growth factor (HGF) release from its tethering in the extracellular matrix in a nitric oxide-dependent manner (Tatsumi and Allen 2004; Tatsumi et al. 2002). Once released, HGF binds to the c-met receptor which is located on the plasma membrane of the satellite cells. This interaction initiates a cascade of signalling events that lead to DNA synthesis, and, thus, to satellite cell proliferation.

Structural adaptation to strain perturbation

It has long been known that muscles adapt to a new functional length by adding or removing sarcomeres in series at the ends of the existing myofibrils (Dix and Eisenberg 1990; Griffin et al. 1971; Tabary et al. 1972; Williams and Goldspink 1971). Immobilisation at long muscle length results in an increase in the number of sarcomeres in series. Conversely, immobilisation at short muscle length leads to a decrease in the number of sarcomeres in series. Furthermore, remodelling of the connective tissue following immobilisation has been demonstrated multiple times in mice, rats, rabbits, and cats (Goldspink 1985; Tabary et al. 1976; Tardieu et al. 1977, 1982; Williams and Goldspink 1984). However, both the occurrence and the extent of remodelling seem to depend on the connective tissue type (series elastic element and/or parallel elastic element), species, age, muscle length during immobilisation, and time period of immobilisation.

In exercise physiology, serial sarcomere number modulation has been a neglected topic so far (Morgan and Proske 2004). Only recently has serial sarcomere number modulation been investigated in the context of exercise. Direct evidence for exercise-induced modulation of serial sarcomere number has come from treadmill-trained rat vastus intermedius muscles, the postural knee extensors (Lynn and Morgan 1994; Lynn et al. 1998). Rats were trained by running on a climbing or descending treadmill for approximately 10 min day⁻¹ for 5 days. The latter had previously been shown to cause muscle damage. Subsequently, serial sarcomere analysis for single fibres was performed by laser diffraction. As a result, the descending-trained rats had the largest sarcomere count, the climbing-trained rats had the smallest count, and

sedentary rats had intermediate counts, although closer to the climbing group. In another series of experiments, rat vastus intermedius muscles were tested mechanically while still *in situ*, i.e. attached to the bones, but with all other muscles about the knee joint removed. As a result, in descending-trained rats, the knee angle for optimum torque generation corresponded to longer muscle lengths than in climbing-trained rats. It follows from these results that eccentric exercise (lengthening contractions) leads to accretion of serial sarcomeres. Conversely, exercise comprising only shortening contractions leads to a decrease in the number of serial sarcomeres. Due to the finding that 38% of the difference in sarcomere numbers between decline- and incline-trained rats does not appear as a difference in optimum angle, the authors suggest that it has been taken up by shortened tendons (Lynn et al. 1998).

Recently, the contraction type-dependent differential serial sarcomere number adaptation has been confirmed by measuring vastus intermedius and vastus lateralis muscle fibre dynamics of up- and downhill-running rats *in vivo* (Butterfield et al. 2005). It was shown in that study that vastus intermedius and vastus lateralis muscles of uphill-walking rats undergo repeated concentric contractions, and therefore they suffer no contraction-induced injury. Conversely, the vastus intermedius and vastus lateralis muscles during downhill walking undergo repeated eccentric contractions (Butterfield et al. 2005). Accordingly, short muscle lengths for uphill concentric-biased contractions result in a loss of serial sarcomeres, while long muscle lengths for downhill eccentric-biased contractions result in a gain of serial sarcomeres (Butterfield et al. 2005).

In humans, the optimum angle for torque generation can be measured reliably, e.g. by isokinetic dynamometry. By that means, an angle-torque curve is measured during maximum voluntary contraction with constant velocity shortening. As determined by this measure, a series of eccentric contractions (“hamstring lowers”) of human hamstring muscles produced a significant shift of approximately 7° in optimum knee angle for torque generation to longer muscle lengths (Brockett et al. 2001). The shift in optimum knee angle for torque generation was paralleled by delayed-onset muscle soreness (DOMS) in the hamstrings. The shift occurred immediately after exercise and persisted 8 days postexercise, consistent with a training effect. The mechanism by which eccentric exercise produces muscle damage, DOMS, and increased optimum length for torque generation has been postulated in the “popping sarcomere hypothesis” (Morgan 1990). The popping sarcomere hypothesis states that stretch-induced muscle damage results from nonuniform lengthening of sarcomeres, when active muscle is stretched beyond optimum length. If sarcomeres are beyond optimum length, then the longest sarcomeres will be the weakest and, so, will be stretched more rapidly than the others. Thus, they will become weaker, until rising passive tension compensates for falling active tension. For at least some muscles this

corresponds to lengths beyond filament overlap. The term “popping” is used to describe the uncontrolled, virtually instantaneous lengthening of a sarcomere from a length commensurate with its passive length to a length where passive structures primarily support the tension. Because the weakest sarcomeres are not at the same point along each myofibril, this nonuniform lengthening leads to a shearing of myofibrils, exposing membranes, especially T-tubules, to large deformations. This is thought to lead to loss of Ca^{2+} homeostasis, and, hence, damage, either through tearing of membranes or opening of stretch-activated channels (Allen et al. 2005). Support for the nonuniform lengthening of sarcomeres comes from a recent study of myofibrils from rabbit psoas muscle and left ventricles of guinea pig during activation and relaxation (Telley et al. 2006a). However, these authors also show that albeit half-sarcomeres of contracting single rabbit psoas myofibrils lengthen to different extents during a stretch, rapid elongation of individual sarcomeres beyond filament overlap (popping) does not occur. Moreover, in contrast to predictions of the popping sarcomere hypothesis, they postulate that a stretch rather stabilises the uniformity of half-sarcomere lengths and sarcomere symmetry (Telley et al. 2006b).

With respect to eccentric exercise, it is postulated that the structural adaptation consists of an increase in the number of sarcomeres in series so that a given muscle length corresponds to a shorter sarcomere length (Morgan and Talbot 2002). Whether and how serial sarcomere adaptation in humans following eccentric exercise is paralleled by changes in tendinous and/or muscle belly connective tissue remains to be established.

A consequence of the popping sarcomere hypothesis is that the unloaded shortening velocity of muscle fibres should increase with eccentric training. The reason for this is that the unloaded shortening velocity of a fibre is the sum of the velocities of its sarcomeres. Thus, the more sarcomeres in series, the faster the unloaded shortening velocity, provided that no alterations in MyHC composition occur. However, this will have to be demonstrated in future experiments, especially with respect to the MyHC isoform gene switching associated with stretch and force production (Goldspink et al. 1991). Another consequence of the popping sarcomere hypothesis is that significant muscle damage also can occur with endurance exercise, provided that the duration (marathon running) or mode (downhill running) of exercise is extreme. Therefore, it is predicted that under certain circumstances, endurance exercise can lead to serial sarcomere accretion with concurrent increase in unloaded fibre shortening velocity. On the contrary, “conventional” endurance exercise, which is associated with a bias towards shortening contractions at short muscle lengths, will lead to a decrease in the serial sarcomere number and, thus, to shorter muscle lengths. Short muscle lengths come with a reduction in functional range of motion (ROM). In general, a reduction in ROM is not desirable in health-based settings that aim at increasing

musculoskeletal and cardiovascular function. Hence, in order to increase or preserve a functional ROM by serial sarcomere number modulation, eccentric resistance exercise covering the functional articular range might be the method of choice.

In conclusion, there is a substantial body of evidence that muscle fibres and satellite cells can sense changes in length. Accordingly, active and passive excursions from resting length are transduced into a molecular and cellular response with subsequent structural adaptation. However, with respect to active excursions from resting length, the response at the molecular, cellular, and structural level is dependent on the contraction mode. It follows directly that muscle length change as well as contraction mode are two mechano-biological determinants of exercise-induced skeletal muscle length adaptation. Therefore, these two mechano-biological determinants, among others (described below), need to be specified in reports coming from investigations into the plasticity of skeletal muscle following (resistance) exercise. As a measure of muscle length change we suggest to specify ROM (x_{11} , Table 1) during exercise and the number of length excursions [i.e. the number of repetitions (x_2 , Table 1)]. It must be pointed out, however, that ROM might not always be indicative of fascicle length excursion. The reason for this is that length changes of the muscle-tendon units do not necessarily correspond to the length changes in the muscle fascicles (Hoyt et al. 2005). This means that the muscle-tendon-unit may lengthen, while the contracting muscle is shortening or isometric. As a measure of contraction mode we propose to report the fractional distribution of the three contraction types [shortening (concentric), isometric, lengthening (eccentric)] per repetition in terms of occurrence and temporal requirement (x_7 , Table 1). Also, the number of contractions should be reported (x_2 and x_3 , Table 1). For example, did the exercise comprise one set of several, only lengthening contractions or was one repetition composed of one shortening, one isometric, and one lengthening contraction? How much time did it take to perform one repetition and how was this time distributed over the respective contraction modes? The importance of the latter point for inducing muscle hypertrophy and gains in strength has been demonstrated in studies where the effect of fast lengthening contractions versus slow lengthening or slow shortening contractions has been investigated (Farthing and Chilibeck 2003; Shepstone et al. 2005).

