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Upregulation of the creatine synthetic pathway in skeletal muscles

of mature *mdx* **mice**

Warrren C. McClure^a, Rick Rabon^a, Hirofumi Ogawa^b, and Brian S. Tseng^a

a Departments of Pediatrics, Neurology and Cell & Developmental Biology, University of Colorado Denver Health Science Center, The Children's Hospital Fitzsimons Campus, Mail Stop 8108, PO Box 6511, Aurora CO 80045

^b Department of Biochemistry, Toyama Medical and Pharmaceutical University, Faculty of Medicine, 2630 Sugitani Toyama 930-0194, Japan

Abstract

Duchenne muscular dystrophy (DMD) is a fatal neuromuscular human disease caused by dystrophin deficiency. The *mdx* mouse lacks dystrophin protein, yet does not exhibit the debilitating DMD phenotype. Investigating compensatory mechanisms in the *mdx* mouse is important. This study targets two metabolic genes, guanidinoacetate methyltransferase (GAMT) and arginine:glycine amidinotransferase (AGAT) which are required for creatine synthesis. We show that GAMT and AGAT mRNA are up-regulated 5.4 and 1.9-fold respectively in adult *mdx* muscle compared to C57. In addition, GAMT protein expression is up-regulated at least 2.5-fold in five different muscles of *mdx* vs. control. Furthermore, we find GAMT immunoreactivity in 80% of mature *mdx* muscle fibers in addition to small regenerating fibers and rare revertants; while GAMT immunoreactivity is equal to background levels in all muscle fibers of mature C57 mice. The up-regulation of the creatine synthetic pathway may help maintain muscle creatine levels and limit cellular energy failure in leaky *mdx* skeletal muscles. These results may help better understand the mild phenotype of the *mdx* mouse and may offer new treatment horizons for DMD.

Keywords

mdx; Duchenne muscular dystrophy (DMD); creatine; skeletal muscle; guanidinoacetate methyltransferase (GAMT); arginine:glycine amidinotransferase (AGAT)

1. Introduction

Duchenne Muscular Dystrophy (DMD) is the most common, severe and lethal progressive muscular dystrophy to affect children. It is also the most common inherited lethal defect worldwide (1/3,500 liveborn males) [1]. There are a number of animal models of DMD including dog, cat, and mouse [2–4]; which all display varying degrees of histopathologic muscle features consistent with muscular dystrophy. The *mdx* mouse has been a valuable animal model of DMD because it lacks dystrophin protein [1], has elevated serum creatine kinase (CK)'s and elevated intracellular calcium [5,6] which are all similar to that of human

^{*}Corresponding author fax 303 724-3420, E-mail address: E-mail: brian.tseng@UCHSC.edu.

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DMD. In addition, *mdx* skeletal muscle has abnormal contractile properties [7,8] and exhibits pathologic dystrophic features [9,10] particularly in the diaphragm [11].

Despite the histopathologic similarities between the *mdx* and humans with DMD, we postulate the *mdx* mouse has made unique compensatory adaptations to dystrophin-deficiency to enable a relatively benign phenotype. *Mdx* mice exhibit cage activity indistinguishable from control mice [12], live a near normal life span [13] and can run in voluntary wheels at distances near those run by age matched control mice [13–16]. In spite of the histopathology seen in the diaphragm at 6 months of age [16], *mdx* mice do not suffer respiratory failure at that age and can still exercise at levels equivalent to the levels of control mice even up to 11 months of age [13,15].

The question of how the *mdx* mouse achieves this relatively benign functional phenotype is extremely important, when considering the severe phenotype displayed in human DMD where the same protein, dystrophin, is absent. Some explanations for the phenotypic variations in species such as differences in gait and life-span are readily apparent. It seems likely that there are many other compensatory molecular mechanisms contributing in combination. Since the *mdx* mouse is not severely crippled [12], has less fibrosis and more central nuclei than human DMD [10,17,18], we postulated that there may be additional compensatory molecular pathways or modifier genes in the *mdx* mouse that warrant further investigation. Diseasemodifying factors implicated in the *mdx* mouse have been described [19], such as extra-cellular matrix alterations [20–23], naturally occurring and experimental up-regulation of utrophin [24–27], myostatin inhibition [28,29], calcium protein-handling protein(s) [30–32] and enhanced satellite cell/regeneration functions [33–37]. While some of these modifiers are likely important in making the *mdx* phenotype relatively benign, many of these same *mdx* changes are occurring in parallel with DMD [2,12,38,39] and cannot fully account for the clear discordant phenotypic severity. Our overarching hypothesis is that there are compensatory pathways activated via modifier genes expressed in the *mdx* mouse that are not activated in the boys with DMD.

