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The mERG1a Channel Modulates Skeletal Muscle MuRF1, but not MAFbx, Expression

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Running Title: MERG Modulates MuRF Expression

The MERG1a Channel Modulates Skeletal Muscle MuRF1, but not MAFbx, Expression

ABSTRACT

Introduction. The mechanism by which the mERG1a K⁺ channel increases ubiquitin proteasome proteolysis (UPP) was investigated. Methods and Results. Atrophic gastrocnemius muscles of hindlimb suspended mice express mERG1a, MuRF1 and MAFbx genes. Electro-transfer of mERG1a into non-suspended mouse muscle significantly decreases muscle fiber size (12.6%) and increases UPP E3ligase MuRF1 mRNA (real time PCR; 2.1 fold) and protein (immunoblot; 23.7%), but does not affect MAFbx E3ligase expression. Neither mERG1a-induced decreased fiber size nor mERG1a-induced increased MuRF1 expression is significantly curtailed by coexpression of inactive HR-FOXO3a, a gene encoding a transcription factor known to induce MAFbx expression by binding directly to its promoter. Discussion. The mERG1a K⁺ channel significantly increases expression of MuRF1 but not MAFbx. We explored this expression pattern by ectopically expressing an inactive FOXO3a and showing that it is not involved in mERG1a-mediated expression of MuRF1. These findings suggest that mERG1a does not modulate *MuRF1* expression through the AKT/FOXO pathway.

Key Words: mERG1a; Skeletal Muscle Atrophy; Ubiquitin Proteasome Proteolysis; MuRF1;

MAFbx/Atrogin1

INTRODUCTION

Skeletal muscle (SKM) comprises 35-45% of the human body mass and is necessary for movement, posture, support and temperature regulation. SKM atrophy, a loss in muscle mass and strength, can be induced by numerous stimuli: disease (e.g., cancer cachexia, sepsis, HIV/AIDS, diabetes), injury (spinal cord damage, denervation), immobilization, fasting, ageing and glucocorticoid treatment^{1,2}. Muscle loss is attributable to either a decreased protein synthesis, increased protein degradation or some combination of both of these events. The extent to which each contributes to muscle loss varies with animal model, the evoking stimulus, study time course and muscle fiber type^{2,3,4}. The protein degradation that produces atrophy results primarily from the activity of three proteolytic systems: calpains, cathepsins and the ubiquitin proteasome pathway (UPP). It is reported that the UPP is the primary proteolytic system involved⁵ and is purported to be responsible for as much as 75% of the protein degradation that occurs during SKM atrophy^{1,6,7}. The UPP is a multistep pathway requiring activation of an ubiquitin protein by an ubiquitin-activating enzyme (E1) and ATP hydrolysis. The activated ubiquitin is transferred to the ubiquitin-carrier protein (E2) which binds to an ubiquitin protein ligase (E3 ligase) carrying a protein substrate. The E3 ligase then transfers the ubiquitin to the targeted substrate. Once a substrate carries (a minimum of four) ubiquitin molecules, it is degraded by the 26S proteasome^{1,6,7}. The basic mechanistic nature of the UPP has been described, but the specific players vary as do the signaling pathways that lead to transcription and translation of these players. To date, two muscle specific ubiquitin E3 ligases have been described: Muscle atrophy F-box (MAFbx/Atrogin1) and Muscle RING Finger-1 (MuRF1). MAFbx/Atrogin-1 is a member of the SCF (Skp1, cdc53/Cullin and F-box protein) subfamily of ligases, containing an F-box domain which serves to bind the E3 complex to a targeted

protein^{8,9,10}. MuRF1 belongs to the RING Finger E3 ligase subfamily which has a canonical Nterminal RING domain followed by a conserved region characteristic of the MuRF1 family and a zinc-finger domain^{8,11}. Expression of these E3 ligases has been shown to be variably modulated by both the PI3K/AKT/FOXO and the IKK-β/IκB-α/Nfkappa B (NF-κB) pathways, dependent upon animal model and the stimulus inducing the atrophic state^{1,2,6,7}. It has recently been shown that these two pathways are responsible for roughly half each of the muscle wasting that occurs in immobilization-induced SKM atrophy, demonstrating that they are the main pathways involved in this type of atrophy 12 .

The ether-a-gogo-related gene 1a (ERG1a) K⁺ channel (Kv11.1, KCNH) produces the I_{Kr} current which is partially responsible for late phase repolarization of the cardiac action potential. Mutations in this channel have been linked with Long QT Syndrome 2 (LQT2), a cardiac disorder characterized clinically by a prolonged QT interval, torsade de pointes and sudden cardiac death¹³. Two alternative splice variants of *ERG1* have been cloned from human (HERG1A and $1B^{14}$) and mouse (mERG1a and $1b^{15}$) cDNA libraries and high levels of ERG1a and 1b expression have been detected in heart and brain of various mammals, including rats, mice and humans 14,15,16. In previous studies with mice, we showed that the mERG1a homomultimeric channel is linked to SKM atrophy induced by hindlimb suspension (HS; i.e., an unloading model) and cancer cachexia¹⁷. Specifically, we showed that: 1) mERG1a channel protein level is upregulated in the gastrocnemius muscles (GM) of hindlimb suspended mice experiencing atrophy relative to matched muscles in weight bearing (WB) control mice; 2) ectopic expression of the wildtype (WT) mERG1a splice variant in SKM of WB mice induced atrophy while co-expression of the WT and a dominant negative mERG1a subunit (DNmERG1a, G628S¹⁸) blocked atrophy in these mice; 3) ectopic expression of the DN-merg1a

mutant in hindlimb suspended mice inhibited atrophy; 4) pharmacological block of mERG1a inhibited atrophy in hindlimb suspended mice and increased muscle fiber cross sectional area (CSA) in WB mice; and 5) ectopic expression of the WT mERG1a channel in mouse GM increased UPP activity.

