Molecular Pathogenesis of Genetic and Inherited Diseases

Long-Term Blocking of Calcium Channels in mdx Mice Results in Differential Effects on Heart and Skeletal Muscle

Louise H. Jørgensen,*† Alison Blain,* Elizabeth Greally,* Steve H. Laval,* Andrew M. Blamire,‡ Benjamin J. Davison,* Heinrich Brinkmeier,§ Guy A. MacGowan,*¶ Henrik D. Schrøder,‖ Kate Bushby,* Volker Straub,* and Hanns Lochmüller*

From the Institute of Human Genetics,* International Centre of Life, and the Institute of Cellular Medicine and Newcastle Magnetic Resonance Centre, Newcastle University, Newcastle upon Tyne, United Kingdom; the Institute of Clinical Research,† and Department of Clinical Pathology,¶ University of Southern Denmark, Odense C, Denmark; the Institute of Pathophysiology,§ Ernst Moritz Arndt University of Greifswald, Karlsruhe, Germany; and the Department of Cardiology,¶ Freeman Hospital, Newcastle upon Tyne, United Kingdom

The disease mechanisms underlying dystrophin-deficient muscular dystrophy are complex, involving not only muscle membrane fragility, but also dysregulated calcium homeostasis. Specifically, it has been proposed that calcium channels directly initiate a cascade of pathological events by allowing calcium ions to enter the cell. The objective of this study was to investigate the effect of chronically blocking calcium channels with the aminoglycoside antibiotic streptomycin from onset of disease in the mdx mouse model of Duchenne muscular dystrophy (DMD).

Treatment in utero onwards delayed onset of dystrophic symptoms in the limb muscle of young mdx mice, but did not prevent degeneration and regeneration events occurring later in the disease course. Long-term treatment had a positive effect on limb muscle pathology, reduced fibrosis, increased sarcolemmal stability, and promoted muscle regeneration in older mice. However, streptomycin treatment did not show positive effects in diaphragm or heart muscle, and heart pathology was worsened. Thus, blocking calcium channels even before disease onset does not prevent dystrophy, making this an unlikely treatment for DMD. These findings highlight the importance of analyzing several time points throughout the life of the treated mice, as well as analyzing many tissues, to get a complete picture of treatment efficacy. (Am J Pathol 2011, 178:273–283; DOI: 10.1016/j.ajpath.2010.11.027)

Duchenne muscular dystrophy (DMD) is a devastating neuromuscular disease affecting 1 in 3500 live male births. The disease is caused by mutations in the gene encoding dystrophin, a protein localized to the cytoplasmic surface of the sarcolemma. Here, dystrophin is believed to exert a dual role, both as structural protein important for membrane integrity and as a scaffolding protein for ion channels and various signaling pathways.1 Loss of dystrophin results in sarcolemmal instability, which causes the muscle to undergo continuous cycles of degeneration and regeneration. The regenerative capacity is not maintained, however, and this results in early, progressive weakening of skeletal muscle leading to loss of ambulation and respiratory failure.2 Dystrophin is also important in cardiac muscle, and patients frequently develop a dilated cardiomyopathy. One key finding in the pathology of DMD is an elevated intracellular calcium level,3 and several studies have explored the potential of pharmacological modulation of calcium influx as a therapeutic option in this disease.4–8

One of the earliest suggestions that calcium is important in DMD was based on cytosolic calcium accumulation in nonnecrotic fibers from DMD patient biopsies not observed in other dystrophies or myopathies.3 Calcium accumulation has also been reported in cultures of dys-

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Address reprint requests to Professor Hanns Lochmüller, MD, Institute of Human Genetics, International Centre for Life, Newcastle University, Central Parkway, Newcastle upon Tyne, NE1 3BZ, United Kingdom. E-mail: Hanns.Lochmuller@newcastle.ac.uk.
trophin-deficient myotubes from DMD patients and dystrophic mdx mice.9–11

The simplest explanation for excessive calcium influx is entry through microtubers in the sarcolemma, which are more abundant in the dystrophic muscle.2 However, calcium influx may also occur through the non-voltage gated calcium channels, such as the calcium leak channels12 or transient receptor potential (TRP) channel family members.13,14 The TRP channels have been proposed to be implicated in regulated calcium entry of skeletal muscle. Specifically, members of this family have been suggested to either directly form calcium channels or indirectly regulate calcium influx. It has been proposed that the nature of these channels could constitute store-operated calcium entry and/or stretch-activated calcium entry.15 Of these, the best characterized in skeletal muscle are TRPC1 and TRPC3. TRPC1 associates with α-syntrophin, a protein connected to the dystrophin-glycoprotein complex (DGC) of the sarcolemma.16 Disruption of the DGC complex in Duchenne muscular dystrophy could therefore alter these protein complexes and change the activity level of TRPC1, causing excessive calcium influx into dystrophic myofibers. TRPC3 has been shown to associate with the type 1 ryanodine receptor and other triadic proteins.17 This channel protein is not involved in store-operated calcium entry, but it has been suggested that TRPC3 could form a link between calcium entry and gene regulation in skeletal muscle via the calcineurin-nuclear factor of activated T cells signaling pathway.18

Recent studies have suggested a direct relationship between calcium dysregulation and dystrophin-deficiency. Transgenic overexpression of TRPC3 in mice is sufficient to induce muscular dystrophy without concomitant muscle membrane fragility,19 and inhibition of TRP channels in dystrophin-deficient mdx mice drastically reduces both calcium influx and dystrophy.19,20 Dystrophin suppresses spontaneous sarcoplasmic reticulum elementary Ca2+ release events, and this inhibitory control, mediated via mechanosensitive pathways, is disrupted in mdx mice.21 In addition, myofiber degeneration alone, as exemplified in the α-dystrobrevin null mouse, does not cause functional muscle defects, and so the development of the DMD/mdx dystrophic pathology cannot be explained solely by membrane fragility.22

The precise molecular nature of the channel or channels involved is currently unclear, and there is controversy as to whether the non-voltage gated calcium channels constituted by TRPC family members initiate the disorder by triggering a cascade of pathological events.23

Regardless of entry mechanisms, calcium overload in skeletal muscle fibers results in activation of calcium-dependent proteolytic enzymes (calpains), which cause necrosis and degeneration24,25 and which even degrade the compensatory protein utrophin.26 In addition, calcium overload has a detrimental impact on mitochondrial structure and function, resulting in apoptosis, decreased activity and mislocalization.27 This chronic damage eventually exhausts the normal muscle repair mechanisms mediated by the resident satellite cells28 and leads to end-stage disease.

