Myostatin negatively regulates satellite cell activation and self-renewal

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Satellite cells are quiescent muscle stem cells that promote postnatal muscle growth and repair. Here we show that myostatin, a TGF-β member, signals satellite cell quiescence and also negatively regulates satellite cell self-renewal. BrdU labeling in vivo revealed that, among the Myostatin-deficient satellite cells, higher numbers of satellite cells are activated as compared with wild type. In contrast, addition of Myostatin to myofiber explant cultures inhibits satellite cell activation. Cell cycle analysis confirms that Myostatin up-regulated p21, a Cdk inhibitor, and decreased the levels and activity of Cdk2 protein in satellite cells. Hence, Myostatin negatively regulates the G1 to S progression and thus maintains the quiescent status of satellite cells. Immunohistochemical analysis with CD34 antibodies indicates that there is an increased number of satellite cells per unit length of freshly isolated Mstn−/− muscle fibers. Determination of proliferation rate suggests that this elevation in satellite cell number could be due to increased self-renewal and delayed expression of the differentiation gene (myogenin) in Mstn−/− adult myoblasts. Taken together, these results suggest that Myostatin is a potent negative regulator of satellite cell activation and thus signals the quiescence of satellite cells.

Introduction

Muscle satellite cells are a distinct lineage of myogenic progenitors responsible for postnatal muscle growth. Satellite cells are mitotically quiescent and are known to express several myogenic markers. When stimulated by damage to the muscle, satellite cells are activated to reenter the cell cycle and express myogenic regulatory factors. The resulting myoblasts subsequently differentiate and fuse to form new replacement myofibers (Bischoff, 1989). The total number of quiescent satellite cells remains constant over repeated cycles of degeneration and regeneration, suggesting that the steady-state satellite cell population is maintained by self-renewal (Yoshida et al., 1998; Schmalbruch and Lewis, 2000; Heslop et al., 2001).

Several different growth factors, including HGFs (Maina et al., 1996) and FGFs (Flanagan-Steet et al., 2000), have been shown to affect the postnatal muscle growth by regulating satellite cell activation. Nowhere is the influence of growth factors on skeletal muscle development more dramatically demonstrated than by the remarkable widespread skeletal muscle mass of myostatin-deficient animals (Kambadur et al., 1997; McPherron and Lee, 1997; McPherron et al., 1997).

Myostatin is a secreted growth and differentiation factor (GDF-8) that belongs to the TGF-β superfamily and shares several features with other members. The myostatin-null mice and cattle with mutation in myostatin showed increased muscling due to generalized hyperplasia and, to a lesser extent, hypertrophy of muscle. The phenotype of myostatin mutant cattle and mice has clearly established that Myostatin negatively regulates muscle growth. Consistent with this, work from our laboratory as well as other published results suggest that Myostatin inhibits myoblast proliferation (Thomas et al., 2000; Zhu et al., 2000; Taylor et al., 2001). Investigations into mechanisms revealed that one of the targets of Myostatin signaling is the cell cycle. In cell culture studies, Myostatin inhibited the progression of myoblasts at G1 and G2 phases of the cell cycle. This cell cycle arrest was shown to be mediated through p21, which was the only Cdk inhibitor to be up-regulated by Myostatin. In addition, Myostatin treatment also specifically reduced levels of Cdk2, the combination of which resulted in markedly reduced activity of Cdk2. This, in turn, led to the hypophosphorylation of retinoblastoma protein and thus inhibition of progression into S phase of the cell cycle (Thomas et al., 2000).
Several studies have implicated a role for Myostatin postnatally in muscle misuse and wasting. Serum concentrations of Myostatin have been reported to be increased in HIV-infected men with weight loss, suggesting that increased Myostatin levels may contribute to the pathophysiology of muscle wasting during HIV infection (Gonzalez-Cadavid et al., 1998). Similarly, a study by Wehling et al. (2000) and Carlson et al. (1999) indicated that atrophy-related muscle loss due to hind limb suspension in mice was associated with increased Myostatin levels in m. plantaris. One of the explanations for the elevated levels of Myostatin in hind limb unloading is that Myostatin may function as an inhibitor of satellite cell proliferation. Indeed, this is supported by the fact that myostatin-null mice display muscle hypertrophy and increased postnatal muscle growth, both of which have been related to increased satellite cell activity (Grounds and Yablonka-Reuveni, 1993; Mesires and Doumit, 2002). This circumstantial evidence led us to speculate that Myostatin may have a role in postnatal muscle growth by controlling the satellite cell quiescence. Here we report that Myostatin is expressed in satellite cells and that lack of myostatin in mice leads to increased number of satellite cells per unit length of muscle fiber. We also show that in myostatin-null mice, a higher percentage of the satellite cells are in an activated state.

