Glycogen synthase kinase 3β represses MYOGENIN function in alveolar rhabdomyosarcoma

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MYOGENIN is a member of the muscle regulatory factor family that orchestrates an obligatory step in myogenesis, the terminal differentiation of skeletal muscle cells. A paradoxical feature of alveolar rhabdomyosarcoma (ARMS), a prevalent soft tissue sarcoma in children arising from cells with a myogenic phenotype, is the inability of these cells to undergo terminal differentiation despite the expression of MYOGENIN. The chimeric PAX3-FOXO1 fusion protein which results from a chromosomal translocation in ARMS has been implicated in blocking cell cycle arrest, preventing myogenesis from occurring. We report here that PAX3-FOXO1 enhances glycogen synthase kinase 3β (GSK3β) activity which in turn represses MYOGENIN activity. MYOGENIN is a GSK3β substrate in vitro on the basis of in vitro kinase assays and MYOGENIN is phosphorylated in ARMS-derived RH30 cells. Constitutively active GSK3β(S9A) increased the level of a phosphorylated form of MYOGENIN on the basis of western blot analysis and this effect was reversed by neutralization of the single consensus GSK3β phosphoacceptor site by mutation (S160/164A). Congruently, GSK3β inhibited the trans-activation of an E-box reporter gene by wild-type MYOGENIN, but not MYOGENIN with the S160/164A mutations. Functionally, GSK3β repressed muscle creatine kinase (MCK) promoter activity, an effect which was reversed by the S160/164A mutated MYOGENIN. Importantly, GSK3β inhibition or exogenous expression of the S160/164A mutated MYOGENIN in ARMS reduced the anchorage independent growth of RH30 cells in colony-formation assays. Thus, sustained GSK3β activity represses a critical regulatory step in the myogenic cascade, contributing to the undifferentiated, proliferative phenotype in alveolar rhabdomyosarcoma (ARMS).

Cell Death and Disease (2014) 5, e1094; doi:10.1038/cddis.2014.58; published online 27 February 2014
Subject Category: Cancer

Rhabdomyosarcoma (RMS) is the most common pediatric soft tissue sarcoma, accounting for 5% of all childhood cancers and approximately 50% of soft tissue sarcomas.¹ ³ There are two main subtypes: embryonal and alveolar RMS and although embryonal RMS is more common, alveolar RMS is considered to carry a worse prognosis. A gene fusion resulting in the t(2;13)(q35;q14) somatic cell chromosomal translocation fuses PAX3 and Foxo1 to create a potent transcription factor (PAX3-FOXO1) which is a predominant causative genetic lesion for the development of alveolar rhabdomyosarcoma (ARMS).¹ ARMS is a highly malignant mesenchymal tumor that has properties of immature striated muscle tissue resulting in dense aggregates of poorly differentiated cells that are separated by fibrous membranes resulting in a loss in cellular cohesion.² ³ PAX3 is a key determinant of somatic myogenesis and, is involved in the migration of progenitor cells to the dermomyotome region of the somite where they grow and divide in the presence of growth factors.⁴ PAX3 is also required to activate the myogenic determination gene, MYOD.⁵ MYOD is one of four myogenic regulatory factors (MRFs, which include MYF-5, MRF4 and MYOGENIN) from the basic helix-loop-helix superfamily of transcription factors which interact with myocyte enhancer factor-2 (MEF2) proteins in the hierarchical control of muscle-specific gene expression.⁶ ⁹ Two kinases that potently exert effects on this myogenic regulatory cascade are p38 mitogen activated protein kinase (MAPK) and glycogen synthase kinase 3β (GSK3β). p38 MAPK is a key regulator of skeletal myogenesis that critically interacts with and activates MEF2 in the somite myotome during development.⁷ ⁹⁻¹⁰ Conversely, GSK3β activation leads to a repression in skeletal and cardiac muscle differentiation, in part by antagonizing p38 MAPK-mediated activation of MEF2.¹⁰ ¹¹ GSK3β usually targets proteins that have already been phosphorylated by another kinase at a ‘priming’ serine or threonine residue located four amino acids C-terminal to a consensus (S/T)XXX(S/T)-PO₄ motif.¹² ¹³ Regulation of MEF2 and the MRFs leads to morphological changes including epithelial to mesenchymal transition, cell alignment and fusion to form multinucleated myotubes that eventually develop into

