Validity and reliability of an ultrasound measurement of the free length of the Achilles tendon

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ABSTRACT

INTRODUCTION: Valid length measurements of the different segments of the Achilles tendon are needed in order to investigate if differential elongation of the Achilles tendon takes place after rupture. The purpose of this paper was to present data concerning the accuracy and reliability of an ultrasound measurement of the free part of the Achilles tendon.

METHODS: Both legs of 19 non-injured subjects were examined by magnetic resonance imagining (MRI) and ultrasound. The length from the distal tip of the soleus muscle to the tendon insertion on the calcaneus was measured by three independent ultrasound examiners. Repeated ultrasound measurements were performed and compared with MRI measurements. Intra-rater and inter-rater reliability and the agreement between MRI and ultrasound were determined. Data were evaluated using the intraclass correlation coefficient (ICC), the standard error of the measurement (SEM) and the minimal detectable change (MDC).

RESULTS: The measurement showed excellent intra-rater reliability (ICC = 0.94 (95% confidence interval (CI): 0.91-0.96), SEM = 5 mm and MDC = 13 mm) and inter-rater reliability (ICC = 0.96 (95% CI: 0.93-0.97), SEM = 4 mm and MDC = 11 mm). On average, ultrasound measurements exceeded the MRI measurements by 2 mm (non-significant); resulting in a measurement error of 5%.

CONCLUSIONS: The ultrasound measurement of the free part of the Achilles tendon showed good reliability and accuracy. For comparison between groups of non-injured subjects, differences of > 5 mm can be detected. For repeated assessment of individual subject differences ≥ 13 mm can be detected.

FUNDING: none.

TRIAL REGISTRATION: Institutional Review Board of Zealand, Denmark, Ref. no: SJ-318.

Validity of the ultrasound measurement of the Achilles tendon was assessed in a study on non-injured subjects. The measurement showed excellent reliability and agreement with MRI measurements. The ultrasound method was considered applicable and easy to perform in the acute phase of rupture.

Achilles tendon rupture is a devastating injury causing many patients to suffer from chronically impaired locomotor function, and only 30-66% resume previous sports activities [1, 2]. The debilitating effect of such injury is suggested to be associated with elongation of the Achilles tendon, leading to ineffective working length [3, 4]. This leads to reduced plantar flexion force of the ankle, thereby impairing a powerful lift-off [5].

The Achilles tendon provides linkage between the calcaneus and the three muscle bellies of the triceps surae: the soleus, the medial and the lateral head of gastrocnemius [6, 7]. It is often considered one tendon moving “en bloc” [5, 7], but studies suggest that the separate collagen fibre bundles can move independently of each other, allowing for differential displacement of the three parts of the tendon during normal muscle contraction [6, 8, 9]. If this is the case, the three muscle bellies might retract differently after rupture due to differentiated elongation of the three parts of the tendon. The medial head of the gastrocnemius muscle shows the greatest amount of contraction during normal locomotion [8]. It might therefore be expected that this part of the tendon elongates the most after rupture. A differentiated elongation of the Achilles tendon might result in altered walking and running patterns due to alterations in force transmission.

Valid length measurements of the different parts of the tendon are needed to investigate if rupture causes differential elongation of the Achilles tendon. An ultrasound (US)-based measurement of the length from the calcaneus to the medial head of the gastrocnemius muscle has previously been investigated at our clinic (Figure 1) [10] and showed excellent validity and reliability. This study also included measurements of the length from the calcaneus to the distal tip of the soleus muscle (Figure 1), but those data were not included in the paper as the Achilles tendon was thought to be moving “en bloc”. The aim of the paper was to develop a measurement that could be used in the acute phase of rupture, and the distal tip of the soleus was thought to be an unreliable anatomical landmark in the acute phase of rupture due to the proximity of the rupture haematoma.

The purpose of the present paper is to present data concerning the accuracy and reliability of a clinically applicable and easy to perform US measurement of the free part of the Achilles tendon (Figure 1).

METHODS

The study was performed in the same setting as previously described for the development and validation of an US measure from the calcaneus to the most distal tip of the medial gastrocnemius head [10]. Measurements were performed on the same subjects by the same investigators. The study population was first scanned by
three independent US investigators; this was followed by MRI scans that were used as the gold standard to test concurrent validity.

