A model for creating a single stretch injury in murine biarticular muscle

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Abstract

Background: We developed a single stretch injury model to create damage near the musculotendinous junction (MTJ) of the gastrocnemius muscle in mice. Our hypothesis was that magnitude of muscle injury could be controlled by stepped shortening of the Achilles tendon (AT) prior to a lengthening contraction. Increased shortening would result in a greater isometric torque deficit and morphological damage 24 hours post-injury.

Methods: Sixteen mice were randomly assigned to sham or injury predicated on stepped increases in AT shortening. The AT was exposed and placed in a customized stainless steel roller-clamp system to achieve a specific level of shortening; 0 mm (resting length), 0.7 mm or 1.4 mm. Plantar flexors were stimulated to tetany with a needle electrode and then actively lengthened at 450°/sec from neutral to 75° of dorsiflexion. Passive and isometric torques were measured pre- and immediately post-injury. Isometric torque was measured again 24 h post-injury. Peak isokinetic torque was recorded during eccentric injury.

Results: Injury resulted in decreased passive and immediate absolute isometric torque only when induced with AT shortening. The percentage of pre-injury isometric torque was significantly lower in the AT shortened groups immediately and 24 h post-injury, but was unaffected by the level of shortening. Relative isometric torque deficits were noted in the 0 mm group only 24 h post-injury. Peak isokinetic torque during injury was similar in all groups. Histological evaluation 24 h post-injury revealed increased morphological damage near the MTJ in the AT shortened groups.

Conclusion: Single stretch with AT shortening created morphological damage near the MTJ and isometric torque deficits immediately and 24 h post-injury, but the magnitude of damage could not be titrated with stepped increases in AT shortening. This model provides an opportunity to utilize transgenic mice in order to elucidate inflammatory mediators that promote regeneration and inhibit fibrosis in order to optimize therapeutic interventions for complete functional recovery.

Keywords: Eccentric contraction, Experimental model, Muscle injury

Background

Muscle strains are among the most common injuries sustained at work, accounting for 49% of outpatient visits in the military, and comprise 30% of a typical sports medicine practice [1,2]. Incomplete functional recovery and susceptibility to recurrent injury are common sequelae of acute muscle strains. Treatment for muscle strain injuries has traditionally included rest, ice, compression and elevation (RICE). Promising new trends to promote complete recovery include novel therapies such as growth factors, gene and stem-cell based therapies [3]. Elucidating effective treatment measures requires a model by which a reproducible, standardized injury can be created that mimics acute muscle strains seen clinically. In this way, various inflammatory mediators that promote regeneration and inhibit fibrosis can be better understood and ultimately optimized for therapeutic intervention.

Acute strains should be distinguished from delayed onset muscle soreness (DOMS) which occurs after exercise involving repeated eccentric contractions and is characterized by microdamage to the muscle. Acute muscle strain injuries manifest as severe disruption at the musculotendinous
junction (MTJ) primarily in biarticular muscles [1,4]. Fast twitch or Type II fibers appear to be more vulnerable, possibly due to metabolic profile [5,6], higher tensions [7] or relatively shorter optimal lengths of the motor unit [8]. It is well established that muscle strains occur during lengthening of an activated high force generating muscle. Mechanical factors identified as contributing to injury include muscle fiber strain [9-12], peak force [13-15], the product of force and strain [9], contraction velocity [9,14], and initial muscle length [12,16]. However, there is not a consensus as to the relative contribution of these mechanical factors contributing to injury.

Models that have elucidated the mechanical events in eccentric injury have incorporated repetitive lengthening contractions in situ [13,15,17] and in vivo ranging from 100 to 900 stretches [6,18,19]. For example, 225 stretches to 10% beyond optimal length resulted in 65% force deficit mouse extensor digitorum longus [18] and 900 stretches at 12.5% strain caused a 40% deficit in rabbit TA [11]. It is difficult, however, to induce disruptive muscle damage at or near the MTJ that characterizes acute muscle strains with repetitive eccentric contractions or exercise [20-25]. Furthermore, assessment of damage is confounded with repetitive contractions as force decline is reflective of injury as well as fatigue [11,13].