DMD fetuses display both ultrastructural disturbances and increased calcium content of the skeletal muscles.29,30 In the mdx mouse model of DMD, the dystrophic process also begins early in embryonic development and includes stem cell loss and structural muscle changes.31 Knockdown of dystrophin in adult mice does not cause any evident dystrophy,32 indicating that the disease has a developmental onset. Based on the involvement of calcium homeostasis and stretch-activated calcium channels in dystrophic pathology and the embryonic/fetal prepathological disturbances, the objective of the present study was to test whether blocking calcium influx from embryonic development can prevent initiation of pathology or alleviate the pathology to a greater extent than has been shown previously using short-term treatment regimes.33–35 We started our treatment regimen when female mice were fertilized, as detected by a vaginal plug, and continued the treatment for up to 6 months after birth. The strategy adopted was to nonspecifically block calcium channels by treating our mice with the aminoglycoside antibiotic streptomycin in the drinking water, as previously described.8

Long-term treatment starting in utero of mdx mice did alleviate limb muscle pathology, but had adverse effects on cardiac and diaphragm tissue. This discordant tissue effectiveness could relate to variations in pathological severity of the analyzed tissues and suggests that the pathology is more complex than previously believed. Any preclinical drug trial conducted in the mdx mouse should therefore include analysis of a range of time points and a variety of tissues. Specifically, because DMD patients succumb to respiratory and cardiac failure, cardiac tissue and diaphragm should be included in any preclinical studies on treatments for DMD.

Materials and Methods

Animal Experiments

C57BL/10-mdx (mdx) and C57BL/10 (BL/10) mouse breeding pairs were plug-checked every day. When a vaginal plug was observed, the mother was treated with streptomycin (Sigma-Aldrich, St. Louis, MO) in the drinking water (4 g/L); the bottles were changed twice per week. Streptomycin is known to cross the placental membrane.35 On giving birth, the mother was treated on streptomycin in the water; the pups, when weaned, were continued on the same dose in their drinking water. The treated pups were analyzed after 6 weeks (n = 3–5), 10 weeks (n = 4–6), and 6 months (n = 3–5 for histology and Western blots, n = 5–11 for magnetic resonance imaging [MRI]) of continued postnatal treatment. Male mice were used for MRI scanning and Evans Blue dye uptake, because extensive control data were available in male mice for these measurements and to be able to correlate heart membrane damage with heart function. To minimize animal usage, females were used for all other analyses.

Toxicity of streptomycin was tested on mouse urine samples. Samples were collected by scruffing the mice.
and allowing them to urinate into an Eppendorf tube. The samples were then analyzed using a dipstick test for proteinuria and ketonuria (Bayer Multistix GP diagnostic strips; MediSupplies, Dorset, UK). Urine samples were additionally analyzed for proteinuria using a bicinchoninic acid kit for protein determination (Sigma-Aldrich).

All animal experiments were in accordance with the Home Office Guidelines, United Kingdom, and were performed under license no. PPL60/3445.

**Human Biopsy Material**

Human fetal tissue was obtained during autopsy of aborted fetuses aged 13 weeks (DMD) and 15 weeks (control). Diagnosis of DMD was made on the basis of DNA analysis. Biopsies were obtained and used according to the guidelines of and with permission from the Regional Ethics Committee for Southern Denmark #15879.

**Histology**

Diaphragm, tibialis anterior (TA), and heart ventricular tissue was harvested and processed for histology by snap-freezing in isопентane cooled in liquid nitrogen and stored at −80°C. Cryosections were cut (10 μm thick) and were stained with hematoxylin and eosin (H&E) for general pathology, Sirius Red for detecting fibrosis, and Alizarin Red for detecting large accumulations of calcium/necrotic fibers, using standard procedures.

Muscle regeneration was assessed by counting the number of myofibers with centrally localized nuclei. Two random areas covering a total of 300–500 fibers were counted from 3 to 5 animals at 6 weeks and at 10 weeks of age. Sections were analyzed and photographed using a Zeiss Axioscan microscope.

**Immunohistochemistry/Immunofluorescence**

The 10-μm cryosections were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) for 15 minutes. Sections were washed and endogenous peroxidase blocked in 1.5% H2O2/Tris-buffered saline (TBS) followed by washing and blocking of endogenous biotin using avidin-biotin (DakoCytomation, Glostrup, Denmark). This was followed by three washes in TBS-Tween20 (TBS-T); finally, the slides were blocked with 4% bovine serum albumin (BSA)/TBS. Primary antibodies were used mouse-anti-human desmin 1:25 (DakoCytomation, D33) labeled with biotinylated-F(ab')2 fragments using the ARK research kit (DakoCytomation) as previously described,36 rat-anti-CD45 1:50 (MCA1388; AbD Serotec, Kidlington, UK) as previously described,37 and rabbit-anti-TRPC3 (Alomone Labs, Jerusalem, Israel) 1:500. TRPC3 was detected using donkey anti-rabbit Alexa594 1:200 (Invitrogen, Carlsbad, CA) and mounted using Vectashield with 4',6-diamidino-2-phenylindole (DAPI) for detection of nuclei (Vector Laboratories, Burlingame, CA). CD45 was detected using rabbit anti-rat biotinylated antibody 1:200, followed by streptavidin-horseradish peroxidase and development using the DAB+ kit (Vector Laboratories), nuclei were counterstained in hematoxylin and slides were mounted with Aquatex (Merck Biosciences, Darmstadt, Germany). Desmin was detected similarly to CD45, except for omitting the secondary antibody step.

Desmin- and CD45-positive areas marking regenerating and inflamed areas, respectively, were semiquanti-fied using a scoring system as follows: no staining = 0, weak staining with few cells/structures/areas = 1, moderate expression with small number of areas/structures = 2, strong expression, vast areas, many cells/structures = 3, and massive expression all over the tissue = 4. Sections were analyzed and photographed using a Zeiss Axioscan microscope.

**Western Blotting**

Two different preparations were used, either crude muscle extracts, used for the 6-week time point (10 mmol/L Tris-HCl, 1% Triton X-100, 0.5 μL/ml phenylmethylene-sulfonilfluoride [Roche], and protease inhibitor cocktail [Roche]) or the total membrane fraction isolated using a plasma membrane protein extraction kit (BioVision, Mountain View, CA), used for the 10-week and 6-month time points. Only total membrane fractions were extracted and analyzed using the kit.