**Description of the ultrasound measurement**

The length of the free part of the Achilles tendon was defined as the distance between the most proximal point of the posterior border of the calcaneus and the distal tip of the soleus muscle-tendon junction. The measurement was carried out in two steps; first the anatomical landmarks were identified and marked on the skin, and then the distance between them was measured on the skin with a tape measure [10]:

1) Positioning: Participants were positioned in a prone position with the knee flexed 10 °. The feet rested on a triangular-shaped foam pad as shown in Figure 2A. Using a goniometer, the ankle joint was positioned in 10 ° of plantar flexion by adjustment of the foam pad. B. With a marker, the point where the ultrasound probe and the needle crossed was marked on the skin. C. The distance between landmarks was measured with a tape measure following the curves of the leg.

**Study population**

The study was performed on the same nineteen non-injured subjects who participated in the previous study [10]. The subjects were aged 26-63 years and had no prior Achilles tendon problems. Investigations were performed in the period November 2012 to January 2013. All participants received oral and written information and their written consent was obtained. The study was approved by the Institutional Review Board of Zealand, Denmark, Ref. no: SJ-318.
Evaluation protocol
The novel US measurement was evaluated by three independent US investigators with 2-5 years of experience within musculoskeletal US. Three different US scanners were used: GE Healthcare Logiq S8, Logiq 9 and logiq P5. The frequency was set to 15 MHz and the focus was dynamically adjusted by the US operator. After each scan, the marks on the skin were removed in order to secure blinding of results between investigators.

Inter-rater and intra-rater reliability and agreement for the novel US measurement were determined for the three independent US investigators. Repeated measurements were performed at a three-week interval between scans. Subsequently, MRI examinations were performed within two months. MR images were evaluated by two independent investigators, both specialised in musculoskeletal MRI. All investigators were blinded to the results of the other investigators and to their own previous results. Participants were positioned identically for US and MRI investigations using the same triangle-shaped foam pad. The position of the ankle and knee joint was controlled using a goniometer.

Magnetic resonance imaging setup
MR imaging was performed on a 1.5T system (Magnetom Avanto, Siemens, Erlangen, Germany). Axial 2D sequences covering both entire lower legs were performed as follows: T1-weighted without fat saturation (TR/TE, 485/12 ms; flip angle, 90°; pixel matrix, 384 × 256; voxel size, 1.2 × 0.9 × 6.0 mm; slice thickness, 6.0 mm; distance factor, 30%; field of view (FOV), 350 × 312 mm; slices, 58) and T2-weighted without fat saturation (TR/TE, 4,090/78 ms; flip angle, 150°; pixel matrix, 448 × 355; voxel size, 1.0 × 0.8 × 6.0 mm; slice thickness, 6.0 mm; distance factor, 20%; FOV, 370 × 367 mm; slices, 60). Sagittal 2D sequences oriented in a plane perpendicular to the axis of the talocrural joint covering each entire lower leg separately were performed as follows: T1-weighted without fat saturation (TR/TE, 555/22 ms; flip angle, 180°; pixel matrix, 448 × 238; voxel size, 1.2 × 0.9 × 3.0 mm; slice thickness, 3.0 mm; distance factor, 10%; FOV, 410 × 291 mm; slices, 37) and T2-weighted without fat saturation (TR/TE, 4740/72 ms; flip angle, 150°; pixel matrix, 512 × 512; voxel size, 1.7 × 1.7 × 4.0 mm; slice thickness, 4.0 mm; distance factor, 20%; FOV, 440 × 440 mm; slices, 26).

T1-weighted slices were used for all measurements. Length measures were performed on the slice that showed the distal end of the soleus muscle, covering the distance from the distal tip of the soleus muscle to the distal end of the Achilles tendon, which was defined by the axial plane intersecting the most cranial aspect of the tuber calcanei.

### Statistical methods
In this exploratory study we a priori decided to obtain a sample size of 19 test persons using both legs. Standard procedures were used for descriptive statistics. Reliability was determined using the intercorrelation coefficient (ICC). ICC2.2 Two-way random, absolute agreement was used for evaluation of intra-rater reliability, whereas ICC2.3 Two-way random, absolute agreement was used for inter-rater reliability. Differences of the mean and standard error of the measurement (SEM) = standard deviation of the crude measurements × √ (1–ICC) were calculated to assess agreement between groups of data [11]. The minimal detectable change (MDC) was calculated to assess agreement between data on the individual patients (MDC = 1.96 × √2 × SEM) [11]. A paired t-test was used for comparison of means as the data showed a normal distribution and the scale was considered to be continuous. Correlation between US and MRI was assessed using Pearson’s correlation. Statistical analyses were performed using the Statistical Package for Social Sciences (SPSS, version 22.0 for Windows, SPSS Inc., Chicago, Ill).

