Testosterone Improves the Regeneration of Old and Young Mouse Skeletal Muscle

Carlo Serra, Frances Tangherlini, Sara Rudy, Daniel Lee, Gianluca Toraldo, Nicolae Lucian Sandor, Anqi Zhang, Ravi Jasuja, and Shalender Bhasin

Department of Medicine, Section of Endocrinology, Diabetes and Nutrition, and the Boston Claude D. Pepper Older Americans Independence Center for Function Promoting Therapies, Boston University School of Medicine, Boston Medical Center, Boston, Massachusetts.

Address correspondence to Carlo Serra, PhD, Section of Endocrinology, Diabetes, and Nutrition, Boston Medical Center, 670 Albany Street, Boston, MA 02118. Email: cserra@bu.edu

MUSCLE aging is characterized by reduced muscle mass, strength, and by lower muscle regenerative potential. The progressive loss of muscle mass and functional capacity with aging predisposes to increased risk of falls, disability, and dependency in the elderly person (1,2). The growing geriatric population has prompted efforts to develop therapeutic interventions to mitigate muscle wasting (3,4). Many factors contribute to the multifactorial pathophysiology of muscle loss and functional decline, such as reduced physical activity, nutritional deficits, oxidative stress, inflammation, as well as age-related decline in anaerobic stimuli, like growth hormone (5), insulin-like growth factor-1 (IGF-1) (6), and testosterone (7).

Satellite cells are responsible for postnatal skeletal muscle growth and regeneration. Muscle repair recapitulates several aspects of the early steps of developmental myogenesis. There are two main stages of muscle regeneration: an initial degenerative phase, characterized by recruitment of inflammatory cells and necrosis of damaged muscle fibers, and the subsequent regenerative phase, characterized by the formation of new muscle fibers (8). Signals released either by necrotic fibers and inflammatory cells activate the satellite cells (8). Newly formed muscle fibers appear by 3–6 days postinjury, and by 10 days postinjury, most of the morphological appearance of damaged muscle is restored.

The early stage of muscle regeneration is characterized by the activation and proliferation of quiescent Pax7+ satellite cells that enter the cell cycle to generate new satellite cells and myoblasts that, in turn, migrate and fuse to form new myofibers (9,10). During muscle regeneration, satellite cells activation and proliferation are regulated by several extracellular stimuli, such as follistatin (11), IGF-1 (12), and hepatocyte growth factor (13).

Aged skeletal muscles show reduced satellite cell number due to the lower level of stimuli that sustain their self-renewal and activation to the higher level of cell apoptosis (14,15); these limits partially explain the lower regenerative capacity of old muscle (16–19). However, aged skeletal muscles reacquire a regular regenerative capacity when exposed to a “young” environment or when Notch signaling is experimentally reactivated (16,20).

Testosterone induces skeletal muscle hypertrophy due to protein accumulation and myonuclear accretion (21–23). This suggests that androgens stimulate satellite cell proliferation and myoblast fusion to preexisting fibers to increase the myonuclear number due to the growing need for protein synthesis (23,24). This hypothesis is supported by studies showing that testosterone treatment increases satellite cell number in human (21,25), in the androgen-dependent “levarter ani” muscle in rodents (26) and in vitro (27–29), and by the evidence that the androgen receptor (AR) plays a role on muscle cell proliferation and differentiation in vivo and in vitro (30–33). In addition, AR knockout mice showed more pronounced reduction of limb muscle mass compared with...
Muscle fiber–specific AR knockout mice (34–36), further supporting the idea of a direct effect of testosterone on satellite cells. Androgen deprivation induces systemic loss of muscle mass, strength, and voluntary running in young and old rodents, whereas subsequent testosterone supplementation reverses these effects (37–39).