The preparations (200–500 ng total membrane fraction or 25 μg crude extracts) were mixed in a 30-μL volume containing loading buffer (Invitrogen), reducing agent (Invitrogen) and water. Samples were incubated at 95–100°C for 5 minutes, briefly spun down and loaded onto Bis-Tris 4–12% gradient gels (Invitrogen). Gels were run for 45 minutes at 200V. Proteins were transferred to a PVDF membrane (GE Healthcare, Waukesha, WI) for 1 hour at 350 mA. Loading and transfer were analyzed using Ponceau staining (Sigma-Aldrich). Membranes were washed in TBS-T and blocked for 1 hour at room temperature or overnight at 4°C in TBS-T with 5% nonfat milk. Primary antibodies were incubated overnight at 4°C except for neural cell adhesion molecule (NCAM), which was incubated for 2 hours at room temperature, in the following concentrations: rabbit-anti-TRPC3 1:4000 (Sigma-Aldrich), rabbit-anti-TRPC3 1:5000 (Alomone Labs), mouse-anti-β-dystroglycan 1:50 (43DAG; Novo-casta, Newcastle upon Tyne), mouse anti-dystrophin 1:100 (DYB; Novo-casta), and rabbit anti-NCAM 1:8000 (Millipore Bioscience Research Reagents, Temecula, CA). Membranes were washed three times for 10 minutes with TBS-T and incubated with secondary goat anti-mouse-horseradish peroxidase (HRP) 1:5000 or goat anti-rabbit-HRP 1:5000 for 1 hour at room temperature. Immunoreactive bands were detected using an enhanced chemiluminescence kit (GE Healthcare).

Semiquantification of Western blots (TRPC3 and β-dystroglycan) was performed using Adobe Photoshop, CS5, and standardized to the ponceau stain band of approximately 80–90 kDa.

**Evans Blue Dye Uptake**

Evans Blue dye (0.25 mg per 10 g body weight in PBS) was injected intraperitoneally 16–24 hours before MRI.
After in vivo cardiac measurements by MRI, muscles were dissected out for histological analysis. Atrial tissue was removed from the hearts, which were then washed in PBS and snap-frozen in isopentane cooled in liquid nitrogen along with TA and diaphragm muscle. Transverse sections (8–10 μm thick) were cut and incubated in ice-cold acetone at −20°C for 10 minutes, washed three times for 10 minutes with PBS, and mounted with Vectashield mounting medium (Vector Laboratories).

All sections were examined and photographed using a Zeiss Axioplan fluorescence microscope. Evans Blue dye staining shows a bright red emission. An Evans Blue dye uptake score was obtained as follows, performed blinded by two individual observers who scored all sections in parallel and decided on an agreed scoring: 0, no uptake; 1, fewer than three areas of uptake; 2, greater than three areas of uptake and at least one area >100 μm in diameter; 4, greater than three areas of uptake and with all areas showing massive uptake of >100 μm in diameter.

Magnetic Resonance Imaging

Mice were anesthetized using 5% isoflurane, and anesthesia was maintained at 1.5% in oxygen with a flow rate of 0.5 L/min. The tail vein was cannulated and the mouse was placed on a sled with cutaneous ECG electrodes, respiration pillow, and temperature detector (Ozai Research Instruments, Toronto, ON) and connected to SA instrument monitoring equipment (SA Instruments, Stony Brook, NY).

Mice were scanned in a 39-mm-diameter quadrature birdcage volume coil (Rapid Biomedical, Rimpar, Germany).

Images were acquired on a horizontal 7-T microimaging system (Varian; Agilent Technologies, Santa Clara, CA) equipped with a 12-cm microimaging gradient insert (maximum gradient, 40 mT/m).

After power calibration and global shimming, a series of four pilot transverse images were acquired over the heart. Single-slice coronal and sagittal images were then obtained to view the apex and mitral valve planes. These images were used to plan for the true short-axis plane.

Up to 10 contiguous short-axis slices were acquired to cover the entire left ventricle using a spoiled gradient-echo cine sequence: TR = 5 ms, TE = 1.42 ms, flip angle = 15 degrees, field of view = 30 × 30 mm, data matrix = 128 × 128, slice thickness = 1 mm. Images were ECG-triggered to the R-wave with a cine delay of 15 ms, and typically 30 phases were acquired distributed through the cardiac cycle. Images were zero-filled to a matrix size of 256 × 256.

Scans were analyzed using the freely available analysis software program Segment v1.8 (http://segment.heiberg.se) to give left ventricular functional parameters.

Statistics

Student’s unpaired two-tailed t-test with an α-value of 0.05 was performed using GraphPad Prism 5 (GraphPad, La Jolla, CA). Cardiac MRI data were compared using one-way analysis of variance.

Results

Prepathological Calcium Accumulation Indicates Involvement of Calcium Dysregulation in DMD/mdx Pathogenesis

Skeletal muscle from DMD fetuses displays ultrastructural changes including degenerative fibers, suggesting that onset of DMD occurs already in utero. It is unclear what comparison can be made between late gestation in humans and the early days after birth in the mouse; however, the onset of pathology is later in the mdx mouse, beginning about 3 weeks postpartum. We therefore wanted to analyze if we could detect calcium disturbances in relation to this early onset of DMD pathology and also extend the analysis to include detection of the earliest time point for accumulation of calcium in the mdx mouse model of DMD. This would prove our hypothesis that calcium accumulation immediately precedes the onset of histologically defined pathology.

We analyzed human DMD (13 weeks) and control (15 weeks) fetal skeletal muscle biopsies in comparison with mdx and C57/BL10 control mouse (BL/10) embryos at embryonic age 9.5, 11.5, and 18.5 days (data not shown) and mouse postnatal muscle at 12–13 days (data not shown) and 3 weeks old (Figure 1, A and B). We detected calcium accumulation in the human DMD fetal biopsy (Figure 1A), compared with the control fetal muscle. We did not detect any calcium accumulation in mdx muscles until age of known onset, which is around 3 weeks postnatally (Figure 1B). Centrally located nuclei indicative of ongoing muscle development were

![Figure 1. Pre-onset accumulation of calcium in dystrophin-deficient skeletal muscle suggests early implication of calcium dysregulation in DMD pathology. A: Calcium uptake (Alizarin Red positive fibers, red) in skeletal muscle from a fetus (13 weeks old) with DMD, compared with skeletal muscle from a control fetus (15 weeks old), which shows no uptake. B: Calcium uptake in tibialis anterior (TA) of C57/BL10 (BL10) and mdx mice was not detectable until postnatal week 3. Central nuclei were detectable in both C57/BL10 control (BL10) and mdx skeletal muscle (TA), confirming ongoing muscle development at this stage. Insets correspond to the higher-magnification image to the left. Scale bars = 100 μm.]
detected in both dystrophic and control mouse muscle, whereas in the dystrophic muscle an inflammatory infiltrate in the area positive for calcium uptake was observed as well (Figure 1B, H&E stain).