**Figure 1.** Myostatin is expressed in satellite cells. M. tibialis anterior was serially sectioned and immunostained with antibodies specific for (A) Myostatin and (B) Pax7. The myonuclei were stained with DAPI. The same muscle was used for in situ and probed for Myostatin (C) and Pax7 transcripts (D). Arrows indicate the stained satellite cells, and DAPI-stained myonuclei are also shown in the insets (B and D). (E) Micrograph showing in situ hybridization performed with myostatin antisense probe on muscle sections of myostatin-null mouse. (F, i) Agarose gel electrophoresis of RT-PCR products derived from satellite cell total RNA. Primers specific for 3′ region of myostatin amplify the expected 515-bp product in a combined RT-PCR reaction (Mstn). Amplicons are not detected in the absence of template (Negative control). Pax7 was amplified with primers designed to produce a 571-bp product (Pax7), and CD34 splice variants were also PCR amplified from the same RT reaction (CD34 splice variants). 1-kb plus DNA ladder is shown. (F, ii) Western blot showing the presence of the full-length 52-kD Myostatin protein in satellite cell protein extract. Bars, 10 μm.
state as compared with wild-type mice. Therefore, the observed hypertrophy phenotype and increased postnatal muscle growth in myostatin-null mice could be due to increased number and activation of satellite cells.

Results
Myostatin is expressed in satellite cells and in the adult myoblasts
To assess if Myostatin is present in satellite cells, we performed immunocytochemistry and in situ hybridizations on serial cross sections of m. tibialis anterior. Satellite cells on fibers were identified by positive immunostaining with anti-Pax7 antibodies (Fig. 1 B). In the serial sections, the Pax7-positive satellite cells were also positive for Myostatin immunostaining (Fig. 1 A). Similarly, in situ hybridization studies with myostatin and Pax7 digoxigenin-labeled probes revealed that Pax7-positive satellite cells (Fig. 1 D) also express abundant levels of myostatin mRNA (Fig. 1 C). Expression of myostatin mRNA is not detected in the muscle sections derived from myostatin knockout mice, confirming the specificity of the myostatin probe (Fig. 1 E). Therefore, it is convincing that the satellite cells attached to the muscle fibers express Myostatin. To further prove that myostatin is present in satellite cells, RT-PCR and Western blot analysis were performed on the total RNA and protein extracts, respectively, of quiescent satellite cells isolated by Percoll gradient. The PCR primers designed to specifically amplify the processed portion of myostatin do indeed amplify the expected 515-bp product from the cDNA pool derived from satellite cell RNA (Fig. 1 F, i). Furthermore, anti-Myostatin antibodies specifically recognized the full-length Myostatin in the protein extracts, confirming that the satellite cells do express Myostatin (Fig. 1 F, ii). To demonstrate that the cells isolated by Percoll gradient techniques are indeed satellite cells, independent PCR reactions were performed with Pax7 and CD34 primers (satellite cell–specific markers). The results show that RT reactions indeed contained Pax7 and CD34 splice variant cDNAs (Fig. 1 F, i).

To gain insight into the role of Myostatin in satellite cell activation and proliferation, low passage number primary cultures were isolated from hind limb muscles of 8-wk-old myostatin-null and wild-type mice to generate highly purified satellite cell–derived cultures and preclude the inclusion of neonatal myoblasts (see Materials and methods). Previous work has demonstrated that proliferating primary adult myoblasts derived from satellite cells express myoblast markers such as MyoD, Desmin, and adhesion molecules, but not terminal differentiation markers such as Myogenin or Myosin heavy chain. Therefore, to characterize the cultures used in this experiment, we immunostained the adult myoblast cultures with MyoD- and M-cadherin–specific antibodies. Over 95% of proliferating primary cells derived from the hind limb muscles of wild-type and myostatin knockout mice expressed M-cadherin and MyoD, indicating that the isolation procedure generated highly purified cultures of myogenic cells from both wild-type and myostatin knockout mice (Fig. 2, A and B). Proliferating myogenic precursor cells in vivo, and myoblasts in vitro, express the intermediate filament Desmin, whereas, satellite cells do not express Desmin (George-Weinstein et al., 1993). Hence, primary cultures were immunostained with antibodies reactive with Desmin to assess their developmental status. As shown in Fig. 2 C, uniform expression of Desmin is detected, indicat-
ing that the primary cultures are all activated myogenic precursor cells. It has been recently demonstrated that expression of CD34 protein is associated with skeletal muscle precursors but not with the differentiated fibers. To rule out any myogenic differentiation during the isolation procedure of primary cultures, the cells were immunostained with anti-CD34 antibodies. As shown in Fig. 2 D, almost all of the cultured cells were immunopositive for CD34, indicating that under the culturing conditions, the myogenic precursor cells are in active proliferation status (Beauchamp et al., 2000). The negative control did not show any nonspecific background (Fig. 2 E).

