Activity-Dependent Gene Regulation in Conditionally-Immortalized Muscle Precursor Cell Lines

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Abstract Skeletal muscle contractile activity has been implicated in many aspects of muscle cell differentiation and maturation. Much of the research in this area has depended upon costly and labor-intensive cultures of isolated primary muscle cells because widely available immortalized muscle cell lines often do not display a high level of either spontaneous or stimulated contractile activity. We sought to develop conditionally-immortalized skeletal muscle cell lines that would provide a source of myofibers that exhibit robust spontaneous contractile activity similar to primary muscle cultures. Using a tetracycline-regulated retroviral vector expressing a temperature-sensitive T-antigen to infect primary myoblasts, we isolated individual clonal muscle precursor cell lines that have characteristics of activated satellite cells during growth and rapidly differentiate into mature myotubes with spontaneous contractile activity after culture in non-transformation-permissive conditions. Comparison of these cell lines (known as rat myoblast-like tetracycline (RMT) cell lines) to primary cell cultures revealed that they share a wide variety of morphological, physiological, and biochemical characteristics. Most importantly, the time-course and extent of activity-dependent gene regulation observed in primary cell culture for all genes tested, including subunits of the nicotinic acetylcholine receptor (nAChR), muscle specific kinase (MuSK), and myogenin, is reproduced in RMT lines. These immortalized cell lines are a useful alternative to primary cultures for studying muscle differentiation and molecular and physiological aspects of electrical activity in muscle fibers. J. Cell. Biochem. 91: 821–839, 2004. © 2004 Wiley-Liss, Inc.

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Activity-dependent gene regulation has been implicated in many aspects of skeletal muscle differentiation and maturation. During myoblast differentiation, the family of myogenic transcription factors (MTF) MyoD, Myf-5, MRF4, and myogenin is differentially expressed, but as myotubes mature, become innervated and functionally active, expression levels are suppressed [Duclert et al., 1991; Buonanno et al., 1992; Adams et al., 1995; Buckingham et al., 2003]. Similarly, nicotinic acetylcholine receptor (nAChR) expression is high during the initial stages of myotube development, but is suppressed once innervated fibers become functionally active [Goldman et al., 1988; Witzemann et al., 1991]. In contrast, expression of myosin isoforms change on multiple occasions during muscle fiber maturation [Whalen et al., 1981], but ultimately are controlled by the degree of muscle activity via nerve stimulation [Buller et al., 1969; Pette and Vrbova, 1992]. These components of skeletal muscle as well as metabolic enzymes and sarcoplasmic reticulum constituents show clear evidence of the importance of contractile activity in regulating muscle specific gene expression.

The vast majority of activity-dependent gene regulation research has been performed on isolated primary muscle cells in vitro or whole...
muscles in vivo. Although these model systems remain the gold standard by which experimental data is and should be evaluated, they present significant technical and scientific drawbacks. Isolation of primary muscle cells is time consuming, costly, and often unpredictable with respect to yield and contamination with non-muscle cells. Similarly, in vivo muscle research typically requires a large number of animals which can be costly, and time consuming and introduces an additional level of expertise with respect to surgical technique. Moreover, in both cases, cell heterogeneity often contributes to difficulty in the interpretation of results.

An alternative to isolation of primary muscle cells for in vitro experimentation is the use of an immortalized muscle cell line of which there are a number [Yaffe, 1968; Kimes and Brandt, 1976; Linkhart et al., 1980; Blau et al., 1983; Mulle et al., 1988]. Although these cell lines have been extremely useful in gaining insight into cell cycle regulation and myotube differentiation, in our hands, they have not proven optimal for the study of activity-dependent gene expression because they do not display a high level of contractile activity. To further our studies of muscle development, maintenance, and in particular, activity, we sought to develop skeletal muscle cell lines from rat primary muscle cell cultures. Of particular interest to us were cell lines that demonstrated rapid cell proliferation followed by differentiation into mature, active, myofibers that reproduced the features of activity-dependent gene regulation observed in primary cell cultures and whole muscle in vivo.

We elected to use the SV40 large T-antigen (Tag) [Aaronson and Todaro, 1968; Henderson and Livingston, 1974] as a transforming agent. Tag has proven very effective for generating immortalized transformed cell lines from a variety of tissues and species including mesoderm-derived tissues such as bone, cartilage, and muscle [Iujvidin et al., 1990; Yang et al., 1992; Simon et al., 1996; Hicok et al., 1998; Robbins et al., 2000]. Temperature-sensitive Tag variants have the added benefit of permitting reversal of transformation through inactivation of the Tag protein at a non-permissive temperature of 37–39°C [Tenen et al., 1977; Brown et al., 1986; Jat and Sharp, 1989; Iujvidin et al., 1990]. An additional level of control over the immortalized state was introduced through the use of a retroviral vector known as NIT-Tag to infect the primary rat myoblasts. In addition to permitting a reduction of Tag-mediated transformation by thermodenaturation of the Tag protein, the NIT-Tag vector also allows blocking of Tag protein expression through the introduction of tetracycline-analogs into the culture medium and inactivating a “tet-off” tetracycline-transactivator (TTA) protein [Gossen and Bujard, 1992; Hoshimaru et al., 1996] expressed from the NIT vector. The combination of both inhibitors was predicted to very tightly and rapidly regulate the transformation status of infected cells.

