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Author(s): Finni Juutinen, Taija; Bernabei, Michel; Baan, Guus C.; Noort, Wendy; Tijs, Chris; Maas, Huub

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Nonuniform displacement and strain between the soleus and gastrocnemius subtendons of rat Achilles tendon

Running head: Nonuniformities within Achilles tendon

1Taija Finni, 2Michel Bernabei, 2Guus C. Baan, 2Wendy Noort, 3Chris Tijs, and 2Huub Maas

1Neuromuscular Research Center, Faculty of Sport and Health Sciences, University of Jyväskylä, Finland

2Department of Human Movement Sciences, Faculty of Behavioural and Movement Sciences, Vrije Universiteit Amsterdam, Amsterdam Movement Sciences, The Netherlands

3Department of Organismic and Evolutionary Biology, Harvard University, Concord Field Station, Bedford, MA, USA.

Corresponding author

Taija Finni, Ph.D.

Neuromuscular Research Center

Faculty of Sport and Health Sciences

University of Jyväskylä

PO Box 35 (VIV227)

FI-40014 University of Jyväskylä

Finland

email: taija.finni@jyu.fi
Abstract

Achilles tendon (AT) comprises of three subtendons arising from the soleus (SOL) and the lateral (LG) and medial (MG) heads of the gastrocnemius muscle. While recent human studies show differential displacement within AT, these displacement have not been attributed to specific subtendons. We tested the hypothesis that the SOL and LG subtendons show differential displacement and strain during various combinations of SOL, LG and MG excitations. Movement of knots, sutured onto SOL and LG subtendons of 12 Wistar rats, was videotaped while the muscles were stimulated intramuscularly and ankle torque was assessed. When SOL only was stimulated, the plantar flexion torque was the smallest among the different conditions (P<0.001). In this condition, from passive to active state, the displacement (0.57 vs. 0.47 mm, P=0.002) and strain (8.4 vs. 2.4%, P<0.001) in the SOL subtendon was greater than in LG subtendon. When LG only was stimulated, a higher ankle torque was measured as compared to SOL stimulation (P<0.001); the displacement was similar in both subtendons (~0.6mm) while the strain was greater in LG than in SOL (4.7 vs. 1.7%, P<0.001). When all three muscles were stimulated simultaneously, ankle torque was highest and the displacement (0.79 vs. 0.74 mm, P=0.002) and strain (7.7 vs. 4.4%, P=0.003) were greater in SOL than in LG. These data show that the different subtendons of AT can experience relative displacement and differential strains. Together with anatomical dissections, the results revealed that such uniformities may be due to a lower stiffness of SOL subtendon compared to LG.

Keywords: anatomy, deformation, three dimensional model, ankle torque, Achilles tendon, triceps surae
Introduction

The Achilles tendon (AT) bears high loads during everyday movements yet AT is vulnerable to disorders. AT is typically considered as one entity but it consists of three distinct subtendons contiguous with soleus (SOL), and lateral (LG) and medial gastrocnemius muscles (MG), which may not always be activated to the same extent. Consequently, one of the etiological factors behind Achilles tendon injuries may be non-uniform stress brought by heterogeneous loading within the tendon. While direct measures of stresses within the tendon are invasive, information on non-uniform loading can be achieved based on measures of tendon strain. Strain is typically calculated from tissue displacements assessed by kinematic methods.

Recently, displacements within human tendon have been measured using non-invasive ultrasound-based methods using e.g. commercial speckle tracking and radiofrequency data with 2D cross-correlation. It was found that the anterior (deep) portion of AT experienced greater displacement during passive ankle movement and eccentric loading of the calf muscles. Furthermore, the anterior portion of AT experienced greater displacement but less elongation than the posterior (superficial) tendon during human walking.