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Keywords: alveolar rhabdomyosarcoma; PAX3-FOXO1; MYOGENIN; GSK3β; cell proliferation; tumorigenicity

Abbreviations: GFP, green fluorescent protein; GSK3β, glycogen synthase kinase 3β; Luc, luciferase; MAPK, mitogen activated protein kinase; MCK, muscle creatine kinase; MEF2, myocyte enhancer factor-2; P38K, phosphoinositide 3-kinase; PKB, protein kinase B

Received 05.11.13; revised 20.1.14; accepted 21.1.14; Edited by A Stephanou
functional, contractile muscle fibers. In particular, cells that express MYOD and MYOGENIN are typically fusion competent\textsuperscript{14,15} with the exception of ARMS cell types. To date, lack of myogenic differentiation of PAX3-FOXO1 expressing ARMS cells has been attributed to their inability to upregulate p57\textsuperscript{Kip2} activity, hence destabilizing the DNA binding affinity of MYOD transcription complexes.\textsuperscript{16} Dysfunctional MYOD/E-protein complex association and transcriptional control is a common feature between ARMS and the non-PAX3-FOXO1 expressing embryonal rhabdomyosarcoma (ERMS). Subsequent restoration of the MYOD/E12 complex has been shown to switch ERMS cells from an arrested myofibroblast phase to a more differentiated state.\textsuperscript{17} Similarly p38 MAPK activity can potentiate myogenic differentiation in ERMS cells by enhancing MYOD trans-activation properties.\textsuperscript{18} Therefore, it is fairly clear that in both rhabdomyosarcoma subtypes the ability of MYOD to potentiate transcription is compromised. However, the role of MYOGENIN in RMS is more equivocal. For normal myogenesis to occur, \textit{both in vitro and in vivo}, an absolute requirement for MYOGENIN is evident. Thus, MYOGENIN activity constitutes a pivot point for irreversible commitment to terminal differentiation.\textsuperscript{19,20} The combination of data from gene targeting studies of the MRFs\textsuperscript{21,22} supports the prevailing consensus that while the other three MRFs can compensate each other’s functional roles,\textsuperscript{23–26} MYOGENIN is absolutely essential for skeletal muscle fiber formation.\textsuperscript{20} Despite its expression in RMS, the paradox as to why MYOGENIN cannot mediate competence for differentiation is unknown.

Here, we examined the posttranslational regulation of MYOGENIN in ARMS. On the basis of the \textit{in silico} prediction of a single consensus phosphorylation site for GSK3\textbeta on the MYOGENIN protein and also high levels of GSK3\textbeta activity in these cells, we determined that MYOGENIN function is potently repressed by GSK3\textbeta activity in ARMS. Moreover, pharmacological inhibition of GSK3\textbeta results in a profound decrease in size and, to a certain extent, number of RMS colonies in a colony-formation assay. This effect is mimicked by introduction of MYOGENIN bearing neutralizing mutations in the GSK3\textbeta consensus site. In combination, these data reveal MYOGENIN as a key target of GSK3\textbeta activity in ARMS.

**Figure 1** MYOGENIN protein expression and GSK3\textbeta activity are both maintained in ARMS: (a) C2C12 myoblasts were transfected with HA-PAX3-FOXO1 or pcDNA3.1 control plasmid for 1 day before extraction or serum withdrawal and then extraction at 1 day increments for up to 4 days as indicated. Protein levels were compared with protein extracts from PAX3-FOXO1 expressing RH30 cells 1 day in growth media (GM) and 4 days in differentiation media (DM). The results show that despite the expression of PAX3-FOXO1, RH30 cells also express MYOGENIN. On the other hand, HA-PAX3-FOXO1 overexpression in C2C12 inhibits MYOGENIN expression and subsequent myogenic differentiation. (b) C2C12 myoblasts were transfected with CMV-dsRed2, MCK-eGFP and, either HA-PAX3-FOXO1 or pcDNA3.1 control plasmid. HA-PAX3-FOXO1 overexpression repressed the formation of multinucleated myotubes. (c) Endogenous GSK3\textbeta protein levels and phosphorylation at serine 9 were compared in C2C12 myoblasts, RH30 and ERMS RD cells. Although GSK3\textbeta is expressed in all three cell types, it is predominantly phosphorylated and hence inactive in C2C12 myoblasts and RD cells but not PAX3-FOXO1 expressing RH30 cells. (d) C2C12 myoblasts were transfected with HA-PAX3-FOXO1 or pcDNA3.1 control plasmid for 1 day before extraction. Overexpression of HA-PAX3-FOXO1 resulted in decreased phosphorylation of GSK3\textbeta at serine 9 indicating its activation.
ARMS, indicating that pharmacologic manipulation of this signaling axis may provide an opportunity for therapeutic intervention.