Here we investigate the mechanism by which mERG1a modulates the UPP. Using a mouse HS model of atrophy, our studies begin with a time course which demonstrates that our HS model indeed induces increased expression of the *mERG1a* gene and also genes encoding the E3 ligases MuRF1 and MAFbx while, interestingly, electro-transfer^{17,19} of mERG1a into mouse GM induces expression of the MuRF1, but not the MAFbx gene. These results are confirmed with immunoblot studies showing that MuRF1 and Atrogin1 (MAFbx protein product) protein levels are increased in response to HS, but that only MuRF1 protein levels increase in response to ectopic expression of mERG1a. We confirm that mERG1a expression is not modulating MAFbx transcription by ectopically co-expressing mERG1a and a MAFbx/Atrogin1 luciferase reporter²⁰ and showing that mERG1a expression does not increase MAFbx promoter-driven luciferase activity levels. Additionally, we co-expressed mERG1a and FOXO3a, a transcription factor shown to induce expression of both MAFbx and MuRF1 genes in some models of atrophy^{20,21}, and demonstrate that mERG1a expression does not modulate MuRF1 (or MAFbx) transcription through *FOXO3a* expression.

MATERIALS AND METHODS

Animals. All procedures were approved by the Purdue Animal Care and Use Committee. Seven to eight week old ND4-Swiss Webster male mice (Harlan-Sprague; Indianapolis, IN) were used in all procedures. Animals were housed in Purdue University facilities on a 12 hour light/dark cycle, monitored by lab animal veterinarians and provided food and water *ad libitum*.

Hindlimb Suspension (HS). Custom suspension cages were constructed as described previously²². Mice were placed in these cages resting in approximately a 30° head down tilt with their hindlimbs elevated so that they were unable to place any load on their hindlimbs. Control mice were kept in commercial mouse cages in a normal weight bearing state.

Tissue Sectioning, Staining and CSA Determination. GMs were embedded, cryo-sectioned (12 μ m) and stained for β –galactosidase (lacZ) activity as described earlier¹⁷. Images of sections were captured with a Leaf Micro–Lumina digital camera (Scitex; Tel-Aviv, Israel). The CSA (μ m²) of each muscle fiber was determined using Image J (NIH; Bethesda, MD).

Plasmids. The *mERG1a* clone in pBK/CMV¹⁵ and the dominant negative *mERG1a* (*DN-mERG1a*) clone in pBK/CMV¹⁸ were generous gifts from Dr. Barry London (Cardiovascular Institute, University of Pittsburgh, PA). The CMV-*lacZ* in pNL vector was purchased from the Center Commercial de Gros (Toulouse, France). The phRL synthetic *Renilla* luciferase reporter vector was purchased from ProMega (Madison, WI). The *MAFbx*/Atrogin1 luciferase reporter was generously provided by Dr. Stewart Lecker (Beth Israel Deaconess Medical Center, Boston, MA²⁰). The *HR-FOXO3a* (*H212R*) plasmid, a mutated form of *FOXO3a* which is inactive because of its inability to bind DNA, was developed in the laboratory of Dr. Naoya Fujita (Japanese Foundation for Cancer Research, Japan²³).

Electro-transfer. Mice were anesthetized with 10 μl/g body weight of a solution of xylazine (1 mg/ml) and ketamine (9 mg/ml) in sterile saline. GMs of shaved hindlimbs were injected with expression plasmids in 50 µl sterile saline and electroporated with 8 pulses at 200V/cm for 20 ms at 1 Hertz with an ECM 830 ElectroSquare Porator (BTX; Hawthorne, NY). This method has been shown to result in gene transcription and translation in SKM in our laboratory^{17,19}.

Real time PCR. Trizol reagent (Invitrogen; Carlsbad, CA) was used to extract total RNA from SKM according to manufacturer's instructions. The extraction was followed sequentially by phenol/chloroform extraction and ethanol precipitation. Any contaminating DNA was degraded by two 10 min treatments with DNase I (ProMega; Madison, WI). DNase was then heat inactivated. SYBR Green Supermix with Rox (Applied Biosystems; Foster City, CA) was added to the PCR reaction (per manufacturer's instructions) and finally primers (see Table 1) for the gene of interest were added to the samples (in triplicate) while primers for an appropriate "housekeeping" gene (the 18S ribosomal subunit) were added to duplicate samples (in triplicate). A 7300 Real Time PCR System (Applied Biosystems) was used to detect SYBR Green fluorescence as a measure of amplicon. Sample C_T values were normalized to (subtracted from) the C_T values of the "housekeeping" gene and the number 2 was raised to a power equal to the difference between the sample C_T values of the 18S subunit and the gene of interest.

Dual Luciferase Reporter Assay. The Dual-Luciferase Reporter Assay Kit (Promega; Madison, WI) was used in accordance with manufacturer's instructions. Firefly luciferase and Renilla luciferase activities were measured with a TD-20/20 Luminometer (Promega, WI).

Western Blot. GMs were homogenized in Tris buffer (10 mM, pH 7.4) containing 1 mM EDTA and protease inhibitors (0.5 mM Pefabloc, 1 uM pepstatin A and 1 mM of each benzamidine and

iodoacetamide; Sigma-Aldrich, St. Louis, MO). The homogenates were centrifuged at 1000xg for 10 minutes and the supernatants were collected and the protein contents were determined using a DC Protein Assay Kit (Bio-Rad; Hercules, CA) according to manufacturer's instructions. Samples were immunoblotted basically as described earlier^{16,17}: protein samples (40 µg) were electrophoresed through 4-20% acrylamide gels, transferred to PVDF membrane (BioRad; Hercules, CA) and immunoblotted using Atrogin1 and MuRF1 antibodies (ECM BioSciences; Versailles, KY) and an ImmunStar Western Chemiluminescent Kit (Bio-Rad). Optical densities of protein bands were determined using ImageJ (NIH) software.

Study Designs.