Single stretch models performed in situ [9,26] and in vitro [27,28] have also provided powerful insights into the mechanisms for acute strain injuries. Most notably, a minimally invasive standardized single stretch injury method using tendon shortening of the tibialis anterior (TA) muscle in rabbits was developed by Best et al. [29]. Similar in vivo single stretch models have been subsequently adapted by other investigators for use in rats [30,31]. Plantar flexion without superimposed tendon shortening produced damage in the TA muscle belly but not at the MTJ [32], which is a hallmark of acute muscle strain injury [1,4,33]. Damage to the gastrocnemius MTJ was observed using ankle excursion in dorsiflexion rather than tendon shortening to manipulate injury [31].

The aim of this study is to evaluate a model for creating a standardized single stretch injury to the biarticular gastrocnemius muscle in mice using Achilles tendon (AT) shortening to control magnitude of injury. A customized stainless steel roller-clamp system with a bifurcated pin was used to shorten the AT. While the fast twitch fiber composition makes the TA an attractive target muscle, we sought to examine a biarticular muscle as they are more often injured compared to monoarticular muscles [1]. Prior work using tendon shortening [29] provided the unique opportunity to explore the role of inflammatory mediators in exacerbating damage following muscle strain in rabbit TA [34,35]. In the present study, we have sought to adapt this model to the murine species in order to capitalize on knowledge that can be gained through transgenic animals. A single stretch murine injury model would be instrumental in facilitating the exploration and optimization of novel treatments such as stem cell therapies for muscle injury [36].

**Methods**

**Instrumentation and experimental apparatus**

This system consists of a geared electric motor, torque sensor, angular position sensor, custom designed closed loop control system and nerve stimulator (Figure 1). Torque is measured with a custom fabricated cruxiform torsional load cell with a full scale range of 100 mN·m. Angular position is measured using a custom modified precision potentiometer (Vishay/Spectrol model 138-0-0-103, Vishay Americas, Shelton, CT). The motor is a Faulhaber Model 3863A024C + 38/2S43:1 + X0744 low inertia gear motor (MICROMO, Clearwater, FL). Command signals for the closed loop control system are supplied by a Wavetek arbitrary waveform generator (Model 75, Wavetek Corp, San Diego, CA). Three channels of data (torque, angular displacement and time) are recorded on a Dell computer using a 16 bit Data Translation model DT322 data acquisition board and Data Translation Measure Foundry data acquisition software (Data Translation Inc., Marlboro, MA).

**Experimental injury protocol**

This study was performed according to a protocol approved by the University of Wisconsin Institutional Animal Care and Use Committee. Sixteen C57BL/67 male 12 week old mice (26.1 ± 1.7 g, Jackson Laboratory, Bar Harbor, Maine) were assigned to one of four conditions (n = 4 mice, 8 limbs in each group); sham control group or injury group according to one of three incremental levels of AT shortening. The plantar flexor muscles were subjected to a single lengthening contraction without shortening (0 mm), AT shortening of 0.7 mm or 1.4 mm, corresponding to approximately 10% and 20% of resting AT length, respectively [37], consistent with the percentage of TA shortening in the rabbit model [29,38]. Tendon shortening was achieved using a bifurcated pin system connected to a precision potentiometer. To calibrate the shortening, a fine thread was inserted between the two pins and connected to a precision rectilinear potentiometer. The bifurcated pin potentiometer shaft was then rotated, while its output voltage and the rectilinear potentiometer output voltage were acquired with a data acquisition system. Collected data were converted into a displacement (shortening) versus voltage (theta) curve. A sixth order polynomial was then fit to the curve over the range of 0 to 4 mm with a resultant R^2 of 0.9999.