Conditionally-immortalized myogenic cell lines expressing temperature-sensitive T-antigen have been produced by a number of laboratories and from a number of species using both T-antigen-expressing transgenic mice and viral vectors [Iujvidin et al., 1990; Yang et al., 1992; Simon et al., 1996]. These reports are variable in their depth of characterization of the differentiation status of generated lines (in large part because many of the markers of myogenic differentiation have only been characterized within the last few years) and either make no mention of contractile activity, or present no data regarding the molecular, biochemical, or physiological changes accompanying it. Furthermore, since a major goal of this study is to determine to what degree these conditionally-immortalized cell lines can reproduce activity-dependent gene regulation observed in primary cell cultures, we felt that in order to directly correlate with earlier work from the laboratory, the lines should derive directly from the primary cell cultures that we routinely generate.

We have generated cell lines using the NIT-Tag retrovirus that are readily perpetuated in culture and have been subcloned and characterized as individual clonal lines. We have used molecular markers to characterize these lines during active growth, contact inhibition, and after differentiation in a combination of culture conditions. Many of the lines differentiate into myofibers with features of mature muscle including contractile activity. We have compared these cells to primary cell cultures and despite the obvious differences in purity and uniformity of the starting population, most morphological, molecular, developmental, and contractile characteristics examined are highly similar. The data presented indicate that these immortalized cell lines will be a useful alternative to primary cell cultures for laboratories...
studying muscle development and differentiation and laboratories studying questions related to contractile activity and electrical responses of muscle fibers.

MATERIALS AND METHODS

Viral Vectors, Infection, and Clonal Selection

The NIT-Tag vector was produced by insertion of a BamHI fragment encoding a full-length temperature-sensitive SV40 Tag into the tetracycline-regulated MMLV-based, replication-defective retroviral vector NIT. The NIT-Tag retrovirus was packaged in HEK293T cells using triple plasmid co-transfection as described for other MMLV-based retroviruses [Suhr et al., 2001]. Rat primary myoblasts, isolated as described previously [Goldman et al., 1991], were plated on 35 mm collagen-coated culture dishes at a density of approximately $1 \times 10^5$ cells/plate and allowed to grow at $37^\circ C/8\% CO_2$ in growth medium (DMEM supplemented with 10% fetal bovine serum (FBS) and 10% horse serum). After growth for 18 h, plates of adherent cells were infected by the addition of NIT-Tag virus conditioned medium at an estimated MOI of 0.1 and transferred to $32^\circ C$. After an additional 24 h of growth, the infected cells were passaged to 150 mm culture plates at a density of 10^5 cells/plate and placed under selection in G418 (500 μg/ml) for 10 days. After selection was complete, 91 G418-resistant colonies that were well separated from neighboring colonies were picked and transferred to 24-well plates for additional growth. Of these colonies, 42 displayed robust growth at low-density with continued selection and were passaged in duplicate to a second set of 24-well plates for further analysis. These five lines numbered 22, 46, 59, 62, and 63, as well as five fibroblast-like lines, 8, 34, 55, 64, and 70 were expanded, tested for the presence of helper virus (found to be negative), and cryogenically preserved in multiple aliquots for further analysis. Each of the myotube-forming lines was subject to further clonal selection by limiting-dilution plating to 96-well plates. These individual clones were then expanded and analyzed individually to compare growth rate, morphology, differentiation, and other characteristics. DNA blot analysis using standard methods revealed an identical banding pattern of the integrated NIT-Tag provirus within individual clones isolated by limiting dilution, but a different pattern comparing between parental lines, suggesting that the parental lines tested arose from independent infections. A single integration was observed in all lines examined except for clones from line 46 that displayed two bands suggesting two proviral integrations (data not shown). The first clonal line of each group was expanded and used for further characterization. These are myoblast-like lines 221, 461, 5951, 6211, and 631, and for purposes of comparison, the fibroblast-like line 8. Myoblast-like myotube-forming lines are heretofore designated RMT cells for rat myoblast-like tetracycline-responsive cells and fibroblast-like lines are designated RFT cells for rat fibroblast-like tetracycline-responsive cells.

Electrical Stimulation

Rat primary myoblasts were isolated and plated as described above. Between 48 and 96 h post-plating primary cell cultures became confluent and the media was changed to DM to induce myotube formation. At this time, cells were treated with 3 μg/ml cytosine arabinoside for 48 h to inhibit fibroblast proliferation. RMT cell lines were plated at a density of approximately 10^5/plate in growth medium and allowed to proliferate at $32^\circ C$ until confluent. Upon
reaching confluence, the media was changed to DM with dox and the plates were shifted to 37°C. Discernible differentiation and myotube formation of RMTs typically commenced 3 days after switching to DM. Primary and RMT myotubes were continuously treated with 1 μg/ml tetrodotoxin (TTX) (Oretek, Inc., Fremont, CA) from the onset of myotube formation until time of electrical stimulation (7–10 days). Prior to experiments in which myotubes were electrically stimulated, cultures were rinsed three times with TTX free media, and then returned to the incubator briefly before checking for spontaneous contractile activity. Myotubes were then electrically stimulated to contract for up to 48 h using conditions described previously [Macpherson et al., 2002]. Data are presented for myotubes that were electrically stimulated for 6 h.

Transfections

To obtain agrin condition media RMT461 myotubes were co-transfected with plasmids containing expression vectors for either soluble or membrane bound forms of agrin [Ferns et al., 1993] and green fluorescent protein (GFP). pCMV-cAg4,8 and pCMV-Ag12,4,8 (generously provided by Dr. M. Ferns) harbor soluble and membrane bound forms of the agrin gene, respectively, which are down stream of the cytomegalovirus promoter. pCS2EGFP, containing the cytomegalovirus promoter driving GFP expression, was used to identify transfected cells and determine transfection efficiency. RMT461 myotubes were co-transfected with 0.5 μg of pCS2EGFP and either 1.5 μg of pCMV-Ag12,4,8 or pCMV-cAg4,8. Total DNA was adjusted to 3 μg/plate using pBSK (Stratagene, La Jolla, CA). Transfections were performed by calcium phosphate precipitation as previously described [Chahine et al., 1992].