Three independent techniques, RT-PCR, Western blotting, and immunocytochemistry, were used to ascertain the expression of myostatin in adult myoblasts. As shown in Fig. 3 A, the PCR reaction amplified the expected size fragment of 1.2-kb myostatin ORF. In addition, anti-Myostatin antibodies specifically recognized Myostatin on the Western blot performed on the protein extracts derived from the adult myoblasts (Fig. 3 B). To further confirm the expression of Myostatin, primary cultures were immunostained with Myostatin antibodies. As shown in Fig. 3 C (i and ii), virtually all the cultured myoblasts showed staining with Myostatin antibodies. Thus, it is clear from these results that Myostatin is expressed in satellite cells and in the adult myoblasts derived from satellite cells.

**Increased activation of satellite cells in the absence of Myostatin in vivo**

Using single-cell PCR analysis, Cornelison and Wold (1997) have previously speculated that Myostatin is an important regulator of the quiescent state of satellite cells. Thus, to determine the involvement of Myostatin in satellite cell quiescence, the cell cycle phase of the satellite cells attached to myofibers in vivo in wild-type and myostatin knockout mice was assessed. Both wild-type and myostatin knockout mice were pulsed with BrdU, and satellite cells from m. tibialis anterior muscle were isolated by Percoll gra-
Characterization of satellite cells with M-cadherin and CD34 immunostaining revealed that the cultures were >95% pure and that there is no difference in the yield or purity between wild-type and myostatin knockout mice (unpublished data). BrdU-positive satellite cells were identified by immunostaining. As shown in Fig. 4 A (i), ~18% of satellite cells isolated from 4-wk-old wild-type mice were labeled with BrdU. However, in myostatin knockout mice, a significantly higher number (33%) of satellite cells was activated to enter S phase. In 8-wk-old wild-type muscle, 8% of the satellite cells isolated were BrdU positive, whereas in the myostatin-null muscle, 15% of the satellite cells were BrdU positive. When the same experiment was repeated on older mice (6 mo old), only 3.2% of the wild-type satellite cells were labeled with BrdU.
had incorporated BrdU. A threefold higher number, ~10% of Myostatin-deficient satellite cells, had incorporated BrdU (Fig. 4 A i), indicating that irrespective of the age, lack of functional Myostatin results in an increased number of activated satellite cells in uninjured muscle. To investigate the role of Myostatin in inhibiting satellite cell activation ex vivo, recombinant Myostatin was added to single isolated fibers in varying concentrations, and satellite cells were left to detach from the fiber and migrate to the well. Fig. 4 A (ii) graphically demonstrates that on average, 80% of the wells had identifiable satellite cells, which were subsequently differentiated in low serum to show they were myogenic. In comparison, in media containing 1 μg/ml of Myostatin, only 37% of fibers had migrated myogenic precursor cells in the well, while media with increased amounts of Myostatin, 2 and 5 μg/ml, had 12 and 0% migrated myogenic cells, respectively. These results confirm that a lack of myostatin expression leads to increased activation of satellite cells in vivo, and overexpression of myostatin inhibits activation.

**Steady-state satellite cell numbers are increased on Myostatin-deficient muscle fibers**

A constant pool of satellite cells is maintained on muscle fibers during repeated cycles of degeneration and regeneration. A portion of myoblasts derived from the activated satellite cells returns to quiescent status, thus maintaining a constant number of satellite cells per unit length of muscle fiber by the self-renewal process. Adult myoblasts derived from the myostatin-null mice tend to proliferate faster (see the results below), suggesting the possibility of increased self-renewal of the satellite cells in these knockout mice. Thus, the steady-state number of satellite cells per fiber of the muscle fibers isolated from both wild-type and myostatin-null mice was measured. As shown in Fig. 4 B (i), there is a significantly higher number of steady-state satellite cells per unit length of muscle fiber in myostatin knockout mice as compared with the wild-type mice. An example of CD34-positive satellite cells on the muscle fiber is shown in Fig. 4 B (ii).

To further demonstrate this increase in satellite cell number per fiber, myofiber explant culture (Fig. 4 C, ii) was used (Hawke et al., 2003). Results showed that a significantly higher number of myogenic precursor cells had migrated from the myostatin-null fiber (83 ± 8; n = 31) as compared with the wild type (51 ± 6; n = 31). The difference in total number of satellite cells between myostatin-null and wild-type fibers was significant (P < 0.05) (Fig. 4 C, i).