Several large-scale expression profiling or microarray studies of the mature *mdx* mouse hindlimb muscles have been published [23,40–43]. These studies were reviewed in an effort to find the most reproducible gene expression differences and then compare to three human DMD studies [44–46]. Genes that were up or down-regulated in the *mdx* mouse were compared to the DMD studies (manuscript in preparation). Genes where expression moved in parallel or the same direction (up or down) in both *mdx* mouse and human DMD were eliminated leaving only the genes that were differentially expressed. Of these genes, two in the same metabolic pathway guanidinoacetate methyltransferase (GAMT) and arginine:glycine amidinotransferase (AGAT) were found in multiple microarray studies to be up-regulated in *mdx* vs control mice while both down-regulated in DMD (vs human control). GAMT and AGAT are the only two enzymes required for creatine synthesis [47,48]. We were further intrigued by this novel finding given a prior study reported upregulated creatine kinase (CK) adaptations [49]. We were also struck with magnetic resonance spectroscopy (MRS) studies that showed near normal intramuscular creatine levels in *mdx* mice [50] yet intramuscular creatine levels in boys with DMD were 20% of control boys [51]. This present study reports both GAMT and AGAT upregulation for *de novo* creatine synthesis in mature *mdx* muscle which may help limit the cellular energy failure associated with the absence of dystrophin.

2. Materials and Methods

2.1. Animals, care, specimen collection and preparation

Adult control (C57BL/10ScSn) (n = 10) and *mdx* (C57BL/10ScSn-mdx) mice were obtained from Jackson Laboratories (Bar Harbor, ME). All mice were housed and handled in accordance

C57 ($n = 10$) and *mdx* ($n=12$) mice age-matched at 9–11 weeks were housed with 4 $\frac{1}{2}$ inch running wheels (Super Pet Mini Run-Around) adapted with bicycle odometers (Sigma Sport BC 401). Weekly running distances and weights were recorded over a six-week period. Average daily running distances were determined for each mouse. These values were averaged for *mdx* and control mice for an overall value. Differences between means were analyzed using Student's *t*-tests.

Blood was collected via retro-orbital eye bleed. Muscle and liver specimens were excised and frozen in liquid nitrogen for homogenization. Contralateral muscles were removed, mounted in Tissue-Tek O.C.T. compound (Miles Lab, Elkhart, IN) and frozen in isopentane cooled to −160°C in liquid nitrogen. Eight micron tissue cross-sections were obtained with a Leica cryostat at −18°C. Both C57 and *mdx* tissue sections were collected onto given poly-L-silane coated slides and stained with H&E. Serial sections were used for immunolabelling studies (see below) and succinate dehydrogenase (SDH) histochemical stain as reported [52].

2.2 mRNA Analysis

(100 mg/kg).

Total RNA was acid/phenol extracted from age 14–16 week old male control and *mdx* gastrocnemious mouse muscle using Trizol Reagent (Invitrogen). The RNA concentrations and purity from each sample were determined by fluorimetric analysis (Beckman DU 640 Spectrophotometer). Oligos (dT12–18) were used to synthesize cDNA from equal concentrations of RNA using SuperScript III Rnase H-Reverse Transcriptase (Invitrogen). Real time quantitative PCR was performed using Taq-Man reagents and ABI 7000 Prism Sequence Detection System. PCR Applied Biosystems Inc. (#377215) commercially available GAMT primers and probes, and custom AGAT (GenBank Accession # NC000068) primer/probe sets (Biosearch Technologies Inc.) [(FWD primer: 5′d(ATGGCTGACGAACTGTATG)3′; REV primer: 5′d(GGCCAATTTGTGTCTGT)3′ PROBE: 5′ CAL Fluor Orange 560 d (CCAGAATTATCCCATCCATTCCGTGGA); BHQ-1 3′)] were designed to span known introns to detect contaminant genomic DNA template if present. Commercially available primers for (#386832) glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (Applied Biosystems Inc.) served as an internal expression standard [53]. 96 well plates were set up with serial dilutions (1:1, 1:10, 1:100, 1:1000) of pooled control and *mdx* mouse cDNA to establish a standard curve for each primer set. Mean quantities were calculated on individual samples of control and *mdx* cDNA at 1:5 dilutions. Each PCR reaction sample was run in triplicate, and non-template controls were utilized for each primer/probe set. GAPDH, GAMT and AGAT primer pairs amplified the same cDNA stocks to determine mean fold change between control and *mdx* mouse strains and compare relative expression levels of GAMT and AGAT.