RNA Isolation and RNase Protection Assay

Total RNA was isolated by homogenizing cell cultures in Trizol (Gibco BRL, Grand Island, NY) followed by the single step purification method as described by the manufacturer’s protocol. Antisense probes used to detect myogenin and the nAChR α-, β-, δ-, γ-, and ε-subunit RNAs were the same as those described by [Chahine et al., 1993]. The muscle specific kinase (MuSK) probe is a HindIII/ApaI subclone of the rat MuSK cDNA (provided by Dr. S.J. Burden). This probe spans nucleotides 1426–1899 of the published rat cDNA [Valenzuela et al., 1995]. The p21 probe is a clone of the mouse cDNA (provided by Dr. K. Huppi) that when linearized with Smal protects approximately 250 nucleotides of rat p21 mRNA. The retinoblastoma (Rb) probe is a Kpn1/Pst1 subclone of the mouse Rb cDNA (provided by Dr. E. Wang) that when linearized with Nde1 protects approximately 340 nucleotides of rat Rb mRNA. RNase protection assays were carried out as previously described [Adams and Goldman, 1998]. The probe for glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was obtained from Ambion (Austin, TX). GAPDH probes were included in each experiment and served to normalize for differences in the amount of RNA in each of the samples based on the observation that electrical stimulation of skeletal muscle does not regulate GAPDH gene activity [Huang et al., 1992]. The RNA for GAPDH was not regulated by any of the conditions employed in this report. RNase resistant hybrids were analyzed on 6% polyacrylamide, 8 M urea gels. After electrophoresis, gels were dried and exposed to the X-ray film. Probe signals were quantified by scanning densitometry and values were normalized to the RNA signal obtained for GAPDH. The specificity of the protected bands was confirmed by hybridizing probes to tRNA, which resulted in no protected fragments on the gel. Probe integrity was monitored for each experiment by running an aliquot of non-hybridized probe on each gel.

Western Blots

Cell lysates from cultured myotubes were prepared by scraping cells from the dishes in RIPA buffer (50 mM Tris-HCl, pH 8.0, 1% NP-40, 0.5% deoxycholate, 0.1% SDS, 150 mM NaCl, 1 mM EDTA) containing a protease inhibitor cocktail (P8340 Sigma, St. Louis, MO) at a dilution of 1:100. Cells were sheared by passing through a 26.5 gauge needle and centrifuged briefly to remove cellular debris. Protein concentrations were determined using the Bio-Rad DC protein assay (Hercules, CA). Protein samples were subjected to SDS–PAGE (10%) and transferred electrophoretically to Immobilon-P membranes (Millipore, Bedford, MA). After electro-transfer the proteins remaining in the gels were stained with Coomassie brilliant blue and used as an additional control for equilibration of protein loading. After trans-
fer, Immobilon-P membranes were blocked in Blotto buffer containing 5% dry milk in PBST (phosphate-buffered saline/0.2% Tween-20) and then incubated overnight at 4°C with mouse myogenin monoclonal antibody (mAb) supernatant 1:50 (clone F5D, obtained from the Developmental Studies Hybridoma Bank) or at room temperature for 1–2 h with rabbit polyclonal antibodies MyoD, Myf-5, or MRF4 1:1,000 (sc-760, sc-302, sc-784, respectively, Santa Cruz Biotechnologies, Santa Cruz, CA). Immunodetection was done using peroxidase-conjugated secondary antibodies, goat anti-mouse or goat anti-rabbit, 1:1,000 (Jackson ImmunoResearch Lab., West Grove, PA) with subsequent chemiluminescent detection (ECL, Amersham Pharmacia Biotech, Piscataway, NJ).

**Immunohistochemistry**

For all comparative immunocytochemical studies, cells within an experiment were grown, harvested, fixed, and processed identically and in parallel to provide favorable cross-comparison. For immunohistochemical analysis, tissue culture plates or multi-well plates were rinsed with PBS and then blocked with 2% paraformaldehyde for approximately 30 min. After fixation, plates were rinsed 5× in PBS and then blocked with 10% serum containing 0.3% Triton X-100 for 1 h at room temperature. Incubation of monoclonal or polyclonal antibodies was at 4°C overnight. After removing primary antibodies the plates were rinsed 3× in PBS for 5 min and then incubated with the appropriate secondary antibody. To visualize nAChR clusters, plates were incubated with Texas Red conjugated α-bungarotoxin (α-BTX) (1:1,000 dilution) (B-7489, Molecular Probes, Eugene, OR) for 1 h at room temperature. Antibody and other reagent sources were as follows: monoclonal anti-myogenin (1:50 dilution), anti-myosin (1:250 dilution), and anti-Pax7 (1:200) (clones F5D, MF20, and PAX7, respectively) were obtained from the Developmental Studies Hybridoma Bank; rabbit polyclonal antibody against MyoD (1:200 dilution), Myf-5 (1:250 dilution), or MFR4 (1:250 dilution) obtained from Santa Cruz Biotechnologies, Inc.; goat anti-mouse, goat anti-rabbit, donkey anti-mouse used at 1:250 dilution, donkey anti-rabbit used at 1:250 dilution (Jackson ImmunoResearch Lab.); Tag antibody used at 1:500 dilution (MAB986 Chemicon, Temecula, CA), DAPI stain (Sigma).

**Statistics**

Mean ± SD were determined for samples from primary and RMT cultures. Differences in mean values of Rb, p21, myogenin, MuSK and nAChR mRNA were compared within groups (e.g., primary TTX treated vs. primary stimulated) and significant differences were determined by t-tests analysis. The level of significance was set a priori at \( P < 0.05 \).