**Primary and secondary myoblasts that lack Myostatin proliferate faster**

Previously, we have shown that high concentrations of Myostatin inhibit the proliferation of myoblasts by controlling the G1 to S transition during cell cycle (Thomas et al., 2000). Thus, here, we determined the proliferation rate and the doubling time of adult myoblasts isolated from both wild-type and myostatin knockout mice. For this purpose, we plated secondary myoblasts cultured from embryonic day (E) 17 and adult myoblasts from postnatal muscle and determined their proliferation. Primary myoblasts isolated from wild-type and myostatin knockout mice underwent a lag period of 3 d before they became proliferative (unpublished data). Cultured secondary myoblasts, isolated from E17 embryos from myostatin knockout mice, proliferated significantly faster, resulting in an increase in the number of Mstn−/− myoblasts as compared with the wild-type myoblasts (Fig. 5 A). The increased proliferation rate of Mstn−/− myoblasts is also reflected by three times higher cumulative population doublings attained by the myostatin-null myoblasts, compared with the wild-type myoblasts (unpublished data). In addition, the proliferation rate of adult myoblasts derived from hind limbs of 4-wk-old myostatin-null and wild-type mice were also assayed using the methylene blue cell proliferation assay. As seen in Fig. 5 B, myostatin-null myoblasts proliferate significantly faster than wild-type myoblasts after 3 d in culture. The proliferation experiments described above were performed on myoblasts derived from satellite cells that were isolated from mixed muscles that contain both fast and slow twitch fibers. A recent report, however, suggests that the satellite cells isolated from fast and slow fibers are inherently different (Rosenblatt et al., 1996). Thus, to address the role of Myostatin in the proliferation of myoblasts from fast or slow fibers, we cultured adult myoblasts from m. tibialis anterior (fast twitch) or m. soleus (slow twitch) of both normal and myostatin knockout mice. The proliferation assay again indicates that both fast (Fig. 5 C) and slow (Fig. 5 D) fiber–specific adult myoblasts isolated from myostatin knockout mice proliferate much faster as compared with the wild-type adult myoblasts. Thus, regardless of the origin of satellite cells, lack of myostatin appears to increase the proliferation rate of both fast and slow fiber–specific satellite cells.

**Addition of exogenous Myostatin inhibits satellite cell activation and myoblast proliferation**

To show the direct inhibitory effects of Myostatin on myoblast proliferation, tissue-dissociated satellite cells were isolated from both wild-type and myostatin-null mice and used in a methylene blue assay. The same number of wild-type and myostatin-null myoblasts were cultured in the media, and myostatin was added in increasing concentrations to only myostatin-null cultures. After 48 h of proliferation, the cells were fixed with 10% formal saline and stained with the methylene blue. When recombinant Myostatin was added to the media in increasing amounts, the enhanced proliferation rate seen in the myostatin-null myoblasts was reduced to that of the wild-type myoblasts (Fig. 5 E).

**Cell cycle progression of wild-type and Myostatin-deficient satellite cells**

Proliferation assays clearly demonstrate that adult myoblasts that lack Myostatin proliferate much faster than the wild-type myoblasts. Furthermore, our previous results with C3C12 cells indicate that addition of exogenous Myostatin to the growth medium (GM) arrests the proliferation of C3C12 cells in G1 phase of the cell cycle (Thomas et al., 2000). Thus, we next asked if the increased proliferation observed in the myostatin knockout myoblasts is due to deregulated G1 to S phase transition. FACS® analysis of the cell cycle distribution of asynchronous populations of wild-type and myostatin knockout adult myoblasts in proliferation media revealed that myostatin knockout myoblasts had a twofold
increase in S phase cell population (Fig. 6 A). To confirm these results, we then synchronized the adult myoblasts in G0 phase by growing them in methionine-free media and initiated S phase entry by serum stimulation. To monitor the number of cells in S phase, the cells were pulsed with BrdU, and the percent of labeled cells was determined by immunostaining the fixed cells at the end of the assay. The results indicate that within the first hour after serum stimulation, 7% of Mstn−/− myoblasts entered S phase. In contrast, only 2% of wild-type myoblasts were found to be in S phase. Similarly, an enhanced number of BrdU-positive cells was observed up to 8 h after serum stimulation in Mstn−/− myoblasts (Fig. 6 B). These results indicate that lack of Myostatin increases the proliferation of myoblasts by deregulating the S phase entry.