2.3 Western blot analysis

Excised muscles and liver were crushed with a mortar and pestle chilled with liquid nitrogen. Crushed tissue (100mg/ml) was homogenized on ice for 15 seconds using a Fisher Power Gen 125 polytron set on ¾ speed in buffer consisting of 1% Triton X-100, 1% sodium deoxycholate in phosphate buffered saline (PBS) with protease inhibitor cocktail (Roche). Protein concentration was determined using the BioRad DC Protein Assay Kit. Muscle (200 μg) homogenates were spun for 1 minute at 7,200 g. Pellets were suspended in equal volumes of ddH2O and loading buffer then run in 12% SDS PAGE gels before transfer to nitrocellulose for immunoblotting. Liver (60 μg) homogenates served as a positive tissue control. Membranes were incubated in blocking buffer containing 5% nonfat dry milk in tris-buffered saline with

0.1% Tween-20 (TTBS) at room temperature. Affinity purified rabbit anti-pig GAMT polyclonal antibodies previously used in mouse tissues (1:500 dilution) were incubated for 3 hours at room temperature [54]. Protein A conjugated to HRP (Zymed) was used at a 1:2500 dilution in blocking buffer for secondary detection in ECL western blotting solution (Pierce). Blots were exposed to CL-X Posure Film (Pierce). As an internal expression control, rabbit anti-desmin polyclonal antibody 1:500 dilution was applied in blocking buffer (MP Biomedicals, #681221) and secondary detection was carried with donkey anti-rabbit Alexa-594 (Molecular Probes). Images were collected using the Typhoon 9400 imaging system and quantitation of images was done using ImageQuant (GE Healthcare).

2.4 Creatine Concentration

A ninhydrin based biochemical assay was performed as published [55]. For muscle homogenates, 0.2 μg of total protein (BioRad DC Kit) were used for each muscle in each reaction. For serum samples, 20 μl of each serum sample was used per reaction. Reactions were run for 10 minutes in black 96 well plates then fluorescence was measured using a Spectromax Fluorimeter (Molecular Devices). Concentrations were determined based on standard curves with serial dilutions using purified creatine (Sigma-Aldrich). Standard deviations were calculated and Student's *t*-tests compared values of control and *mdx*.

2.5 Indirect Immunofluorescence

10 μm frozen sections were stained with Hematoxylin and Eosin (H&E) while immunofluorescence used cryosections blocked with buffer consisting of 10% goat serum 0.2% Triton-X 100 in PBS. AGAT antibodies are not currently available. Affinity-purified rabbit anti-pig GAMT polyclonal antibody diluted 1:200 in blocking buffer was applied to cryosections for 4 hours at room temperature [54]. As a control for non-specific binding, anti-Green Fluorescent Protein (GFP) antibody (Sigma G1544) diluted 1:100 was applied for 4 hours at room temperature. Secondary antibody (donkey anti-rabbit Alexa-488; Molecular Probes) diluted 1:400 was applied to sections for 1 hour. To identify early regenerating fibers, embryonic myosin heavy chain antibody (EMyHC) F1.652 (DSHB University of Iowa) diluted 1:400 [56] was secondarily labelled with secondary donkey anti-rabbit Alexa 596 (Molecular Probes). Immunofluorescence protocol was carried out using the M.O.M. kit (BMK-2202 Vector Labs) with a 1:20 primary antibody dilution and a 1:400 dilution of streptavidin Alexafluor 488 (S11223) from Molecular probes. DNA of nuclei were counterstained with 0.1% Hoechst (Sigma H6024). Sections were coverslipped with anti-fade mounting media. Images were captured on a Leica SM4000B fluorescent microscope with a Retiga 4000R QImaging CCD Camera (North Central Instruments, Inc.)

GAMT staining intensity, cell counts and cell area determination were carried out by outlining individual cell perimeters then using NIH Image software for analysis.