Few reports to date have examined the effects of androgen administration on the muscle regeneration. They show variable effects depending on the type of injured muscle and the extent of muscle regeneration of castrated and intact rodents (40,41). For example, a recent report showed the beneficial effect of nandrolone decanoate supplementation on the number and size of regenerating fibers and on the level of expression of MyoD and IGF-1 postinjury (42). Because testosterone modulates several pathways involved in satellite cell activation, proliferation and aging, we evaluated here the effect of testosterone supplementation on the early phases of the muscle regeneration of young and old mice.

Methods

Mice, Castration, and Testosterone Treatment

Sexually mature 2- and 24-month-old male C57Bl/6J wild type mice, purchased from the Charles River Laboratories, Boston, MA, were acclimatized for at least 7 days before starting experiments. The mice were kept on a 12-hour light-dark cycle and given food and water ad libitum. The animals were handled according to the National Institute of Health guidelines for the care and use of laboratory animals, using protocols approved by the Committee for the Ethical Care and Use of Laboratory Animals of the Boston University.

The mice were randomly assigned to three experimental groups: eugonadal, sham-operated (Sham) implanted with empty silastic tubing; orchiectomized mice (Cast) implanted with empty silastic tubing; and orchiectomized mice implanted with silastic implants containing testosterone propionate (Tp) (Cast Tp). Surgical orchietomy was performed under anesthesia through a midline scrotal incision allowing bilateral access to the hemiscrotal content.

Testosterone supplementation was administered through 1.0 cm silastic implants (1.47 mm inner diameter, 1.96 mm outer diameter; Dow Corning Corp. Midland, MI) containing 8 mg Tp (Sigma–Aldrich, St Louis, MO, T1875) and sealed with silicone glue (Dow Corning Corp.). The implants were inserted subcutaneously through a small incision at the nape of the neck 12 days after orchietomy. Sham-operated eugonadal and Cast control mice were implanted with empty silastic tubing.

Serum total testosterone was measured using liquid chromatography-tandem mass spectrometry as described (43). For 24 month-old mice, serum T level (ng/dL; mean ± SD) was higher in Tp-treated mice (293 ± 447.9) than in Sham (345.4 ± 132.9) and Cast mice (23.74 ± 8.32; mean values 4 days postinjury). Similar testosterone levels were found in 2-month-old mice: Sham (296.9 ± 114.1), Cast (19.69 ± 3.48), and Cast Tp mice (3589 ± 308.1; mean values 4 days postinjury).

Muscle Injury

After 12 days of testosterone delivery, the left “tibialis anterior” muscle was injured under anesthesia using four 5 μL injections of 10 μM Naja nigricollis cardiotoxin (Accurate Chemical & Scientific Corporation, Westbury, NY) administered along the longitudinal axis of the muscle using a Hamilton syringe (model 725, Hamilton Company, Reno, NV) with a 30½ gauge needle. Contralateral tibialis anterior muscle was left intact as control. At the end of the treatment, mice were injected intraperitoneally with 5-bromo-2′-deoxyuridine (BrdU; 50 mg/kg body weight; Sigma–Aldrich, B5002) dissolved in sterile saline solution 5 hours before sacrifice. The tibialis anterior muscle was isolated, freed of visible connective tissue, and snap frozen in liquid nitrogen–cooled isopentane.

Antibodies

Primary antibodies used: mouse monoclonal to embryonic myosin heavy chain (emb-MyHC; F1.652, 1:100); rat monoclonal to BrdU (Abcam, Cambridge, MA, ab6326, 1:25); rat monoclonal to laminin 2 alpha (Abcam, ab11576, 1:100); chicken polyclonal to laminin (Abcam ab14055, 1:200); and rabbit polyclonal to neural cell adhesion molecule (NCAM; Millipore, Billerica, MA, AB5032, 1:200). F.1.652 hybridoma was developed by Dr Helen Blau and was obtained from the Developmental Studies Hybridoma Bank of the University of Iowa, Iowa City, IA. Secondary antibodies used: goat polyclonal to mouse Cy3 conjugated (Jackson Immunoresearch, West Grove, PA, 115-165-062, 1:400), goat polyclonal to rat FITC conjugated (Jackson Immunoresearch, 112-095-003, 1:200), goat polyclonal to rabbit Cy3 conjugated (Jackson Immunoresearch, 111-165-003, 1:200), and goat polyclonal to chicken fluorescein isothiocyanate (FITC) conjugated (Abcam, Cambridge, ab46969, 1:200).