**RESULTS**

Primary myoblasts were immortalized by the NIT-Tag retrovirus that combines advantages of temperature-sensitive Tag with tetracycline regulation into a single vector that provides tight conditional control over cell immortalization. NIT-Tag shown schematically in Figure 1A, encodes two expression cassettes. The first cassette is expressed from the 5' viral LTR and encodes a G418 (neomycin) resistance gene, an internal ribosomal entry site (IRES), and the TTA protein. TTA transactivates expression of the second expression cassette through a TTA-responsive promoter (To) to produce the Tag gene product. A shift to the non-permissive temperature (37°C) destabilizes the Tag protein conformation resulting in a reduction in transforming function (Fig. 1B). In addition to temperature shift, high-level expression of Tag via TTA transactivation can be blocked by the addition of tetracycline analogs such as dox, further inhibiting the Tag-mediated transformation (Fig. 1C). As described in “Materials and Methods,” 42 neomycin-resistant NIT-Tag infected cell lines were grown in parallel and tested in an initial screen for their capacity to differentiate into multinucleate myofibers. Several lines indicated a high degree of myofiber formation, and the results for one parental line, RMT46 are shown in Figure 1D. RMT46 maintained at 32°C in growth medium showed a healthy, dense monolayer of mononuclear cells after 1 week of continuous proliferation (growing) whereas an identically plated culture that was shifted to DM supplemented with 1 μg/ml dox for the same duration at 37°C displayed numerous multinucleate muscle fibers and virtually no mononuclear cells (differentiated).

We then compared the individual and combined effects of temperature and dox treatment on the growth and differentiation of NIT-Tag infected cell lines cultured for 1 week under the following four conditions: 32°C, 37°C,
For this next level of analysis, clonal lines isolated by limiting dilution of each of the parental cell lines (described in “Materials and Methods”) were used. Differentiation was grossly determined by the formation of multi-nucleated myotubes in the culture plate. All RMT cell lines followed a similar trend toward differentiation and the results are shown for lines RMT461, RMT631, and RFT8 in Figure 2. The RFT8 line was included to show that morphological and biochemical changes characteristic of RMT differentiation are not the result of NIT infection, dox treatment, or even simple changes in the proliferative status of the cell, but are instead likely to arise from the fundamental character or “lineage” of the original infected cell.

As shown in Figure 2, the most permissive growth conditions (32°C) resulted in continuous cell proliferation as suggested by multiple mitotic profiles and exclusively mono-nucleated cells (Fig. 2A,E,I). During active low-density growth, individual RMT lines displayed doubling times ranging from 14 to 21 h (Table I). When cells were grown at 37°C, a condition that should inactivate the Tag protein, mitotic profiles were again observed with no evidence of myotube formation, suggesting that sufficient functional Tag was still present to drive cell division (Fig. 2B,F,J). In contrast, the addition of dox at the permissive growth temperature produced a much more profound effect on cell growth and the formation of myotubes. For the cell lines shown here, growth was dramatically reduced and no clear mitotic profiles were observed in the DAPI stained plates used for this analysis (Fig. 2C,G,K). Approximately 50% of RMT461 cells converted to myotubes with dox treatment alone (Fig. 2C), and RMT631 cells arrayed themselves in fiber-like strands of individual cells with rare myotubes (Fig. 2G). RFT8 cells grew equally well at either 32 or 37°C (Fig. 2I,J), but as with the RMT lines, dox treatment clearly lowered cell density and also resulted in an increasingly flattened morphology (Fig. 2K).

The greatest degree of morphological differentiation was observed in cells with combined temperature shift to 37°C and dox addition. Under these conditions, no mitotic profiles were observed for any of the cell lines and by 1 week, all muscle-like lines had formed myotubes. For RMT461 and RMT631 cells this conversion was >90% (Fig. 2D,H). Differentiated RMT461 and RMT631 cells produce long, multi-nucleated fibers, the majority of which are narrow in diameter, and accommodated only 1–2 nuclei in width for the length of the fiber. A smaller percentage of the fibers are wide near one terminus with 5–50 nuclei, and taper along their length to fibers with widely spaced individual nuclei. As shown in Figure 2L, RFT8 cells did not form any observed myotubes, but they did flatten and distribute with greater distance between nuclei suggesting a differentiated state.
Although these data are consistent with the predicted action of the NIT-Tag provirus, we sought to confirm changes in Tag production through the use of a mAb that recognizes the native form of Tag but not the denatured form of Tag at the non-permissive temperature, 37°C. This mAb was employed to compare the levels of the Tag protein under each of the above conditions. The results for all RMT cell lines and RFT8 followed the same trend, and the results are shown for lines RMT461 and RFT8 in Figures 2M–T. Nuclear localized Tag was clearly observed in essentially all growing cells at 32°C (Fig. 2M,Q). With the shift to 37°C, a clear decrease in staining intensity is observed, although nuclear localized Tag is still readily apparent (Fig. 2N,R). Under the 32°C + dox conditions, Tag staining is dramatically reduced relative to either of the previous conditions (Fig. 2O,S). The low level observed at 32°C is
even further reduced under the most favorable condition for differentiation, 37°C + dox (Fig. 2P,T). The above data provides evidence that the combination of temperature shift and dox treatment very effectively blocks the transforming function of the Tag protein.