Molecular and cellular determinants of radial muscle hypertrophy

Previous work showed that if the tibialis anterior in the mature rabbit was electrically stimulated while held in the stretched position by plaster cast immobilisation, it increased in mass by 35% within 7 days (Goldspink et al. 1992). Thus, if the lengthened (stretched) rodent muscle is additionally subjected to electrical stimulation, it

Table 1 Mechano-biological descriptors of resistance exercise stimuli. In order to better discriminate between the significant stimulatory cues leading to distinct muscular adaptations, we propose to design and describe resistance exercise-related muscular perturbations based on mechano-biological descriptors (a). Example of how inaccurate exercise stimulus description might lead to wrong conclusions (b). Based on the traditional resistance exercise descriptors (x_1 - x_6), two hypothetical subjects (denoted A and B) receive exactly the same resistance exercise stimulus. However, when further missing

mechano-biologically significant descriptors (x_7 - x_{13}) are taken into account, it is obvious that the two conditions differ. These two differing conditions will lead to distinct mechano-chemical signal transductions in the subjects' muscles, even if the two subjects had the same potential to adapt (i.e. the same response matrix characteristics [see Fig. 1]). Thus, the two differing mechano-chemical signal transductions will most likely lead to distinct muscular adaptations and adaptational effects. Consequently, such results could be misinterpreted

a

| | |
|------------|---|
| x_1 , | load magnitude |
| x_2 , | number of repetitions |
| x_3 , | number of sets |
| x_4 , | rest in-between sets ([s] or [min]) |
| x_5 , | number of exercise interventions (per [d] or week) |
| x_6 , | duration of the experimental period ([d] or weeks) |
| x_7 , | fractional and temporal distribution of the contraction modes per repetition and duration [s] of one repetition |
| x_8 , | rest in-between repetitions ([s] or [min]) |
| x_9 , | time under tension ([s] or [min]) |
| x_{10} , | volitional muscular failure |
| x_{11} , | range of motion |
| x_{12} , | recovery time in-between exercise sessions ([h] or [d]) |
| x_{13} , | anatomical definition of the exercise (exercise form) |

Classical set of descriptors

New set of descriptors

Complete set of mechano-biological descriptors

b

| | x_1 | x_2 | x_3 | x_4 | x_5 | x_6 | x_7 | x_8 | x_9 | x_{10} | x_{11} | x_{12} | x_{13} |
|----------|---------|-------|-------|-------|------------|----------|---|-------|---------|----------|----------|----------|----------|
| A | 75% 1RM | 6 | 1 | - | 2 per week | 10 weeks | 2 s shortening 0 s isometric 2 s lengthening | - | 24+5 s | no | 60% | 24 h | no |
| B | 75% 1RM | 6 | 1 | - | 2 per week | 10 weeks | 10 s shortening 2 s isometric 4 s lengthening | - | 96+10 s | yes | 100% | 72 h | yes |

increases in girth as well as length. The distinct role of active tension in generating radial growth is evidenced by this finding. Conversely, when muscle contractile activity is reduced by means of immobilisation (e.g. casting) or unloading (bed rest, space flight), rapid muscle loss (atrophy) occurs (Booth and Kelso 1973; Thomason and Booth 1990). Muscle loss is accentuated when immobilisation occurs at short muscle length and attenuated when immobilisation occurs at long muscle length, i.e. in a stretched position (Dupont Salter et al. 2003). Therefore, muscle growth and muscle atrophy are two opposing phenomena that are mechanistically linked. Either the activity or inactivity of a common set of molecules controlling a few cellular pathways determines whether the skeletal muscle tissue will respond to defined stimuli with increased protein synthesis and stimulation of cell growth or with increased protein breakdown and reduced cell proliferation (Glass 2003a, b, 2005; Nader 2005; Rennie et al. 2004; Sartorelli and Fulco 2004). In

essence, the maintenance of skeletal muscle mass is the result of the dynamic balance between muscle protein synthesis and muscle protein degradation. Thus, these two opposite processes are believed to hold the key to the understanding of the mechanisms involved in the regulation of skeletal muscle mass.

Resistance exercise in humans and relevant animal models such as functional overload via synergist ablation can produce a significant increase in the mass of the overloaded muscles. In contrast to endurance exercise, resistance exercise is associated with high-intensity-short-duration workloads. The high-intensity-short-duration workloads placed on skeletal muscle during resistance exercise are at or near maximal capacity, and as such produce significant perturbations to the skeletal muscle fibres and the associated extracellular matrix. These perturbations can lead to significant muscle damage, especially if lengthening contractions (eccentric exercise) with supramaximal loads are performed. How-

ever, while both resistance and endurance exercise can result in muscle injury, resistance exercise is more likely to be associated with increases in fibre cross-sectional area and mass. The reasons for the response's specificity point to differences in the integration of hormonal, metabolic, mechanical, neuronal, and immune responses, which are all likely involved in the distinct transcriptional responses that characterise endurance and resistance training.

Molecular determinants of skeletal muscle hypertrophy and atrophy

Exercise-induced hypertrophy mediators upstream of Akt

Resistance exercise-induced muscle hypertrophy results when muscle protein synthesis exceeds muscle protein degradation. In contrast, muscle atrophy is the result from increased muscle protein degradation over muscle protein synthesis. The integration of both anabolic and catabolic signals that lead to the increase or decrease in skeletal muscle mass (Fig. 2) is believed to occur at the molecular nodal point Akt (Nader 2005). Thus, activated (phosphorylated) Akt is both an effector of anabolic signals and a dominant inhibitor of catabolic signals. Activation of Akt is mediated by the insulin-like growth factor 1 (IGF-1)/phosphatidylinositol-3 kinase (PI3K) pathway. The IGF-1/PI3K pathway is triggered by increased muscle loading and subsequent expression of the gene encoding IGF-1 in both animal models (DeVol et al. 1990) and humans (Bamman et al. 2001). On the basis of their mRNA transcripts, three human muscle IGF-1 isoforms have been identified so far: IGF-1Ea ("liver type" isoform), IGF-1Eb, and IGF-1Ec (MGF) (Goldspink 2005; Hameed et al. 2003). In overloaded rodent muscle there are two clearly identified transcripts, IGF-1Ea and IGF-1Eb, of which IGF-1Eb has been termed MGF (Goldspink 2005). Rodent MGF differs slightly from the human MGF sequence as it contains a 52 base pair insert in exon 5 (Goldspink 2005). Other terms such as "mIGF-1", which corresponds to the IGF-1Ea isoform, have also been used to describe the different isoforms (Musaro et al. 2001). However, only MGF appears to be activated by mechanical signals (Yang et al. 1996).

These muscle-specific isoforms of IGF-1 are believed to be sufficient to induce hypertrophy through either autocrine or paracrine mechanisms (DeVol et al. 1990). Transgenic mice engineered to overexpress systemic or liver type IGF-1 under the control of a muscle-specific promoter have skeletal muscles that are twofold greater in mass than those seen in normal mice (Coleman et al. 1995; Musaro et al. 2001). Binding of the cytokine IGF-1 induces a conformational change in the muscle IGF-1 receptor (IGFR) tyrosine kinase, resulting in its *trans*-phosphorylation and the subsequent phosphorylation of insulin receptor substrate 1 (IRS-1). In turn, this results in the activation of PI3K. Finally, activation of PI3K results in the production of phosphatidylinositol-3,4,5-