Given that Myostatin inhibits the G1 to S phase progression of myoblasts, we speculated that Myostatin may affect one or more of the cell cycle control proteins. Of particular interest was the Cdk inhibitor p21, which has previously been shown to be involved in cell cycle arrest in the G1 phase. In addition, growth arrest by Myostatin signaling has been shown to influence p21 expression (Thomas et al., 2000; Joulia et al., 2003). To examine this possibility, actively growing adult myoblasts derived from satellite cells were cultured with or without Myostatin in the media for 24 h before harvesting. Western analysis with p21-specific antibodies revealed that Myostatin treatment up-regulated p21 expression (Fig. 6 C). To determine the direct effect of Myostatin on satellite cell proliferation, adult myoblasts were isolated from wild-type and myostatin-null mice. Exogenous Myostatin was added to the proliferating myostatin-null myoblast cultures. As the concentration increased, the enhanced proliferative potential of the myostatin-null myoblast decreased to that of the wild type (E).

Delayed differentiation potential of Myostatin-deficient myoblasts

Myogenic differentiation of myoblasts occurs upon withdrawal of actively growing myoblasts from cell cycle into G0 and subsequent expression of myogenic markers. As the adult myoblasts derived from the Mstn−/− muscle fibers...
myoblasts were synchronized to G1 phase, and once released into S phase, they were BrdU labeled and the percentage of BrdU-positive cells was counted at various time points. Data presented are an average of three replicates. **, P < 0.01. At least a total of 1,000 cells were counted in each of three replicates. Data presented are an average of three animals. (C) Western blot showing the levels of p21 and Cdk2 protein in primary myoblasts cultured with (+) or without (−) Myostatin for 24 h. p21 protein was detected using anti-p21 antibodies, and Cdk2 protein was detected using Cdk2 antibody. Tubulin protein levels are included to show equal loading.

have deregulated S phase entry, we next evaluated the ability of Mstn−/− myoblasts to withdraw from the cell cycle upon mitogen withdrawal. Adult myoblasts derived from both wild-type and myostatin knockout mice were switched to DM, and protein extracts were made from differentiating myoblasts at 0 (majority quiescent), 6, 12, 24, 48, and 72 h after switching to the low-serum media. The levels of differentiation-specific genes were quantified by Western and densitometric analysis. As expected, MyoD protein levels peak immediately after switching to the DM (within 6 h) in both wild-type and myostatin knockout myoblasts. However, in the myostatin-null myoblasts, the peak levels of MyoD continue until 12 h and then slowly decline to basal levels at ~48 h after switching to DM (Fig. 7 B). Whereas in the wild-type myoblasts, a sharp decline in the levels of MyoD was observed at 12 h of differentiation. Similar to MyoD, the levels of the early differentiation marker Myogenin peak within 6 h into differentiation in wild-type myoblasts. However, in the Myostatin-deficient myoblasts, the peak expression of Myogenin is delayed and is seen at 24 h into differentiation (Fig. 7 C).

Discussion

Investigations into the molecular basis of Myostatin function revealed that Myostatin functions by controlling the myoblast number during embryonic myogenesis (Thomas et al., 2000). Several lines of evidence, including the expression of Myostatin in postnatal muscle (Sharma et al., 1999), increased postnatal muscle growth in myostatin-null mice (McPherron et al., 1997), and expression of Myostatin in nascent myotubes after muscle regeneration (Kirk et al., 2000), compelled us to evaluate the role of Myostatin in adult myogenesis. The results presented here clearly demonstrate that Myostatin is expressed in satellite cells and that Myostatin regulates satellite cell quiescence and self-renewal.

Before delineating the role of Myostatin in adult myogenesis, we attempted to demonstrate the expression of myostatin mRNA and protein specifically in satellite cells, which are the precursors of adult muscle. Immunocytochemistry with anti-Myostatin antibodies on muscle transverse sections revealed intense Myostatin immunoreactivity in cells that are juxtaposed to the surface of muscle fibers. Further, immunostaining with antibodies against Pax7 (satellite cell marker) on serial sections established that the cells positive for Myostatin immunoreactivity are indeed the satellite cells (Seale et al., 2000) (Fig. 1, A and B). Myostatin expression
was further confirmed in Pax7-positive satellite cells by in situ hybridization (Fig. 1, C and D). In addition, purified satellite cells appear to express myostatin mRNA and protein, confirming that Myostatin is present in satellite cells. These results thus are in agreement with an earlier report, where Cornelison et al. (2000) have shown the presence of myostatin mRNA in the total RNA isolated from the single satellite cell attached to the fiber. In the same study, they also showed that, on activation of satellite cells, the levels of Myostatin (GDF-8) were significantly down-regulated, suggesting that high levels of Myostatin might be involved in satellite cell quiescence.