2.6 Evan Blue Vital Dye

Evans Blue Dye (EBD) is a vital dye that binds to serum albumin and is excluded from the intracellular space in healthy muscle fibers. EBD penetrates fibers with membrane disruptions and can be readily visualized under fluorescence [57]. Sixteen week old C57 and *mdx* mice were given an IP injection of 2% solution EBD (50 μl/10 g body weight) dissolved in normal saline, 4 hours prior to euthanasia [57]. EBD in sections is visualized using excitation/emission \approx 590 nm/617 nm) filters.

3. Results

3.1 Voluntary wheel running distances

Based on previous reports, *mdx* mice run distances nearly as well as control mice in voluntary exercise wheels [13–16]. In our Colorado facility at mild altitude (5280 feet above sea level), no significant difference in running distance was observed over a six week period in mature 10–15 week old C57 (n=10) and *mdx* mice (n=12) (Figure 1A). While there is some individual variation from week to week, overall C57 mice averaged 5.7 km/24 hour while *mdx* averaged 5.2 km/24 hour (Figure 1B). These summary values are consistent with previous reported distances run by C57 and *mdx* mice [13–16].

3.2 mRNA analysis

To validate mRNA-based microarray and bioinformatics screening results of up-regulated GAMT and AGAT in *mdx* vs. control, Real-Time quantitative PCR was performed on mRNA from gastrocnemius muscles of *mdx* vs. control adult mice. Figure 2 shows that GAMT RT-PCR products are 5.4-fold higher in *mdx* vs. C57 (p = 0.003) and AGAT RT-PCR products in *mdx* mice are 1.9-fold higher vs. control ($p = 0.047$). GAPDH served as internal expression control as previously reported [53]. This quantitative mRNA based assay demonstrates that both GAMT and AGAT mRNA expression relative to GAPDH levels are upregulated in *mdx* mice vs. C57.

3.3 Western blots

After verifying GAMT mRNA upregulation in gastrocnemius muscles of 16 week old *mdx* mice, GAMT immunoblots were performed on *mdx* and C57 homogenates of 5 different skeletal muscles. AGAT antibodies are not available currently. GAMT protein has a calculated MW = 26kD and liver, which expresses highest levels of GAMT protein, served as a positive tissue control [54]. In Figure 3A, every *mdx* muscle group revealed at least a 2.5-fold higher GAMT protein vs. C57. The top half of the blots were immunoblotted with desmin antibody to verify equal protein loading and internal expression controls (RF and TA blots are shown in Figure 3B). Additional gels were run and stained with Coomassie blue to also verify equivalent loading of total protein in the gastrocnemius muscle (data not shown).

In figure 3C with western blot analysis of skeletal muscle, we show that 2 week and 4 week old C57 mice have elevated levels of GAMT signal which gradually decreases by 5 weeks of age and becomes undetectable by 16 weeks of age (Figure 3A). In contrast, 2 week, 4 week and 5 week old *mdx* mice have increased levels of GAMT (Figure 3C) which continued to show elevation GAMT protein through at least 16 weeks of age (Figure 3A).

3.4 Creatine Concentration

Creatine concentrations were determined in serum and homogenates of five different muscles and no statistically significant difference was found comparing control vs. *mdx.* There was a trend toward higher total serum creatine in *mdx* vs. control mouse serum but not significant (Table 1). Concentrations in both serum and muscle homogenates were comparable to levels previously reported in mouse [58].

3.5 Indirect Immunofluorescence

GAMT protein cellular localization in hindlimb muscle cryosections was performed using *insitu* indirect immunofluorescence. Increased anti-GAMT immunofluorescence (green) in the gastrocnemius muscle was visualized in the cytoplasm in the majority (80%) of muscle fibers of *mdx* mice while only background levels were seen in all fibers of control muscles (Figure 4A). Dystrophin immunofluorescence (data not shown) revealed <3% revertant fibers which

is consistent with previous reports [12] so the great majority of GAMT expression+ fibers are visualized in widespread distribution far beyond the rare revertant fibers.

Single fiber characteristics were further investigated by performing more detailed image analysis. In Figure 4B, single muscle fiber cross-sectional area was plotted against GAMT fluorescence intensity for 300 fibers. *Mdx* fibers were further grouped into central or noncentral nucleated fibers, both of which can express high levels of GAMT protein. Control fibers have a much smaller range of fiber sizes ranging mostly from 3000 to 7000 μm² while *mdx* fibers range in size from 500 to $14,000 \mu m^2$. Further analysis shows that control fibers (independent of size) have background levels of GAMT immunofluorescence whereas the *mdx* fibers show higher immunofluorescence in all fiber sizes with the greatest intensity being in the smaller caliber fibers with central nuclei \langle <2000 μ m²). Marked GAMT immunofluorescence intensity is also found in many of the mid-size (3000 to 7000 μ m²) *mdx* myofibers which should be considered mature fibers when compared to control fibers of the same size Figure 4A [59].