**Results**

**MYOGENIN is expressed in PAX3-FOXO1 expressing RH30 cells.** Serum (10% FBS) contains growth factors that repress the transcriptional activity of MRFs and also stimulate cell cycle progression hence rendering C2C12 myoblasts proliferative. In tissue culture, serum withdrawal (2% HS) results in activation of MEF2 and MRFs causing cell alignment and fusion to form multinucleated myotubes. Initially, in order to investigate the effect of PAX3-FOXO1 on this differentiation program, proliferating C2C12 myoblasts were transiently transfected with CMV-dsRed2, MCK-eGFP, and either HA-PAX3-FOXO1 or pcDNA3.1 control vector. Growth media (GM) was replaced with differentiation media (DM) 19 h after transfection and cells were allowed to differentiate for 96 h. SDS-PAGE samples were prepared from populations of myoblasts that either expressed or did not express PAX3-FOXO1, (a) before serum withdrawal (time = 0; GM = 10% FBS) and (b) at 24 h increments upon serum withdrawal (days 1–4; DM = 2% HS). Protein expression levels of these samples were then compared with protein samples from PAX3-FOXO1 expressing RH30 cells in GM and DM, by western blotting. These data indicate that despite the expression of PAX3-FOXO1, MYOGENIN protein expression is maintained in human ARMS-derived RH30 cells (Figure 1a). In addition, PAX3-FOXO1 repressed myotube formation in C2C12 myoblasts (Figures 1a and b). Detection of myogenic differentiation using an MCK promoter driving GFP expression revealed GFP expressing, multinucleated myotubes in the controls but not in cells expressing PAX3-FOXO1 (Figure 1b).

It is well documented that MRFs and MEF2 proteins are highly sensitive to pro-myogenic kinases such as p38 MAPK\(^{28-30}\) and also kinases such as GSK3/β which are

![Figure 2](image-url)
repressive to myogenesis.\textsuperscript{10,31} Therefore we tested for GSK3\(\beta\) activity under conditions when myogenesis is suppressed. As GSK3\(\beta\) is constitutively active until it is repressed by phosphorylation at serine 9 (by PKB), we assessed both total GSK3\(\beta\) protein expression levels and S9 phosphorylation levels using appropriate antibodies as indicated. We document that GSK3\(\beta\) is expressed in proliferative C2C12 myoblasts, PAX3-FOXO1 expressing ARMS cells (RH30) and, non-PAX3-FOXO1 ERMS cells (RD). However only in PAX3-FOXO1 expressing RH30 cells, is GSK3\(\beta\) predominantly in its unphosphorylated form (at serine 9) and, hence fully active state (Figure 1c). In addition, ectopic expression of PAX3-FOXO1 resulted in reduced phosphorylation of GSK3\(\beta\) at serine 9 (Figure 1d).

**MYOGENIN trans-activation function is repressed by GSK3\(\beta\).** To assess the effect of GSK3\(\beta\) activity on MYOGENIN function, trans-activation of a 4x E-box Luciferase construct was measured in proliferating C2C12 myoblasts that were transfected with different combinations of constitutively active GSK3\(\beta\)(S9A) and MYOGENIN as indicated in Figure 2a. The data indicate that MYOGENIN potentiates the 4x E-box Luc reporter gene and that GSK3\(\beta\)(S9A) abrogates this effect (\(P<0.001\)) indicating repression of MYOGENIN by active GSK3\(\beta\) (Figure 2a, left panel) without affecting the MYOGENIN protein expression levels (Figure 2a, right panel).