Three independent techniques, RT-PCR, Western blotting, and immunocytochemistry, were used to ascertain the expression of myostatin in adult myoblasts. As shown in Fig. 3 A, the PCR reaction amplified the expected size fragment of 1.2-kb myostatin ORF. In addition, anti-Myostatin antibodies specifically recognized Myostatin on the Western blot performed on the protein extracts derived from the adult myoblasts (Fig. 3 B). To demonstrate the expression of myostatin in actively proliferating cells, primary cultures were immunostained with Myostatin antibodies. As shown in Fig. 3 C, virtually all the cultured myoblasts showed staining with Myostatin antibodies and did not detect Myostatin in myoblasts isolated from myostatin-null mice. Thus, it is clear from these results that Myostatin is expressed in satellite cells and in the adult myoblasts derived from satellite cells.

If Myostatin negatively regulates the activation of satellite cells, we reasoned that in myostatin knockout mice there would be increased activation of satellite cells. Consistent with this hypothesis, a BrdU labeling experiment showed that in myostatin knockout animals, there is a significant increase in the pool of activated satellite cells as compared with wild-type mice. Although there is a significant reduction in the pool of activated satellite cells with age, myostatin-null muscle fibers exhibit considerably more activated satellite cells even at 6 mo (Fig. 4 A, i), which is consistent with the
persistent postnatal muscle growth and lack of muscle wasting observed in myostatin-null mice (McPherron and Lee, 2002). Recently, the breeding of the myostatin-null mouse with the mdx mouse resulted in the amelioration of the symptoms (Wagner et al., 2002). Also, the injection of a specific antibody to Myostatin into the mdx mice alleviated the severity of the phenotype; in fact, the mice gained muscle mass and increased the production of the muscle protein creatine kinase, indicating improvement of muscle strength (Bogdanovich et al., 2002). Whitemore et al. (2003) also injected inhibitory Myostatin antibody into wild-type mice and demonstrated increased skeletal muscle mass and increased grip strength. These results further indicate that Myostatin acts postnatally as a negative regulator of satellite cell activation and skeletal muscle growth. In addition to the increased muscle growth through hyperplasia, myostatin-null mutant mice also display hypertrophy, which has been related to increased satellite cell activity. One explanation for the increased number of myonuclei, and thus hypertrophy of myostatin-null muscle fiber, is the observed spontaneous activation of satellite cells in the absence of Myostatin. Conversely to this, the addition of Myostatin to isolated myofibers decreased the activation of satellite cells, when normally they would be induced to activate (Fig. 4 A, ii).

For activation, it is essential that the quiescent satellite cells either in G0 or G1 should reenter the S phase to undergo cell cycle progression. Hence, negative regulators of satellite cell activation, such as Myostatin, could possibly regulate satellite cell activation by controlling the G1 to S phase transition. Indeed, the results from this study (Fig. 6 C) suggest that high levels of Myostatin can inhibit G1 to S progression of the cell cycle by down-regulating Cdk2 and up-regulating p21, which would result in hypophosphorylation of pRb. Hence, it is quite possible that the higher levels of Myostatin observed in quiescent satellite cells (Cornelison et al., 2000) maintain the G0 status by regulating the G1/S check point. In the myostatin knockout mice, we also observed an increase in the number of CD34-positive satellite cells per unit length of fiber as compared with wild-type mice (Fig. 4 B, i). A recent report suggested that in order to meet the demand and maintain a constant pool of satellite cells on muscle fibers, satellite cells must self-renew like stem cells. The self-renewal process is accomplished by returning a pool of satellite cell–derived myoblasts to the quiescent status on muscle fibers (Seale and Rudnicki, 2000). An increased number of satellite cells per unit fiber thus means that in the absence of Myostatin, there is increased self-renewal of satellite cells, possibly by increased proliferation of satellite cell–derived myoblasts. This theory is supported by the observation that the adult myoblasts derived from the satellite cells of myostatin knockout mice appear to have a shorter doubling time and proliferate much faster as compared with the wild-type adult myoblasts (Fig. 5). This increased proliferation rate of Mstn 