Transcription Factor Immunostaining

To further establish the identity of these cell lines both in the proliferating and differentiated state we examined the expression of Pax7 and the MTFs (MyoD, Myf-5, MRF4, and myogenin). Pax7 has been characterized as specific to undifferentiated mononuclear muscle precursors known as satellite cells [Seale et al., 2000]. MyoD and Myf-5 are predominantly associated with the commitment of differentiating mononuclear cells to myogenic determination while MRF4 and myogenin are believed to be involved in myoblast terminal differentiation [Buckingham et al., 2003]. Analysis of these markers is shown in Figure 3. In Figure 3A–D, RMT461 cells in different growth conditions were processed for Pax7 (red) and Myf-5 (green) immunostaining. In Figure 3A,B, respectively, proliferating and confluent RMT461 cells are strongly nuclear positive for both Pax7 and Myf-5 (co-localization indicated by yellow color). After differentiation (Fig. 3C,D), Pax7 intensity is significantly decreased and appears to be lost altogether in the nuclei of differentiated myotubes, whereas Myf-5 continues to be observed at a high-levels in the nuclei, and, to a lesser extent, the cytoplasm, of both mono- and multinucleated cells. Each of the RMT lines followed the same pattern of marker expression as the RMT461 line.

<table>
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<th>Name/rep line</th>
<th>No. of SC</th>
<th>Morph</th>
<th>Het</th>
<th>Doub (in h)</th>
<th>Df. time (in days)</th>
<th>Fib/mon</th>
<th>Act.</th>
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<td>5–7</td>
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<td>++</td>
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<td>21</td>
<td>3–4</td>
<td>++++</td>
<td>+++</td>
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<tr>
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<td>13</td>
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Table I. Gross Characterization of Cell Lines

Name, line name of the original isolate; rep line, representative subclone obtained by limiting dilution and used in most later analyzes; no. of SC, number of individual subclones obtained by limiting dilution and subject to analysis. RMT8 was not subcloned, but was selected based on comparison to other fibroblastic RMT lines 44, 55, 58, 84, and 70 not described. Morph, morphological appearance while growing; Het, differences in characteristics such as morphology and differentiation between subclones within a line. (−) Indicates no apparent differences; (+) indicates differences between clonal lines. Doub, approximate doubling time of the original isolate; Df time, number of days following initiation of differentiation to obtain maximal cell fusion, myofiber formation, and onset of spontaneous activity; Fib/mon, approximate number of cells participating in fiber formation versus cells that remain mononuclear at maximal differentiation. (−) No fiber formation, (+) <20% nuclei in fibers, (++) 20–50%, (+++) 50–90%, (+++++) >90%. Act, spontaneous activity following majority fiber formation. (+) Very rare active fibers, (+++) occasional active fibers, (++++) frequent active fibers, (++++++) many active fibers.

*Because of proximal colonies in original selection of the isolate, lines 59 and 62 underwent an intermediate round of clonal selection prior to subcloning by limiting dilutions.
A similar analysis of MRF4 (green) with accompanying nuclear DAPI stain (blue) is shown in Figure 3E–H. Proliferating RMTs displayed no detectable expression of MRF4 (Fig. 3E). At high cell density, we observed rare cells that were positive for MRF4 (Fig. 3F and inset). In contrast, after differentiation all of the RMT lines displayed essentially uniform MRF4 positive staining in the nuclei and weak expression the cell body of myotubes and mononuclear cells (Fig. 3G,H). Again, each of the RMT lines followed a similar pattern of marker expression as the RMT461 line.

The expression of MyoD and myogenin were analyzed together and the results are shown for RMT461, RMT631, and RMT221 cell lines (Fig. 3I–Q). MyoD-positive signal (green) was observed in the nucleus and to a lesser extent, the cytoplasm, of all RMT cells at all phases of growth (Fig. 3I–Q). Unlike MyoD, during active proliferation, myogenin-positive immunostaining (shown in red) was a rare occurrence.

Fig. 3.
for any of the RMT lines (Fig. 3I,L,O). In confluent RMT cultures, however, notable differences were observed in myogenin expression that were not seen with other MTF's described above. As shown in Figure 3J,M, the RMT461 and RMT631 lines displayed a significant increase in myogenin expression, while other lines, like RMT221, showed rare positive cells (Fig. 3P). After 1 week of differentiation, the overwhelming majority of RMT461 and RMT631 mononuclear cells and fibers were positive for nuclear myogenin signal (Fig. 3K,N). The RMT221 line, which was slower to differentiate, also displayed many cells and fibers that were strongly positive for myogenin at the 1-week time-point (Fig. 3Q).

Taken together, these data suggest that RMT cells possess the characteristics of a muscle precursor cell known as an “activated satellite cell” that progresses through a predictable course of differentiation to become a mature myotube. It is noteworthy that a parallel analysis of transcription factor expression was performed on RFT cells, and under each growth condition, no positive signal was observable (data not shown).

After extensive testing of each of the RMT cell lines under various conditions, we could assign a generalized “differentiation” ranking to the cell lines that combined speed of differentiation, extent of morphological maturation (mononuclear cells vs. myotubes), and degree of differentiation-marker expression. The ranking is as follows with the most “differentiatable” first: RMT461 > RMT631 > RMT6211 > RMT221 > RMT5951. Many of the features of each of these cell lines are summarized in Table I.

Primary Versus RMT461 Immunostaining

During low-density growth, both primary muscle cells and RMT461 express abundant MyoD (Fig. 4). In contrast, few cells appear to be myogenin positive in either primary cell cultures or RMT461 cells, consistent with observations of RMT lines above. Upon differentiation, both primary muscle cells and RMT461 cells fuse to form MyoD positive myotubes and myogenin expression is dramatically increased relative to non-differentiated cells and appears to be present in essentially all of the myonuclei.