GSK3\(\beta\) directly phosphorylates MYOGENIN in vitro. In order to determine whether MYOGENIN is a substrate for GSK3\(\beta\), an in vitro kinase assay was performed using GST-MYOGENIN (1–225), purified GST-GSK3\(\beta\) and \(\gamma\)-\(^{32}\)P ATP. Bands were resolved using SDS-PAGE and subsequent autoradiography showed \(^{32}\)P labeled bands for MYOGENIN, autophosphorylated GSK3\(\beta\) and MyBP (positive control, Figure 2b). In addition, Coomassie Blue staining revealed a lower mobility band indicative of phosphorylation (Figure 2b). To further test the idea that the lower mobility band is hyperphosphorylated, we used calf-intestinal phosphatases on RH30 cell lysates and found that the low mobility band was eradicated (Figure 2c). Collectively these data suggest that MYOGENIN is a GSK3\(\beta\) substrate in vitro.

Pharmacologic manipulation of GSK3\(\beta\) activity alters MYOGENIN properties. To further investigate the effect of GSK3\(\beta\) on MYOGENIN, COS7 cells were co-transfected with MYOGENIN and GSK3\(\beta\)(S9A) and, then treated with or without 10 \(\mu\)M GSK3\(\beta\) inhibitor, AR-A014418, as indicated in Figures 3a and b. Western blot analysis revealed two predominant forms of MYOGENIN, a low mobility hyperphosphorylated isoform and a high mobility, hypophosphorylated isoform (Figure 3a, lane 2). The lower mobility, hyperphosphorylated band is reduced upon pharmacological treatment with AR-A014418 as indicated (Figure 3a, lane 3). This corresponded with a significant increase in trans-activation of an E-box cis element driven reporter gene (\(P<0.001\), Figure 3b). In contrast, constitutively active GSK3\(\beta\)(S9A) without pharmacological inhibition resulted in an increase in the low mobility, hyperphosphorylated band (Figure 3a, lane 4) which corresponded to a decrease in E-box luciferase activity in reporter gene assays (\(P<0.05\), Figure 3b).

Mutation of a consensus GSK3\(\beta\) phosphoacceptor site on MYOGENIN (S160/164A) prevents GSK3\(\beta\)-mediated repression. By in silico analysis, MYOGENIN contains a highly conserved putative GSK3\(\beta\) consensus phosphoacceptor site (Table 1), which we targeted by neutralizing site-directed mutagenesis. We observed that although wild-type MYOGENIN is sensitive to the repressive effects of constitutively active GSK3\(\beta\)(S9A), MYOGENIN
MYOGENIN neutralizing phosphomutant (S160/164A) is resistant to GSK3β repression of transcription activity as well as an increased slower migrating, hyperphosphorylated MYOGENIN band. (**a**) 4x E-box Luc activity was assessed in C2C12 myoblasts that were transfected with either wild-type MYOGENIN or MYOGENIN (S160/164A) and co-transfected with HA-GSK3β(S9A) or pcDNA3.1 control plasmid as indicated. HA-GSK3β(S9A) repressed MYOGENIN trans-activation of the 4x E-box promoter region (P < 0.001) but had no effect on mutated MYOGENIN (S160/164A) transcriptional activity. (**b**) Western blot analysis of the same samples revealed a decrease in a slower migrating, hyperphosphorylated band for overexpressed MYOGENIN (S160/164A, lane 2) with respect to overexpressed wild-type MYOGENIN (lane 1). Co-transfected HA-GSK3β(S9A) caused an increase in the slow migrating, hyperphosphorylated MYOGENIN band (lane 3) but not with overexpressed mutated MYOGENIN (S160/164A, lane 4). (**c**) Independent analysis of E-box Luc activity in C2C12 myoblasts with different combinations of overexpressed MYOGENIN, mutated MYOGENIN (S160/164A), HA-GSK3β(S9A) or pcDNA3.1 control plasmid as indicated. ***P < 0.001, **P < 0.05, #, ns

(S160/164A) was not (Figure 4a). Western blot analysis revealed that MYOGENIN (S160/164A) mutations correspond with a decrease in the low mobility, hyperphosphorylated upper band (Figure 4b, lane 2) and that this effect was not altered by ectopically expressed HA-GSK3β(S9A). Together these data indicate that S160/164A mutations in MYOGENIN render it insensitive to the repressive effect of GSK3β. GSK3β(S9A) expression resulted in an increase in the low mobility, hyperphosphorylated form of wild-type MYOGENIN (Figure 4b, lane 3) and this corresponded with decreased E-box luciferase activity (P < 0.001, Figure 4a). Although trans-activation of the skeletal muscle gene E-box cis-element by mutated MYOGENIN (S160/164A) is marginally less potent than wild-type MYOGENIN (P < 0.05, Figure 4c); it is resistant to inhibition by activated GSK3β (P < 0.001, Figure 4c).