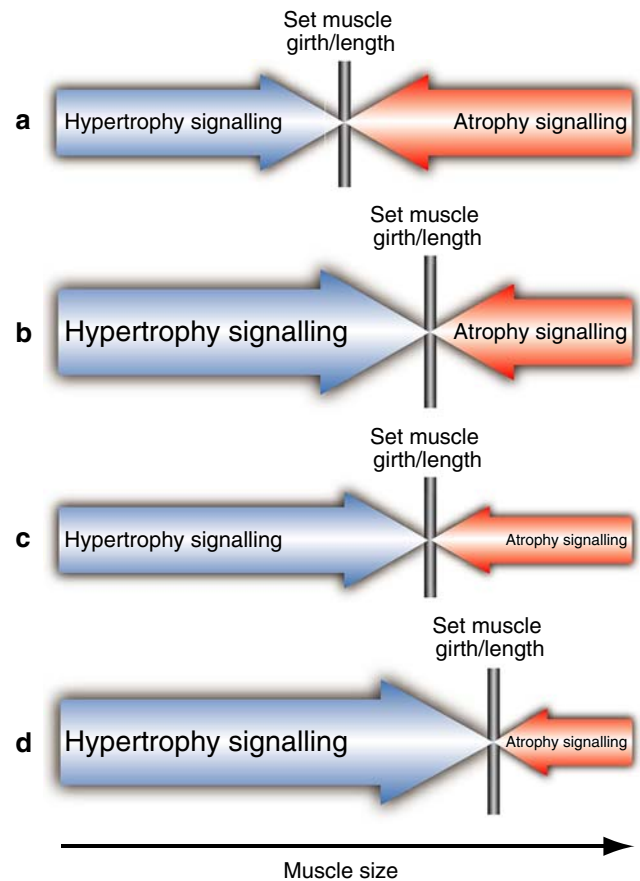


Fig. 2 Simplified model for the relationship between muscle fibre size and the balance between anabolic and catabolic stimuli. Muscle size (girth and/or length) is set by the balance between activity-induced hypertrophy (anabolic) (*blue*) and counteracting atrophy (catabolic) (*red*) signals. In normal muscle, subjected to some amount of tear and wear, hypertrophy and atrophy signals are in balance (**a**). Resistance exercise perturbs the balance by inducing hypertrophy signals over atrophy signals (**b**), or by inhibiting atrophy signals (**c**), or both (**d**), thus driving hypertrophy. This model does not take into account changes in the contractile and metabolic profile that may occur following resistance exercise

triphosphate and activation of Akt via 3'-phosphoinositide-dependent protein kinase 1 (PDK1). However, whether IGF-1 acts as an extracellular cue in muscle biology depends on its availability for muscle IGFR. Indeed, the availability of IGF-1 for muscle IGFR is controlled by IGF-1-binding proteins (IGFBPs). Binding of IGF-1 to IGFBPs can lead either to stimulation or inhibition of IGF-1 effects.

Hypertrophy mediators downstream of PI3K and Akt

Two pathways downstream of PI3K and Akt are believed to mediate hypertrophy (Glass 2005; Nader 2005): the Akt/mammalian target of rapamycin (mTOR) pathway, and the Akt/glycogen synthase kinase 3 beta (GSK3 β) pathway. Both pathways lead to marked hypertrophy through activation of the protein synthetic machinery. Activation of mTOR by phosphorylated Akt

results in an increase in protein translation by two mechanisms: first, mTOR activates 70 kDa ribosomal S6 protein kinase (S6K1/p70^{S6k}), a positive regulator of protein translation; second, mTOR inhibits the activity of PHAS-1 (also known as 4E-BP1), a negative regulator of the protein initiation factor eIF-4E. Conversely, phosphorylation of Akt results in the inactivation of GSK3 β . GSK3 β blocks protein translation initiated by the eIF2B protein. Therefore, GSK3 β inhibition may induce hypertrophy by stimulating protein synthesis independent of the mTOR pathway.

Other growth-signalling pathways in skeletal muscle

Other signal transduction pathways shown to be activated in response to various forms of muscle contraction include those involving the mitogen-activated protein kinase (MAPK) signalling pathways (Long et al. 2004). The MAPK-signalling pathways constitute a network of phosphorylation cascades that link cellular stress to changes in transcriptional activity. Relevant to the present review is the observation that exercise leads to the activation of at least three MAPK-signalling pathways, i.e. extracellular signal-regulated kinases (ERK)1/2, p38 MAPK, c-JUN NH₂-terminal kinase (JNK), in skeletal muscle (Aronson et al. 1998; Boppart et al. 1999; Widgren et al. 2000). Further, AMP-activated protein kinase (AMPK) activity has been shown to be increased during contractions and exercise both in rodents and humans (Winder 2001). However, the relevance of the AMPK-signalling pathway has recently been questioned (Brooks 2005; Wadley et al. 2006). With respect to calcineurin signalling in working skeletal muscle, the reader is referred to chapter “[Molecular and cellular determinants of contractile and metabolic tuning](#)”.

Atrophy signalling

As with protein synthesis, degradation of cellular proteins is an essential process for the maintenance of myocellular homeostasis. However, in some specific situations, when protein degradation exceeds protein synthesis, skeletal muscle mass loss occurs. This process of mass loss is termed atrophy. Skeletal muscle atrophy is a serious consequence of various conditions such as microgravity, hindlimb suspension, immobilisation, and numerous diseases, including cancer and AIDS (Baracos 2001; Booth and Kelso 1973; Miro et al. 1997; Thomason and Booth 1990). Muscle loss is paralleled by profound transcriptomic (Bey et al. 2003; Cros et al. 2001; St-Amand et al. 2001; Stein et al. 2002; Stevenson et al. 2003; Wittwer et al. 2002) and proteomic (Isfort et al. 2000, 2002a, b; Toigo et al. 2005) changes. Over the years, several studies have identified at least five different systems involved in the degradation of proteins during muscle atrophy (Jackman and Kandarian 2004; Kandarian and Jackman 2006). These are the lysosomal system, the calpain system, the caspase or apoptotic protease system, the ubiquitin proteasome system, and the nuclear factor

kappa B (NF- κ B) system. At present, it remains unclear what the relative contribution of these systems to the atrophy process are, and which specific roles they may play during each particular disease state or context in which muscle atrophy develops. However, among the various systems involved in muscle proteolysis during atrophy, the ubiquitin-proteasome system is thought to play a major role (Jagoe et al. 2002). In addition to ubiquitin, three distinct enzymatic components are required, an E1 ubiquitin-activating enzyme, an E2 ubiquitin-conjugating enzyme, and an E3 ubiquitin-ligating enzyme (Glickman and Ciechanover 2002). The E3 ubiquitin ligases are the components which confer substrate specificity. In multiple models of skeletal muscle atrophy, the expression levels of two genes increased significantly: Muscle Ring Finger 1 (MuRF1) (Bodine et al. 2001a) and Muscle Atrophy F-box (MAFbx) (Bodine et al. 2001a) [also called Atrogin-1 (Gomes et al. 2001)]. Both MuRF1 and MAFbx/Atrogin were shown to encode E3 ubiquitin ligases and to be specifically expressed in skeletal muscle (Bodine et al. 2001a). However, the upregulation of MuRF1 and MAFbx/Atrogin requires the nuclear translocation and activity of a family of transcription factors termed Forkhead box O (FOXO). Indeed, in the context of skeletal muscle atrophy, an increase in FOXO1 mRNA in addition to several other atrophy-related genes was reported (Lecker et al. 2004). Also, FOXO3 activation was demonstrated to be sufficient to induce atrophy (Sandri et al. 2004). However, recent evidence shows that FOXO transcription factors are excluded from the nucleus when phosphorylated by Akt, and translocate to the nucleus upon dephosphorylation. Thus, muscle atrophy is prevented by FOXO inhibition through nuclear exclusion by phosphorylation through Akt. This finding highlights Akt's role as a molecular checkpoint for the integration of both anabolic and catabolic signals that lead either to the increase or decrease in skeletal muscle mass. However, although there is a distinct set of genes which are inversely regulated by hypertrophy and atrophy (Latres et al. 2005), distinct transcriptional pathways are activated in skeletal muscle atrophy. These distinct transcriptional pathways are not necessarily the converse of those seen during hypertrophy. Thus, it seems that atrophy is not simply the converse of hypertrophy.

Muscle mass enhancement by inhibition of negative regulators

Myostatin, also known as growth and differentiation factor 8 (GDF-8), is a transforming growth factor- β (TGF- β) family member. It inhibits the progression of myoblasts from G1- to S-phase of the cell cycle through upregulation of p21, the only cyclin-dependent kinase 2 (Cdk2) inhibitor (McCroskery et al. 2003). Myostatin also inhibits myoblast differentiation by downregulation of MyoD/Myogenin expression (Langley et al. 2002). Consequently, myostatin acts as a negative regulator of skeletal muscle mass in (1) cattle, (2) mice, and (3) humans. (1) Naturally occurring mutations in the myo-

statin gene are responsible for the “double-muscling” phenotype, which is characterised by a dramatic increase in muscle mass of certain breeds of cattle (McPherron and Lee 1997). (2) Myostatin-null mice show an increase in muscle mass due to muscle hyperplasia and hypertrophy (McPherron et al. 1997). (3) Recently, a child with muscle hypertrophy was found to have a loss-of-function mutation in the myostatin gene (Schuelke et al. 2004). This individual showed a quadriceps cross-sectional area 7.2 standard deviations above the mean for age- and sex-matched controls and the ability to hold two 3 kg dumbbells in “horizontal suspension with arms extended” at the age of 4.3 years (Schuelke et al. 2004). As suggested, other less dramatic changes in the myostatin gene (or heterozygosity for the splice site mutation) may confer enhanced athletic prowess in a less conspicuous manner (McNally 2004). However, the child’s mutation has not been found in any other individual, and is therefore not a polymorphism-driving normal human variation (Gordon et al. 2005). Furthermore, genetic association studies with myostatin polymorphisms have consistently failed to demonstrate any statistically significant relationship with any human muscle trait (Ferrell et al. 1999; Ivey et al. 2000; Thomis et al. 2004).