Study 1. Eight groups of seven mice each were hindlimb suspended²² for either 1,2,3,4,5,7,10 or 14 days while another group of 7 weight bearing animals were used as day 0 controls (n=63). After the assigned control or suspension duration, each group of mice was killed according to the approved protocol and the GMs were harvested and flash frozen in liquid nitrogen. The left GMs from all mice were prepared for real time PCR and thus assayed for expression of mERG1a, MuRF1 and MAFbx. Normalized sample C_T values for the Day 0 control were averaged and the fold increase in gene expression for each gene per mouse was determined by calculating the ratio of each daily sample mouse value to the average Day 0 value.

Study 2. The left legs of 30 (6 groups of 5) mice were injected with 10 µg lacZ and 20 µg empty control expression plasmid whereas the right legs were injected with a combination of 10 µg lacZ and 20 µg mERG1a expression plasmid and all legs were subjected to electrotransfer. A group of 5 mice each was killed at 1,2,3,4,5 and 6 weeks each after electrotransfer and the GMs from all legs were harvested and flash frozen in liquid nitrogen. The muscles were cryosectioned and stained for lacZ activity as a marker for gene expression. Because a greater abundance (2X) of mERG1a than lacZ cDNA was injected, we assume that all myofibers staining for lacZ reporter activity also express *mERG1a* and that those myofibers not staining for lacZ activity are not expressing plasmid.

Study 3A. The left GMs of 49 mice (7 groups of 7) were injected with plasmid encoding *lacZ* (10 μg) and a second plasmid encoding mERG1a (30 μg) while the right GMs were injected with the *lacZ* plasmid (10 µg) and an empty (control) plasmid (30 µg), followed by electro-transfer. Seven mice were killed immediately to give a Day 0 data point. Seven more mice were killed at days 1,2,3,4,5 and 7 each post electro-transfer. RNA was extracted from GMs and real time PCR was used to determine expression levels of mERG1a, MuRF1 and MAFbx. C_T values were normalized (see methods) and the left (mERG1a treatment) to right (control) leg gene expression ratio was determined for each mouse as an indicator of treatment effect. Expression ratios for the Day 0 control were averaged and the fold increase in gene expression for each gene was determined by calculating the ratio of each daily sample mouse value to the average Day 0 value (baseline).

Study 3B. Ten mice were anesthetized and (as control) the GMs of both legs were injected with: 1) plasmid encoding the *Renilla* luciferase reporter (10 µg), 2) the *MAFbx*/Atrogin1 luciferase reporter plasmid (20 µg), and 3) an empty plasmid (20 µg). The right GMs of another 10 mice were injected as above while the left legs received the same mixture except that mERG1a plasmid (20 µg) was injected instead of the empty plasmid. Seven days after electrotransfer, the GMs were harvested and flash frozen in liquid nitrogen. These were later homogenized and

assayed for Renilla and firefly luciferase activities using a Dual Luciferase Reporter Kit (ProMega).

Study 4. Hindlimb suspension. Two groups of 3 mice each (n=6) were: 1) allowed to remain weight bearing; or 2) hindlimb suspended. After 7 days of treatment (day 7 is when our model induces significant levels of myofiber CSA decrease¹⁷), the mice were killed and the GMs were harvested and used to prepare muscle protein samples (both muscles of one mouse composed a sample). Electro-transfer. The GMs of a group of 6 mice (n=6) were injected with DNA (left = 10 µg lacZ and 30 µg mERG1a; right leg = 10 µg lacZ and 30 µg empty control plasmid) and the legs were electroporated. Two muscles were combined to make a sample; that is, two left legs made one mERG1a treated sample and two right legs composed a control sample, etc. GMs of mice from both designs were homogenized and centrifuged as described. Aliquots of each protein sample were electrophoresed and immunoblotted with either MuRF1 or Atrogin1 antibody.

Study 5. Mice (n=56) were randomly assigned to 4 groups (of 16 each), anesthetized and the left GMs were injected with expression plasmid: 1) lacZ (10 μ g) and empty plasmid (40 μ g); 2) lacZ(10 µg), empty (20 µg) and mERG1 (20 µg); 3) lacZ (10 µg), empty plasmid (20 µg) and HR-FOXO3a (20 μg); and 4) lacZ (10 μg), mERG1 (20 μg) and HR-FOXO3a (20 μg). All right GMs received *lacZ* (10 µg) and control plasmid (20 µg) and all GMs were electroporated. After 7 days, the muscles of 28 mice were assayed for mERG1a, HR-FOXO3a, MuRF1 and MAFbx expression by real time PCR and the fold increase in gene expression of left leg over right leg was calculated for each mouse. The muscle fiber CSAs were measured in the GMs of the remaining 28 mice.

Statistics. Data were analyzed using either ANOVA or a Student's t-test as reported in the figure legends. When ANOVA was used, data were analyzed for a completely randomized design. When significant differences were found, means were separated by Fisher's Protected Least Significant Difference. All data were analyzed using the General Linear Model Procedure of SAS (SAS Institute Inc.; Cary, NC). Statements of significance are based on p levels as noted in legends.

RESULTS

Study 1. Hindlimb suspension induces mERG1a, MuRF1 and MAFbx gene expression in the GMs of mice. Rationale. To aid in design of an optimal time course for our mERG1a ectopic expression model, we determined the time course of expression for mERG1a, MuRF1 and MAFbx genes in response to HS. Data. Real time PCR results show that mERG1a, MuRF1 and MAFbx genes were all expressed in response to HS (Fig. 1). Increases in MAFbx mRNA levels are noted as early as Day1 and continue to increase steeply to reach a maximum at Day 4. MAFbx mRNA levels begin to decline after Day 4 and expression of this gene is back to Day 1 levels between Days 7 and 10. mERG1a mRNA levels show a humble rise at Day 3 (Fig. 1, inset) just prior to a rise in MuRF1 mRNA levels, which increase sharply between Days 3 and 4 (Figure 1, inset). MuRF1 mRNA levels peak at Day 4 and then drop to initial levels between Days 7 and 10 while mERG1a mRNA levels peak at Day 5 before dropping to initial levels also between Days 7 and 10.