These observations of non-uniform displacements within AT have been attributed to the hierarchical structure of tendon that allows relative sliding of the tendon fascicles and to the unique structure of the AT that, in contrast to the most frequent view, contains distinguishable subtendons of SOL, LG and MG muscles. Differences in moment arm lengths, muscle slack lengths, and activation levels between these muscles as well as orientation of calcaneus may contribute to nonuniform AT behavior.

Previous human studies have also shown relative sliding between SOL and LG muscles, suggesting that AT subtendons are capable - to a certain degree – of moving independently. These studies have used selective activation of muscles within the triceps surae group; thus suggesting that the nonuniform behavior of AT subtendons could be associated with individual muscle activation. However, assessment of the deformation of individual AT subtendons is not yet possible in human studies, due to imaging resolution and selectivity of muscle activation.

Therefore, the purpose of this study was to investigate whether displacement and strain differ between SOL and LG subtendons during various activation conditions in the triceps surae.
muscles of the rat. It was hypothesized that the subtendons will undergo differential
displacement and strain and that these nonuniformities would be most pronounced when SOL
or LG muscles are stimulated individually. We also performed an anatomical study to
evaluate if AT anatomy in rat is similar to that of human and to reveal potential factors
affecting relative displacement and strain.

Methods

Animals

Experiments were performed on 12 adult male Wistar rats (235.5±16.5 g). Surgical and
experimental procedures were in strict agreement with the guidelines and regulations
concerning animal welfare and experimentation set forth by the Dutch law and were
approved by the institutional Committee on Ethics of Animal Experimentation at the Vrije
Universiteit Amsterdam. According to standard procedures ²⁰, the rats were deeply
anesthetized by intraperitoneal injection of urethane solution (1.2 ml/100g body mass, 12.5%
urethane solution), as judged by complete absence of withdrawal reflexes. If withdrawal
reflexes could be elicited, supplemental doses (0.3–0.5 ml/time) were administered. To
prevent hypothermia, animals were placed on an electrical heating pad to keep their core
temperature at approximately 37°C. Rats were continuously monitored for body temperature
and respiration. After completion of the experiments, the animals were euthanized with an
overdose of pentobarbital sodium (Euthasol 20%) injected intracardially, and double-sided
pneumothrax. The left (experimental) hindlimb was stored in -40°C for later anatomical
study.

Surgery and preparation

The left hindlimb was shaved and the skin covering the lower leg was removed. The femur
was exposed, biceps femoris and gracilis muscles were removed. Anatomical landmarks for
axis of rotation of ankle and knee joints were identified and marked with a permanent marker.

The AT was freed from all connective and fat tissues leaving the insertion to calcaneal bone
intact. For kinematic analysis, four knotted suture markers (Mariderm Schwarz, thread
thickness 7/0 USP, Catgut GmbH, Germany) were sutured into each of the LG and SOL
subtendons (Fig. 1). All exposed tissues were regularly irrigated with saline to prevent
dehydration.
Following procedures by Tijs et al.\textsuperscript{21}, the foot was attached to a 6 degrees-of-freedom load cell (Mini40-E, ATI, Apex, NC, USA) and the femur was rigidly secured. During all measurements, the ankle was set at an angle of 120° and the knee at an angle of 90°.

Wire electrodes for intramuscular stimulation were placed near the neuromuscular junctions of SOL, LG MG muscles. The location of the neuromuscular junctions was tested by stimulating the muscle superficially with a set current and identifying a location for greatest response. Then the wires were inserted around this location intramuscularly in bipolar configuration. Stimulation intensity was defined as the intensity resulting in the highest twitch torque.

**Experimental procedures**

A trial consisted of two conditioning twitches with 1 s in between followed by a 1 s tetanus (stimulation frequency 100 Hz, pulse width 100μs, Digitimer DS3, Digitimer Ltd., Hertfordshire, England). These trials were performed for five conditions: stimulation of SOL only (SOLs), LG only (LGs), MG only (MGs), LG and MG simultaneously (LGs+MGs), as well as SOL and LG and MG simultaneously (SOLs+LGs+MGs). Stimulation order was randomized between animals. Two minutes rest was given between trials. To monitor changes in force producing capacity, control trials with SOLs+LGs+MGs and ankle and knee angles at 90° were performed before and after the experiment. The second control trial was not significantly different from the first but showed individual variation (-0.3 %, SD 9.7).