In summary, skeletal muscle mass depends on the dynamic balance of protein synthesis versus protein breakdown. Whether muscle fibre protein synthesis outweighs protein degradation depends on the activity of intracellular hypertrophy- and atrophy-inducing mediators (Fig. 2). The activity of intracellular hypertrophy- and atrophy-inducing mediators is coordinated at molecular checkpoints within the myofibre. These molecular checkpoints integrate anabolic and catabolic signals that are triggered by (patho-)physiological conditions. Resistance exercise is a physiological condition that aims at inducing hypertrophy signalling while repressing atrophy signalling (Fig. 2), finally leading to myofibre hypertrophy. Resistance exercise is associated with high active tension that is imposed on skeletal muscle. As shown, active tension through muscular contraction is per se a potent anabolic stimulus for myofibre hypertrophy. However, the levels of active tension required to induce graded hypertrophic effects or to prevent atrophy are most likely to differ. Therefore, such graded tensional effects must be investigated at the molecular and cellular level, if specific exercise regimens, e.g. for the prevention or treatment of sarcopenia, are to be developed. Consequently, the level of active tension that is imposed on skeletal muscle during resistance exercise is a further significant mechanobiological determinant of skeletal muscle size adaptation (x_1 , Table 1). As such, it should be quantified in resistance exercise reports. However, the quantification of the load magnitude poses some problems, since usually, load magnitude is reported in terms of the one-repetition-maximum (1RM), e.g. 75% 1RM. It is beyond the scope of this review to discuss issues related to the 1RM. Suffice it to say that in a scientific setting we do not consider the 1RM an appropriate measure to determine the

magnitude of the tensional load for exercise. In a scientific setting, we suggest to construct maximal voluntary torque (MVT)-angle curves, whenever possible. Based on these MVT-angle curves, the respective choices with respect to tension magnitude can be legitimated. Moreover, MVT intramuscular imbalances (joint angles of disproportionate torque) can be detected and pre-/post-MVT-angle curves can be compared with respect to MVT as well as optimum angle for torque generation (see “Structural adaptation to strain perturbation”). In most other settings it might be more practical to report the load magnitude in terms of 1RM [e.g. % 1RM (x_1 , Table 1)]. Importantly, information about the 1RM should always be combined with information about the time under tension (TUT) (x_9 , Table 1) until failure. That is, how many seconds the exercise can maximally be sustained prior to volitional failure (x_{10} , Table 1). This will additionally give important information about the metabolic changes occurring with training (see “Molecular and cellular determinants of contractile and metabolic tuning”). However, load magnitude per se is not a measure of muscular loading. Only an anatomically perfect technique will allow the efficient “delivery” of the load to the muscle under investigation. It follows that a sound anatomical definition of the exercise in terms of joint positions, movement velocity (movement control), etc. should be an integral part of the exercise stimulus descriptions (x_{13} , Table 1). It is imperative to know if the muscle was under permanent tension and how much of the load effectively “reached” the target muscle.

Cellular determinants of muscle hypertrophy and atrophy

As mentioned above, satellite cells are lineage-committed adult muscle stem cells, located between the basal lamina and the sarcolemma of myofibres. Satellite cells contribute to postnatal muscle growth and muscle regeneration after injury (Charge and Rudnicki 2004; Dhawan and Rando 2005; McKinnell et al. 2005; Wagers and Conboy 2005). Upon myotrauma, quiescent satellite cells become activated, proliferate, and ultimately fuse to existing damaged muscle fibres or among themselves to form new myofibres. Satellite cells are activated in response to hypertrophic stimuli, such as those occurring during muscle mechanical overload (Darr and Schultz 1987; Moss and Leblond 1971; Schiaffino et al. 1976). In several animal models of compensatory hypertrophy (Hanzlikova et al. 1975; Snow 1990) or after resistance training in humans (Kadi et al. 1999a, b, 2004; Roth et al. 2001), the total number of activated satellite cells is substantially increased. The mechanisms leading to satellite cell activation during muscle hypertrophy are not entirely understood. It is postulated that extensive physical activity, such as resistance training or muscle overloading (chronic stretch, agonist muscle ablation, tenotomy), inflicts muscle injury (Allen et al. 2005; Armstrong et al. 1991; Faulkner et al. 1993; Gibala et al. 1995). Consequently, muscle injury, similar

to more severe muscle damage, may initiate a process of regeneration. An indirect proof of muscle damage after mechanical stress is given by an increase of serum markers such as muscle creatine kinase, an enzyme that is usually restricted to the myofibre cytosol. Muscle injury initiates an inflammatory response with the attraction of nonmuscle mononucleated cells, such as neutrophils and macrophages, into the damaged zone (Fielding et al. 1993). Subsequently, several growth factors are released either by the infiltrating cells or by the damaged myofibres themselves. These growth factors may ultimately regulate satellite cell proliferation and differentiation. Indeed, several cytokines have been described that modulate proliferation and differentiation of satellite cells *in vitro* or during regeneration after (exercise-induced) muscle injury (Charge and Rudnicki 2004; Vierck et al. 2000). As mentioned above, HGF is considered to be a key regulator of satellite cell activity during muscle regeneration (Allen et al. 1995; Bischoff 1997). HGF is secreted by damaged tissue during the early phase of muscle regeneration in amounts proportional to the extent of muscle injury (Sheehan and Allen 1999; Tsumi et al. 1998). It seems that HGF directly regulates satellite cell activation. As described, a large body of evidence supports the importance of IGF-1 in the genesis of skeletal muscle hypertrophy. IGF-1 can promote both proliferation and differentiation of cultured satellite cells, and these findings have been confirmed in animal models (Charge and Rudnicki 2004). Experiments showed that muscle-localised expression of IGF-1Ea ("mIGF-1") prevented, through an increase of the regenerative potential of satellite cells, the age-related loss of muscle mass (Musaro et al. 2001). Also, satellite cells derived from mice overexpressing IGF-1Ea display an increased proliferative potential (Chakravarthy et al. 2000b). Increased proliferative potential seems to be mediated by activation of the IGF-1/PI3K/Akt pathway, which results in the inactivation (phosphorylation) of FOXO1 (Machida et al. 2003). Inactivation of FOXO1 downregulates the activation of the p27^{Kip1} promoter (Chakravarthy et al. 2000a). Therefore, the molecular pathways activated by IGF-1 in the muscle fibres to promote increased protein translation appear also to be activated in satellite cells. However, IGF-1 action on satellite cells seems to be IGF-1 isoform-specific with apparently different expression kinetics (Goldspink 2005). After exercise and/or damage, the IGF-1 gene is first spliced towards MGF but after a day or so becomes completely spliced towards the systemic IGF-1 isoforms, which in human muscle are IGF-1Ea and IGF1-Eb (Goldspink 2005; Haddad and Adams 2002; Hill and Goldspink 2003; Hill et al. 2003; Yang and Goldspink 2002).