To determine if fibers expressing GAMT are only restricted to small regenerated fibers, anti-Embryonic Myosin Heavy Chain (EMyHC) was utilized on serial sections of gastrocnemeus muscles. Pathologic areas with degeneration-regeneration features are seen in the mature *mdx* mice and a great majority of GAMT immunofluorescence is seen in non-pathologic areas. There are small fibers with both elevated GAMT levels and EMyHC+ immunolabelling in these pathologic areas and comprise less than 5% of total fibers (Figure 5A). However, GAMT + expression is not restricted to nascent small fibers that express EMyHC as GAMT immunofluorescence is observed in a greater majority of larger fibers that do not co-label with EMyHC (Figure 5B).

In an effort to eliminate the possibility that mdx muscle fibers are prone to non-specific immunolabelling because of pathologic/inflammatory features, anti-GFP antibody was incubated with serial sections and showed no affinity to the sections (Figure 5C). Finally sections were also incubated without primary antibody followed by secondary antibody to look for non-specific binding showing little to no background fluorescence (Figure 5D).

16 week old C57 gastrocnemius cryosections did not reveal immunoreactivity to GAMT nor EMyHC antibodies above background levels; nor with secondary antibody alone (data not shown).

3.6 Evans Blue Dye

Finally, we assessed the co-localization of GAMT immunofluorescence with Evans Blue Dye (EBD), a vital dye for identifying leaky cells, in single fiber analysis on triple-stained cryosections, (GAMT - green, EBD - red & Hoechst - blue) (Figure 6). Serial sections of a rare hyper-contracted fiber from mature *mdx* RF of the quadriceps muscle group are shown (40X objective lens). A loss of GAMT immunofluorescence (panel B) corresponds to the fiber of interest (X) shown in the H&E stain from Panel A. Panel C shows presence of EBD (red) in the same fiber signifying membrane leakage. DNA in nuclei are stained with Hoechst (blue). This fiber is also highly oxidative as determined by succinate dehydrogenase (SDH) activity visible as dark blue. Some EBD+ fibers are also GAMT positive but these fibers did not necessarily appear pathologic (arrows). The great majority of *mdx* fibers with elevated GAMT immunofluorescence were not found to be hyper-contracted or severely damaged (H&E criteria) and the few pathologic fibers (hyalinized or hypercontracted) that were identified exhibited minimal to no GAMT immunofluorescence. Central nuclei are a pathologic but relatively minor feature compared to this hypercontracted appearance. We have seen at least ten severely pathologic fibers with these same features of EBD+, SDH+ and GAMT negative.

Other *mdx* fibers that exhibited minimal GAMT immunofluorescence were the largest fibers $\left($ <7000 μm²) in cross-section (Figure 4) and tended to show traces of EBD uptake/fluorescence.

4. Discussion

Previous metabolic studies in *mdx* mouse skeletal muscle have revealed reductions in glycolytic and oxidative enzyme activities [60,61]. Similarly human DMD studies have reported also reductions in glycolytic, glycogenolysis and oxidative enzymes [62–64]. However, one *mdx* study suggested that there may be discrete metabolic adaptations occurring in the phosphocreatine synthetic energy pathway through up-regulation of creatine kinase enzyme activity and protein content [49]. Nuclear magnetic resonance (NMR) studies [65,66] and a biochemical study [67] of *mdx* leg muscles have shown 80–90% of creatine in *mdx* vs control mice whereas NMR studies of boys with DMD showed only 20% of muscle creatine compared to controls [51] while urine creatine is 17-fold higher in boys with DMD vs. control [68].

Creatine in skeletal muscle acts to buffer the "cytosolic phosphorylation potential" for large fluctuations in energy demands yet creatine is not thought to be synthesized in skeletal muscle but taken up from serum after synthesis in the liver or oral intake [48]. Therefore aim of this study was to determine if there is an up-regulation of the creatine synthetic pathway in *mdx* skeletal muscle that could help maintain physiologic intracellular creatine concentrations despite leakage into the serum.