Mice were anesthetized with isoflurane and a 2-mm incision made to expose the AT. A custom splint was placed over the femur and tibia to ensure the knee joint angle
remained at 0° throughout the test. The animal was placed side lying in a metal half cylinder warming unit to maintain body temperature at 37°C. Care was taken to ensure that the animal was positioned so that the axis of ankle joint rotation was aligned with the axis of rotation of the loading frame. The foot was secured to a foot plate and locked in a neutral position (tibiotarsal angle of 90°) perpendicular to the tibia. Side lying was selected to eliminate the effect of gravity on joint torque. The ankle was aligned between two pins immediately anterior and posterior to the axis of rotation at the ankle joint. Brass cylinders of various sizes were machined to fit over the pin to allow the ankle to remain secured between the pins without impeding motion throughout the physiological range.

**Pre-injury passive peak and isometric torque measurements**

With the animal secured in the apparatus, the ankle was rotated from neutral through a 75° arc into dorsiflexion at an angular velocity was 450°/sec and the peak passive torque recorded (Additional file 1: Figure S1). The load frame was then fixed so that isometric torque could be recorded. The plantar flexors were stimulated to tetany (2.8 mA pulse train, 100Hz, 2.0 ms pulse width and triangular wave) with a needle electrode inserted subcutaneously in the popliteal fossa. To determine the simulating frequency, plantar flexor muscles were stimulated at 50, 75, 100 and 150Hz. Isometric torque did not increase from 100 to 150Hz and therefore 100Hz was selected. Two isometric torques were recorded with 3-min rest intervals between tests and the average torque was calculated.

**Isokinetic torque measurements**

The AT was visualized and placed in a customized stainless steel roller-clamp system and left at resting length (0 mm, no shortening), or shortened 0.7 mm or 1.4 mm. The bifurcated pin was tapered to allow ease of accepting the AT into the roller-clamp initially while prohibiting slippage once the tendon is turned to wrap back onto itself, shortening it 0.7 mm or 1.4 mm from resting length. A 1.0-s trigger delay between the initial muscle stimulation and ankle rotation ensured the achievement of complete muscle tetany. Muscle stimulation ceased immediately upon the footplate returning to neutral. In all groups, the angular velocity was 450°/sec and the ankle was rotated from neutral through a 75° arc into dorsiflexion and returned to neutral, at which time stimulation ceased. Torque-time plots were analyzed for ankle joint torque produced during muscle tetany and the peak joint torque at the end range of dorsiflexion (Figure 2). A sham group was included whereby the AT was exposed and ankle rotation occurred without stimulation or AT shortening (passive lengthening).
Post-injury peak passive and isometric torque measurements
The tendon was released from the roller-clamp system following injury isometric torque was immediately assessed as described above. The needle electrode was removed, the footplate unlocked, and peak passive torque was recorded. Incisions were closed with Vetbond and animals were returned to their cages for 24 h. Isometric torque measurements were again recorded 24 h post-injury in the absence of AT shortening.

Histology
The AT was transected and the triceps surae muscle resected. The gastrocnemius was isolated, dissected and frozen in isopentane cooled in liquid nitrogen and stored at −80°C. Serial 10 μm cross-sections immediately adjacent to the tendon were obtained using a cryostat (Leica/ Jung CM 1800 model, IMEB, Inc, San Marcos, CA) and subsequently stained with hematoxylin and eosin (H&E). Injury was identified as the region with greatest disruption of fibers and nuclei accumulation. Damage was characterized in a semi-quantitative manner by counting multiple nuclei accumulation (expressed as nuclei/mm²) 24 h post-injury. The cellular response to injury infers an increase in nuclei above baseline, but is not specific for myonuclei, neutrophils, macrophages and fibroblasts. In order to establish the location and extent of injury, damage was investigated in a spatially sensitive manner in the 1.4 mm shortening group. Tissue was carefully cut into 1 mm segments (0-1 mm, 1-2 mm, 2-3 mm and 3-4 mm) starting at the MTJ and continuing proximally through the muscle belly. As damage was localized immediately adjacent to the MTJ, only this area was viewed for subsequent groups. Micrographs were collected using a camera-assisted microscope (Nikon Eclipse microscope model E6000, Nikon Instruments, Inc., Melville, NY with an Olympus camera, model DP70, Olympus Imaging America, Inc., Center Valley, PA). Five sections from each muscle were viewed and two computer images (0.25 mm x 0.25 mm) were captured per section and viewed using Image J.