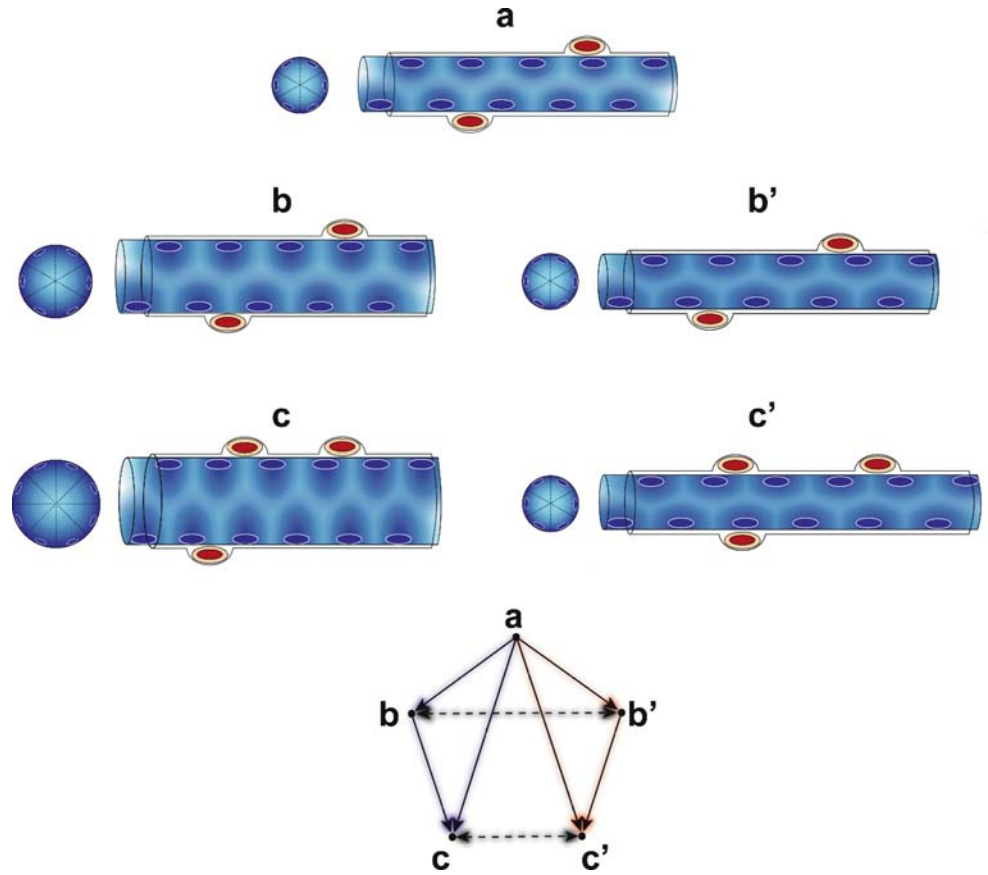
Concepts of myocellular enlargement

Muscle fibres, i.e. multinucleated muscle cells, develop during embryonic differentiation, when mononucleated myoblasts first proliferate and then fuse to form myotu-

bes that become innervated. Following myoblast fusion, no further mitotic divisions occur within the myotubes or muscle fibres. Thus, under normal biological conditions, adult skeletal muscle is an extremely stable tissue with little turnover of nuclei (Decary et al. 1997; Schmalbruch and Lewis 2000). These findings about the postmitotic and multinucleated nature of muscle fibres have led to the concept of a DNA unit or myonuclear domain (Allen et al. 1999; Cheek 1985; Hall and Ralston 1989). The myonuclear domain is the theoretical amount of cytoplasm supported by a single myonucleus. However, the concept of a myonuclear domain is a theoretical one since regulation of the expression and distribution of individual proteins within the muscle fibre is dependent on a number of different variables related to the nature of each protein. Nonetheless, since each muscle fibre is made up of many myonuclear domains, muscle fibre radial or longitudinal hypertrophy could conceivably result from either an increase in the number of domains (by increasing myonuclear number) or by an increase in the size of existing domains (Edgerton and Roy 1991) (Fig. 3). Research to date has strongly supported the former concept by showing that satellite cell activation is required for muscle hypertrophy. The requirement of satellite cell activation was first demonstrated by an approach in which mild γ -irradiation was employed to block satellite cell proliferation. In response to functional overload, myonuclear number or muscle size was not increased in irradiated rat and mice muscles (Adams et al. 2002; Rosenblatt and Parry 1992). However, recent reports indicate that the size of myofibres can increase without the addition of new myonuclei (Kadi et al. 2004; Wada et al. 2003; Zhong et al. 2005). It was found that following 30 and 90 days of resistance exercise, the fibre area controlled by each myonucleus gradually increased throughout the training period and returned to pretraining values during detraining (Kadi et al. 2004). No alterations in the number of myonuclei were detected. Moreover, it has been shown that under normal physiological conditions myonuclear domain size might vary throughout mouse lifespan (Wada et al. 2003) and that myonuclear domain size is not constant during rat soleus muscle atrophy (Zhong et al. 2005).

Whether the increase in mass during hypertrophy results from an increase in the size of each fibre (hypertrophy) or by an increase in fibre number (hyperplasia), has been under debate. Although the evidence has been somewhat contradictory, there has been some suggestion that an increase in fibre number may occur in some animals under certain experimental conditions. Indeed, a review of several investigations into skeletal muscle growth concluded that in several animal species certain forms of mechanical overload increases muscle fibre number (Kelley 1996). However, it has been suggested that some reports have misinterpreted the intricate arrangements of elongating fibres as increases in fibre number (Paul and Rosenthal 2002). Indeed, studies reporting an increase in the number of muscle fibres used avian or cat muscles (Kelley 1996). Both avian and cat

Fig. 3 Hypothetical model of skeletal muscle fibre cytoplasmic enlargement. Schematically depicted are satellite cells (red/orange) lying beneath the basal lamina and sarcolemma of multinucleated (dark blue) skeletal muscle fibres (cross and longitudinal sections). According to the concept of the myonuclear domain (see text for details), muscle fibre hypertrophy could conceivably result from either an increase in the size of existing domains (dark blue shading) ($a \rightarrow b$ and $a \rightarrow b'$ for radial and longitudinal hypertrophy, respectively) or by an increase in the number of domains by addition of new myonuclei (dark blue) provided by the satellite cells (red) ($b \rightarrow c$ / $a \rightarrow c$ and $b' \rightarrow c'$ / $a' \rightarrow c'$ for radial and longitudinal hypertrophy, respectively)



muscles have multiple endplate bands and fibres that do not insert into both tendons but terminate intrafascicularly (Paul 2001). Thus, it remains to be determined whether the radial growth of muscles with intrafascicularly terminating fibres in larger mammals arises from new fibre formation as assumed previously, or from elongation of existing fibres as recently proposed (Paul and Rosenthal 2002). However, most human muscle fascicles, despite their great length, consist of fibres that extend continuously from one tendon to the other with a single nerve endplate band. Therefore, muscle hypertrophy in the adult human apparently can be accounted for predominantly by hypertrophy of existing fibres via addition of newly constructed myofibrils to the contractile apparatus. Accordingly, it has been suggested that hyperplasia does not occur in humans following resistance exercise (MacDougall et al. 1984; McCall et al. 1996). However, hyperplasia still remains a thinkable mechanism of muscle enlargement. Thus, more conclusive evidence might come from future investigations into the plasticity of human muscle fibre number. Fibre splitting or branching is also a characteristic feature of muscle regeneration (Charge and Rudnicki 2004). Fibre splitting is commonly observed in muscles from patients suffering neuromuscular diseases, in hypertrophied muscles, and in ageing mouse muscles, all of which are associated with abnormal regenerative capacity (Bockhold et al. 1998; Charge et al. 2002; Schiaffino et al. 1979). It has been hypothesised that fibre splitting occurs due to

the incomplete fusion of fibres regenerating within the same basal lamina (Blaivas and Carlson 1991; Bourke and Ontell 1984).

Summarising, mechanical stress through high-intensity resistance exercise (especially, but not exclusively supramaximal eccentric exercise) inflicts myotrauma. Upon myotrauma, quiescent satellite cells become activated: (1) through anabolic cytokines that are released by the perturbed extracellular matrix; (2) by infiltrating cells involved in the inflammatory response; (3) by the damaged myofibres; (4) in an autocrine manner by the satellite cells themselves. Following activation, the satellite cells proliferate, and ultimately fuse to existing muscle fibres or among themselves for tissue repair/regeneration. Thus, besides myofibre hypertrophy due to increased protein synthesis/decreased protein degradation (see “Cellular determinants of muscle hypertrophy and atrophy”), satellite cell-based myoplasmic enlargement is a further mechanism in adult skeletal muscle hypertrophy. It is consistent with the concept of the myonuclear domain, where the satellite cells provide the additional DNA for the establishment of additional myonuclear domains during myoplasmic enlargement. It follows that muscle damage is a further significant mechano-biological determinant of skeletal muscle size adaptation. Thus, load magnitude (as a measure of extracellular matrix and satellite cell perturbation) as well as the number of lengthening contractions (as a measure or eccentric damage) need to be specified in resistance exer-

cise reports (x_1 , x_2 , x_7 , Table 1). Additionally, successful recovery (repair) from muscle injury depends on the balance between the degenerative and regenerative processes. Importantly, degenerative and regenerative processes are time-dependent. Thus, if resistance exercise perturbations are delivered at very short time intervals, the degenerative processes prevail. If the degenerative processes prevail, muscle mass is lost. As a consequence, studies in which subjects are resistance-trained over a determinate period of time (x_6 , Table 1) should report recovery times in between the exercise sessions (x_{12} , Table 1) and number of exercise interventions per week (x_5 , Table 1), as a function of exercise-induced muscle damage (exercise intensity). The importance of these three factors (x_6 , x_{12} , x_5 , Table 1) has been demonstrated in studies, in which the duration of elevated protein turnover following resistance exercise sessions was investigated (Chesley et al. 1992; MacDougall et al. 1995; Phillips et al. 1997).