**Recordings and data analysis**

Torque measured by the load cell was sampled at 1000 Hz and stored. Three-dimensional gravity-corrected ankle torques were calculated following previously described procedures\textsuperscript{21}. Passive torque was obtained from a 50 ms time window in relaxed condition (after the second twitch). This was subtracted from the torque obtained from a 50 ms window during tetanus to obtain the active torque used in the statistical analysis.

All trials were videotaped at 50 frames/s (HC-V720, Panasonic, UK) for two-dimensional kinematic analysis of the tendon markers (see Fig. 1). A ruler was placed below the tendon from which 5 mm length was digitized and used for calibration. The camera was placed orthogonal to the sagittal plane with focus on the Achilles tendon which yielded an average resolution of 0.023 mm/pixel (range 0.022-0.024).
An image captured in relaxed condition (after the second twitch corresponding to the passive
torque) and an image during tetanus (corresponding to the time of active torque) were
extracted for further analysis of tendon marker displacements. From these two images, the
tendon markers were digitized manually by selecting the centroid of the knot (Fig. 1), and
[x,y] coordinates were extracted. The image analysis was done using custom written software
in MATLAB (R2014b, Mathworks, Natick, MA, USA).

From the [x,y] coordinates data, displacement (D) of each of the markers (n) from passive to
active state (Δx and Δy) was calculated. Then the average displacement (D) was calculated as
the mean of the four SOL and four LG marker displacements, respectively.

\[
D_n = \sqrt{\Delta x_n^2 + \Delta y_n^2} \quad n = 1,2,3,4
\]

\[
\bar{D} = \frac{D_1 + D_2 + D_3 + D_4}{4}
\]

Strains (ε) based on a linear 3-segment model connecting the four markers in SOL and LG
subtendons, respectively, were calculated as \( \varepsilon = (L-L_0)/L_0 \), where L was the 3-segment length
in an active state and \( L_0 \) the 3-segment length in the passive state.

LG muscle-tendon unit (MTU) lengths were estimated using an ankle-knee geometric model
by Ettema \(^{22}\), while SOL MTU lengths were estimated by adjusting the model for ankle joint
angle displacement only \(^{23}\). The following anatomical parameters, required in the model,
were measured from the animals of the present study: ankle lever arm length (mean of 6.8
mm used in the model) and tibia length (38.3 mm).

**Anatomical study**

Using a precision caliper, SOL and LG tendon lengths were calculated as the distance
between their myotendinous junction and calcaneus while keeping the ankle and knee joint
angles at 90°. The distance of the most proximal suture from the myotendinous junction was
also measured for complete anatomical information. For all animals, dissection of AT
subtendons confirmed that the markers were sutured into either SOL or LG subtendons (Fig.
2B).

In two animals, an exploratory anatomical study was performed immediately after the
stimulation protocol, but before euthanizing the animal. After removing plantaris MTU (Fig.
2A), SOL, MG and LG muscle bellies were dissected from proximal to distal until the
tendon. To separate SOL, MG and LG subtendons careful blunt dissection along the orientation of tendon fascicles were done while minimizing tissue damage. In the most distal part, the separation of SOL, LG and MG tendons continued by removal of tissue around the calcaneus and the tendon until the insertion sites were fully exposed (Fig. 2C).