Short-Term Treatment with Streptomycin Improves mdx Limb Skeletal Muscle Pathology

Streptomycin treatment was started as soon as female mice were plugged, to expose the offspring in utero. At 6 weeks of age, an improvement in pathology in the TA muscle was detectable (Figure 2), consistent with previous short-term streptomycin treatments of mdx mice.6,8 A significant drop in the percentage of centrally localized nuclei (Figure 2A) (P < 0.05) and less inflammation (CD45 semiquantification 1.6 ± 0.55 in mdx S vs. 2.7 ± 0.57 in mdx C, P < 0.05) was observed in the treated animals (Figure 2D).

Active regeneration was observed as desmin-positive myotubes in dystrophic mice, regardless of treatment, suggesting that streptomycin could not completely prevent onset and progression of pathology. This was supported by the presence of necrotic foci regardless of treatment as judged by Alizarin Red staining (Figure 2D). TRPC3 expression overall was very patchy, even in C57BL/10 control muscle (BL/10 C), with localization to the membrane and cytosol without any obvious difference in expression level. However, Western blot confirmed that 6-week-old mdx C mice showed increased expression of TRPC3, compared with BL/10 C (Figure 2B).

Intriguingly, TRPC3 was detected in the nuclei of newly regenerating fibers (Figure 2D). This staining pattern was generally not observed in nuclei of BL/10 C muscle, un-injured mdx muscle, or nuclei in heart transverse sections from both mdx and BL/10 mice. A negative control staining, omitting primary antibody, showed no nuclear staining (Figure 2D, inset); staining was abolished by competition with the appropriate peptide (data not shown), and the antibody specificity has been verified previously.40 An identical staining pattern was observed with an alternative antibody raised against TRPC3 (data not shown). The increased expression of TRPC3 observed in mdx mouse muscle (Figure 2B) could therefore be a consequence of ongoing regeneration, rather than a primary pathological feature.

Streptomycin is an aminoglycoside, and some aminoglycosides read through premature stop codons.41 Because the mdx mutation is a stop codon in exon 23,42 it appeared possible that read-through had occurred and caused any improvement seen in our study. However, the improvement observed at 6 weeks of age was not caused by read-through of the mdx stop-codon mutation, because dystrophin protein was not detected (Figure 2C).

Continued Treatment up to 10 Weeks of Age Failed to Prevent Progressive Pathology in Limb, Heart, and Diaphragm Muscle

Although the 6-week time-point data were supportive for unspecific calcium channel blockade as a potential treat-
Abnormality in dystrophin in both control and treated mdx mice confirmed that there was no read-through activity at 10 weeks of age (Figure 3B).

TRPC3 expression showed the same overall features as the 6-week time point, with patchy expression and presence of TRPC3 protein in the nuclei of regenerating fibers as well (Figure 3C). Semi-quantification of TRPC3 Western blot (Figure 3D) showed that TRPC3 protein was increased in mdx C (1.62 ± 0.24) vs. BL/10 C (0.89 ± 0.59) (*P < 0.05). After treatment with streptomycin, TRPC3 protein expression level did not change in BL/10 mice (0.97 ± 0.31, *P > 0.05) BL/10 C vs. BL/10 S), but appeared to decrease in treated mdx mice (0.95 ± 0.45); however, this could not be confirmed by semi-quantification (*P > 0.05, mdx C vs. mdx S).

Semi-quantification of β-dystroglycan protein expression in mdx C (1.36 ± 0.31) vs. BL/10 C (2.46 ± 0.31) (*P < 0.05) showed down-regulation in dystrophic mice, as expected (Figure 3E). Expression of β-dystroglycan did not appear to change in BL/10 mice after streptomycin treatment (BL/10 S 1.90 ± 0.19, *P > 0.05 vs. BL/10 C). In dystrophic mice, treatment appeared to increase expression level of β-dystroglycan, but this could not be confirmed statistically (mdx S 1.90 ± 0.47, *P > 0.05 vs. mdx C).

We also examined heart muscle pathology at this time point, using Sirius Red staining for fibrosis followed by semi-quantitative analysis of the histological images (Figure 3F). There was a trend toward increased fibrosis in the hearts of treated mdx mice (mdx S % fibrosis 0.68 ± 0.47), compared with controls (mdx C % fibrosis 0.14 ± 0.071), but this did not reach statistical significance (*P > 0.05). No fibrosis was detected in hearts from BL/10 C or BL/10 S mice (data not shown).

**Long-Term Treatment Alleviates Limb Muscle Pathology but Cardiac and Diaphragm Pathology Is Worsened**

The final time point to be analyzed was after continued treatment until the age of 6 months. To examine whether long-term treatment with streptomycin induced kidney damage, total protein urine content was analyzed. All mice had similar levels (0.3 g/L). In addition, we analyzed the urine samples for ketones, and all were negative. Histological analysis of kidneys and livers in H&E-stained sections from paraffin-embedded tissue did not demonstrate histological changes in any of the groups (data not shown), suggesting that there were no adverse side ef-
fects from the treatment. Ototoxicity was tested by performing a simple hearing test in which fingers were snapped behind the mice and the Preyer reflex observed. All groups behaved similarly.

On routine histological examination of H&E stainings, an improvement in limb muscle pathology was observable in treated mdx mice. Specifically, large areas with ongoing necrosis were still detectable in mdx C (Figure 4A), whereas in mdx S mice (Figure 4A) the only sign of pathology was occasional small repaired areas with myotubes and centrally located nuclei without any necrosis. However, in both treated and control mdx mice >95% of the limb muscle fibers had centrally located nuclei (data not shown), suggesting that most muscle fibers had gone through at least one round of degeneration and regeneration. Diaphragm muscles presented with severe necrosis combined with large areas of fibrosis and regenerative fibers, suggesting that streptomycin treatment had no effect in this more severely affected tissue (Figure 4A). These histological observations were confirmed by Sirius Red staining for fibrosis (Figure 4B) showing that TA muscles from treated mdx mice (Figure 4B) appeared to have reduced thickening of the endo- and perimysium, compared with control mice (Figure 4B), whereas there was no obvious difference in the diaphragm muscles (Figure 4B).