Study 2. Electrotransfer of expression plasmid into mouse GM yields gene transcription and translation with significant results lasting up to one month. Rationale. To support the validity of our studies reporting responses (over time) to electrotransfer of expression plasmid into skeletal muscle, we conducted a long term study of changes in GM CSA in response to mERG1a expression. Data. Indeed, as reported earlier¹⁷, the myofiber CSA was significantly decreased in the myofibers of the *mERG1a* injected legs expressing plasmid relative to the myofibers in the same leg not expressing plasmid and to both the expressing and non-expressing myofibers in muscles injected with *lacZ* and empty plasmid (Fig. 2). The decrease in fiber size is attributable to mERG1a expression and is significant for 4 weeks; the mERG1a expressing myofiber CSAs are no longer significantly smaller than controls at 5 and 6 weeks after electrotransfer.

Study 3. MuRF1, but not MAXbx, gene expression occurs within the GMs of mice in response to ectopic expression of mERG1a. A. Rationale. To isolate responses to mERG1a expression, we ectopically expressed mERG1a in GMs of mice and assayed for mRNA of interest over time to ensure that we did not miss changes in expression of the E3 ligase genes. Data. Real time PCR results show that indeed mERG1a was expressed in response to mERG1a plasmid electrotransfer in the GMs (Fig. 3A). Importantly, the MuRF1 E3 ligase gene was also expressed in response to the mERG1a expression and, although no significant differences from baseline transcript levels were detected at Days 1 or 2 post electrotransfer (Fig. 3A), the data reveal that mERG1a and MuRF1 transcript levels both increased above Day 1 levels significantly by Day 3 and peaked at Day 4. Transcript levels for both genes declined toward baseline levels by day 7. The data indicate that MuRF1 expression is increased by ectopic mERG1a expression. Most interestingly, there is no significant increase in MAFbx gene expression over the 7 days during which mERG1a was expressed, suggesting that ectopic expression of mERG1a did not activate this potential route for protein degradation. B. Rationale. To confirm that mERG1a expression does not affect MAFbx expression, we co-expressed mERG1a, a MAFbx luciferase reporter and a Renilla luciferase reporter (as control for differences in transfection efficiencies) and assayed for dual luciferase activities 7 days later. Data. The ratio of firefly to Renilla luciferase activities was determined for each leg and the ratio of luciferase activity in the left to right legs was calculated for each mouse (Fig. 3B). Although mice from a set of positive controls showed that mERG1a expression decreases levels of ubiquinated firefly luciferase (data not shown), in concert with expectations, MAFbx/Atrogin1 reporter luciferase activity was not significantly affected by mERG1a co-expression (Fig. 3B). These studies confirm that mERG1a expression does not affect MAFbx expression.

Study 4. MuRF1 protein levels increase in response to HS and ectopic expression of mERG1a while Atrogin1 protein levels increase in response to HS only. Rationale. Because increased mRNA expression is not always an indicator of augmented protein synthesis (nor can it be equated with increased protein activity), we assessed levels of MuRF1 and Atrogin1 proteins in GMs from both hindlimb suspended mice and those ectopically expressing mERG1a using western blots. Data. Both visual observation and optical density data show that MuRF1 protein (~37 kDa) levels increase over controls in the GMs of both HS mice and those ectopically expressing mERG1a (Fig. 4A and C, respectively); however, Atrogin1 protein (~41 kDa) levels increase only in response to HS (Fig. 4B,D). The Coomassie stained membranes confirm that equal amounts of protein were loaded into the lanes. These data further demonstrate that mERG1a expression induces transcription and translation of MuRF1, but not MAFbx.

Study 5. Block of FOXO3a DNA binding does not inhibit mERG1a up-regulation of MuRF1. Rationale. To test if MERG1a modulates MuRF1 expression through FOXO3a, we expressed a mutant form of FOXO3a (HR-FOXO3a, unable to bind DNA²³) $\pm mERG1a$ in WB mice and then assayed for atrophy. If the mERG1a K⁺ channel modulates MuRF1 expression through the FOXO3a transcription factor, then inhibition of active FOXO3a DNA binding will lower MuRF1 mRNA levels and diminish the changes in myofiber CSA shown to occur in response to mERG1a expression¹⁷. Data. The fold increase in mERG1a and FOXO3a mRNA levels in muscles injected with each respective expression plasmid shows that transcripts were made (Fig. 5A). The data also show that the MuRF1 mRNA level was significantly increased in response to mERG1a expression and confirm that MAFbx levels are not affected by mERG1a expression or by the levels of MuRF1 expression reached in this experiment. It further shows that mERG1a expression did not result in increased levels of FOXO3a transcription because our primers would

have detected this native mRNA. Although statistically insignificant, the fold increases in MAFbx transcript do "creep" above 1.0, hinting that the HR-FOXO3a could be inhibiting some basal level of MAFbx transcription. Most interestingly, there was no significant change in the levels of mERG1a-induced MuRF1 mRNA transcribed in the presence of HR-FOXO3a as would be expected if mERG1a modulated MuRF1 transcription through FOXO3a (Fig. 5A). Muscle fiber CSA data demonstrate that, indeed, muscle fiber size decreases in response to mERG1a expression; however, this decrease in muscle fiber CSA is not affected by expression of HR-FOXO3a (Fig. 5B). These sets of data suggest that mERG1a does not modulate MuRF1expression through FOXO3a activity. Verification: To ensure the validity of experiments performed with HR-FOXO3a, we determined that the HR-FOXO3a plasmid functions as a dominant negative when ectopically expressed in mouse GM. We ectopically expressed empty control plasmid (40 µg) in the right legs of 20 mice. The left legs of 10 of these mice received the same plasmid as control while another group of 10 received HR-FOXO3a (40 µg) in their left legs. All mice were hindlimb suspended. After 4 days, we harvested the GMs and determined the MAFbx mRNA expression levels using real time PCR and analyzed the data to yield ratios of MAFbx mRNA expression in left to right legs. As expected, HS resulted in increased MAFbx expression over weight bearing controls. The ratio of MAFbx mRNA expression in the control mice (left over right leg) was 1.06 ± 0.06 while the ratio was 0.8 ± 0.08 (p ≤ 0.05) in the mice receiving the HR-FOXO3a plasmid in their left legs, indicating the inactive HR-FOXO3a interfered with transcription of MAFbx in muscle of HS mice as would be expected.