For three dimensional AT reconstruction, the tendon of one animal was prepared in the following manner: AT was first fixed in buffered 4% paraformaldehyde, preserved in buffered 20% sucrose solution and then taken out of the solution and dried before staining the three subtendon surfaces with different colors (Staedler Permanent Dry Safe, Lumocolor (AP), Germany). Then, AT was embedded in a 4% gelatin solution and frozen before cutting into 1 mm cross-sections. These cross-sections were photographed and CSAs were measured using ImageJ (imagej.nih.gov) by manually segmenting five slices along each of the subtendons. Cross-sectional images were uploaded to Reconstruct (synapseweb.clm.utexas.edu) to create a 3D reconstruction of the AT. Finally, the reconstructed tendon was imported in Anim8or (anim8or.com) and modified to include the insertion onto calcaneus (Fig.3D).

**Statistics**

After confirming normality with Saphiro-Wilk test, by comparing mean and median values and by checking values of skewness and kurtosis, parametric statistics were used. The effects of stimulation condition on ankle torque were tested using repeated measures ANOVA while two factor ANOVA (stimulation*subtendon) was used to examine differential displacements and strains between the SOL and LG subtendons. Greenhouse-Geisser correction was used when assumption for sphericity was not met and Bonferroni was used as post hoc for pairwise comparisons. Level of significance was set at $P \leq 0.05$.

**Results**

*Tendon anatomy*

Mean tendon length was 7.62 mm (SD 0.33) for SOL and 11.22 mm (SD 0.34) for LG. In SOL, the most proximal marker was located 0.26 mm (SD 0.55) distally from the myotendinous junction and for LG this distance was 3.89 mm (SD 0.55). Mean resting lengths used in strain calculations (i.e. between proximal and distal markers) were 1.56 mm (SD 0.25) for SOL and 1.68 mm (SD 0.35) for LG corresponding to about 20% (SOL) and 15% (LG) of the total tendon lengths. The calculated MTU lengths of SOL and LG were
29.36 mm and 37.46 mm, respectively. LG tendon was 27% and SOL tendon 23% of the estimated MTU length.

The attachment site of the tendons on the calcaneus is shown in figure 2D. We were not able to identify separate SOL and LG entheses or insertion sites onto the calcaneus (Fig. 2C). The attachment area of SOL+LG was slightly smaller than that for MG, the latter being located laterally to the attachment site of SOL+LG. The model of AT illustrates that the MG travels from medial to postero-lateral side while LG travels medially when traveling distally (Fig. 3). Mean cross-sectional area of the subtendon was 0.14 mm$^2$ for SOL, 0.34 mm$^2$ for LG and 0.48 mm$^2$ for MG.

**Ankle joint torque**
Stimulation condition had a main effect on ankle torque (p<0.001) with all other conditions differing from each other except LGs and MGs. Ankle torque was smallest when SOL was stimulated and highest when all three muscles were stimulated (Fig. 4A). LG ankle torque (16.9±5.6 mNm) was over four times higher and MG torque (12.8±2.9 mNm) over three times higher than SOL torque (3.9±2.4 mNm).

**Displacement of subtendons**
Both subtendons showed displacements even if their respective muscle was not stimulated. Displacements of SOL and LG subtendons differed for some, but not all, stimulation conditions (Fig. 4B). Main effects of stimulation condition (p=0.001) and subtendon (p=0.003) on tendon displacements were found as well as a significant stimulation*subtendon interaction effect (p=0.002). Post hoc pairwise comparisons revealed differences between subtendons (Fig 4B) and between the following stimulation conditions: LGs and SOLs+LGs+MGs (on average 24% greater displacement with SOLs+LGs+MGs, p=0.016), MGs and LGs+MGs (43%, p=0.003) and MGs and SOLs+LGs+MGs (52%, p=0.002).

**Subtendon strain**
There was a main effect of stimulation (p<0.001) and subtendon (p=0.017) on SOL and LG subtendon strain with significant interaction (p<0.001). Post hoc pairwise comparisons showed that SOL and LG subtendon strains were different in each stimulation condition (Fig. 4C). Overall, subtendon strains in SOLs condition differed from LGs (p=0.022), MGs
(p<0.001) and LGs+MGs conditions (p<0.001). Strain also differed between LGs and MGs (p<0.001), LGs and LGs+MGs (p=0.021), MGs and LGs+MGs (p<0.000), MGs and SOLs+LGs+MGs (p<0.001) and LGs+MGs and SOLs+LGs+MGs (p=0.003) conditions. Note that in both conditions when SOL was stimulated, SOL tendon was stretched to a greater extent than LG tendon.