In adult control mice, tissue blots have previously shown GAMT protein is predominantly expressed in liver and testis, not skeletal muscle [54]. However, GAMT and AGAT protein are both found in the skeletal muscles of rat embryos and postnatal 21 day old control mice suggesting de novo creatine synthesis may have an important role in the development of embryonic and perinatal skeletal muscle [58,69]. We believe that these features are also seen in small regenerative fibers ($\langle 500 \text{ µm}^2 \rangle$) of adult *mdx* mice with the novel observation that GAMT and AGAT mRNA are both significantly higher in adult *mdx* vs. control muscles (Figure 2). Figure 3 shows that five different muscles have 2.5–5 fold greater GAMT protein immunoreactivity in adult *mdx* compared to control.

Figure 3C demonstrates that GAMT protein expression is high in both *mdx* and control mice at 2 weeks of age. By 4 weeks and 5 weeks of age GAMT protein expression drops in both control and *mdx* mice, but *mdx* mice maintain higher GAMT protein levels from 4 weeks on and certainly at 16 weeks of age (Figure 3A). GAMT immunofluorescence is detected in 80% of all skeletal fibers from the gastrocnemius in cross-section (Figures 4A and 5A) while GAMT/AGAT mRNA levels are also upregulated (Figure 2). It may be possible to reconcile the long-held mystery of why does the mdx mouse appear to undergo a necrotic crisis phase exclusively at 3–6 weeks of age [5] and this decrease in GAMT expression could perhaps account for some aspect of this temporal relationship and recovery after 6 weeks of age. Other crisis factors may contribute including the following: weaning from nutrition of mother's breastmilk to standard rodent chow, increased weight-load, increased motor activity, etc.

These results suggest that *de novo* creatine synthesis can occur in small regenerating fibers but what about stably regenerated mature myofibers from adult *mdx*? To confirm that the mature fibers are not in an actively regenerating, serial sections of adult *mdx* mice were stained for EMyHC and found that fewer than 5% of all fibers stained positive and were always of small caliber ($\textless 500 \ \mu \text{m}^2$) as seen in Figure 4B. Fibers that are EMyHC+ do indeed also co-express GAMT at high levels, while other larger caliber mature fibers express marked GAMT without co-expression of EMyHC as seen in Figure 5A and 5B.

Considering the diaphragm has been shown to be the most severely affected muscle in *mdx* mice [11], one could postulate GAMT/AGAT would be decreased and not confer a

compensatory role. However we find the *mdx* diaphragm with western blot analysis shows an upregulation in GAMT protein. This may be due to several possibilities. One interpretation is different ages of *mdx* mice and technical differences employed in relevant studies. This study investigated 4 month old (16 week old) *mdx* mice which do not have profound respiratory, exercise or histologic deficits while other studies have shown severe pathologic alterations in the *mdx* diaphragm starting at 24 weeks of age (6 months) and histologically devastated by 16 month old *mdx* mice [12]. Another explanation may be that only small DIA fibers and regenerating fibers are expressing very high levels of GAMT but are shut down in the larger vulnerable fibers. Detailed single fiber diaphragm muscle immunolabelling analysis at different time points of mature and old mdx mice will be necessary to clarify this relationship if causal or incidental.

The up-regulation of GAMT protein expression in larger caliber (>2000 μm²), *mdx* mature fibers can be seen in figures 4 and 5. Since half of the fibers ranging from 2000 to 7000 μm² express higher levels of GAMT protein (compared to C57) we believe GAMT plays more than just a development/regeneration role in the *mdx* muscle fibers. Increased GAMT may help maintain creatine concentration in *mdx* mature fibers making those fibers more resistant to the metabolic challenges of leaky membranes.

The difference in GAMT immunofluorescence between large and small caliber fibers may offer insights into regulation of GAMT expression that prove valuable in understanding its expression. The largest *mdx* fibers have very little GAMT immunoreactivity. If GAMT expression is important to limit energy failure and support muscle cell survival, this finding may offer an explanation to previous work that reports small caliber fibers do not suffer necrosis compared to larger fibers in *mdx* mice [70,71]. It seems that GAMT up-regulation in adult *mdx* skeletal muscle may support both regeneration requirements of nascent fibers and enhanced metabolic support of established mature fibers including many stably regenerated fibers with central nuclei. The data presented here demonstrates that AGAT mRNA plus GAMT mRNA and GAMT protein are upregulated in multiple skeletal muscles. We believe this strongly implicates the novel finding that de novo creatine synthesis can occur in skeletal muscles of mature mdx mouse. Substrate availability for this pathway however, has not been determined in this study. Dietary experiments are currently underway in which mdx mice are fed modified arginine and glycine diets.