DISCUSSION

In earlier work, we showed that the mERG1a K⁺ channel is up-regulated in atrophic GM from hindlimb suspended mice in contrast to muscle from weight bearing control animals and that mERG1a up-regulates UPP activity¹⁷. The mechanism(s) by which mERG1a modulate(s) UPP activity are unknown. We hypothesize that mERG1a acts upstream of pathways already known to modulate the UPP in SKM atrophy. For example, it has been shown that deactivation of the PI3K/AKT1/FOXO pathway and/or activation of the IKK-β/IκB-α/NF-κB pathway can upregulate UPP activity in atrophic SKM mainly by modulating expression of "atrogenes" (i.e., genes up-regulated during muscle loss). Particularly, these pathways have been shown to upregulate genes which encode the SKM specific E3 ligases, MuRF1 and Atrogin 1^{1,2,7}. In fact, recent studies demonstrate that both FOXO and NF-κB transcription factors are up-regulated in many atrophy models (including unloading)^{20,24-26} and one particularly elegant study by Reed and colleagues demonstrates that each of these transcription factor families is responsible for roughly 50% of the muscle loss occurring in immobilization-induced atrophy¹². Here we report that hindlimb suspension induces atrophy and up-regulates expression of mERG1a, MuRF1 and MAFbx genes and, interestingly, that ectopic expression of the mERG1a K⁺ channel in mouse GM increases expression of the MuRF1 gene, but not that of the MAFbx gene. We hypothesize, therefore, that although mERG1a could act on MuRF1 gene expression through a novel pathway, it likely modulates MuRF1 expression by a pathway currently known to modulate E3 ligase expression.

PI3K/Akt/FOXO: In general, deactivation of PI3K (phosphoinositide 3-kinase) results in reduced Akt phosphorylation and, thereby, deactivation of this enzyme. Once deactivated, Akt no longer phosphorylates FOXO transcription factors; the dephosphorylated/activated FOXO

(three isoforms have been described in SKM: FOXO1, 3 and 4) then moves to the nucleus and promotes transcription of MAFbx and/or MuRF-1 and other known atrogenes 1,2,7. Research with diabetes-induced muscle atrophy suggests that UPP activity is increased by suppressed PI3K and Akt activities with concomitant increased FOXO1 phosphorylation and resultant increases in MAFbx and MuRF1 mRNA levels²⁷⁻²⁹. In dexamethasone treated C2C12 myotubes, it has been shown that FOXO1 does not increase MuRF1 or MAFbx transcription directly, but does so indirectly by inhibiting the IGF-1 block of their upregulation³⁰. It has been demonstrated that FOXO3a expression results in increased levels of MuRF1 mRNA³¹ in cardiomyocytes and also induces the transcription of $MAFbx^{20}$. In immobilization studies, upregulation of MAXbx and MuRF1 expression occurs in numerous species including mice³² and rats^{8,33}. Unloading studies in humans also reveal changes in MuRF1 and MAFbx expression³⁴ and show that these changes differ according to which muscle is monitored and suggest, therefore, that E3 ligase expression may be fiber type specific^{4,35}. Nonetheless, the mechanism(s) responsible for these changes in E3 ligase abundances are not well understood.

Here we show that hindlimb suspension induces expression of mERG1a, MuRF1 and MAFbx. In fact, transcription of all the assayed genes occurs early in HS, prior to Day 7, at which time muscle fiber CSA is significantly smaller than in control WB mice¹⁷. However, while MuRF1 mRNA levels are correlated with mERG1a expression, MAFbx mRNA levels begin to rise prior to any noted increase in mERG1a mRNA levels, suggesting that at least initial MAFbx gene expression is likely not linked to HS induced mERG1a expression. Importantly, we demonstrate that ectopic expression of mERG1a indeed induces expression of MuRF1 in mouse GM. In fact, MuRF1 expression mirrors that of electro-transferred mERG1a very closely, with mERG1a levels rising at day 3 prior to a sharp increase in MuRF1 mRNA levels between Days 3 and 4.

These data suggest that the increase in MuRF1 expression is likely subsequent to the day 3 rise in mERG1a transcript. Interestingly, data also show that mERG1a expression does not induce MAFbx transcription. This is most interesting because this isolation of mERG1a-expression effects reveals a separation of E3 ligase modulation at the level of a membrane protein. Because it has been shown that FOXO3a induces expression of MAFbx specifically by binding directly to the MAFbx promoter in mouse muscle²¹, our data suggest that mERG1a does not modulate FOXO3a gene expression or protein activity. To test if mERG1a affects MuRF1 expression through FOXO3a, we co-expressed mERG1a and an inactive FOXO3a (HR-FOXO3a). We found that inhibition of FOXO3a activity (i.e., DNA binding) did not diminish changes shown to occur in response to mERG1a expression; that is, it did not lower MuRF1 mRNA levels or prevent the decrease in muscle fiber CSA seen in *mERG1a* treated controls. Further, our studies reveal that mERG1a expression does not modulate FOXO3a mRNA levels. Therefore, we conclude that MERG1a does not induce MuRF1 transcription through up-regulation of FOXO3a expression. Further, we suggest that it is not likely that MERG1a induces FOXO3a activity (increased inhibition of its phosphorylation or increased dephosphorylation) mainly because MAFbx mRNA levels do not increase with mERG1a expression as would be expected if FOXO3a activation were augmented and translocation to the nucleus were up-regulated. (Of course, our data does not rule out the possibility that mERG1a up-regulation of MAFbx transcription requires some MAFbx promoter specific co-factor not present in this model, but present during HS, which would allow FOXO3a to bind the MAFbx promoter.) It still remains to be definitively determined if the mERG1a channel modulates MuRF1 expression through the FOXO transcription factor family. We suspect that mERG1a does not affect FOXO factors because: 1) it does not modulate MAFbx expression (and it has been shown that FOXO 1 and 4