We also examined the effect on cardiac pathology. Fibrosis was observed (Figure 4C) in hearts from both treated and control mdx mice, and on quantification the percentage of fibrosis was significantly increased in treated hearts (mdx C 3.27 ± 0.34 vs. mdx S 4.28 ± 0.28, P < 0.05) (Figure 4D). In addition, there was severe necrosis in the treated hearts, which was not observed in control dystrophic hearts (Figure 4C).

Magnetic resonance imaging scanning of treated and control BL/10 and mdx mice did not reveal any difference in left ventricular function between dystrophic mice, regardless of treatment, and both groups demonstrated well-compensated function, as previously reported using the conductance catheter technique. Cardiac pathology in dystrophic mice has previously been characterized by increased left ventricular mass. Magnetic resonance imaging confirmed that there was a significant increase in left ventricular mass in mdx C, compared with BL/10 C and that this was not improved in mdx S mice (Figure 4D). Overviews from the heart scans are presented in Figure 4D (long-axis and short-axis images taken from a healthy BL/10 C mouse).

No damage was observed in TA, diaphragm, or heart from BL/10 S, compared with controls, as judged from histological analysis (data not shown).

**Membrane Damage Is Reduced in Limb Muscle after Long-Term Treatment with No Effect on Diaphragm and Cardiac Muscle**

Membrane damage, as detected by Evans Blue dye uptake, was significantly reduced in TA muscle after 6 months treatment in dystrophic mice (Figure 5, A and B) (P < 0.01), compared with controls; however, there was no detectable difference in membrane damage in diaphragm and cardiac muscle (Figure 5, A and B).

Using Western blotting analysis of both heart tissue and TA, we confirmed that there was still no detectable read-through activity in the dystrophin gene by the aminoglycoside that could account for the positive effect streptomycin treatment (Figure 5C). There was increased...
protein expression of NCAM in TA of treated mdx mice, compared with all other groups, suggesting that there was increased muscle regeneration or stalled myofiber maturation in these mice at this time point (Figure 5D). Both treated and control mdx mice appeared to have a reduced expression of TRPC3 compared with the BL/10 groups. However, TRPC3 protein expression level appeared to be higher in treated mdx mice, compared with controls, again indicative of TRPC3 protein being actively transcribed and expressed by regenerating fibers, because increased NCAM was detected in the treated mdx mice as well (Figure 5D). Finally, there appeared to be increased expression of β-dystroglycan in the treated dystrophic mice, which appeared to have normalized toward wild-type levels (Figure 5D).

Figure 5. Membrane fragility is improved in limb muscle of mdx mice after 6 months of treatment with streptomycin, but heart and diaphragm pathology is not improved. A: Representative images of Evans Blue Dye (EBD) uptake in TA, diaphragm, and heart tissues harvested from the same mouse for mdx C and mdx S, respectively. There is great variability in uptake both in different tissues examined from the same mouse and in the same tissue types examined from different mice. B: Quantification of EBD uptake in TA, diaphragm, and heart tissue from mdx C and mdx S. Data are from 4–5 mice, expressed as means ± SD. **P < 0.01 for EBD uptake in TA in mdx C vs. mdx S. There were no significant differences in diaphragm and heart tissue, which both show a highly variable uptake pattern. C: Western blot detection of dystrophin (427 kDa) protein expression in total membrane fractions from 6-month-old TA muscle from BL/10 C, mdx C, BL/10 S, and mdx S. Representative results from two mice per group are shown. Ponceau staining was used as loading control (protein band of 80–90 kDa is shown). D: Protein expression pattern Tibialis Anterior.

Discussion

The understanding of the pathophysiological mechanisms of DMD has become increasingly complex since the discovery of the underlying genetic defect, and still no cure exists, more than 20 years later. The only available treatment to ameliorate skeletal muscle weakness and wasting is glucocorticoids, but this treatment is associated with potentially severe side-effects, such as weight gain and osteoporosis. It is therefore essential to continue testing new treatment strategies.

In the present study, we focused our attention on the hypothesis that calcium dysregulation could be a primary event in developmental onset of pathology, based on several recent studies suggesting that calcium channel proteins might be involved in disease initiation.

Our confirmation of early calcium accumulation in DMD fetal muscle is important for determination of the pathophysiological mechanisms, and also for defining the optimal time-point for initiation of treatment. This is supported by a previous study in which transgenic expression of utrophin in mdx mice from developmental onset reduced dystrophic pathology to a much greater extent than expression initiated at birth or postnatally, indicating that it is important when a therapeutic intervention is started. Based on the defects observed before histopathological onset in DMD/mdx, we wanted to analyze whether blocking calcium channels in the dystrophic mice from embryonic development and throughout adulthood could prevent pathological onset or progression. We observed a complex pathological pattern in which treatment outcome varied with age, ongoing disease progression, and tissue. Long-term treatment for 6 months, which may be the most relevant time point for comparison with the human condition, resulted in improved limb muscle pathology, but worsened cardiac pathology.

Previous studies have suggested a direct role for the TRP channel proteins in DMD pathology, and recent data show that transgenic expression of TRPC3 could induce a dystrophic pathology without concomitant membrane fragility. During this study we therefore recorded expression and localization pattern of TRPC3 to see if calcium channel blockade induced changes in expression pattern and whether this correlated with pathology.

TRPC3 expression was increased in mdx mouse muscle of young mice, but appeared to be reduced in older mice. Expression appeared stronger and localized both to the membrane and to cytoplasm and nuclei in regenerating fibers, compared with uninjured fibers in younger mice. The intracellular localization is in agreement with a recent study in which transgenic expression of TRPC3 with GLUT4 in the transverse tubular system. The nuclear localization of TRPC3 is unexplained, but has also been observed in a recent study, and regulated expression of TRPC3 during myotube development in vitro supports a correlation between TRPC3 expression and regeneration.