In Figure 7 we offer a model showing that de novo creatine synthetic upregulation in *mdx* mice could offer metabolic advantage for muscle cells which lack dystrophin and offset the sequelae of leaky membranes including metabolic compromises e.g. creatine kinase and creatine efflux. Maintaining normal ATP levels is essential for all normal muscle function including calcium pumps, myosin ATPases and membrane potentials. In addition to this model GAMT may have an important role in normal development and regeneration of skeletal muscle. We have begun studies breeding *mdx* mice with GAMT knockout mice. The GAMT null mice are small but viable and active [58]. Our double knockout studies should offer insights for possible GAMT loss-of-function effects. Initial observations of these double knockout mice have shown that they struggle to thrive prior to weaning (unpublished observations). Studies with these mice should provide insight of the potential role of GAMT up-regulation and *de novo* creatine synthesis in skeletal muscles of mature *mdx* mice.

As the creatine synthetic pathway is being investigated in the *mdx* mouse, one may wonder why human DMD oral creatine supplementation studies have not found statistically significant benefit with functional activities, activities of daily living or modified manual muscle testing [72,73]. Supra-physiological levels of supplemental creatine (10 –14% of diet from birth) resulted in no clear benefit in one *mdx* study [67] while a second oral creatine supplementation *mdx* study found histologic improvements in the EDL, not soleus [74]. We conclude oral

creatine supplementation studies have not shown striking functional improvements in human DMD nor *mdx* mouse functional parameters so one may further explore avenues to enhance oral creatine as a treatment modality.

No direct study has ever demonstrated the existence of a functional creatine transporter (CrT) at the membrane of a dystrophin-deficient skeletal muscle although it has been shown to localize on the membranes of control embryonic rat skeletal muscle in addition to brain and intestines [69]. Dystrophin-deficiency can lead to disrupted membrane complexes, such as those involving dystroglycans [75] and freeze-fracture scanning electron-microscopy studies have shown a profound loss of intramembranous particle density in *mdx* skeletal muscles [76,77]. It stands to reason that transmembrane proteins and transporters such as the CrT may also have aberrant localization or impaired function in dystrophin-deficient muscles. Efficacy of oral creatine supplementation may be undermined if insufficient CrTs are at the muscle membranes of *mdx* and DMD. If *mdx* muscle cannot efficiently transport plasma creatine into the muscle cell comparable to controls, and *mdx* muscles maintain 80+% [67] (Table 1) of normal creatine levels, one could deduce that *de novo* creatine synthesis in skeletal muscle with the increased GAMT expression would be advantageous. The downregulation of GAMT expression [45,63,78] coupled with an impaired CrT function may explain why human DMD muscle only contains 20% of normal creatine levels [51]. Detailed studies are warranted to characterize the localization and function of CrT in the *mdx* mouse and in humans with DMD.

The work described in this manuscript uncovers a novel facet to possibly explain why the *mdx* mouse is not crippled. We do believe there are likely to be many differential genes and combined compensatory mechanisms in the *mdx* mouse. This study adds a new aspect to that complex and evolving story. The work done here demonstrates discrete creatine synthesis capabilities in mature skeletal muscle fibers not just small nascent fibers of the adult *mdx* mouse. Future promoter studies may help elucidate the apparent divergent signaling mechanisms that influence GAMT/AGAT gene regulation in mouse versus human skeletal muscle. In future efforts, these findings could be important directions to seek new targets and potential treatments for human DMD.

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Figure 1.

Average distances run in a 24 hour period were measured daily over a six week interval. In panel A, C57 mice (n=10) are denoted with a solid black box and *mdx* mice (n=12) are denoted with an open diamond. In panel B, bar graph summarizes the average daily voluntary wheel run distances of mature *mdx* and C57 mice. No statistical difference is seen.

Figure 2.

Quantitative real time Taq-man RT-PCR in gastrocnemius muscles of 16 week old *mdx* vs C57 mice. In mdx , both GAMT and AGAT mRNAs are significantly elevated, with $p = 0.003$ and $p = 0.047$, respectively. *Mdx* mice ($n = 5$) are denoted by black and white diagonal bars while C57 ($n = 5$) are denoted by black and gray diagonal bars.

McClure et al. Page 16

Figure 3.