can modulate MAFbx expression in some models^{30,36}); and 2) increasingly research suggests that MAFbx expression is modulated by FOXO transcription factors while MuRF1 expression is regulated by NF-kB factors^{20,21,36,37}. Nonetheless, further studies with mutant forms of FOXO1 and FOXO4 should be performed to determine definitively that mERG1a is (or is not) inducing MuRF1 expression through these transcription factors. If FOXO family members are implicated, then studies determining if Akt and PI3K are involved will be required.

Other recent work reports that treatment of rat skeletal muscle with PPAR- α agonist yields decreased PI3K/Akt signaling activity, increased FOXO1 transcription factor activity and concomitant increases in MuRF1 and MAFbx expression³⁸. If mERG1a is increasing MuRF1 production by inactivating some point of the PI3K/Akt pathway, then it could be acting through effects of PPAR-α agonists on the membrane-bound PPAR-α receptors. Of course, the work does not address potential effects on other pathways (e.g., NF-kB factors), so it does not rule out that PPAR- α activation is increasing MuRF1 (or MAFbx) expression through other activity(-ies). PI3K/Akt independent mechanisms also have been shown to modulate FOXO activity. For example, Nemo-like kinase (NLK) has been shown to bind and phosphorylate FOXO1, thereby inhibiting its transcriptional activity through the PI3k/Akt independent transforming growth factor-beta-activated kinase (TAK1)-NLK pathway³⁹. Further, Atrogin1 expression has been shown to be induced by a p38MAPK dependent pathway in cardiac myocytes, independent of the Akt/FOXO pathway⁴⁰. These routes represent further possibilities for mERG1a modulation of MuRF1 expression.

IKK-β/IκB-α/NF-κB: Activation of NF-κB family members by upstream IKK-β activation leads to production of E3 ligases^{2,24,37,41}. In brief, some factor triggers activation of IKK-B

(kinase) by phosphorylation. In cachectic atrophy, evidence suggests that activation of tumor necrosis factor alpha receptors triggers this step, but this does not appear to be the case with atrophy induced by unloading. In both cases, however, active IKK-β phosphorylates IκBα which is then ubiquitinated and degraded, releasing bound NF-κB factors. Different NF-κB factors are released dependent upon what triggers the atrophy¹. In unloading atrophy, it has been shown that levels of p50 (NF-κB factor) and Bcl-3 (an IκB family member) increase with atrophic conditions and, in fact, the decrease in SKM size caused by hindlimb suspension is ameliorated in both p50 and Bcl-3 knockout mice²⁶. Once released, the NF-κB proteins then translocate to the nucleus and cause transcription of numerous genes, including MuRF1, but interestingly, not $MAFbx^{37,41}$. We suggest, in fact, that because ectopic expression of mERG1a induces an increase in MuRF1 (and not MAFbx) mRNA and protein levels, that mERG1a would modulate (increase) the expression and/or activity of upstream *NF-κB* transcription factors. Nonetheless, given that mERG1a expression causes a significant increase in MuRF1 mRNA and protein levels and the fact that NF-κB factors are known to be involved in MuRF1 expression, the data suggest that mERG1a could modulate NF-κB factor expression and/or activity. Identifying if/which NF-κB factors are modulated by mERG1a and the mechanism(s) by which this occurs clearly requires study.

In summary, we present data which show that MuRF1 gene expression is significantly increased in response to mERG1a expression and confirms that MAFbx expression is not affected by mERG1a expression or by the levels of MuRF1 reached in these experiments. That is, we have shown what appears to be a complete separation of MuRF1 and MAFbx/Atrogin1 E3 ligase modulation at the level of a membrane protein. Further, we demonstrate that MuRF1 modulation by mERG1a is not occurring through FOXO3a. Considering our data, this latter conclusion is

not surprising given the facts that NF- κB is basically required for disuse muscle atrophy 27,37,42 and that MuRF1 transcription, and not MAFbx, is increased by NF-κB factors³⁷. Obviously, the mechanism by which the mERG1a K⁺ channel up-regulates MuRF1 expression in SKM needs further investigation along with the factors inducing up-regulation of the K⁺ channel itself. Investigation of the mechanism(s) initiating SKM atrophy is important because SKM atrophy is coincident with many pathological conditions and is related to increased disability, morbidity and mortality and understanding these pathways will suggest specific, more effective therapies for abatement of this debilitating condition.

ABBREVIATIONS

SKM - skeletal muscle

UPP - ubiquitin proteasome proteolysis

E3 ligase - ubiquitin protein ligase

MAFbx - Muscle atrophy F-box

MuRF1 - Muscle RING Finger-1

SCF - Skp1, cdc53/Cullin and F-box protein

NF-κB – Nfkappa B

ERG1a – ether-a-gogo-related gene 1a

mERG1a - mouse ether-a-gogo-related gene 1a

HERG1A - human ether-a-gogo-related gene 1A

HS - hindlimb suspension

GM – gastrocnemius muscle

WB – weight bearing

WT - wildtype

DN-mERG1a - dominant negative mERG1a

CSA - cross sectional area

 $lacZ - \beta$ -galactosidase enzyme

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Target Gene	TM °C	Forward Primer	Reverse Primer	Amplicon
mERG1a	60	5'-CGC AGA ACA CCT	5'-GCA GAA GCC GTC	119 bp
		TCC TCG ACA C-3'	GTT GCA GTA G-3'	
MuRF1	55	5'-GGG CTA CCT TCC	5'-ACT CCT CCT CCT	183 bp
		TCT CAA GTG-3'	CCT CAT CTG-3'	
MAFbx	55	5'-CGT CGC AGC CAA	5'-ATC CAG GAT GGC	171 bp
		GAA GAG AAA G-3'	AGT CGA GAA G–3'	
FOXO3a	70 f;	5'-AAA TGT TCG TCG	5'-GTC GCC CTT ATC	153 bp
	60 r	CGG CGG AAC-3'	CTT GAA GTA-3'	
18S	57 f;	5'-CGG CTA CCA CAT	5'-GCT GGA ATT ACC	123 bp
(ribosomal)	58 r	CCA AGG AA–3'	GCG GCT-3'	

Table 1. Forward and reverse primers used for real time PCR.