In the present study, localization of TRPC3 to the nuclei was observed only in regenerating myofibers of skeletal muscle; we did not detect this nuclear pattern in uninjured skeletal muscle or ventricular myocytes (data not shown). Our observations are in accordance with a re-
cent study in which localization of TRPC3 to the intercalated disks and intracellular membranes of ventricular myocytes was shown.52 Because the function of TRPC3 appears to be different in adult muscle versus developing muscle,14 our observations lead us to propose that the up-regulation of TRPC3 protein in young dystrophic mouse muscle could relate to ongoing regeneration. Nonetheless, dysregulated TRPC3 in early development caused by lack of dystrophin could also be a specific pathological feature. This needs to be investigated further in specific models of muscle regeneration.36,37

TRPC1 may form part of the putative calcium channels implicated in DMD pathology, because inactivating TRPC1 expression in mdx muscle fibers abrogated a calcium leak channel13 and because TRPC1 is directly associated with the dystrophin-glycoprotein complex.16 Specifically it has been proposed that the TRPC1 protein might constitute a stretch-activated calcium channel.53 However, recent observations in TRPC1 knockout mice suggest that this channel is not mechanosensitive, nor is it involved in store-operated calcium entry54; thus, the actual gating mechanism remains unknown. Nonetheless, TRPC1 appears to be indispensable for muscle activity, and TRPC1 knockout mice undergo muscle fatigue much faster than control mice.54 The role of TRPC3 and TRPC1 in relation to DMD remains elusive.

In the present study, reducing calcium influx in limb muscle via blockade of the calcium channels resulted in reduced signs of fiber damage in 6-week-old mice, in accordance with a previous study.48 The number of central nuclei then increased in both treated and untreated mdx mice with continued treatment and was no longer significantly different at 10 weeks of age. This suggests that there is an early delay in pathological progression after treatment.

Streptomycin has also been shown to have a directly harmful effect on mitochondrial ribosomes;56; thus, cumulative damage on mitochondria may present later as accelerated degeneration (observed here as increased numbers of centrally localized nuclei). Another explanation for the transitory effect might be that regeneration has occurred, not how many times each individual myofiber has undergone a round of necrosis followed by regeneration. It can therefore be speculated that, if the muscles have undergone fewer cycles of degeneration and regeneration, the satellite cell pool would benefit, because it will not become exhausted. The satellite cells will then still be available for repair of muscle fibers damaged by membrane fragility-induced events. This would result in the observed pattern of regeneration (increased NCAM expression) with reduced necrosis.

We observed a positive effect on membrane stability in the limb muscles of older, treated mdx mice, as ascer-
uptake might delay onset, but without concomitant correction of the membrane fragility defect, microtare, and nonspecific calcium uptake; eventually, necrosis, degeneration, and end-stage fibrosis will occur. This protection may be more pronounced in limb muscles, compared with diaphragm and heart, simply due to differences in physical activity. There are two possible scenarios to consider. First, the drug might simply reduce daily activity of the mice. This could relate to their behavior or to the mice feeling sick. It is important, because reduced or altered movements would directly reduce activity-induced necrosis of mdx mice. Therefore, introducing mechanical stress (eg, using forced treadmill running) and hence inducing necrosis of the muscle might counter the early improvement observed. We did not notice any obvious behavioral changes, and the mice appeared to be active and well-groomed, but this possibility should be tested using specific behavior studies and exercise regimens. Second, the relative activity level in limb muscle versus diaphragm and heart may in part explain differential effects of the drug. Physical activity level of the mice is more relevant for limb muscle than for heart and diaphragm, as the latter are in constant motion throughout life and undergo the same level of stress in both control and treated mice. Based on the present findings, systemic calcium channel blockade may not be a useful therapeutic strategy for DMD, as it may result in an aggravation of heart pathology. Calcium accumulation and structural disturbances before clinical onset in DMD/mdx needs careful consideration in designing future preclinical drug trials in the mdx mouse, because the type of muscle changes present at the start of intervention might influence the outcome considerably.