Panel A shows GAMT immunoblot analysis of five different muscle groups of 16 week old *mdx* and C57 mice. Lanes 1–4 are *mdx* and lanes 5–8 are C57. Lane 9 is liver used as a positive control tissue for GAMT protein. GAMT protein has a calculated MW= 26kD. GAST is gastrocnemius; RF is rectus femoris; TA is tibialis anterior; HAM is hamstring; and Dia is diaphragm. $N=8-12$ in all groups have been tested with this representative blot showing $n=4$ per group. Two GAMT bands seen in the TA and HAM of C57 have unclear significance. Two alternative transcripts been reported in NCBI database and a human case report from lymphoblastoid tissue suggested the smaller GAMT species lacks a terminal exon and may be non-enzymatic [79]. The predominate GAMT band in the *mdx* mice is the larger 26 kd species.

The function of this smaller GAMT species in C57 is not known but does not undermine the observation that the mdx mouse highly upregulates 26 kD GAMT unlike C57. Panel B shows the upper half of blot membrane in Panel A probed with desmin antibody to verify relative equal loading and internal expression control. Expected molecular weight of desmin = 53 kD. (C) Gastrocnemius immunoblot analysis for GAMT in mdx and C57 mice ages postnatal age 5 weeks, 4 weeks and 2 weeks. Desmin immunoblotting demonstrated equivalent protein loading and internal expression control for each sample.

Figure 4.

Panel A. Indirect immunofluorescence of GAMT expression in transverse gastrocnemius cryosections of C57 and *mdx* mice. Sections were treated identically on the same slide and images were captured using identical exposure parameters. Panel B. Fiber cross-sectional area plotted against relative GAMT expression in 100 fibers of each C57 (O) and *mdx* gastrocnemius fibers grouped into central (\blacktriangle) or non-central nucleated fibers (\triangle).

Figure 5.

Indirect immunofluorescence on serial cryosections to show both GAMT and EMyHC protein expression in gastrocnemius muscles of 16 weeks old *mdx* mice. Panel A is immunostained for GAMT (green) and the area within the yellow line is a pathologic area. Panel B is a serial section of panel A, which shows the same region with fibers staining for EMyHC in red, circled with blue. Yellow stars denote fibers that are positive for GAMT and negative for EMyHC. White Xs denote fibers of a similar size that are negative for both GAMT and EMyHC. Adjacent cryosections incubated with a polyclonal antibody against green fluorescent protein in Panel C for non-specific antigenicity labeling and Panel D for secondary antibody only control for non-specific labeling. 16 week old C57 gastrocnemius cryosections are not immunoreactive to GAMT nor EMyHC antibodies (data not shown). Bar = 50 microns.

Figure 6.

Panel A – D are serial sections showing an isolated hyper-contracted (pathologic) fiber, denoted with an X, from mature *mdx* rectus femoris. Panel A. H&E staining. Bar = 40 microns. Panel B. Indirect immunoflourescence of GAMT protein appears green with Alexa 488 plus nuclear DNA (blue) with Hoechst stain. Panel C is the same section as panel B using different filter set to show Evans Blue Dye (red) is leaking into a subset of cells. Panel D shows succinate dehydrogenase (SDH) enzymatic activity with the greatest activity in the darkest fibers. White and black arrows denote fibers that are leaking (EBD positive), expressing GAMT, have high SDH activity but do not appear pathologic on H&E features. Images were captured using a 40X objective lens.

Figure 7.

We propose that leaky *mdx* muscle fibers are able to maintain near normal creatine concentrations by synthesizing creatine *de novo* through up-regulation of both GAMT and AGAT. Our data shows that creatine levels are similar in the serum and muscle homogenates of both *mdx* and control mice. The up-regulation of GAMT and AGAT for *de novo* synthesis of creatine would allow for discrete *mdx* metabolic adaptations to offset leaky membranes and excess loss of intracellular creatine. Limiting energy failure in muscle cells, *mdx* mice may be divergent in a unique metabolic adaptation compared to muscles from boys with DMD. Further studies including human analysis are warranted.

Creatine Concentration in C57 vs mdx Muscle and Serum Creatine Concentration in C57 vs *mdx* Muscle and Serum

-Serum creatine concentration (μ M) is not statistically different in WT vs. mdx (p=0.1). -Serum creatine concentration (μ M) is not statistically different in WT vs. mdx (p=0.1).

-There is no difference in WT and mdx muscle creatine concentrations. -There is no difference in WT and mdx muscle creatine concentrations.

-A minimum of 5 homogenates were tested per animall. -A minimum of 5 homogenates were tested per animall.