After differentiation, sarcomeric myosin is expressed throughout the cytoplasm of primary and RMT461 myotubes in a similar manner (Fig. 4). Typically, both primaries and RMT461s begin to spontaneously contract within 24 h of myotube formation. One notable difference between primary myotubes and the immortalized cell lines in general, is that the cell lines tend to make smaller myotubes that contain fewer myonuclei (Fig. 4 compare myogenin stained primary myotubes with 461 myotubes). Varying differentiation media conditions does not seem to alter this characteristic and it is currently not known whether this is an intrinsic property of the cell lines or is due to heterogeneity of the cell population in the primary cell preparation.

nAChR Clustering

nAChRs are ligand-gated ions channels that transmit excitatory signals from motor neurons to stimulate skeletal muscle contractile activity. During early muscle development nAChRs are expressed throughout the membrane, but as myotubes mature, receptors begin to aggregate (cluster) spontaneously. This process of nAChR clustering is dramatically increased with the arrival of pre-synaptic motor terminals which secrete a neural specific form of the synapse organizing protein, agrin. Since nAChR clustering is a fundamental precursor to the processes of activity-dependent signaling at the neuromuscular junction, we sought to further characterize the maturation of RMT myotubes by evaluating their ability to cluster nAChRs. When RMT461 myotubes were fluorescently-labeled with α-BTX, these mature myotubes formed nAChR clusters that were comparable in size and number to primary myotubes (Fig. 5A). This observation suggests that the signaling pathway for the formation of neuromuscular junctions remained intact in RMT461 cells.

As a further test of the integrity of this synapse forming pathway we treated RMT461
myotubes with conditioned media from cells expressing either a secreted or a membrane bound form of agrin. Compared with control non-conditioned myotubes, those that were exposed to conditioned media containing secreted agrin (cAg4-8) displayed visibly increased numbers of nAChR clusters (Fig. 5B) whereas myotubes treated with conditioned media from cells expressing membrane bound agrin (Ag12-4-8) showed no evident increase in nAChR clustering. Taken together these data indicate that the components necessary for the initial stages of synapse formation are intact in the immortalized cell lines.

**Activity-Dependent Regulation**

In differentiating cultured primary myotubes and RMT cell cultures, contractile activity is readily observed; however, this characteristic is difficult to quantify. Although direct comparative measurements of activity are problematic for a variety of reasons, it is possible and also of perhaps greater utility to quantify the down stream effects of activity on myofiber biology. A number of genes and their protein products have been found to be rapidly and quantifiably changed in response to electrical stimulation both in vitro and in vivo. To better assess the usefulness of the RMT461 cell line as an alternative model system for studying muscle biology, we examined several genes with activity-dependent regulation by RNase-protection assay and Western blot analysis. Among the gene products examined were cell cycle regulators, Rb protein and p21, the four MTFs, the five subunits (α, β, δ, γ, and ε) of the nAChR, and MuSK, a muscle specific tyrosine kinase required for motor endplate formation.

We have previously shown that 6 h of electrical stimulation is sufficient to produce significant
changes in gene transcription in primary myotubes [Macpherson et al., 2002]. In the present study, RNase protection assays show that after 6 h of electrical stimulation transcripts for Rb and p21 are not acutely affected by activity in primary myotubes, but decrease in RMT461 myotubes (Fig. 6A,B). Interestingly, preliminary data from our lab indicate that mRNA for both Rb and p21 increase significantly in denervated whole muscle, and these increases appear to be suppressed by electrical stimulation (unpublished data). Consequently, the observation that the levels of Rb and p21 in primary myotubes are less responsive to the effects of electrical stimulation than in whole muscle or RMT461 myotubes may reflect the heterogeneity of the cell population in primary cultures.

While the role of cell cycle regulators in activity-dependent gene regulation remains to be determined, many reports have implicated myogenin as an important MTF that not only is regulated by muscle activity, but is also involved in regulating nAChR gene expression in response to muscle depolarization [Neville et al., 1992; Mendelzon et al., 1994; Gundersen et al., 1995; Tang et al., 2001]. Six hours of electrical stimulation produced significant decreases in myogenin mRNA in primary myotubes, but only slightly decreased myogenin mRNA in RMT461s (Fig. 7A). This was also true for at least two of the other RMT cell lines (not shown). In contrast, myogenin protein was regulated in a similar manner in primary and RMT461 myotubes (Fig. 7B). These data suggest that although muscle activity can reduce myogenin expression in both primary and immortalized muscle lines, the mechanisms contributing to this reduction may differ between primary and immortalized cell lines.

Since changes in the levels of protein expression, rather than RNA, are more likely to impact downstream targets of transcription factors we also checked for changes in the protein levels of the other MTFs (MyoD, Myf-5, and MRF4). When protein levels for each of these proteins were incubated for 24 h with unconditioned medium (control), conditioned media from myotubes transfected with a secreted form of agrin (c-Ag4-8), or media from myotubes transfected with a membrane bound form of agrin (Ag12-4-8), Agrin induced clustering was evaluated qualitatively by comparing the relative level of clustering in four randomly selected fields for each condition tested. Images are at 100x magnification.

Fig. 5. nAChR clustering in primary and RMT461 myotubes. A: After 1 week of differentiation, primary and RMT461 myotubes were incubated with Texas red α-bungarotoxin (α-BTX) to visualize nAChR clustering. Brightfield images are included to indicate the relative density of myotubes on each plate. Images are at 200x magnification. B: Agrin induced nAChR clustering in RMT461 myotubes. After 1 week of differentiation, myotubes were incubated for 24 h with unconditioned medium (control), conditioned media from myotubes transfected with a secreted form of agrin (c-Ag4-8), or media from myotubes transfected with a membrane bound form of agrin (Ag12-4-8). Agrin induced clustering was evaluated qualitatively by comparing the relative level of clustering in four randomly selected fields for each condition tested. Images are at 100x magnification.
was assessed after 6 h of electrical stimulation, no substantial effect was observed in either rat primary or RMT461 myotubes (Fig. 7C–E). Whether these proteins are affected by longer periods of muscle activity will require further experimentation.