Immunohistochemistry Analysis

Cryosections of the tibialis anterior, 6- to 8-μm-thick, tibialis anterior were fixed in 4% paraformaldehyde for 25 minutes at 4°C, permeabilized with 0.5% Triton X-100 for 15 minutes, and blocked in 5% normal goat serum (NGS) for 1 hour. Samples were incubated with primary antibodies overnight at 4°C in 1% bovine serum albumin (BSA)/1% NGS and then with secondary antibodies for 1 hour at room temperature. For the identification of BrdU+ nuclei, after fixation and permeabilization, muscle sections were incubated in 1 N HCl on ice for 10 minutes, 2 N HCl at 60°C for 5 minutes and then at room temperature for 15 minutes, washed with 0.1 M borate buffer for
12 minutes, incubated in 1% Triton X-100, 1 M glycine and 5% NGS for 45 minutes, in goat anti-mouse IgG (H + L) Fab fragment (Jackson Immunoresearch, 115-007-003, 1:100) in 5% NGS for 30 minutes, and then overnight at 4°C with primary antibodies diluted in 1% BSA and 1% NGS. Nuclei were counterstained with 4',6'-diamidino-2-phenylindole. Haematoxylin and Eosin staining was performed using a standard protocol. Pictures were acquired using a Nikon Eclipse TE2000-E microscope (Nikon Instruments Inc., Melville, NY). Regenerating area was identified as the region of the section showing the infiltrate of inflammatory cells and the centro-nucleated/emb-MyHC⁺ fibers and was measured using the SPOT imaging analysis software (Diagnostic Instruments, Sterling Heights, MI).

Data Analysis
Results are means ± SEM. One-way analysis of variance (ANOVA) was used in experiments with more than two independent groups. If overall ANOVA revealed significant difference, Student’s t test was used to analyze differences between groups. Chi-square test was used to compare frequency distribution of fiber cross-sectional area (CSA) among groups. p values ≤ 0.05 were considered statistically significant.

Results
Testosterone Administration Is Associated With Improved Muscle Regeneration in 24-Month-Old Mice
To evaluate the effect of testosterone supplementation on the muscle regeneration of aged skeletal muscle, 24-month-old C57BL/6J mice, described in Methods section, were sacrificed after BrdU injection 2, 4, and 9 days after cardiotoxin injury, to evaluate the rate and the extension of muscle regeneration and to capture the different phases of the regeneration process. Necrotic area and inflammatory infiltrates were found in the tibialis anterior 2 days after the cardiotoxin injury, indicating the induction of muscle regeneration in all groups of aged mice (Figure 1A). Double immunostaining for BrdU, an established marker of cell proliferation, and for NCAM, a recognized marker of satellite cells (44), revealed that 2 days after cardiotoxin injury, the control orchiectomized mice that were implanted with empty silastic implants had a lower number of proliferating BrdU⁺/NCAM⁺ satellite cells when compared with sham-operated mice (Figure 1B and C). However, the number of proliferating satellite cells was restored in orchiectomized mice treated with testosterone (Figure 1C). Similar data were obtained 4 days after cardiotoxin-induced injury in that the number of BrdU⁺/NCAM⁺ satellite cells in orchiectomized mice was still lower than in sham-operated mice, whereas testosterone rescued this deficit (Figure 1D and E).