Discussion

Typically, the AT is considered as one entity but it consists of three distinct subtendons contiguous with SOL, LG and MG muscles. This study investigated to what extent SOL and LG subtendons displacement and strain differ during various activation conditions of the triceps surae muscles. We found differential displacements for the SOL and LG subtendons in some, but not all, conditions and different strains when the triceps surae muscles were activated individually or simultaneously. This is in agreement with human imaging studies reporting nonuniform displacements within AT (e.g. 9,11-12) and confirms the earlier notion that a single measure of strain may not provide an accurate representation of the AT behavior as a whole 24. In spite of the observed nonuniform strain, our findings suggest that the distinct subtendons are not completely independent because activation of any combination of the muscles produced displacements of similar order of magnitude in both SOL and LG subtendons (Fig. 4B).

We hypothesized that the differences in SOL and LG subtendon behavior would be the greatest when their respective muscles were stimulated individually. This was true when only SOL was stimulated; both the displacement and strain in the SOL subtendon were significantly higher than in the LG subtendon. However, when only LG was stimulated the displacement was similar in both subtendons while the strain was greater in LG. This result may indicate that activation of a single muscle results in a pulling force on the neighboring subtendon without stretching it much. The observed asymmetries in deformations of SOL and LG subtendons may come from the differences in anatomy and properties of the MTU and/or asymmetries in the connective tissue linkages between these muscles.

As shown in figure 4, when SOL was activated individually it produced the smallest ankle torque but strained the SOL subtendon by ~8% while activation of SOL together with LG and MG produced much greater ankle torque but a similar strain in SOL subtendon. The LG
subtendon, on the other hand, strained ~4% in the conditions when its muscle was stimulated individually or simultaneously with other muscles. When all muscles were stimulated simultaneously, SOL subtendon strain was nearly twofold compared to LG strain. The results show convincingly that tendons lengthen and store elastic energy even in small animals.

Using the torque and strain values of figure 4, we may conclude that the SOL subtendon has a lower stiffness than that of LG. However, considering that the LG tendon had a larger CSA, their Young’s modulus can still be quite similar. Previously, considerable variation in tendon material properties has been reported in various tendons of lower limb of turkeys but with rather similar maximum strain of ~10%.

Considering that in the current, supposedly submaximal activation condition, the SOL subtendon strained up to ~8%, the observation raises question about the susceptibility of SOL subtendon to injuries in tasks requiring maximal effort.

Based on the previously reported nonuniform AT displacements from human experiments, the following question has been asked: why would the free AT, representing the tendon length from the distal end of SOL muscle belly to the calcaneus (i.e. SOL subtendon), show a greater elongation than the longer tendon of the gastrocnemius muscle? The present results suggest that the tendon properties may vary within the AT. While anatomical and functional differences between human and rat triceps surae MTU must be considered, in both species the SOL subtendon is located anteriorly (Fig. 3). In agreement with our results in the rat AT, the anterior part of the human AT has been shown to elongate and strain more than posterior tendon where the gastrocnemius subtendons are located.