Activity-dependent regulation of the nAChR subunit genes has been well documented both in whole muscle and in primary muscle cells (Goldman et al., 1988; Witzemann et al., 1991; Chahine et al., 1993; Dutton et al., 1993; Bessereau et al., 1994). After 6 h of electrical stimulation, we observe suppression of the mRNA for each of the nAChR subunit genes in the RMT461 cell line (Fig. 8A–E). Furthermore, as can be seen for the gamma and alpha subunits, the levels of suppression that are achieved in the cell line are similar to those in primary myotubes (Fig. 8D,E). As a further test of activity-dependent regulation in the RMT lines we probed for changes in MuSK, which has also been shown to be regulated by muscle activity (Bowen et al., 1998). Similar to the nAChRs, 6 h of electrical stimulation caused a significant decrease in the level of MuSK mRNA in both primary and RMT461 myotubes (Fig. 8F).

Collectively, these data indicate that morphological, physiological, and biochemical characteristics of primary myoblast cultures and RMT cells are very similar and that RMT cells are likely to be a useful alternative model system to study muscle differentiation, maintenance, and activity in vitro.

**DISCUSSION**

The goal of this study was to produce a conditionally immortalized muscle cell line for the purpose of investigating processes of activity-dependent gene regulation. With the use of a temperature-sensitive, tetracycline-regulated retroviral vector we successfully generated a number of these cell lines from embryonic rat muscle tissue that display both the morphological and contractile characteristics of isolated primary myotubes. These lines have the capacity to proliferate under a variety of growth conditions and yet assume intrinsic muscle properties after stimulation to differentiate. Several of these RMT cell lines and the RMT46-derived lines in particular, have demonstrated reproducible contractile activity through passage 15 even when combined with limited-dilution subcloning or re-infection and selection.

The NIT vector has two potential advantages over many other tetracycline-regulated vectors in that it is more rapidly regulated by the addition or removal of Tet-analogs than many other vectors and NIT has no evident toxicity in cultured cells with repeated passage. This may stem from a lowered level of TTA protein indirectly expressed from the viral LTR via the IRES element (generally 2–5% of the protein encoded in the up stream position) that may
both reduce toxicity and speeds responsiveness to Tet-analogs. Even though the level of TTA-transactivator production is low, it is still likely to be saturating with respect to activation of the To response promoter. In the absence of tetracycline analogs, the NIT vector typically provides higher levels of transgene expression at low copy than direct expression from strong viral promoters such as the hCMV, sCMV, or RSV promoters (unpublished observations).

Two general morphologies were observed in the original isolated colonies: the spindle-shaped colonies that produced the RMT cell lines, and more numerous fibroblast-like colonies.

**Fig. 7.** Changes in myogenic transcription factors (MTF) after short-term electrical stimulation. **A:** RNase protection assays were used to assay for changes in levels of myogenin (Mgn) mRNA in primary myotubes (black bars) and RMT461 myotubes (gray bars) after 6 h of electrical stimulation. Changes in RNA levels are compared to non-stimulated myotubes that were maintained in media containing TTX. Representative RNase protections are presented at the left of the graph. Values presented are the average of two experiments and each experiment was performed in duplicate. For each protection assay, levels of Mgn mRNA were normalized to the level of expression of GAPDH. Bars represent mean ± SD. *P < 0.05, significantly different from within group controls (TTX). **B–D:** Representative Western blots of changes in myogenin (B), MyoD (C), Myf-5 (D), and MRF4 (E) protein after 6 h of electrical stimulation. Coomassie blue stained gels are shown as an additional control for total protein loading (∼20 μg/lane). For each figure, samples from control (TTX) and electrically stimulated (Stim) primary myotubes are presented in the first two lanes, respectively and samples from RMT461 myotubes are presented in the second two lanes.
nies such as RFT8. Although the RFT cell lines are described as “fibroblast-like” and do not form myotubes under the conditions used in this report, it should not be assumed that these cells are fibroblasts. They may instead derive from any of the other cell types in the original preparation or may be a muscle-resident cell that is not yet committed to the muscle lineage.

With regard to RMT cell lines such as RMT461, the evidence presented in this report strongly suggests that it displays characteristics of committed muscle progenitor cells known as “activated” satellite cells. Pax7 is a hallmark of satellite cell specification in general [Seale et al., 2001], and co-expression of the MTFs Myf-5 and MyoD places these cells squarely within the activated category since “quiescent” satellite cells are generally considered negative for MTF expression [Seale et al., 2001]. An interesting feature of the RMT cell lines is that differentiation from mononuclear “activated satellite” cell to mature myofiber is marked by an apparent step-wise progression that is not as clearly delineated in other cell lines or primary cultures. One example of this is the onset of widespread myogenin expression in confluent RMT461 and RMT631 cultures that can be trypsinized, re-passaged, apparently resume mononuclear growth, and later, differentiate in a normal manner. MRF4 expression, on the other hand, does not increase substantially in the majority of dense RMT461 and RMT631 cells but is instead observed at high levels only after fusion is well underway. This difference