PAX3-FOXO1 activation of GSK3β antagonizes muscle creatine kinase promoter activation. To further examine the functional significance of our findings, we used MCK promoter activity, as a key indicator of the activation of myogenic differentiation, in C2C12 myoblasts that were transfected with or without the PAX3-FOXO1 oncogene (Figure 5a). These data depict that PAX3-FOXO1 represses MCK promoter activity in myoblasts that have been co-transfected with MYOGENIN (P < 0.01) and this effect is not only abrogated by pharmacological inhibition of GSK3β, but further activated (P < 0.001, Figure 5a). Interestingly, in PAX3-FOXO1 expressing, human ARMS-derived RH30 cells, ectopically expressed MYOGENIN had no effect on MCK promoter activity unless it was coupled with pharmacological inhibition of GSK3β using AR-A014418 (P < 0.001, Figure 5b). Conversely, mutated MYOGENIN (S160/164A)
muscle-specific genes by repressing the transcriptional activity of MYOGENIN. When coupled with pharmacological inhibition of GSK3β, expression of MYOGENIN enhanced MCK-Luc activity as expected. This was repressed by co-expression of PAX3-FOXO1, indicating and then treated with either 10 μM AR-A014418 or DMSO solvent. MYOGENIN enhanced MCK-Luc activity as expected (P < 0.001) and this effect was repressed by co-expression of PAX3-FOXO1 (P < 0.01). Pharmacological inhibition of GSK3β not only reversed the effect of PAX3-FOXO1 but resulted in a super-activation (P < 0.001). To assess the importance of these findings in human-derived ARMS, RH30 cells were transfected with either MYOGENIN or mutated MYOGENIN (S160/164A) and MCK-Luc promoter activity was assessed. The data shows that wild-type MYOGENIN could not trans-activate the MCK promoter region unless it was coupled with pharmacological inhibition of GSK3β (P < 0.001). This was in contrast to mutated MYOGENIN (S160/164A) and MCK-Luc promoter activity was assessed. The data shows that mutant MYOGENIN could not trans-activate the MCK promoter region unless it was coupled with pharmacological inhibition of GSK3β (P < 0.001). This was in contrast to wild-type MYOGENIN (S160/164A), which could potentiate MCK promoter activity (P < 0.001) regardless of GSK3β inhibition. A summary of our findings: GSK3β activity promotes the tumorigenicity of RH30 cells and that this effect is neutralized by expression of MYOGENIN bearing mutations that render it insensitive to GSK3β.

Electrical stimulation of ARMS-derived RH30 cells reduces GSK3β activity through Akt (PKB). Electrical stimulation of skeletal muscle cells in cell culture has been shown to induce phenotype alterations and differentiation. Given that rhabdomyosarcoma shares properties of the skeletal muscle lineage, we electrically stimulated cultured RH30 cells for 4 h/day (5 Hz) for up to 4 days with the idea that it might promote differentiation by affecting the Akt/GSK3β signaling pathway. Stimulation of these cells resulted in an increase in pAktT308 to levels that were 3.00 ± 0.72-fold higher than those in non-stimulated cells after 4 days of stimulation (Figures 7a and b). Concomitantly, pGSK/S9 was also increased 2.25 ± 0.37 fold following 4 days of stimulation (Figures 7a and c). These increases in kinase activity corresponded with increased E-box promoter activity in stimulated cells compared with controls (Figure 7f). Collectively, these data indicate that electrical stimulation suppresses GSK3β activity and correspondingly activates MRF activity supporting our previous findings and also highlighting the possibility of using electrical stimulation as a therapeutic intervention in ARMS patients.