The orchiectomized mice exhibited a smaller CSA of regenerating fibers immunostained for emb-MyHC, a marker of regenerating muscle fibers 4 days after cardiotoxin-induced injury (Figure 2A and B). The control orchiectomized mice implanted with empty silastic implants had significantly lower mean CSA of emb-MyHC⁺ muscle fibers than testosterone-treated orchiectomized mice and intact sham-operated mice. Consistent with these findings, the distribution of the muscle fibers in testosterone-treated orchiectomized mice (Figure 2C) is shifted toward the right compared with control orchiectomized mice implanted with empty silastic implants (chi-square p value for trend <.001). Thus, testosterone supplementation in orchiectomized mice rescued the CSA of regenerating muscle fibers to that seen in sham-operated intact mice. However, even though testosterone-treated orchiectomized mice showed increased number of emb-MyHC⁺ muscle fibers, the number of emb-MyHC⁺ muscle fibers was not significantly different between orchiectomized placebo-control mice (implanted with empty drug delivery device) and intact sham-operated mice (Figure 2D). These data suggest that testosterone predominantly enhances the formation of new muscle fibers that have larger CSA than those in orchiectomized mice.

By 9 days after cardiotoxin-induced injury, the muscle regeneration was nearly complete. At this time, we quantitated the number of centro-nucleated fibers (CNFs) in the regenerating tibialis anterior muscle. We found that neither the number (Figure 3A) nor the CSA (Figure 3B) of CNFs differed significantly among the three groups of mice. These data suggest that neither orchiectomy nor testosterone supplementation affects the late steps of muscle regeneration.

Testosterone Administration Is Associated With Improved Muscle Regeneration in 2-Month-Old Mice
To confirm the effect of testosterone on the muscle regeneration of old skeletal muscle we found, we repeated the same experimental protocol that was used for the aged mice in 2-month-old mice. Two days after cardiotoxin-induced injury, the mice of all experimental groups showed consistent muscle regeneration (Figure 4A). Consistent to what shown in aged mice, orchiectomized mice that received empty silastic implants had fewer proliferating satellite cells in comparison to sham-operated eugonadal controls and testosterone-treated orchiectomized mice (Figure 4B and C). Similarly, 4 days after injury, orchiectomized mice had a lower number of BrdU⁺/NCAM⁺ satellite cells than sham-operated eugonadal controls and testosterone-treated orchiectomized mice (Figure 4D and E). However, in both cases, the differences were not statistically significant.

The CSA of emb-MyHC⁺ muscle fibers was significantly smaller in the control orchiectomized mice implanted with empty drug delivery device than in testosterone-treated orchiectomized mice and sham-operated eugonadal controls (Figure 5A and B). In addition, similar to aged mice, the distribution of the muscle fiber diameters was shifted.
Figure 1. Effect of testosterone supplementation on satellite cell proliferation in aged skeletal muscle 2 and 4 days postinjury. Hematoxylin & Eosin staining of the tibialis anterior muscle 2 days postinjury; necrotic areas are present in damaged muscle (A). Representative image showing BrdU (green) and neural cell adhesion molecule (NCAM; red) staining 2 days postinjury (B). Testosterone supplementation in orchiectomized mice increased the number of the BrdU+/NCAM+ satellite cells when normalized for the extension of regenerating area, indicating a positive effect of androgens on satellite cell activation and/or proliferation in the aged skeletal muscle 2 days postinjury (C). Similar results are showed 4 days postinjury, when again castration is associated with reduced number of proliferating satellite cells, a deficit rescued by testosterone administration (D-E). Nuclei were counterstained with 4′,6′-diamidino-2-phenylindole (blue). In (B) and (D), arrows indicate BrdU+/NCAM+ satellite cells. Results are means ± SEM (n = 4 per group). *p < .05. Representative critical values for t test statistic: Figure C, Sham versus Cast: t = 3.117, df = 4.

toward right in testosterone-treated orchiectomized mice compared with control orchiectomized mice receiving empty silastic implants (Figure 5C). The number of emb-MyHC+ muscle fibers per unit regenerating area was greater in testosterone-treated orchiectomized mice than in castrated mice implanted with empty silastic implants (Figure 5D), although values do not reached statistical significance. As in the aged mice, CNFs number and mean CSA of CNFs were not significantly different among the three treatment groups 9 days after injury (Figure 6A and B).