Statistics
Means and standard errors were calculated. The paired Students t test was used to compare the passive torques before and immediately after injury. Dunnett’s multiple comparison test was performed when absolute isometric torques immediately and 24 h post-injury were compared with pre-injury torques. Two-way analysis of variance (ANOVA) and the Bonferroni post hoc test were used to compare the percentage of pre-injury isometric torque between the groups at both time points. Statistical analyses were carried out using KaleidaGraph, version 4.03 (Synergy Software, Inc., Reading, PA).

Results
Passive torque
The peak passive torque during 75° of dorsiflexion was measured in the absence of tendon shortening in all groups prior to and immediately following injury. Passive torque was compared in a pairwise fashion with the pre-injury value. Significant decreases were observed in the 0.7 mm (P < 0.01) and 1.4 mm (P < 0.05) AT shortened groups (Figure 3).

Isometric and peak isokinetic torques
Isometric torque measures were recorded pre-injury and repeated immediately and 24 h post-injury. As shown in Table 1, multiple-comparisons of absolute isometric torque were significantly lower immediately and 24 h post-injury...
in the 0.7 mm (P < 0.01) and 1.4 mm (P < 0.001) AT shortening groups. The 0 mm group showed a trend for a torque deficit at 24 h (P = 0.07). The relative difference in isometric torque (percentage of pre-injury torque) between groups was examined immediately and 24 h post-injury and found to be significant (Figure 4). A substantial decrease in the percentage of isometric torque was observed in both the 0.7 mm and 1.4 mm AT shortened groups immediately (P < 0.01) and 24 h post-injury (P < 0.001) compared to all sham time points. The 0 mm injury group showed a trend for relative isometric torque deficit immediately (P = 0.07), and was significant at 24 h from all sham time points (P < 0.05). Peak isokinetic torque occurred at the end range of dorsiflexion and reflects the passive and active torques. Peak torque was unaffected by AT shortening and reached 40.06 ± 11.21, 39.49 ± 9.89 and 39.32 ± 9.67 mN·m in the three injured groups, respectively (mean ± SD). The isometric to peak isokinetic torque ratio is consistent with those reported by Best et al. in rabbit TA [29] using a similar model.

**Histology**

Damage was characterized by disruption of fibers and described in a semi-quantitative manner by counting multiple nuclei accumulation 24 h post-injury. Damage between groups was assessed immediately adjacent to the MTJ and found to be significant in injured groups with AT shortening (Figure 5). Nuclei accumulation in the injured gastrocnemius without AT shortening (0 mm) was 565 ± 38 nuclei/mm² and not different from uninjured controls (594 ± 20 nuclei/mm²). AT shortening to 0.7 or 1.4 mm resulted in increased cellularity above uninjured and 0 mm groups to 748 ± 86 and 888 ± 28 nuclei/mm², respectively, but there was not a further increase with the stepped increase in AT shortening (P < 0.05) (Figure 6).

**Discussion**

We have developed a single stretch injury model for murine gastrocnemius via shortening the AT that results in damage near the MTJ, similar to the location of damage in acute muscle strains noted clinically. Injury was defined by morphological damage and isometric torque deficits immediately and 24 h post-injury. A murine model for single stretch may provide a useful tool for utilizing transgenic animals to elucidate the role of various inflammatory mediators in order to facilitate regeneration and minimize fibrosis.