Myoblast differentiation is a highly orchestrated step wherein the withdrawal of myoblasts from the cell cycle to G0 should occur before the committed myoblasts postmitotically differentiate into myotubes. Molecular analysis indicates that low mitogen medium triggers the up-regulation of MyoD, which in turn induces the expression of the p21 gene. Concomitant up-regulation of p21 and decrease of G1 cyclins and Cdk2 results in the withdrawal of myoblasts to G0 and terminal differentiation. Thus, the rate of withdrawal of myoblasts is an important step in the differentiation of myoblasts into myotubes. The studies with BrdU labeling clearly indicated that the wild-type myoblasts immediately withdraw from the cell cycle, whereas, the myostatin-null myoblasts appear to undergo the cell cycle longer,
resulting in delayed withdrawal from the cell cycle (Fig. 7A). The molecular basis for delayed withdrawal appears to be prolonged expression of MyoD and delayed peak expression of a differentiation-specific marker, such as Myogenin, in myostatin-null myoblasts (Fig. 7, B and C). These results certainly point out that Myostatin, by regulating MyoD expression, plays a key role in adult myoblast differentiation. Immunostaining with Myosin heavy chain antibody (unpublished data) and gene expression analysis revealed that loss of Myostatin results in delayed withdrawal from the cell cycle and differentiation of primary myoblasts derived from satellite cells. Consistent with this hypothesis, Rios et al. (2002) and Langley et al. (2002) very recently demonstrated that excess Myostatin in DM blocks the differentiation of C2C12 myoblasts by inhibiting MyoD expression. Eventually, however, the Myostatin-deficient myoblasts do differentiate, albeit delayed.

In conclusion, we propose that Myostatin blocks the activation of satellite cells and also negatively regulates self-renewal of satellite cells (Fig. 8). Increased activation of Mstn−/− satellite cells, and subsequent enhanced proliferation with delayed differentiation of myostatin-null myoblasts, could thus be the primary reason for the increased postnatal muscle growth and hypertrophy in myostatin-null animals. To our knowledge, this is the first report of a secreted growth factor that is expressed in satellite cells and maintains the satellite cell quiescence.

Materials and methods

Animals

myostatin-null mice (C57BL/10 background) were obtained from S.-J. Lee (The Johns Hopkins University, Baltimore, MD). Wild-type mouse strain C57BL/10 was bred at the Ruakura Small Animal Colony.

BrdU labeling of satellite cells in vivo

To determine the activated satellite cells in vivo at a given time point, BrdU labeling was used. Three C57BL/10 or Mstn−/− mice at ages 4 wk, 8 wk, or 6 mo were intraperitoneally injected with BrdU (30 mg/kg) as a single pulse 2 h before killing. Satellite cells from m. tibialis anterior were isolated using Percoll gradient as per the published protocol (Allen et al., 1997; Partridge, 1997; Yablonska-Reuveni et al., 1999). To detect activated satellite cells, 5,000–10,000 satellite cells were fixed using 0.25% glutaraldehyde and immunostained for BrdU using the cell proliferation kit (Amersham Biosciences). A minimum of 1,000 immunostained satellite cells were counted to calculate the percentage of BrdU-positive cells.

Determination of satellite cell number on muscle fiber

M. tibialis anterior was excised from six C57BL/10 or Mstn−/− mice, and individual muscle fibers were isolated by the method of Rosenblatt et al. (1999). Isolated fibers were fixed in an eight-well Lab-Tek chamber slide (Nunc) with 0.25% glutaraldehyde for 1 min and rinsed with PBS for immunocytochemistry. Satellite cell detection with CD34 antibodies (Santa Cruz Biotechnology, Inc.) and counting were performed according to the published protocol of Beauchamp et al. (2000). The total number of satellite cells was counted after 72 h incubation of the wild-type and myostatin-null tibialis anterior muscle fibers in fiber medium (FM) at 37°C, 5% CO2.

Effect of Myostatin on satellite cell migration

To determine the effect of Myostatin on the migration of satellite cells from the fiber, isolated muscle fibers were either placed in FM or in FM with tritrating amounts of Myostatin for 96 h at 37°C, 5% CO2. Wells were analyzed for satellite cell migration, and satellite cells were counted. Replicates of at least 24 fibers were used for statistical analysis.

Cell cycle analysis of adult myoblasts

Adult myoblasts were cultured according to the published methods (Allen et al., 1997; Partridge, 1997; Yablonska-Reuveni et al., 1999). FACS® analysis of primary myoblasts was done according to the method of Thomas et al. (2000). To monitor the cell cycle progression, adult myoblasts were synchronized at the G1/S boundary and released to S phase by the method of Kitzmann et al. (1998). The myoblasts in S phase were pulsed with 0.1 mM BrdU for 15 min, fixed, and processed for immunocytochemistry. To determine the cell cycle withdrawal of adult myoblasts, proliferating adult myoblasts were switched to DM and pulsed with 0.1 mM BrdU at 0, 6, 12, 24, or 48 h time points. The myoblasts were fixed and immunostained for BrdU.