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References

1. Pilgram GS, Potikanond S, Baines RA, Fradkin LG, Noordermeer JN: The roles of the dystrophin-associated glycoprotein complex at the synapse. Mol Neurobiol 2010, 41:1–21
2. Deconinck N and Dan B: Pathophysiology of Duchenne muscular dystrophy: current hypotheses. Pediatr Neurol 2007, 36:1–7
3. Bedensteiner J, Nic B and Engel AG: Intracellular calcium accumulation in Duchenne dystrophy and other myopathies: a study of 567,000 muscle fibers in 114 biopsies. Neurology 1978, 28:439–446
4. Bertorini TE, Palmieri GM, Griffen JW, Igarashi M, McGee J, Brown R, Nutting DF, Hinton AB, Karas JG: Effect of chronic treatment with the calcium antagonist diltiazem in Duchenne muscular dystrophy. Neu- rology 1988, 38:609–613
5. De Luca A, Nico FR, Rolland JF, Cozzioli A, Burdi R, Mangieri D, Giannuzzi V, Lianot A, Cipponio V, De Bellis M, Nicchia GP, Camerino GM, Frigeri A, Svelto M, Camerino DC: Gentamicin treatment in exercised mdx mice: identification of dystrophin-sensitive pathways and evaluation of efficacy in work-loaded dystrophic muscle. Neurobiol Dis 2008, 32:243–253
6. Matsumura CY, Fertile A, Albuquerque TC, Santo Neto H, Marques MJ: Diltiazem and verapamil protect dystrophin-deficient muscle fibers of MDX mice from degeneration: a potential role in calcium buffering and sarcocellular stability. Muscle Nerve 2009, 39:167–176
7. Pernice W, Beckmann R, Ketelszik UP, Frey M, Schmidt-Redemann B, Haap KP, Roehren R, Sauer M: A double-blind placebo-controlled trial of diltiazem in Duchenne dystrophy. Klin Wochenschr 1988, 66:565–570
8. Yeung EW, Whitehead NP, Suchyna TM, Gottlieb PA, Sachs F, Allen DG: Effects of stretch-activated channel blockers on [Ca2+]+ and muscle damage in the mdx mouse. J Physiol 2005, 562:367–380
9. Mongini T, Ghigo D, Doriguzzi C, Bussolino F, Pescarmona G, Pollo B, Biffler D, Bosia A: Free cytoplastic Ca2+ + at rest and after cholinergic stimulus is increased in cultured muscle cells from Duchenne muscular dystrophy patients. Neurology 1988, 38:476–480
10. Bakker AJ, Head SI, Williams DA, Stephenson DG: Ca2+ levels in myotubes grown from the skeletal muscle of dystrophic (mdx) and normal mice. J Physiol 1993, 460:1–13
11. Turner PR, Fong PY, Denetclaw FW, Steinhardt RA: Increased calcium influx in dystrophic muscle. J Cell Biol 1991, 115:701–712
12. McCarter GC and Steinhardt RA: Increased activity of calcium leak channels caused by proteolysis near sarcosomal renales. J Membr Biol 2000, 176:169–174
13. Vandebrouck C, Martin D, Colson-Van Schoor M, Debaix H, Gailly P: Involvement of TRPC in the abnormal calcium influx observed in dystrophic (mdx) mouse skeletal muscle fibers. J Cell Biol 2002, 158:1089–1096
14. Krüger J, Kunert-Keil C, Bisping F, Brinkmeier H: Transient receptor potential cation channels in normal and dystrophic mdx muscle. Neuromuscul Disord 2008, 18:501–513
15. Whitehead NP, Yeung EW, Allen DG: Muscle damage in mdx (dystrophic) mouse: role of calcium and reactive oxygen species. Clin Exp Pharmacol Physiol 2006, 33:657–662
16. Vandebrouck A, Sabourin J, Rivet J, Balghi H, Sebille S, Kitzis A, Raymond G, Cognard C, Bourmeyster N, Constantin B: Regulation of capacitative calcium entries by alpha1-syntrophin: association of TRPC1 with dystrophin-complex and the PDZ domain of alpha1-syntrophin. FEASE B 2007, 21:608–617
17. Lee EH, Cherednichenko G, Pessah IN, Allen PD: Functional coupling between TRPC3 and RyR1 regulates the expressions of key triadic proteins. J Biol Chem 2006, 281:10042–10048
18. Rosenberg P, Hawkins A, Stiber J, Shelton JM, Hutcheson K, Bassel-Duby R, Shin DM, Yan Z, Williams RS: TRPC3 channels confer cellular memory of recent neuromuscular activity. Proc Natl Acad Sci USA 2004, 101:9397–9392
19. Millay DP, Gonnasekera SA, Sargent MA, Maillet M, Aronow BJ, Molkentin JD: Calcium influx is sufficient to induce muscle dystrophy via a TRPC-dependent mechanism. Proc Natl Acad Sci USA 2009, 106:19023–19028
20. Iwata Y, Katanosaka Y, Arai Y, Shigekawa M, Wakabayashi S: Dominant-negative inhibition of Ca2+ influx via TRPV2 ameliorates muscular dystrophy in animal models. Hum Mol Genet 2009, 18:824–834
21. Teichmann MD, Wegner FV, Fink RH, Chamberlain JS, Launikonis BS, Martinez B, Friedrich O: Inhibitory control over Ca(2+) sparks via mechanosensitive channels is disrupted in dystrophic deficient muscle but restored by mini-dystrophy expression. PLoS One 2008, 3:e3644
22. Bunnell TM, Jaeger MA, Fitzsimons DP, Prins KW, Ervasti JM: Dystrophin-deficient muscle from mdx dystrophic mice: role of calcium and reactive oxygen species. Clin Exp Pharmacol Physiol 2006, 33:657–662
23. Squire E, Raymakers JM, Vandebrouck C, Potter A, Tinsley J, Fisher R, Gillis JM, Davies KE: Prevention of pathology in mdx mice by expression of utrophin: analysis using an inducible transgenic expression system. Hum Mol Genet 2002, 11:3333–3344
24. Spencer MJ, Croall DE, Tidball JG: Calpains are activated in necrotic fibers from mdx dystrophic mice. J Biol Chem 1995, 270:10909–10914
25. Alport JM, Steinhardt RA: Calcium influx through calcium leak channels is responsible for the elevated levels of calcium-dependent proteolysis in dystrophic myotubes. J Biol Chem 2000, 275:9452–9460
26. Courdier-Fruh I, Brigueat A: Utrophin is a calpain substrate in muscle cells. Muscle Nerve 2006, 33:753–759
27. Kawamata H, Manfredi G: Mitochondrial dysfunction and intracellular calcium dysregulation in A517. Mech Ageing Dev 2010, 131:517–526
28. Charge SB, Rudnicki MA: Cellular and molecular regulation of muscle regeneration. Physiol Rev 2004, 84:209–238

29. Bertorini TE, Cornelio F, Bhattacharya SK, Palmieri GM, Dones I, Dworzak F, Brambati B: Calcium and magnesium content in fetuses at risk and preneurotic Duchenne muscular dystrophy. Neurology 1984, 34:1436–1440

30. Turk R, Sterrenburg E, de Meijer EJ, van Ommen GJ, den Dunnen JT, Dubois C, Figarella-Branger D, Pastoret C, Rampini C, Karpati G, McGeachie JK, Grounds MD, Partridge TA, Morgan JE: Age-related changes in replication of myogenic cells in mdx mice: quantitative autoradiographic studies. J Neurol Sci 1993, 119:169–179

31. Merrick D, Staeder LK, Larner D, Smith J: Muscular dystrophy begins early in embryonic development deriving from stem cell loss and disrupted skeletal muscle formation. Dis Model Mech 2009, 2:374–388

32. Ghahramani Seno MM, Graham IR, Athanasopoulos T, Trollet C, Pohlschmidt M, Crompton MR, Dickson G: RNAi-mediated knockdown of dystrophin expression in adult mice does not lead to overt muscular dystrophy pathology. Hum Mol Genet 2008, 17:2622–2632

33. Gannier F, White E, Lacamapagne A, Garnier D, Le Guennec YJ: Streptomyacin reverses a large stretch induced increase in [Ca2+]i in isolated guinea pig ventricular myocytes. Cardiovasc Res 1994, 28:1193–1198

34. Willems ME, Stauber WT: Streptomycin and EDTA decrease the effects of antibiotics on mitochondrial calcium channel blockers in isolated guinea pig ventricular myocytes. Cardiovasc Res 1990, 24:168

35. Rubin A, Winston J, Rutledge ML: Effects of streptomycin upon the human fetus. AMA Am J Dis Child 1951, 82:14–16

36. Roulet E, Birnbaumer L, Gailly P: Role of TRPC1 channel in skeletal muscle from dystrophic mice. J Neuro 2005, 281:H74–H88

37. Jørgensen LH, Jensen CH, Wewer UM, Schrøder HD: Transgenic mouse: a point mutation. Science 1989, 244:1578–1580

38. Heiberg E, Engblom H, Engvall J, Hedström E, Ugander M, Arheden H: Semi-automatic quantification of myocardial infarction from delayed contrast enhanced magnetic resonance imaging. Scand Cardiovasc J 2005, 39:267–275

39. Fayssoil A, Nardi O, Orlikowski D, Annane D: Cardiomyopathy in patients with Ca2+ channel blockers. Eur Heart J 2003, 24:960–964