**DISCUSSION**

We show here that testosterone supplementation modestly stimulated several early steps in the muscle regeneration that followed muscle injury created by the injection of cardiotoxin. Concordant changes were observed in the 2-month-old mice and the 24-month-old mice. Lowering of testosterone concentrations by surgical orchiectomy was associated with fewer BrdU+/NCAM+ proliferating myoblasts than in intact eugonadal age-matched controls. Conversely, testosterone-treated young and old...
mice had higher numbers of proliferating BrdU+/NCAM+ myoblasts 2 and 4 days after injury in comparison to control orchiectomized mice that received empty implants, although the difference in the number of BrdU+/NCAM+ cells of 2-month-old testosterone-treated mice did not reach statistical significance when compared with the corresponding values of age-matched orchiectomized mice treated with empty drug delivery devices, in part because the number of BrdU+/NCAM+ cells per unit area was small, consistent with the limited regenerating area. Similarly, we found an increased number of emb-MyHC+ regenerating fibers 4 days after injury in testosterone-treated old mice compared with orchiectomized mice treated with the empty silastic tubing, suggesting that the increased number of proliferating BrdU+/NCAM+ myoblasts resulted in greater differentiation rates of new myofibers in response to testosterone administration. Testosterone induced the formation of larger emb-MyHC+ fibers that had greater CSA than those in control orchiectomized mice. In addition, the rightward shift in fiber size distribution shown by testosterone-treated mice indicates that the treatment improved muscle regeneration. Taken together, our data suggest
that testosterone administration contributes to the expansion of myoblasts during the early stages of muscle regeneration, allowing the formation of bigger regenerating myofibers. In addition, testosterone administration induced the hypertrophy of the early regenerating fibers, indicating a potential positive effect of testosterone on muscle protein accretion in this period of muscle regeneration. By 9 days after injury, neither the young and nor the old regenerating muscle revealed statistically significant differences in the indices of muscle regeneration at this late time point; the number and CSA of regenerated CNFs was similar in the three intervention groups. Moreover, the difference in the CSA of the regenerating fibers 9 days postinjury between young and old mice, for example, between the Sham mice of the two classes of age, is in line with the general agreement that a lower level of satellite cell activation and proliferation

Figure 4. Effect of testosterone supplementation on satellite cell proliferation in young skeletal muscle 2 and 4 days postinjury. Hematoxylin & Eosin staining of the tibialis anterior muscle 2 days postinjury (A). Representative image showing BrdU (green) and neural cell adhesion molecule (NCAM; red) staining (B). Testosterone supplementation restored the number of BrdU+/NCAM+ satellite cells, indicating a positive effect of androgens on satellite cell activation and/or proliferation in the young skeletal muscle 2 days postinjury (C). As in aged mice, castrated mice showed a trend toward reduced number of proliferating satellite cells, a deficit rescued by testosterone administration 4 days postinjury (D-E). The number of BrdU+/NCAM+ satellite cells per unit area did not differ significantly between controls and testosterone-treated orchiectomized mice in figures C and E. Nuclei were counterstained with 4′,6′-diamidino-2-phenylindole (blue). In (B) and (D), arrows indicate BrdU+/NCAM+ satellite cells. Results are means ± SEM (n = 4 per group).
reduces and/or delays the overall rate of regeneration of the aged muscle. Thus, in this mouse model of cardiotoxin-induced muscle injury, the effects of testosterone appear limited to the early stages of muscle regeneration.