Molecular and cellular determinants of contractile and metabolic tuning

Muscle fibre has multiple, complex functional gene groupings adapting independently to environmental stimuli (Spangenburg and Booth 2003). The molecular regulation of these functional gene groupings is fibre type-specific and results in the fibre type's phenotypical characteristics. Such characteristics comprise, e.g. contractile protein isoforms, mitochondrial volume, myoglobin levels, capillary density, and oxidative enzyme capacity. Each of these characteristics could be considered a functional gene domain within the respective fibre type (Spangenburg and Booth 2003). Not only can these functional modules be regulated within different myofibres but also MyHC protein expression can be heterogeneous within a single fibre (Pette and Staron 2000; Talmadge et al. 1996), resulting in "hybrid" (Baldwin and Haddad 2001) or "polymorphic" (Caiozzo et al. 2003) fibres. It is believed that contractile activity following neural activation induces changes in common regulatory factors within a subpopulation of genes (i.e. gene "modules") to modify the muscle fibre phenotype (Spangenburg and Booth 2003). More specifically, neural activation of skeletal muscle results in the release of acetylcholine from the neuromuscular junction and depolarisation of the plasma membrane, which activates force production by a process known as excitation-contraction coupling. The frequency and duration of stimulation determine the amplitude and duration of the Ca^{2+} transients and, as a result, the level of force output by the muscle. Thus, both the amplitude and duration of the Ca^{2+} transient in skeletal muscle are determined by the motor unit (MU) firing frequency. The increases in amplitude, as well as the duration for which these amplitudes are achieved, are thought to encode signals that will be recognised by different downstream Ca^{2+} -depen-

dent pathways. The key signalling pathways downstream of the elevation in intracellular Ca^{2+} that translate this signal into a transcriptional response include the Ca^{2+} /calmodulin(CaM)-dependent phosphatase calcineurin (Cn), Ca^{2+} /calmodulin-dependent kinase II (CaMKII), Ca^{2+} /calmodulin-dependent kinase IV (CaMKIV), and Ca^{2+} -dependent protein kinase C (PKC) (Chin 2005). In turn, these Ca^{2+} -dependent key signalling pathways will determine the set of genes expressed, thus providing a mechanism for tightly coupling the extent of muscle excitation to regulation of transcription (i.e. excitation-transcription coupling) (Chin 2005). Many Ca^{2+} -sensitive target genes have been identified in skeletal muscle. Downstream Ca^{2+} -sensitive target genes of varied expression levels between fibres include the nicotinic acetylcholine receptor (nAChR), glucose transporter 4 (GLUT4), sarcoplasmic reticulum (SR) Ca^{2+} ATPase (SERCA1), MyHC isoforms, oxidative enzymes, as well as genes that regulate mitochondrial biogenesis. Apart from the transcriptional regulation, it has been suggested that muscle fibre contractile characteristics might also be regulated by posttranslational modification of contractile proteins (Canepari et al. 2005). In addition to the role of Cn signalling in the determination of muscle fibre type characteristics, this phosphatase is known to play an important role in muscle hypertrophy (Dunn et al. 1999; Michel et al. 2004). Cn dephosphorylates the transcription factor nuclear factor of activated T cells (NFAT), enabling its nuclear translocation and DNA binding. The Cn-NFAT pathway has been linked to Ca^{2+} -induced skeletal muscle hypertrophy, at least in cultured skeletal muscle (Semsarian et al. 1999). With regard to skeletal muscle hypertrophy in animal models, the role of Cn remains controversial. Current thought suggests that hyperactivation of Cn alone is not sufficient to induce skeletal muscle hypertrophy but that the activation of accessory parallel signalling pathways for growth is required (Michel et al. 2004).

The size principle of motor recruitment—consequences for resistance exercise

As mentioned above, amplitude and duration of Ca^{2+} -transients in skeletal muscle as a function of MU firing frequency are decoded at the molecular level, resulting in expression changes of fibre type-specific functional gene modules. For mammals, it has been shown that for many activities there is a graded level of muscle recruitment that is driven by the different thresholds of the motor neurones: "the size principle of motor recruitment" (Denny-Brown and Pennybacker 1938; Henneman et al. 1965, 1974). The size principle predicts that MU recruitment is determined by force requirements. The majority of units are, in fact, recruited voluntarily in the order of increasing size (Monster and Chan 1977; Tanji and Kato 1973). Typically, small MUs are type I units, and unit size increases with progression through the fibre types: I < IIA < IIX (according to the MyHC isoform classification system in humans). Therefore, when low force is

required, only type I MUs will be active. Only when force is high will recruitment demand involvement of the larger MUs. It follows that slow muscle fibres are activated for low-force contractions and fast muscle fibres are additionally activated to supply greater force demands. Several researchers have shown that MU recruitment is completed by $\sim 50\%$ of maximum voluntary contraction (MVC) in small muscles [adductor pollicis (Kukulka and Clamann 1981), and first dorsal interosseus (De Luca et al. 1982a, b, 1996; Milner-Brown et al. 1973)], and 70–80% of MVC in large muscles [biceps (Kukulka and Clamann 1981), deltoid (De Luca et al. 1982a, b), and tibialis anterior (De Luca et al. 1996; Erim et al. 1996)]. The size principle applies when either slow-ramp force is exerted or constant low forces are compared with constant higher forces. The size principle of orderly recruitment is also preserved during exercise (Gollnick et al. 1974a, b; Vollestad and Blom 1985; Vollestad et al. 1984, 1992), whether the comparison is of sustained contractions at different intensities, or at different times during exercise at the same submaximal force sustained until exhaustion (Adam and De Luca 2003).

However, during some forms of voluntary or reflex contraction, motor neurone size may not be the sole factor determining excitation threshold: (1) Specialisation of the synaptic input among the motor neurones, and joint position have been shown to affect recruitment order in some cases; (2) MU rotation strategy has been demonstrated for sustained isometric contraction of biceps brachii at 10 but not at 40% of maximum; (3) In certain very rapid or sudden corrective movements—such as accelerations or sudden changes in direction—high-threshold units that do not participate in walking or even in running might be selectively recruited, as shown for the human short extensor muscle of the toes (Grimby 1984). With respect to ballistic movements, there is some controversy over the extent to which selective recruitment may occur (if at all) (Zehr and Sale 1994); (4) It has been suggested that selective recruitment of large MUs occurs during lengthening contractions (Nardone and Schieppati 1988; Nardone et al. 1989); (5) In certain cases of altered motor tasks, recruitment order might vary. Based on these results, it has been suggested that differential recruitment simply represents the existence of distinct, task-related subpopulations of motor units, rather than “violations” of the size principle (Burke 2002; Cope and Sokoloff 1999). Hence, when the same motor task is undertaken in exactly the same way, the order in which MUs are recruited remains fixed. This is of special importance, since it supports the notion that recruitment order is maintained during muscular fatigue. In particular, it has been shown that submaximal fatiguing contractions in the vastus lateralis muscle of humans lead to the monotonic decline in the recruitment threshold of all MUs and the progressive recruitment of new MUs, without change in the recruitment order. Thus, as the force capacity of continuously active muscle fibres declines progressively, increased excitation is required to keep the muscle output constant. The increased excita-

tion produces the recruitment of additional MUs. The recruited MUs thus become active at a lower torque level than their initial threshold, and the recruitment threshold continues to decrease in subsequent contractions as the force production of the active MUs continues to decrease (Adam and De Luca 2003). Indeed, it could be shown that the reduction in peak tetanic torque is linearly correlated with the decrease in the mean MU recruitment threshold at corresponding endurance times (beginning, middle, and end of fatiguing contractions) (Adam and De Luca 2003). However, these data do not prove a causal relationship between changes in muscle force output and changes in MU recruitment. Nonetheless, it can reasonably be assumed that the drop in peak tetanic torque that comes with muscular fatigue corresponds to an increase in the relative force requirement to sustain the target torque level.

Based on these findings, we propose a theoretical model of muscle fatigue and MU recruitment during resistance exercise progression (Fig. 4). Provided that exercise is performed to volitional muscular failure, the three different load magnitudes in Fig. 4a–c will lead to similar, i.e. “complete” MU recruitment (Fig. 4d), and, thus, to a similar stimulation of protein synthesis. Indeed, preliminary studies aimed at delineating the dose–response relationship between the intensity of exercise and the rates of muscle protein synthesis have shown that when the same total amount of ATP is turned over and recruitment is complete, exercise at 60, 75, and 90% of the 1RM results in exactly the same stimulation of muscle protein synthesis (Bowtell et al. 2003). Additionally, increases in tension above 65% cause no further stimulation in muscle protein synthesis (Bowtell et al. 2003). However, the different TUT until muscular failure imply different MU recruitment dynamics. As a consequence, distinct metabolic loads are inflicted on the exercising muscle in Fig. 4a–c. In Fig. 4a, progression through the fibre types I \rightarrow IIA \rightarrow IIX occurs fast (30 s). In contrast, in Fig. 4c, progression through the fibre types occurs relatively slowly (180 s). Consequently, in Fig. 4c the fibres’ capability in providing ATP through oxidative metabolism is more pronouncedly challenged than in Fig. 4a. This is due to the fact that with low resistive loads in % of the maximum voluntary torque (MVT), the threshold for complete recruitment is reached later. Thus, if a biceps exercise is assumed for the examples in Fig. 4, only in Fig. 4a the recruitment is predicted to be complete at start. In Fig. 4b, c, complete recruitment will be reached later. It follows directly, that the two strategies of MU recruitment and MU firing rate-coding for the increase in force output will vary to different extents for Fig. 4a–c. Consequently, the distinct pattern of MU recruitment and rate-coding in Fig. 4a–c will have a different impact on the previously described excitation–transcription coupling (Chin 2005).