40. Tadros SF, Kim Y, Phan PA, Birnbaumer L, Housley GD: TRPC3 ion channel subunit immunolocalization in the cochlea. Histochem Cell Biol 2010, 133:137–147

41. Welch EM, Barton ER, Zhuo J, Tomizawa Y, Friens WJ, Trilliphis P, Paushkin S, Patel M, Trotta CR, Hwang S, Wilde RG, Karp G, Taka- sugi J, Chen G, Jones S, Ren H, Moon YC, Corson D, Turpoff AA, Campbell JA, Conn MM, Khan A, Amshead NG, Hedrick J, Moloin A, Risher N, Weetall M, Yeh S, Branstrom CA, Colacino JM, Babiak J, Ju WD, Hirawat S, Northcutt VJ, Miller LL, Spatrick P, He F, Kawana M, Feng H, Jacobson A, Peltz SW, Sweeney HL: PTC124 targets genetic disorders caused by nonsense mutations. Nature 2007, 447:97–91

42. Scincis P, Geng Y, Ryder-Cook AS, Barnard EA, Darlisson MG, Barnard PJ: The molecular basis of muscular dystrophy in the mdx mouse: a point mutation. Science 1989, 244:1578–1580

43. Dubois C, Figarella-Branger D, Pastoret C, Rampini C, Karpati G, Rougon G: Expression of NCAM and its polysialylated isoforms during mdx mouse muscle regeneration and in vitro myogenesis. Neuromuscul Disord 1994, 4:171–182

44. Turk R, Sterrenburg E, De Meijer EJ, van Ommen GJ, den Dunnen JT, ‘t Hoen PA: Muscle regeneration in dystrophin-deficient mdx mice studied by gene expression profiling. BMC Genomics 2005, 6:98

45. Jero J, Coling DE, Lalwani AK: The use of Preyer’s reflex in evaluation of hearing in mice. Acta Otolaryngol 2001, 121:585–589

46. Grounds MD, Radley HG, Lynch GS, Nagaraju K, De Luca A: Towards risk: developing standards. AJP January 2011, Vol. 178, No. 1

47. Bauer R, Straub V, Blain A, Bushby K, MacGowan GA: Contrasting effects of steroids and angiotensin-converting enzyme inhibitors in a mouse model of dystrophin-deficient cardiomyopathy. Eur J Heart Fail 2009, 11:463–471

48. Quinlan JG, Hahn HS, Wong BL, Lorenz JN, Wenisch AS, Levin LS: Evolution of the mdx mouse cardiomyopathy: physiological and morphological findings. Neuromuscul Disord 2004, 14:491–496

49. Hofmann EP, Brown RH Jr, Kunkei LM: Dystrophin: the protein product of the Duchenne muscular dystrophy locus. Cell 1987, 51:919–928

50. Angelini C: The role of corticosteroids in muscular dystrophy: a critical appraisal. Muscle Nerve 2007, 36:424–435

51. McGeachie JK, Grounds MD, Partridge TA, Morgan JE: Age-related changes in replication of myogenic cells in mdx mice: quantitative autoradiographic studies. J Neurol Sci 1993, 119:169–179

52. Frango-Obregón A Jr, Lansman JB: Mechanosensitive ion channels in skeletal muscle from normal and dystrophic mice. J Physiol 1994, 481:299–309

53. Zanou N, Shapovalov G, Louis M, Tajedine N, Gallo C, Van Schoor M, Anguish I, Cao ML, Schakman O, Dietrich A, Lebacci J, Rouegg U, Roulet E, Birnbaumer L, Gaillly P: Role of TRPC1 channel in skeletal muscle function. Am J Physiol Cell Physiol 2010, 298:C149–C162

54. Holliday R: Streptomyacin, errors in mitochondria and ageing. Biogerontology 2005, 6:431–432

55. Grounds MD, Torrisi J: Anti-TNFα (Remicade) therapy protects dystrophic skeletal muscle from necrosis. FASEB J 2004, 18:676–682

56. Acharyya S, Butchbach ME, Sahenk Z, Wang H, Saji M, Carathers M, Ribich MD, Skipworth RJ, Feason KC, Hollandsworth SA, Musarella P, Burghes AH, Rafael-Fortney JA, Guttridge DC: Dystrophic glycoprotein complex dysfunction: a regulatory link between muscular dystrophy and cancer cachexia. Cancer Cell 2005, 8:421–432

57. Pullmann H and Vollmer B: An interaction of aminoglycoside antibiotics with Ca binding to lipid monolayers and to biomembranes. Biochem Pharmacol 1982, 31:3769–3773

58. Fayssoil A, Nardi O, Orlikowski D, Annane D: Cardiomyopathy in Duchenne muscular dystrophy: pathogenesis and therapeutics. Heart Fail Rev 2010, 15:103–107

59. Zhang W, ten Hove M, Schneider JE, Stuckey DJ, Becton MF, Skipworth RJ, Feason KC, Hollandsworth SA, Musarella P, Burghes AH, Rafael-Fortney JA, Guttridge DC: Dystrophic glycoprotein complex dysfunction: a regulatory link between muscular dystrophy and cancer cachexia. Cancer Cell 2005, 8:421–432

60. Fanchauy M, Poliajka E, Jung C, Ogrodnik J, Shirako N, Niggli E: Pathways of abnormal stress-induced Ca2+ influx into dystrophic mdx cardiomyocytes. Cell Calcium 2009, 46:114–121

61. Youm JB, Han J, Kim N, Zhang YH, Kim E, Joo H, Hoon LC, Joon KS, Cha KA, Earm YE: Role of stretch-activated channels on the stretch-induced changes of rat atrial myocytes. Prog Biophys Mol Biol 2006, 90:186–206

62. Sciscius P, Williams IA, Chu Y, Cooper PJ, Ju YK, Allen DG: Stretch-activated channels in the heart: contributions to length-dependence and to cardiomyopathy. Prog Biophys Mol Biol 2008, 97:232–249

63. Hu JF, Gilmer L, Hopkins R, Wolfbarger L Jr, Effects of antibiotics on cellular viability in porcine heart valve tissue [Erratum appeared in Cardiovasc Res 1990;24:168]. Cardiovasc Res 1989, 23:960–964

64. Schirmer M, Kaiser A, Lessenich A, Lindemann S, Fedrowitz M, Gemert M, Löscher W: Auditory and vestibular defects and behavioral alterations after neonatal administration of streptomycin to Lewis rats: similarities and differences to the cingulin (cic2/c2) Lewis rat mutant. Brain Res 2007, 1155:179–195