Previous studies have reported that aging is associated with a reduced intrinsic ability of the skeletal muscle to repair after an injury, presumably due to the reduced levels of circulating factors that regulate satellite cell activation and myoblast proliferation (17,20,45), the accumulation of cytokines that inhibit muscle differentiation, for example, TGFβ (18), as well as the intrinsic changes in the satellite cell niche (46,47). In this scenario, the aging generates a state of lower proliferation for muscle stem cells with the consequent reduced regenerative performance, maybe as an adaptation to reduce the risk of cell transformation.

We did not find major qualitative or significant quantitative differences in the temporal profile of muscle regeneration in response to testosterone administration between the

Figure 5. Effect of testosterone supplementation on muscle regeneration in young skeletal muscle 4 days postinjury. Representative image showing regenerating fibers positive for emb-MyHC (red) and for laminin (green) (A). Testosterone treatment increases the cross-sectional area (B) of emb-MyHC+ muscle fibers 4 days postinjury, when compared with Cast mice, as also corroborated by the right shift showed by the distribution by frequency classes of fiber diameter (C) (p value for chi-square analysis <.001). The number of emb-MyHC+ muscle fibers when normalized for the regenerating area did not differ significantly between testosterone-treated orchiectomized mice and Sham or Cast mice (D). Results are means ± SEM (n = 4/group). *p < .05; **p < .01. Representative critical values for t test statistic: Figure B, Sham versus Cast: t = 4.590, df = 6.

Figure 6. Effect of testosterone supplementation on muscle regeneration in young skeletal muscle 9 days postinjury. As for the aged mice, no significant difference was found in the centro-nucleated fibers number, normalized for regeneration area, and in the cross-sectional area of regenerating fibers (A–B).
2-month-old and the 24-month-old mice. In both the young and the old mice, testosterone administration increased the number of proliferating BrdU+/NCAM+ satellite cells and promoted the formation of larger emb-MyHC+ regenerating fibers. As our experimental animals were healthy and living in a pathogen-free environment, these findings may not apply to the community-dwelling older humans who often have a cluster of comorbid conditions, which may directly or indirectly affect the regenerative response to injury, or to still older mice.

Only a few studies have investigated the role of androgens during muscle regeneration, using different types of muscle injury. In one study, nadroline decanoate administration after a muscle contusion injury in intact male rats was associated with increased muscle twitch and tetanic force 14 days after injury (40) and improved growth of regenerating slow twitch muscle 25 days postinjury (41). However, the early stages of muscle regeneration were not investigated, as we have done in this study. In another study, nadroline decanoate increased the expression of IGF-1, MyoD, and cyclin D1 after bupivacaine-induced muscle injury in the tibialis anterior muscle of castrated mice (42), leading the investigators to suggest that nadroline accelerates healing processes through an IGF-1–mediated mechanism. Intramuscular IGF-1 signaling also plays an important role in mediating testosterone’s anabolic effect on the skeletal muscle (29,36,48). IGF-1 enhances myoblast proliferation and differentiation (49–52) and intramuscular IGF-1 receptor (IGF-1R) signaling mechanisms seem essential for mediating the effects of testosterone on the proliferation of muscle progenitor cells and the differentiation of myoblasts (42). Whether the observed effects of testosterone in the present study are also mediated through an IGF-1R–dependent mechanism requires further investigation.

Testosterone administration dose dependently induces skeletal muscle fiber hypertrophy and increases the number of satellite cells and myonuclei in young and older men. Muscle hypertrophy generated by testosterone administration seems to involve not only muscle protein accretion but also myoblast expansion and fusion with preexisting fibers (21–23,25). The studies in the AR knockout male mice (34–36,53) also support a direct role for androgens in inducing satellite cell activation and myoblast proliferation. We focused our investigation to the satellite cells and to the regenerating muscle fibers. However, it is possible that testosterone may also affect other cell types residing within the skeletal muscle such as the fibroblasts, blood vessels, and neural structures. For example, AR is expressed by muscle fibroblasts, vascular endothelial cells, and in nerve end plates. The role of AR signaling in these cell types should be investigated because, for example, fibroblasts regulate satellite cell activation and muscle regeneration (54). In addition, testosterone modulates the activity of immune cells that play an essential role in muscle regeneration (8). Upregulation of serum TGFβ1 and TGFβ family members in response to muscle injury have been reported to contribute to decreased muscle regeneration in older animals (55–57). Testosterone has been shown to block TGFβ signaling through cross-communication of Wnt/β-catenin signaling through follistatin (58). Whether the effects of testosterone on muscle regeneration are also mediated through the modulation of the TGFβ and Wnt pathways needs further investigation.