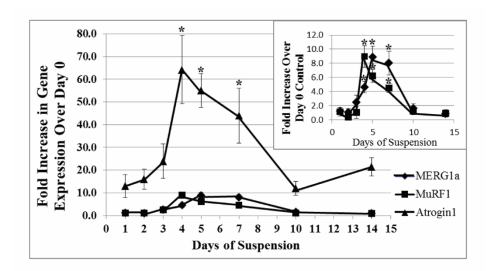


Figure 1. Hindlimb suspension (HS) induces mERG1a, MuRF1 and MAFbx gene expression in the gastrocnemius muscle of mice. Real time PCR demonstrates that HS induces highest expression of mERG1a, MuRF1 and MAFbx genes soon after commencement of suspension. Each daily data point was compared to the average day 0 level (day X/day 0) to yield a fold increase per mouse; fold increase per mouse data were averaged per day. Data points are, therefore, mean fold increase in gene expression (\pm standard deviation) over time. Inset: mERG1a and MuRF1 gene expression is graphed using a smaller y-axis to better show response to time course. Within gene, all data were analyzed by ANOVA (see methods). The "*" indicates a significant (p<0.05) increase in gene expression over the day 0 control within gene. 7 mice/day, n=63 mice.

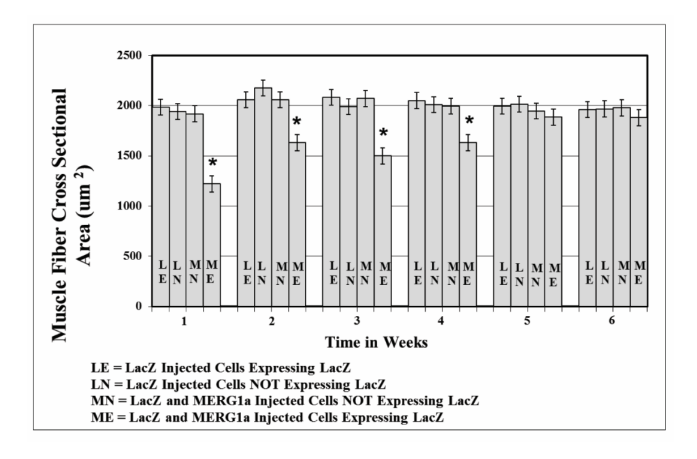


Figure 2. Electrotransfer of mERG1a and the lacZ reporter expression plasmids into gastrocnemius muscle (GM) yields results lasting 4 weeks. Relative to fibers of muscles injected with mERG1a and β -galactosidase (lacZ), but not expressing lacZ activity (MN), the cross sectional areas of the fibers injected with mERG1a and expressing lacZ activity (ME) are significantly decreased in weeks 1 through 4. The areas of the ME fibers are also significantly smaller than lacZ expressing (LE) and non-expressing (LN) fibers of GMs injected with lacZ only. There are no significant differences in fiber cross sectional area after 4 weeks. All data were analyzed by ANOVA (see methods). Statements of significance ("*") were based on p<0.05. n=30 mice.

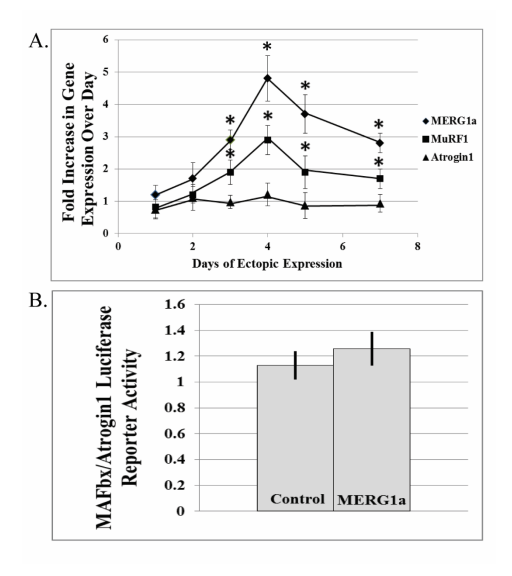


Figure 3. MuRF1, but not MAFbx, gene expression occurs within mouse gastrocnemius muscles (GM) in response to mERG1a expression. A. mERG1a and MuRF1 mRNA levels increase in response to mERG1a electro-transfer. Methods. Left GMs were injected with β -galactosidase (lacZ) and mERG1a plasmids; right GMs received lacZ and empty plasmid. After electrotransfer, GMs from 7 mice were harvested each day at days 0-5,7 and assayed with real time PCR. Data represent mean (± standard deviation) fold increase in daily gene expression over average day 0. Within gene, data were analyzed by ANOVA (see methods). The "*" indicates significant (p<0.05) increase in expression over day 0 within gene. 7 mice/day, n=49. B.

mERG1a expression in GM does not modulate MAFbx. Right GMs (n=20) were injected with: 1) Renilla luciferase reporter, 2) MAFbx/Atrogin1 luciferase reporter and 3) empty plasmid. Left GMs were treated identically except they received *mERG1a* instead of empty plasmid. Seven days post electro-transfer, GMs were assayed using a Dual Luciferase Reporter Kit (ProMega). The ratio of firefly to Renilla luciferase activities was determined per leg and activity ratios (left to right legs) were calculated per mouse. Data were analyzed by Student's T-test (SAS) and represent mean \pm standard deviation; p<0.55.