Discussion

ARMS, unlike ERMS, has a well-characterized cytogenetic basis in the majority of patients resulting from chromosomal translocations between chromosomes 1 and 13 and also 2 and 13 that result in fusion of the DNA binding domains of either Pax7 or Pax3 with the trans-activation domain of the Forkhead (FKHR) transcription factor family member Foxo1. In view of the well-substantiated crucial role of Pax3 and 7 in the development of skeletal muscle, it is therefore not surprising that the signature of ARMS tumor cells is a muscle-like phenotype and the expression of a variety of structural muscle marker genes such as myosin heavy chain and desmin. What is surprising is the sustained expression of MYOD and MYOGENIN in ARMS, and...
which are transcription factors that are intimately associated with the terminally differentiated, non-proliferative phenotype of normal myogenic cells, begging the question as to why they cannot exert this effect in ARMS. In particular, the function of MYOGENIN in the myogenic regulatory hierarchy places it at a pivotal and required step in the terminal commitment of myogenic progenitors to the differentiation program.\textsuperscript{19,20,34}

Thus, our observations reported here, that MYOGENIN function in ARMS is repressed by inappropriate sustained signaling by the kinase GSK3\textsubscript{b}, may be of considerable...
significance for understanding the etiology of this disease. Moreover, as repression of kinase activity is, in many cases, a tractable pharmacologic approach, we now propose targeting GSK3β activity as a tangible therapeutic strategy for ARMS. In support of the above, a recent study showed that ARMS-associated PAX3/7-Foxo1 fusion proteins inhibit MYOD target genes. It was also reported that forced MYOD/E-protein dimer expression could not rescue PAX3/7-Foxo1 repression of myogenic factors. Here, we also report that ectopically expressed PAX3-FOXO1 represses the induction of muscle genes, even when MRFs are expressed. We propose that the posttranslational repression of MYOGENIN

Figure 7  In vitro electrical stimulation of RH30 cells. (a) Western blot analysis revealed that electrical stimulation increased PKB/Akt activity and that this corresponded with increased phosphorylation of GSK3β at S9. Relative increase of phosphorylation at: (b) Akt at T308 and (c) GSK3β at S9, over time. Graphical representation of phosphorylated to total amounts of: (d) Akt and (e) GSK3β. (f) E-box promoter activity decreased with electrical stimulation and this also corresponded with inhibition of GSK3β at S9. *P < 0.01, **P < 0.001.
activity is due to sustained GSK3β activity and, through a cross-talk mechanism, subsequent repression of p38 MAPK (Supplementary Figure 1) as we have previously described.10 p38 MAPK and PKB/Akt are both required for activation of MEF2/MYOD transcriptional control and chromatin remodeling events at crucial myogenic loci for the differentiation program.11,40

In other systems, GSK3β phosphorylation of its protein substrates results in subsequent targeting for proteasomal degradation.12,13 However, GSK3β does not appear to affect MYOGENIN protein stability in our experiments as we observe an increase in a slow migrating, hyperphosphorylated form of MYOGENIN in response to GSK3β signaling that is not reduced in terms of its level of expression suggesting that proteasomal degradation of MYOGENIN is not enhanced by GSK3β. Conversely, neutralizing mutations of the GSK3β consensus enhanced MYOGENIN trans-activation of the muscle creatine kinase promoter, and also reduced the tumorigenic properties of ARMS cells (RH30) in a colony-formation assay. These findings suggest that GSK3β-mediated inhibition of MYOGENIN trans-activation properties impairs MYOGENIN’s ability to promote terminal differentiation in tumorigenic RH30 cells.

Cell cycle control is an essential component of normal growth control and development which goes awry in tumorigenesis. To date several growth-promoting PAX3-FOXO1 target genes have been implicated in RMS such as the IGF-R and c-Met although, while their contribution to proliferation is likely, the extent of their precise involvement in ARMS is still not clear.41 During normal skeletal myogenesis, upregulation of a cyclin-dependent kinase inhibitor, p21, stalls myoblasts in the G2/M phase of the cell cycle thus priming them for differentiation by promoting cell cycle exit, which is a requirement for subsequent muscle-specific gene expression.42 Consistent with the idea that GSK3β activation may contribute to the oncogenic properties resulting from PAX3-FOXO1 expression in ARMS, we observed that the number of proliferative RH30 cells is approximately halved by pharmacological inhibition of GSK3β. So far, the exact mechanism by which GSK3β regulates cell proliferation in ARMS is unknown. However, GSK3β has recently been shown to activate KLF643 and we recently identified that KLF6 enhances cell proliferation in myogenic cells through a TGFβ/Smad3 dependent pathway.44 We therefore speculate that PAX3-FOXO1/GSK3β enhancement of cell proliferation may involve KLF6 as a downstream effector as it is also highly expressed in various RMS cell types.