The regenerative response to different modes of muscle injury and possibly the effects of testosterone on skeletal muscle regeneration in response to different modes of injury may vary (40–42). Our data indicate that testosterone stimulates myoblast proliferation and the formation of larger new fibers in the early phases of regenerative response to cardiotoxin-induced injury in both the young and the old mice. The clinical usefulness of androgens to promote muscle regeneration in humans and the underlying mechanisms by which testosterone modulates the early regenerative response need further investigation.

Funding
This work was supported by grants from the Department of Medicine, Evans Medical Foundation, and the Clinical and Translational Science Institute (grant UL1RR025771) of the Boston University to C.S. and R.J., by the National Institute of Health (grant 5R01DK070534-07 to S.B.), and by the Boston Claude D. Pepper Older Americans Independence Center (grant P30AG031679 to C.S. and S.B.).

References
11. Armand AS, Della Gaspera B, Launay T, Charbonnier F, Gallien CL, Chanoine C. Expression and neural control of follistatin versus myo-
statin genes during regeneration of mouse soleus. Dev Dyn. 2003;227:
256–265.

enhancement of muscle regeneration and prevention of fibrosis. Muscle

13. Allen RE, Sheehan SM, Taylor RG, Kendall TL, Rice GM. Hepato-
cyte growth factor activates quiescent skeletal muscle satellite cells in

14. Renault V, Thornell LE, Eriksson PO, Butler-Browne G, Moulay V.
Regenerative potential of human skeletal muscle during aging. Aging


restoration of regenerative potential to aged muscle. Science. 2003;
302:1575–1577.

17. Carlson ME, Hsu M, Conboy IM. Imbalance between pSmad3 and
Notch induces CDK inhibitors in old muscle stem cells. Nature. 2008;
454:528–532.

Wnt in the systemic regulation and aging of satellite cell responses.

induces the differentiation of myogenic precursors into fibroblastic cells in

TA. Rejuvenation of aged progenitor cells by exposure to a young

21. Sinha-Hikim I, Cornford M, Gaytan H, Lee ML, Bhasin S. Effects of
testosterone supplementation on skeletal muscle fiber hypertrophy and
satellite cells in community-dwelling older men. J Clin Endocrinol
Metab. 2006;91:3024–3033.

22. Kadi F, Bonnerud P, Eriksson A, Thornell LE. The expression of
androgen receptors in human neck and limb muscles: effects of training
and self-administration of androgenic-anabolic steroids. Histochem

increase in muscle size in healthy young men is associated with
muscle fiber hypertrophy. Am J Physiol Endocrinol Metab. 2002;283:
E154–164.

24. Kadi F, Eriksson A, Holmner S, Thornell LE. Effects of anabolic
steroids on the muscle cells of strength-trained athletes. Med Sci

muscle hypertrophy is associated with an increase in satellite cell
number in healthy, young men. Am J Physiol Endocrinol Metab. 2003;

26. Joubert Y, Tobin C. Testosterone treatment results in quiescent satellite
cells being activated and recruited into cell cycle in rat levator ani

27. Powers ML, Fiorini JR. A direct effect of testosterone on muscle cells

28. Dowmit ME, Cook DR, Merkel RA. Testosterone up-regulates andro-
gen receptors and decreases differentiation of porcine myogenic sar-