In conclusion, we deduce that the TUT (x_9 , Table 1), volitional muscular failure (x_{10} , Table 1), number of sets (x_3 , Table 1), rest in-between sets (x_4 , Table 1), and rest in-between repetitions (x_8 , Table 1) are further mechano-

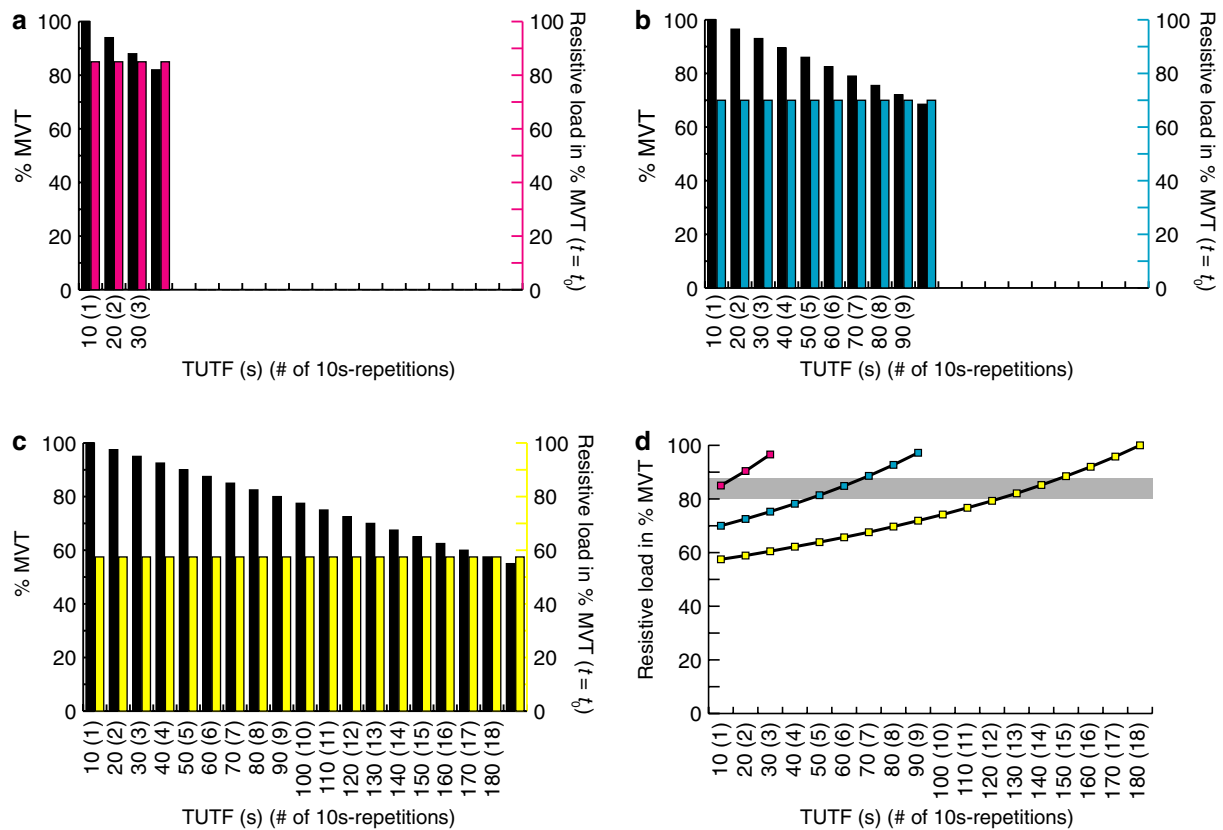


Fig. 4 Hypothetical model of muscle fatigue and its consequences for motor unit recruitment and, thus, for the metabolic load imposed on a skeletal muscle. In this simplistic model (e.g. biceps brachii performing 1 set of resistance exercise to volitional muscular failure), it is assumed that the drop in % of the maximum voluntary torque (MVT) in-between the repetitions (i.e. the “fatigue inroad”) occurs with constant percentage. The magnitude of fatigue inroad per repetition is assumed to be 6, 3.5, and 2.5% for high (magenta bars), intermediate (cyan bars), and low (yellow bars) resistive loads, respectively. The primary ordinates in **a–c** indicate the load magnitude expressed in % of the MVT achieved at a determinate joint angle. The abscissae in **a–d** indicate the total time (s) under tension until muscular failure (TUTF) or the corresponding number of repetitions when 10 s movement time per repetition is assumed. The secondary ordinates in **a–c** indicate the resistive load (i.e. the “training load”) in % of MVT at $t = t_0$. The ordinate in **d** indicates the resistive load in % of MVT through exercise progression, which corresponds to the relative resistive MVT. A high (magenta bars), intermediate (cyan bars), and low (yellow bars) resistive load is assumed in **a–c**, respec-

tively. These high, intermediate, and low resistive loads correspond to high, intermediate, and low % of MVT. Moreover, these resistive loads (“training loads”) remain unchanged during the same exercise set. However, the theoretical MVT (black bars) declines with exercise progression due to muscular fatigue, with a concurrent increase in motor unit (MU) recruitment and MU firing rate (see text for details). The time point of muscular failure corresponds to the point at which the theoretical MVT is below the MVT required to overcome the resistive load. Muscular failure occurs after 30 s (3 repetitions), 90 s (9 repetitions), and 180 s (18 repetitions) in examples **a–c**, respectively. In **d**, the hypothetical increase in relative resistive load (% MVT) for high (magenta), intermediate (cyan), and low (yellow) resistive loads [% MVT ($t = t_0$)] is plotted against the TUTF. The relative load [% MVT ($t = t_0$)] as exercise starts and the steepness with which it increases through exercise progression until voluntary failure, as a function of the muscle’s fibre type distribution, determines MU recruitment. The grey shaded area corresponds to the predicted resistive load in % of MVT at which MU recruitment for biceps brachii muscle is completed (see text for details)

biological determinants that affect MU recruitment and MU firing rate-coding, and, thus, excitation–transcription coupling. In addition, the number of exercise interventions per week (x_5 , Table 1) and the duration of the experimental period (x_6 , Table 1) should be reported for an estimation of the training’s stimulatory effect on muscle protein synthesis.

Molecular sensing of oxygen

Intramyocellular oxygen partial pressure has been suggested to drop with exercise, leading to “local hypoxia” (Gayeski et al. 1985). Local hypoxia, as determined by

measurement of oxygen saturation via assessment of myoglobin desaturation, has been shown to occur within 20 s of onset of exercise in human quadriceps muscle (Richardson et al. 1995). Additionally, oxygen saturation is reduced to a plateau with the onset of exercise (Richardson et al. 2001). Thus, it has been hypothesised that local hypoxic conditions may prevail in muscle even during low intensity exercise in normoxia (Hoppeler et al. 2003).

The partial pressure of cellular oxygen is sensed by a family of prolyl hydroxylases (Wenger et al. 2005). This enzyme family contains three members: prolyl-4-hydroxylase domain 1 (PHD1), PHD2, and PHD3, also known as hypoxia-inducible factor (HIF) prolyl hydroxylase 3

(HPH3), HPH2, and HPH1, respectively. The prolyl hydroxylases covalently modify HIF α subunits. HIF is a heterodimeric transcription factor consisting of one of three different oxygen-sensitive HIF α subunits and a common constitutive HIF β subunit. Under normoxic conditions, HIF α is hydroxylated. HIF α hydroxylation promotes von Hippel–Lindau (VHL) tumour suppressor protein binding to HIF α . Upon binding of VHL, HIF α is targeted for proteasomal destruction. Under hypoxic conditions, however, PHD activity (and thus HIF α hydroxylation) decreases. Thus, under hypoxic conditions, the high turnover rate of HIF α subunits enables the very rapid accumulation of HIF α . Following a further decrease in oxygen availability, the asparagine hydroxylase function of another enzyme, the factor inhibiting HIF (FIH), also becomes impaired, resulting in a decrease in HIF α C-terminal hydroxylation. This decrease in C-terminal HIF α hydroxylation enables the increased recruitment of the p300 and CREB-binding protein (p300/CBP) transcriptional coactivators, leading to the enhanced transcriptional activation of at least 70 different HIF effector genes (Wenger et al. 2005).