While these human studies were not able to specify if the anterior part could be attributed to SOL subtendon, several ultrasound studies have accomplished to separate SOL and MG tendons by assessing displacement of their respective muscle-tendon junctions. During isometric contraction these studies showed strains of 5.2% in SOL vs. 2.6% in MG, 6.6% in SOL vs. 2.6% in MG and 3.8% in SOL vs. 2.7% in MG, which are slightly lower than those in the present rat study. Despite different species, activation type (voluntary vs. intramuscular stimulation) and level of contraction, data from the present and previous studies lead to the conclusion that SOL strains more than LG during isometric contraction.
However, a different behaviour was observed in the intact compartment and with a physiological pattern of activation. During the stance phase of walking in rats, length change up to about 3.1 mm in LG and 0.9 mm in SOL distal tendons have been reported. This corroborates with finding of Franz et al. that the posterior AT had greater elongation than anterior AT during walking. This discrepancy between isometric and dynamic conditions might be explained by the fact that during walking also knee joint angle changes and the deformations of different subtendons may thus differ from purely isometric conditions.

Comparing strain between the subtendons eliminates the differences in absolute length from consideration but observations of their relative movements may also be important. Large relative movements in adjacent tendon fascicles or subtendons may be expected to damage the intrafascicular matrix and smaller relative movements can also lead to injury after repetitive loading e.g.. In the present study the displacement in SOL was slightly but significantly greater than in LG in three out of five stimulation conditions (Fig. 4B). The magnitude of relative movement was in the order of tenth of a millimeter and it may be considered that the displacements were similar between the subtendons. These similarities in displacements may indicate that at least part of the AT acts as a common spring for the muscle compartments of the triceps surae. Similar displacements, even if only one muscle was stimulated, could also be caused by connective tissue linkages between the muscle bellies or subtendons. Future studies should explore what is the degree of dependency between the subtendons and how their behavior is related to subtendon properties, relative muscle length and activation level of the triceps surae muscles.

**Anatomy**

Similar to the human AT, the reconstructed model of the rat AT (Fig. 3) shows rotation of MG subtendon from medial origin to lateral insertion site. In rats, this anatomical organization has been suggested to cause an eversion moment at the ankle joint. In humans, however, MG has been reported to contribute either to eversion or inversion depending on the frontal plane angle of the foot. In the present study, LG attachment was located to lateral aspect of calcaneus together with SOL. In humans, considerable variations in the attachment patterns of the three tendons have been found. Unlike in human AT, the insertion site of rat MG to calcaneus was larger than that of SOL and LG, and these two tendons seemed to share the same enthesis. In rats the joint enthesis of SOL and LG may be a necessity due to the
small cross-sectional area of the SOL tendon, the arrangement providing reinforcement to the insertion site.

The hierarchical structure of tendon with interactions between collagen fibers and the matrix they are embedded in has been a challenge for modeling studies. Because AT injuries are rather common, modeling attempts are important for this specific tendon as well. It has already been shown that subject-specific geometry of total AT may explain rupture location and the next step would be to incorporate the subtendon organization and properties of interfascicular matrix to such models.

Limitations

It is important to note that we did not fully excite these muscles and, thus, ankle torque does not reflect the maximal force capacity of these muscles. While we placed the electrodes as close to the point that yielded largest twitch when stimulated from the surface of the muscle, and used the intensity that elicited maximal twitch, not all the motor units could be recruited. It has been reported that the LGs + MGs peak active ankle moment may be only about 20% when activated intramuscularly as compared to nerve stimulation. However, at least for SOL muscle we may have been closer to 50% of maximal isometric torque since the torque was about 4 mNm in the present study as compared to 8 mNm with supramaximal nerve stimulation. Overall, the torques exerted by the various muscles were in line with what is expected based on muscle sizes and assuming similar moment arms around the ankle for all muscles.

The kinematic measures were done in one plane only. The two-dimensional approach causes some uncertainties in the reported displacement and strain values due to out-of-plane movement and possible rotation of the tendon. Estimating a 10° difference between the imaging and movement planes would have caused about 1.5% error in the displacements reported in this study. When discussing the reported displacements, it needs to be acknowledged that they were tracked from a part of the tendon, not from its entire length. In SOL muscle the tracking was done as proximally as possible whereas the adjacent markers in MG lay in the mid tendon. Potential longitudinal differences in tendon properties could therefore be one reason for the observed differential behavior. While MG tendon deformation could not be measured in the present study due to limitations of the two-dimensional kinematic method used, the effects of MG stimulation on SOL and LG subtendons was
somewhat unexpected. The negative strains observed upon MG activation suggests that the most distal parts of LG and SOL subtendons were pulled proximally relative to the most proximal part, or that this result is due to errors from inability of the two-dimensional analysis to quantify tendon rotation.