29. Serra C, Bhasin S, Tangerlini F, et al. The role of GH and IGF-I in
mediating anabolic effects of testosterone on androgen-responsive

30. Lee DK. Androgen receptor enhances myogenin expression and accelerates
differentiation. Biochem Biophys Res Commun. 2002;294:
408–413.

myoblastoma cell differentiation and proliferation is stimulated by
androgens and associated with a modulation of myostatin and Pax7

32. Sinha-Hikim I, Taylor WE, Gonzalez-Cadavid NF, Zheng W, Bhasin S.
Androgen receptor in human skeletal muscle and cultured muscle
satellite cells: up-regulation by androgen treatment. J Clin Endocrinol
Metab. 2004;89:5245–5255.

33. Singh R, Artaza JN, Taylor WE, Gonzalez-Cadavid NF, Bhasin S.
Androgens stimulate myogenic differentiation and inhibit adipogenesis
in C3H 10T1/2 pluripotent cells through an androgen-receptor-mediated

development and function in male, but not female, genomic androgen

myocytes contributes to the maintenance of muscle mass and fiber
type regulation but not to muscle strength or fatigue. Endocrinology.
2009;150:3558–3566.

controls the strength but not the mass of limb muscles. Proc Natl Acad

37. Axell AM, MacLean HE, Plant DR, et al. Continuous testosterone
administration prevents skeletal muscle atrophy and enhances resis-
tance to fatigue in orchidectomized male mice. Am J Physiol Endor-
clin Metab. 2006;291:E506–E516.

supplementation reverses sarcopenia in aging through regulation of
myostatin, c-Jun NH2-terminal kinase, Notch, and Akt signaling path-

39. bebunco J, Eash JK, Li C, Ma Q, Glass DJ. Voluntary running, skeletal
muscle gene expression and signaling inversely regulated by orchidec-
tomy and testosterone replacement. Am J Physiol Endocrinol Metab.

40. Beiner JM, Joki P, Cholewicki J, Panjabi MM. The effect of anabolic
steroids and corticosteroids on healing of muscle contusion injury. Am

41. Ferré A, Noirel P, Page CL, Salah IB, Daegelen D, Rieu M. Effects of
anabolic/androgenic steroids on regenerating skeletal muscles in the

42. White JP, Baltgalvis KA, Sato S, Wilson LB, Carson JA. Effect of
nandrolone decanoate administration on recovery from bupivacaine-
induced muscle injury. J Appl Physiol. 2009;107:
1420–1430.

43. Sir-Petermann T, Codner E, Perez Y, et al. Metabolic and reproduct-
ive features before and during puberty in daughters of women with
polycystic ovary syndrome. J Clin Endocrinol Metab. 2009;94:
1923–1930.

44. Ila I, Leon-Monzon M, Dalakas MC. Regenerating and denervated
human muscle fibers and satellite cells express neural cell adhesion
molecule recognized by monoclonal antibodies to natural killer cells.

45. Carlson ME, Suetta C, Conboy MJ, et al. Molecular aging and reju-
venation of human muscle stem cells. EMBO Mol Med. 2009;1:
381–391.

46. Carlson ME, Conboy IM. Loss of stem cell regenerative capacity

47. Collins CA, Zammit PS, Ruiz AP, Morgan JE, Partridge TA. A popula-
tion of myogenic stem cells that survives skeletal muscle aging. Stem

48. Lewis MJ, Horvitz GD, Clemmons DR, Fournier M. Role of IGF-1 and
IGF-binding proteins within diaphragm muscle in modulating the
effects of nandrolone. Am J Physiol Endocrinol Metab. 2002;282:
E483–490.

49. Coolican SA, Samuel DS, Ewton DZ, McWade FJ, Florini JR. The
mitogenic and myogenic actions of insulin-like growth factors
utilize distinct signaling pathways. J Biol Chem. 1997;272:
6653–6662.