HIF effector genes regulate, e.g., oxygen supply, cellular metabolism, cell growth, and apoptosis (Wenger et al. 2005). Indeed, it has recently been shown that HIF1 α protein abundance increases in response to the acute exercise-induced increase in oxygen demand in human tissue (Ameln et al. 2005). Concurrently, no changes in HIF1 α mRNA abundance were detected in response to exercise. Thus, it was suggested that the observed increased HIF1 α protein abundance mainly depends on posttranscriptional mechanisms. In parallel with the exercise-induced stabilisation of HIF1 α protein, nuclear translocation of the protein was observed, together with an induced expression of the HIF regulated target genes. In particular, vascular endothelial factor (VEGF) and erythropoietin (EPO) mRNA abundance was increased with exercise (Ameln et al. 2005).

The role of HIF1 α in regulating skeletal muscle function has recently also been investigated with HIF1 α knockout mice (HIF1 α KO) (Mason et al. 2004). It was found that a significant exercise-induced increase in mRNA abundance of VEGF, GLUT4, muscle-specific phosphofructokinase (PFK-M), phosphoglycerate kinase (PGK), and lactate dehydrogenase-A (LDH-A) is missing in HIF1 α KO compared to wild-type (WT) mice. Also, changes in enzymatic activity in a number of key glycolytic enzymes are affected by deletion of HIF1 α . In particular, the activity of one of the key rate-limiting enzymes, PFK, is significantly lower following exercise in HIF1 α KO compared to WT mice. In HIF1 α KO, overall, a metabolic shift away from glycolysis towards oxidation was observed. This metabolic shift away from glycolysis towards oxidation increases swimming and uphill running exercise times in the HIF1 α KO. Conversely, repeated bouts of downhill running (eccentric exercise) results in greatly reduced exercise times and increased muscle damage in HIF1 α KO relative to WT mice. The authors hypothesised that the reduced exercise times with

eccentric exercise is due to the impaired glycolytic capacity in HIF1 α KO (Mason et al. 2004). This assumption was based on the previous findings that primarily fast-twitch glycolytic fibres are activated for eccentric contraction (Nardone and Schieppati 1988; Nardone et al. 1989).

Further evidence for the molecular regulation of skeletal muscle metabolic properties through regulation of fibre type functional modules has recently come from the engineering of a mouse capable of continuous running of up to twice the distance of a WT littermate (Wang et al. 2004). These authors uncovered peroxisome proliferator-activated receptor (PPAR) δ as the first transcription factor able to drive a metabolic fibre reprogramming following the targeted expression of an activated form of PPAR δ . PPAR δ produces profound and coordinated increases in oxidation enzymes, mitochondrial biogenesis, and production of specialised type I fibre contractile proteins—the three hallmarks for muscle fibre type switching. The muscle phenotypes described in that study are remarkably similar to those of transgenic mice expressing either Cn, CaM kinase, or peroxisome proliferator-activated receptor-gamma coactivator 1 α (PGC-1 α) (Lin et al. 2002; Naya et al. 2000; Wu et al. 2002). Thus, it was suggested that PPAR δ could be one of the hypothetical downstream transcription factors of these pathways (Wang et al. 2004). However, according to ligand and gain-of-function transgenic studies, PPAR δ needs to be activated in order to direct the muscle fibre switch. Indeed, in a recent report, simple overexpression of WT PPAR δ in muscle was found not to be sufficient to promote a fibre switch or obesity resistance, although certain oxidation enzymes are increased (Luquet et al. 2003). This supports the notion that the activating signal or ligand, but not the receptor, is limiting (Wang et al. 2004). Thus, PPAR δ activation, rather than merely an increase of PPAR δ levels, is an essential element for fibre switching and its associated functional manifestations. Two models have been proposed for the exercise-induced activation of endogenous PPAR δ (Wang et al. 2004). First, it is possible that exercise generates or increases endogenous ligands for PPAR δ as tissues are undergoing substantial increases in fatty-acid internalisation and oxidation. Fatty acids and their metabolites can activate PPAR δ . A second model is that exercise may induce the expression of PGC-1 α (Goto et al. 2000) and thereby activate PPAR δ . This is consistent with previous work in which it was shown that PGC-1 α physically associates with PPAR δ in muscle tissue and can powerfully activate it even in the absence of ligands (Wang et al. 2003). In line with these findings in rodent muscles, two recent studies show that one-legged knee extensor exercise of different duration and blood supply conditions induce transcriptional activation of the PGC-1 α gene in human skeletal muscle (Norrbom et al. 2004; Pilegaard et al. 2003). Impairment of blood supply following low-intensity exercise with superimposed vascular occlusion has also been shown to stimulate hypertrophy to a greater degree than exercise without occlusion (Abe et al. 2005; Moore et al. 2004; Takarada et al. 2002, 2004).

In summary, it can be derived that decreased oxygen supply to skeletal muscle during resistance exercise affects myocellular oxygen homeostasis. Oxygen supply to skeletal muscle is influenced by the magnitude of active and/or passive tension that is generated during exercise as well as by the modality of exercise (Vedsted et al. 2006). Magnitude of active and/or passive tension dictates to which extent blood flow is reduced. Modality of exercise, i.e. constant versus intermittent tension delivery, is a measure of how long and how many times blood flow is reduced. For example, during “eccentric only” resistance exercise (such as “negative chins” or “negative dips”), only lengthening contractions are performed. Negative chins/dips are performed by climbing into the top position using the legs and by lowering the body back down by the lengthening contractions of the respective target muscles. Consequently, while climbing into the top position, target muscle blood flow is less challenged than during the lowering phase. Thus, we conclude that load magnitude (x_1 , Table 1), TUT (x_9 , Table 1), contraction mode and fractional distribution per repetition (x_7 , Table 1), rest in-between repetitions (x_8 , Table 1), as well as ROM (x_{11} , Table 1), are significant mechano-biological determinants of myocellular oxygen homeostasis. Therefore, they need to be specified in resistance exercise protocols. It must also be emphasised that the total TUT does not necessarily just correspond to the number of repetitions times the time requirement for one repetition. If the exercise is performed to volitional muscular failure, the TUT corresponds to the TUT until failure (TUTF). TUTF includes an isometric contraction of maximal time duration at the end of exercise. Thus, TUTF may exceed the product of the number of repetitions times the time requirement for one repetition.

Recommendations for optimal exercise stimulus design and description

Skeletal muscle is a biological tissue. As such, it adapts to mechano-biological conditions that are present or missing. As described in this paper, these mechano-biological conditions lead to the induction of molecular and cellular determinants for the regulation of skeletal muscle girth, length, and fibre type. Thus, we propose to view (resistance) “exercise” as physiological tissue conditioning, i.e., the targeted delivery of muscular stimuli of quantitative and/or qualitative effect on the muscular phenotype. Whether and to which extent an adaptation occurs is subordinate to the “response matrix” of the respective subject (Fig. 1). Targeted delivery of adaptational stimuli precludes an accurate mechano-biological description of the condition. Unfortunately, most of the past and present resistance exercise prescriptions (Kramer and Ratamess 2004) still use ambiguous stimulus descriptors. This shortcoming makes it a priori virtually impossible to identify which stimuli lead to what kind of

adaptation. Consequently, no exercise prescriptions with specific effect, e.g. for the prevention and/or treatment of sarcopenia and metabolic diseases, exist. In order to allow for a dissection of the relative contributions of exercise stimuli and response matrix to muscular adaptation, we have identified and described a basic set of resistance exercise-related muscular conditions and termed them “mechano-biological determinants of muscular adaptation” (Table 1). While most of these determinants are scientifically tested, some are hypothetical (e.g. rest in-between repetitions). Understanding the relative contribution of these determinants in modulating the skeletal muscle phenotype should provide the basis for the design and prescription of training stimuli with specific quantitative and/or qualitative effect. We are aware of the fact that the use of 13 descriptors might raise some concerns with respect to practicability. However, we consider these determinants mandatory information for scientific experiments that aim at identifying causal connections between stimulus and response. Consequently, we strongly recommend to standardise the design and description of all future resistance exercise investigations by using the herein proposed basic set of 13 mechano-biological determinants (classical and new ones). Finally, although some of the newly proposed descriptors (e.g. TUT) are already employed by some exercise specialists, the identification of graded quantitative and qualitative effects of the most important descriptors should improve the effectiveness of future exercise recommendations.

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