**Trial registration:** Institutional Review Board of Zealand, Denmark, Ref. no: SJ-318.

### RESULTS
Both legs of 19 uninjured persons (eight males and 11 females) were studied. The mean age was 43.4 years (SD = 10.7, range: 26-63 years). The average height was 175 cm (SD = 9, range: 158-192 cm) and the average weight was 76.8 kg (SD = 12.9, range: 58-110 kg). All participants stated their right leg to be dominant.
Reliability
The US measurement of the free part of the Achilles tendon showed excellent intra-rater reliability, good agreement (ICC = 0.94 (95% confidence interval: CI): 0.91-0.96), SEM = 5 mm and MDC = 13 mm) and good inter-rater reliability (ICC = 0.96 (95% CI: 0.93-0.97), SEM = 4 mm and MDC = 11 mm) (Table 1). There was no systematic difference between test days, but a systematic difference between investigators of up to 5 mm (p < 0.01) was seen.

Validity
Validity of the US measurement was tested by comparison with the MRI measurements. The measurements showed good correlation r = 0.82 according to the recommendations of Portney & Watkins [12]. MRI measurements were, on average, 2 mm shorter than the US measurements (non-significant). The average length of the measured Achilles tendons was 42 mm; resulting in a measurement error of 2/42 = 5%.

DISCUSSION
The most important finding of the present study was that the US measurement of the free part of the Achilles tendon showed good reliability and accuracy in non-injured people. For comparison between groups of non-injured people, differences of 5 mm can be detected; and for repeated assessment of individual subjects, differences of ≥ 13 mm can be detected. In comparison with MRI, the US measurement showed an acceptable accuracy with a measurement error of 5%.

Both the inter-rater and intra-rater reliability were fully comparable with measurements using extended field of view (EFOV) US imaging [13]. The advantage of EFOV measurement is its ease and the possibility of saving the pictures for later analysis and/or control. On the other hand, the advantage of the present US measurement is its relative ease and that it is a low-cost technology, making it applicable in any US setting.

The clinical relevance of the US measurement depends on its ability to detect elongations of the tendon. In relation to the average length of the free tendon (42 mm), an elongation of 12% can be detected when comparing groups of patients, whereas the elongation must exceed 31% when assessing individual subjects. If the average elongation after rupture is 2.5-3.5 cm, as described by Silbernagel et al [4], the US measure is fully capable of measuring differences in groups of patients, whereas one should take care in using it for assessment of individual patients.

When using the US measurement, one should be familiar with the anatomical landmarks and be aware not to move the skin while measuring. Our setup was strictly standardised in order to minimise the risk of such measurements errors.

When interpreting the results, it is important to consider the morphology of the distal tip of the soleus, which might change after an Achilles tendon rupture. Therefore, the present findings might not be directly transferable to a population with ruptured Achilles tendons. Particularly, care should be taken in the months following a rupture where the authors’ experience is that the morphology of the distal tip of the soleus muscle is blurred and difficult to identify. Probably a measurement of the free part of the Achilles tendon, using the distal tip of the soleus muscle as a landmark, should not be performed until six months after injury in order for the landmark to become clear and easily identifiable. This needs to be investigated in a sample of patients with Achilles tendon rupture. This study is limited by the lack of a proper sample size calculation. The sample size was determined due to logistic matters and no power calculation was performed. Also, the standardised setup for the measurement might have put the tendon on slack, why the measurement is likely to underestimate the true length of the tendon.

CONCLUSIONS
The US measurement of the free part of the Achilles tendon showed good reliability and accuracy. For comparison between groups of non-injured subjects, differences of > 5 mm can be detected. For repeated assessment of individual subjects, differences of > 13 mm can be detected. The US measurement is a promising clinical tool to be further assessed in the setting of acute Achilles tendon rupture.

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ACCEPTED: 15 January 2018
CONFLICTS OF INTEREST: Disclosure forms provided by the authors are available with the full text of this article at www.danmedj.dk

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