From a clinical perspective, the severity of muscle damage is perhaps best characterized by deficits in force production or torque [13,14,39-41]. Torque is the product of the force produced by a muscle or muscle group and the moment arm of the muscle(s) attachment. Absolute isometric torque measurements recorded using a torsional load cell dropped in AT shortened injured groups immediately post-injury and were consistent with values from other reports [42,43], but were not significant in the 0 mm injured group. AT shortening was prerequisite to elicit significant deficits in absolute isometric torque immediately and 24 h post-injury using multiple comparison analysis. Deficits expressed as the percentage of pre-injury isometric torque between groups revealed that AT shortening was prerequisite to elicit significant deficits immediately post-injury. Without AT shortening, deficits in percentage of isometric torque were only significant 24 h following injury. We propose that the mechanical damage elicited without AT shortening was sufficient to prompt further damage through the inflammatory cascade, resulting in deficits that persisted 24 h post-injury. AT shortening resulted in increased mechanical damage which manifested as immediate isometric deficits and those deficits were magnified 24 h post-injury. Cumulative effects of the initial stretch injury with resultant cascade of chemical and metabolic factors have been proposed to exacerbate mechanical deficits 24 h post-injury [11,13,18,34,35,40]. This study selected 24 h for sacrifice in order to capture the peak time point for mechanical damage; however, a time

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**Table 1 Time course for isometric torque following single stretch injury**

|        | Pre-injury | Immediately post-injury | 24h Post-injury |
|--------|------------|-------------------------|-----------------|
| Sham   | 7.22 ± 0.05| 7.7 ± 0.72              | 7.52 ± 0.55     |
| 0mm    | 6.95 ± 2.18| 4.81 ± 1.34             | 3.94 ± 2.08     |
| 0.7mm  | 6.95 ± 1.92| 2.49 ± 0.93             | 2.21 ± 0.46     |
| 1.4mm  | 7.51 ± 0.79| 3.12 ± 1.01             | 1.65 ± 0.54     |

Values are presented in mN·m as mean ± SD. Isometric torque was measured pre-injury, immediately and 24 h post-injury and analyzed using Dunnett’s multiple comparison test. *P < 0.01, **P < 0.001 versus pre-injury torque.
course to elucidate the duration of mechanical deficits would be an interesting contribution.

Persistent force deficits have been documented up to three days following a single stretch injury without superimposed AT shortening by modulating range of motion from 10° to 25° of dorsiflexion in rat gastrocnemius muscle [31]. One possibility for this disparity in damage between species and models may be the degree of fiber strain during stretch. In a recent elegant in vivo study by Butterfield and Herzog [12], excursion of the muscle tendon unit (MTU) was held constant while the timing of activation and starting length were altered in rabbit TA muscle for 50 lengthening contractions. Maximal fiber strain was measured directly and was found to increase with preactivation and starting muscle length. The magnitude of fiber strain was shown to be more important than muscle force or MTU dynamics in contributing to injury. While the precise relationship between fiber strain and MTU strain has not been established, increased compliance of the contractile element has been shown [44] discrediting the direct one to one relationship assumed in in vitro and in situ injury models. MTU lengthening occurs paradoxically with fiber shortening during ambulation in cat soleus muscle, indicating that MTU strains exceed muscle fiber strains [45]. By shortening the AT prior to lengthening contraction, the ability of the MTU to minimize fiber strain might have been attenuated, thus creating muscle injury.

The viscoelastic nature of muscle must be considered in minimizing the variability of mechanical testing [46]. Passive properties of muscle have been shown to contribute nearly half of total force of plantar flexion in mice [9]. Our data suggests that the passive component contributed to more than half of peak torque during active lengthening.
In permeabilized rat soleus fibers, passive tension reached 80% of maximum isometric force during a first passive stretch, and this decreased to approximately 50% with preconditioning [47]. Our data agrees with passive tension contributing to the majority of peak isokinetic torque, but this did not decline following one passive stretch to precondition the muscle.