In summary, MYOGENIN normally activates genes that regulate cell fusion and terminal differentiation of skeletal muscle. In PAX3-FOXO1 expressing ARMS cells, our data indicate that sustained GSK3β activity represses MYOGENIN function, contributing to the transformed, proliferative phenotype of these cells. On the basis of this evidence, we propose that pharmacologic targeting of GSK3β kinase activity may constitute a tractable therapeutic strategy for ARMS.

Materials and Methods

Plasmids. E-box, MYOGENIN and MCK reporter constructs in pGL3 and expression vectors for MYOGENIN in EMSV were used in reporter gene assays. HA-tagged PAX3-FOXO1 was cloned into pcDNA3.1 and kindly donated by Dr. Malkin at MaRS, Toronto. HA-tagged GSK3β(S9A) was cloned in pcDNA3 ORF 995–2305.

Antibodies. Anti-MYOGENIN and anti-HA mouse monoclonal antibodies as well as anti-MEF2A rabbit polyclonal antibody were produced with the assistance of the York University Animal Care Facility; anti-PAX3 (1:250, Cell Signaling, Whitby, ON, Canada) GSK3β; phospho-GSK3β (1:1000; Cell Signaling); actin, MYOD, Myf-5, GFP, dsRed2 (1:2000; SantaCruz, Santa Cruz, CA, USA) were used for immunoblotting experiments.

Cell culture and transfection. C2C12, Cos7 and RH30 cells were maintained in DMEM supplemented with 10% fetal bovine serum (HyClone, Burlington, ON, Canada), 1% l-glutamine and 1% penicillin-streptomycin. Cells were maintained in a humidified, 37 °C incubator with a 5% CO2 atmosphere. For transfections, cells were seeded 1 day before transfection and transfected according to the standard calcium phosphate method previously described. A mixture of 50 μl 2.5 M CaCl2 per 25 μg DNA with an equal volume of 2X HeBS (2.8 M NaCl, 15 mM Na2HPO4, 50 mM HEPES, pH = 7.15) was used and the cells were incubated overnight followed by washing and addition of fresh media. The cells were counted and transferred to pre-gelatin-coated plates.

Protein extractions, immunoblotting and reporter gene assays. Cells were collected following the recovery period throughout the 4 days of the protocol.

Electrical stimulation. Cells were plated onto 0.1% gelatin-coated 6-well plates. The lids of the plates were fitted with two parallel platinum wire electrodes, placed at the opposite ends of each well and extending into the media. The wires from all wells were arranged in parallel and connected to an electrical stimulator (Harvard Apparatus Canada, Saint-Laurent, Quebec, Canada). Cells were stimulated at 5 V and a frequency of 5 Hz for 4 hours/day and allowed a subsequent 20 h recovery period. Cells were collected following the recovery period throughout the 4 days of the protocol.

Soft agarose colony formation assay. Materials: 0.7% (w/v) DNA grade Agarose, 1% (w/v) DNA grade Agar, 0.005% Crystal Violet (Sigma-Aldrich, Oakville, ON, Canada), 2X Media + 20% (v/v) FBS. After 48 h of transfection with MYOGENIN containing the S160/164A mutations or empty vector, RH30 cells were assayed for their capacity to form colonies as previously described.47 A total of 1 x 104 cells were suspended on a layer of 0.35% agarose in DMEM (10% FBS) with or without 10 μM AR-A014418, in 6-well plates. Medium was refreshed every 3-5 days as needed and on the 22nd day, the amount of colonies were
counted using a contrast phase microscope. The relative colony sizes were calculated using ImageJ software (Scion Corporation, Frederick, MD, USA). Four independent experiments were carried out in triplicate.

Conflict of Interest
The authors declare no conflict of interest.

Acknowledgements. We thank Dr. D Malkin from The Research Institute at The Hospital for Sick Children, Toronto, Ontario, Canada, for providing the HA-PAK3-FOXO1 plasmids as well as the RH3D and RD cells. This work was supported by funding provided by Canadian Institutes for Health Research (CIHR) and Natural Sciences and Engineering Research Council of Canada (NSERC).

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Supplementary Information accompanies this paper on Cell Death and Disease website (http://www.nature.com/cddis)