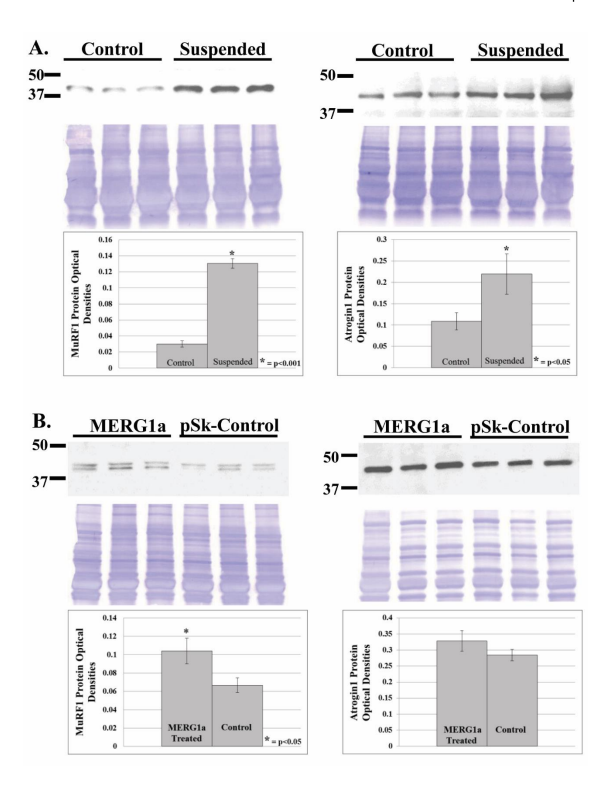
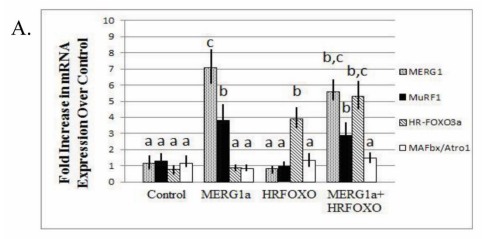


Figure 4. Immunoblots demonstrate that MuRF1 protein levels in gastrocnemius muscles (GMs) increase in response to both hindlimb suspension (HS) and ectopic expression of mERG1a while Atrogin1 protein levels increase in response to HS only. Protein samples (40 µg) from GMs

were electrophoresed through 4-20% acrylamide gels, transferred to PVDF membrane (BioRad; Hercules, CA) and immunoblotted with antibody recognizing either MuRF1 or Atrogin1 protein (ECM BioSciences; Versailles, KY). A and B. In both immunoblots (top), lanes 1-3 contain control GMs while lanes 4-6 contain GMs from mice suspended for 4 days. MuRF1 (A) and Atrogin1 (B) protein levels increased significantly (434% and 222%, respectively) in response to HS. C and D. In both immunoblots (top), lanes 1-3 contain GMs electro-transferred with mERG1a while lanes 4-6 contain GMs electro-transferred with control plasmid. MuRF1 protein levels (C) increased significantly (156%) while the increase (115%) in Atrogin1 protein abundance (D) is not significant. A-D. Coomassie stained membranes (below immunoblots) show that equivalent levels of protein were loaded into the lanes. Data were analyzed by Student's T-test (SAS). Bars represent the mean (± standard deviation) optical density units of proteins as determined by ImageJ (NIH). The p values are noted in the graphs.



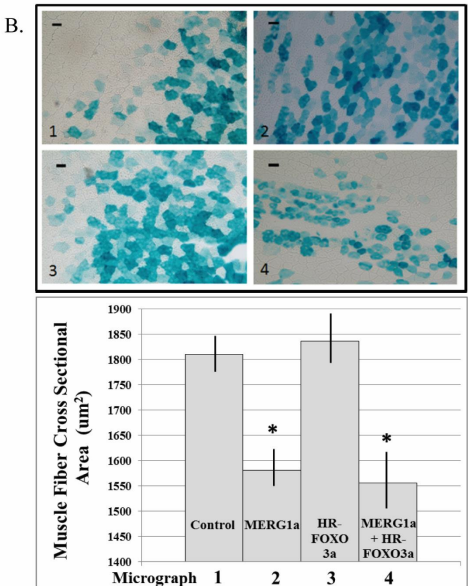


Figure 5. mERG1a-induced up-regulation of MuRF1 expression is not inhibited by block of FOXO3a DNA binding. We expressed HR-FOXO3a (an inactive mutant unable to bind DNA) \pm mERG1a in mouse gastrocnemius muscle (GM) and assayed for atrophy: A. E3 ligase expression; and B. muscle fiber cross sectional area (CSA). Figure A. Real time PCR indicates that the increase in MuRF1 mRNA levels noted in response to mERG1a expression is not curtailed by co-expression of HR-FOXO3a. Further, ectopic mERG1a expression does not affect levels of FOXO3a nor MAFbx mRNA. Data were analyzed by ANOVA (see methods) and are expressed as group mean \pm S.E.M. Different letters indicate significant differences, p \leq 0.05; n=28 mice. Figure B. The decrease in muscle fiber CSA that occurs in response to mERG1a expression is not affected by co-expression of HR-FOXO3a. GMs were injected with expression plasmids: 1. control and β -galactosidase (lacZ); 2. control, lacZ and mERG1a; 3. control, lacZ and HR-FOXO3a; and 4. lacZ, mERG1a and HR-FOXO3a. Seven days after electro-transfer, muscle cryo-sections were stained for lacZ reporter to indicate plasmid expression. Data were analyzed by ANOVA (see methods). Bars represent mean CSA \pm S.E.M. Statistical differences are noted by "*", p<0.01. n=28. Scale bar = 40 μ m.