Passive torque was substantially decreased immediately following injury in groups with AT shortening but was not affected by the level of shortening. Elastic myofilaments contribute to passive stability, with titin serving as a spring activated by calcium binding to increase stiffness during contraction [48]. These findings suggests that passive torque generated by parallel elastic structures such as titin or the extracellular matrix were not incrementally damaged in accordance with AT shortening [49].

Histological verification of damage has been touted as the “gold standard” as it allows visualization of morphological changes that can be appreciated qualitatively [50]. Quantification of histological damage is more difficult, but has been shown to be reliable when there is an established set of criteria to quantify muscle damage [51], such as inflammatory cell infiltration on H&E sections [15]. Disruption of fibers and accumulation of nuclei within the interstitial space or myofibers was not appreciably increased in the 0 mm group over the sham control. AT shortening was prerequisite to elicit damage visualized by morphological changes 24 h following injury, illustrating a similar trend with mechanical deficits. It should be noted that although morphological changes have been correlated with loss of contractile force [19], a causal relationship has not been established for either magnitude or time course [39]. Although persistent isometric torque deficit and morphological damage were only evident upon AT shortening, we were unable to precisely titrate the magnitude of damage with stepped increases in AT shortening.

Developing a murine in vivo model for acute muscle strain that results in damage near the MTJ seen clinically has proven challenging, possibly due to active and passive sarcomere length heterogeneity [47,52]. To our knowledge, this is the first minimally invasive murine model able to create damage near the MTJ with a single stretch of a biarticular muscle while retaining the intact MTU. Pioneering research incorporating single stretch injury models in situ and in vitro have elucidated mechanical factors contributing to damage [9,26-28]. Less invasive injury models that did not require tendon detachment were created for rabbit TA [29] and subsequently adapted for rat TA [30] and gastrocnemius [31]. This model provides an opportunity to utilize genetically engineered mice to expand the exploration of factors involved in tissue healing.

There are several limitations inherent to our study. While shortening of the AT is minimally invasive and an improvement from in situ models whereby the MTJ is detached, it is still a non-physiologic model. Furthermore, we were unable to titrate the level of damage by stepped increases in AT shortening. The contribution of this study is to provide a model for single stretch injury, not to elucidate the mechanical events creating damage. We speculated that shortening of the AT dampened the ability of the MTU to minimize fiber strain, but strain was not measured. In future work, the method of electrical stimulation should be optimized. Isometric tetanic force has been shown to be two times lower with tibial nerve stimulation compared to skin stimulation [31]. A needle electrode does not permit isolation of the target gastrocnemius muscle, but results in the contraction of the triceps surae and deep flexors. Subsequently, isometric torque measurements reflect the contribution of the entire posterior compartment musculature. Finally, while morphological damage confirms injury [13,15], quantitative measures are difficult. Cellular response to injury was non-specific as we only evaluated nuclei using H&E stain. Specific cell type identification using immunohistochemistry or assays to quantify inflammatory genes or proteins would have provided more sensitive information about the inflammatory response.

Conclusions
We have created a minimally invasive murine model for single stretch injury that results in damage localized near the MTJ of a biarticular muscle. Shortening of the Achilles tendon was prerequisite for incurring damage as identified by immediate absolute and relative isometric torque deficits. Morphological damage was noted 24 h post-injury in the AT shortened groups, along with a further decline in relative isometric torque. This model will provide an opportunity to utilize transgenic mice to further elucidate the role of various inflammatory mediators involved in fibrosis and regeneration.

Additional file

Additional file 1: Figure S1. An angle versus time plot illustrating the footplate traveling through 75° of dorsiflexion.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
SB designed the study, performed the injury testing, mechanical data collection, data analysis and manuscript preparation. RM designed the isokinetic device and software. AP performed the histology and helped with software and optimization of the testing. RV provided use of his laboratory. All the authors read and approved the final manuscript.

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