Fig. 8. Activity-dependent changes in RNA levels for nAChR subunits and muscle specific kinase (MuSK) measured by RNase protection assay. Values for RMT461 myotubes are indicated by gray bars and values for primary myotubes are indicated by black bars. “TTX” label indicates non-stimulated cultures and “Stim” label indicates cultures electrically stimulated for 6 h. Representative RNase protections are presented to the left of each graph: A) assay of nAChR ß subunit RNA in RMT461 myotubes; (B) assay of nAChR ð subunit RNA in RMT461 cells; (C) assay of nAChR ε subunit RNA in RMT461; (D) comparative assays of nAChR γ subunit RNA in RMT461 myotubes and primary myotubes; (E) comparative assays of nAChR α subunit RNA in RMT461 myotubes and primary myotubes; (F) comparative assays of MuSK RNA in RMT461 myotubes and primary myotubes. Values presented are the average of two experiments, each performed in duplicate. For each protection assay, the respective level of mRNA was normalized to the GAPDH expression level. Bars represent mean ± SD. *P < 0.05, significantly different from within group controls (TTX).
suggests that MRF4 expression may indeed be an indicator of terminal differentiation whereas myogenin expressing cells, given the proper stimulus, can abort the differentiation program and return to a less differentiated state relatively unchanged. In Figure 9 we have summarized the markers characteristic of each of these steps, and propose a transient developmental stage for the activated cell termed the “queued” cell that represents a cell that has progressed far along the differentiation pathway, but not yet reached the point-of-no-return toward terminal differentiation.

Figure 9 also summarizes changes in the expression of the RNAs and proteins examined in differentiated myofibers with and without electrical activity. We have shown that, with the exception of subtle differences, the cell line RMT461 displays activity-dependent regulation of several muscle-expressed proteins including transcription factors, transmembrane receptors, and other proteins in a manner that suggests many similarities with primary muscle cells. Preliminary data from experiments performed on some of the other RMT cell lines indicate they also maintain contractile regulation similar to RMT461 and primary myotubes.

Possibly the best characterized activity-dependent genes in skeletal muscle are those of the nAChR [Goldman et al., 1988; Witzemann et al., 1991; Chahine et al., 1993; Dutton et al., 1993; Bessereau et al., 1994]. The expression of these genes generally increases with muscle inactivity, and decreases when muscle cells are functionally active. Although a number of signaling pathways have been implicated in regulating these processes [Klarsfeld et al., 1989; Huang et al., 1992; Chahine et al., 1993; Mendelzon et al., 1994; Tang et al., 2001; Macpherson et al., 2002], the physiological details of how the regulatory pathways interact have not been unraveled. One of the problems in resolving these questions is the difficulty in large scale access to the nuclei of mammalian muscle preparations. The development of readily available muscle cell lines that maintain the same functional characteristic as primary cells should prove to be a valuable resource in addressing some of these issues.

**Fig. 9.** Schematic of different stages of RMT461 differentiation, marker expression, and changes in gene expression/protein content with electrical activity. In the left panel, RMT cells are depicted during low- and high-density growth. Marker names below each category of cells indicates gene products found expressed at that developmental stage. Expression levels of nAChRs, MuSK, Rb and p21 were only evaluated in myotubes. At the far right, arrows and dashes associated with gene products indicate expression changes with electrical activity.
One of the differences between primary cells and the immortalized cell lines appears to be in the regulation of Rb and p21 in response to electrical stimulation shown in Figure 6. The observation that Rb and p21 do not appear to be regulated by electrical stimulation in primary cell cultures may simply reflect the differences in the cell populations present in these cultures. Cultures from immortalized cell lines are composed of clonal myogenic cells, whereas primary cultures are heterogeneous in composition with a large percentage of the cells presenting a morphology suggesting fibroblastic origin [Rando and Blau, 1994 and clonal populations this study]. Since fibroblasts and other mononuclear cells do not possess the contractile machinery of myofibers yet do express Rb and p21, the presence of such large populations of cells in the primary muscle preparations would significantly dilute any observable effect of muscle activity. Regardless of the explanation for this difference between primary and RMT cultures, the above observations highlight one of the advantages of performing experiments with a pure population of cells when asking specific questions about the regulation of genes that are expressed in multiple cell types. In addition, because of the purity of the cell lines, they have been used as an enriched protein source to ask questions about protein phosphorylation [Tang et al., in press Cellular Signaling].

The other observed difference in activity-dependent regulation between primary cells and the immortalized cell lines is the regulation of myogenin RNA. Myogenin is thought to be an important mediator of activity-dependent transcription. In primary muscle cells, we observe a decrease in myogenin RNA in response to electrical stimulation that is not as dramatically reflected in the RNA of the RMT cell lines. We do not currently know what mechanism is responsible for this difference, but given that myogenin protein levels are regulated similarly in primary cells and RMT's it appears that in the post-transcriptional pathways mediating myogenin activity, as related to activity-dependent regulation, are likely to be intact in the cell lines.

The activity-dependent data we have presented demonstrates the fidelity with which the cell lines respond to the suppressive effects of muscle activity. Although we have no direct evidence that the cell lines are equally reliable in portraying the inductive signals of muscle stimulation, we assume that this will be the case based on the developmental similarities of the cell lines with primary muscle cells. It will be interesting to determine if proteins that are induced by activity in skeletal muscle, such as some of the contractile proteins [Cerny and Bandman, 1986; Torgan and Daniels, 2001], display a similar phenotype in the cell lines.

In addition to the applications described above, the cell lines appear to be readily transfectable either as myoblasts or myotubes. Consequently, they are amenable to reporter assays, as well as further genetic manipulation. Moreover, given their dual mode of growth regulation they may be less likely to display the tumorgenic characteristics that have been describe for other muscle cell lines when reintroduced in vivo [Iujvidin et al., 1990; Morgan et al., 2002].

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REFERENCES


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Hoshimaru M, Ray J, Sah DW, Gage FH. 1996. Differentiation of the immortalized adult neuronal progenitor cell line HC2S2 into neurons by regulatable suppression of the v-myc oncogene. Proc Natl Sci USA 93:1518–1523.