**PERSPECTIVE**

Achilles tendon comprises of three distinct subtendons that can behave differently. We showed that during isometric contractions the SOL subtendon undergoes substantially greater strain than LG subtendon when their respective muscles are stimulated independently or simultaneously. We conclude that one cause for the nonuniform behavior within the rat AT may be different stiffness of the subtendons, SOL being more compliant than LG. The difference in strain between the subtendons necessitates a degree of nonuniform displacement between the subtendons. While the elastic interfascicular matrix allows relative sliding of the tendon fascicles it can also bear considerable loads. With a stiffness about half of that of tendon fascicles \(^{30,36}\), the matrix can act to distribute forces within the tendon or serve as an alternative pathway for force transmission upon damage to tendon fascicles. Consequently, tendon organization may provide an inbuilt protective mechanism against tendon injuries by shielding tendon from uneven stresses caused by differences in SOL, LG and MG activation levels in combination with differential MTU length changes between the triceps surae muscles during normal physical activities. Future studies are needed to reveal which structures, such as the tendon matrix properties \(^{36}\) or connective tissue links between the muscle bellies \(^{37}\), bring about the mechanical dependency of the subtendons.

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References


Figures & figure legends

Figures

Figure 1. Experimental setup showing the left hindlimb attached to a load cell with stimulating wires going into the muscles. Knot sutures in lateral gastrocnemius (top row of knots) and soleus subtendons (bottom row of knots) are seen as black dots. The scale has 1 mm between lines. The inset shows that during digitizing the middle of the knot was selected.
Figure 2. A) Achilles tendon has been flipped over calcaneus to reveal dorsal view of the tendon. Medial gastrocnemius tendon (MG) has been separated from lateral gastrocnemius (LG) and soleus (SOL) subtendons. Four sutures along the length of LG and SOL subtendons that were used in the analysis are identified. An additional marker was sutured more distally to the LG but it was not used in this study. B) Dorsal view of separated LG and SOL subtendons to confirm that the sutures were sown into the correct subtendons. C) A close-up from the tendon insertion to calcaneus via enthesis (E). While MG had a distinguishably its own enthesis the LG and SOL tendons appeared to share the same enthesis and insertion site. This is also visible in D) where the calcaneus has been fully exposed and tendons detached. The insertion site for MG tendon, marked in red, is larger than the common insertion site for SOL and LG tendons. The scale in A and C has 1 mm between lines.
Figure 3. Three dimensional reconstruction of the rat Achilles tendon (AT). Similarly to human AT, the model illustrates medio-lateral twist of the rat medial gastrocnemius (MG) subtendon (A,B,C). Subtendon of lateral gastrocnemius (LG) travels medially emerging underneath the MG to its lateral attachment site in the calcaneus (D). Contrary to humans, rat soleus (SOL) muscle and its respective subtendon comprise a small cross-sectional area relative to MG and LG (B).
Figure 4. Mean (SD) of ankle joint torque (A), subtendon displacement (B) and strain (C) in the five different stimulation conditions; only soleus: SOLs, only lateral gastrocnemius: LGs, only medial gastrocnemius: MG, LG and MG simultaneously: LGs+MGs and all three muscles together: SOLs+LGs+MGs. Ankle torque differed significantly (p≤0.001) between all conditions except between LGs and MGs.

Differences between subtendons; * p<0.05, ** p<0.01, ***p<0.001.
A

Ankle torque (mNm)

B

Displacement (mm)

C

Strain (%)

Stimulation condition

SOLs  LGs  MGs  LGs+MGs  SOLs+LGs+MGs