Myoblast proliferation assay

Primary myoblasts were cultured from hind limb muscles of three 4-wk-old mice or six to eight E17 mouse embryos according to the published protocols (Chakravarthy et al., 2000; Partridge et al., 1997; Partridge, 1997; Yablonska-Reuveni et al., 1999). Embryonic primary myoblasts and satellite cells isolated from fast or slow fibers were seeded at low density of ~100 cells/cm2 in GM, and cells were counted every 24 h. Adult myoblasts were cultured as described earlier and were seeded at a density of 6,097 cells/cm2 in GM on Matrigel-coated 96-well microtiter plates. Cell counting and methylene blue photometric endpoint assay were performed according to the published protocols (Oliver et al., 1989; Chakravarty et al., 2000).

In situ hybridization

M. tibialis anterior muscle tissue was fixed in 4% paraformaldehyde (wt/vol) in PBS for 16 h at room temperature, washed, dehydrated through an ascending ethanol series, and embedded in paraffin wax. Sections were then cut at 7 μm and mounted on microscope slides (SuperfrostPlus; Fisher Scientific). For the generation of digoxigenin probes, a 326-bp fragment of the mature myostatin cDNA (Lp 903–1231; Genbank/EMBL/DDBJ accession no. NM_010834.1) or a 571-bp fragment of the Pax7 cDNA (Lp 963–1534; Genbank/EMBL/DDBJ accession no. XM_1317582) were cloned into the pGEM-T easy vector and transcribed according to the manufacturer’s recommendation (DIG RNA Labeling Kit; SP6/T7; Boehringer). Paraffin-embedded sections were dewaxed, and in situ hybridizations were performed according to the method of Greiw et al. (2001).

Immunocytochemistry

M. tibialis anterior muscle tissue was removed from C57BL/10 or Mstn−/− mice, fixed, and sectioned according to the method of Sharma et al. (1999). The sections were incubated with either 1:50 diluted Myostatin polyclonal antibody (Sharma et al., 1999) or with 1:100 diluted Pax7 mouse monoclonal antibody (DSHB). Subsequent steps of immunostaining were performed according to the published protocol (Thomas et al., 2000).

Primary cultures fixed with 0.25% glutaraldehyde on eight-well Lab-Tek chamber slides were permeabilized with 0.1% Triton X-100 in PBS at room temperature, and the immunocytochemistry was performed according to the method of Beauchamp et al. (2000). Detailed procedures are available on request. Photographs were taken using an Olympus BX50 microscope (Olympus) fitted with a DAGE-MTI DC-330 color camera (DAGE-MTI, Inc.).

Western blotting

Adult myoblasts were extracted from C57BL/10 or myostatin-null muscle tissue as described above. Percoll gradient–enriched quiescent satellite cells, proliferating myoblasts, or myoblasts changed to DM (DME, 2% [vol/vol] HS) for 0, 6, 12, 24 or 48 h were resuspended in 200 μl of protein lysis solution (50 mM Tris, pH 7.6, 250 mM NaCl, 5 mM EDTA, 0.1% Nonidet P-40; Complete™ protease inhibitor; Boehringer). The suspension was passed through a fine syringe needle to lyse the cells. Protein was measured using protein assay reagent (Bio-Rad Laboratories). Proteins (10 μg) were separated on SDS-PAGE (4–12% gradient; Novex) and then transferred to a nitrocellulose membrane. Western blots for MyoD and Myogenin were performed according to the protocol described previously (Thomas et al., 2000). Western blots for Myostatin were performed as described by Sharma et al. (1999). To detect the changes in the levels of p21 and Cdk2 upon Myostatin treatment, Western blotting was performed. Adult myoblasts were treated with 10 μg/ml Myostatin for 24 h, and the protein extracts from the myoblasts were made as described above. The
Detection of Myostatin in fiber-dissociated satellite cells and adult myoblasts

Satellite cells and adult myoblasts were isolated from 6-wk-old mice using the cell gradient method (Goodell et al., 1996). Total RNA was isolated using TRIzol reagent (Invitrogen) according to the manufacturer’s protocol, and RT-PCR was performed to amplify myostatin as described previously (Kambadur et al., 1997). Pax7 cDNA was PCR amplified using 5′-GCTGCGGAACCTCCTACCACTAC-3′ and 5′-CCACGACACGGGATGGTTCC-3′ primers and the following conditions: 95°C for 15 s, 55°C for 30 s, and 72°C for 30 s for 35 cycles. CD34 was PCR amplified using a published protocol (Beauchamp et al., 2000). Protein extracts from satellite cells were made as described above, and Western blot for Myostatin was performed according to the published method (Sharma et al., 1999).

Statistical analysis

To determine the significance between two groups, comparisons were made using a t test. The data are presented as means ± SEM.

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