Quantitative analysis of patellar tendon size and structure in asymptomatic professional players: sonographic study

Maurizio Giacchino¹ Cristina Caresio² Negar E. Gorji³ Filippo Molinari² Giuseppe Massazza⁴ Marco Alessandro Minetto⁴

¹ Medical Lab, Asti, Italy ² Biolab, Department of Electronics and Telecommunications, Politecnico di Torino, Turin, Italy ³ Division of Endocrinology, Diabetology and Metabolism, Department of Medical Sciences, University of Turin, Turin, Italy ⁴ Division of Physical Medicine and Rehabilitation, Department of Surgical Sciences, University of Turin, Turin, Italy

Corresponding Author: Marco Alessandro Minetto
Department of Surgical Sciences, University of Turin
Corso Dogliotti 14
10126 Turin, Italy
E-mail: marco.minetto@unito.it

Summary

Background: Ultrasonographic abnormalities of the patellar tendon frequently occur in asymptomatic athletes and it is not always clear whether they precede (and may predict) the development of tendinopathy.

Objective: This study aimed to investigate by ultrasonography the prevalence of patellar tendon abnormalities in players of “pallapugno” and to establish whether structural tendon abnormalities predict tendinopathy development.

Methods: Ultrasound B-mode images of the patellar tendon of both sides were acquired in fourteen throwers. Qualitative assessments of tendon structure and neovascularization and quantitative assessments of tendon thickness, cross-sectional area (CSA), and echo-intensity were performed.

Results: Qualitative assessments showed a subclinical tendinopathy of the non-dominant tendon in 5 out of 14 throwers (35% of cases), while quantitative assessments showed abnormalities of the non-dominant tendon in 8 out of 14 players (57% of cases). Echo-intensity and CSA were the quantitative variables most discriminant between asymptomatic players without structural tendon abnormalities and those with tendon abnormalities. Two players (2 out of 8 cases: 25%) developed a clinical tendinopathy after a follow-up of six months.

Conclusion: The prevalence of subclinical tendinopathy in the non-dominant patellar tendon of throwers was high. Patellar tendon abnormalities at baseline seem to increase the risk of development of subsequent patellar tendinopathy.

Level of evidence: II b (individual cohort study).

Key words: patellar tendon, tendinopathy, thrower players, quantitative ultrasonography.

Introduction

Musculoskeletal ultrasonography is an effective technique to visualize normal and pathological tendons and to non-invasively evaluate (qualitative assessment)¹,² or measure (quantitative analysis) parameters reflecting tendon size (length, thickness, and cross-sectional area) and structure (echo-intensity)³-⁶. Patellar tendon ultrasonography is commonly performed in the routine management of athletes of different disciplines to investigate tendon abnormalities that may occur due to repetitive tendon overload⁷-¹⁰. In fact, patellar tendinopathy is a common overuse disorder typically occurring in athletes who participate in sports that require jumping, including volleyball and basketball, hence the label “jumper’s knee”¹¹,¹². However, ultrasonographic tendon abnormalities (such as hypoechoic areas, increased tendon thickness, neovascularization) have been identified in large percentages of asymptomatic athletes⁶,¹⁰,¹³, therefore it is not always clear whether structural tendon abnormalities precede and predict the development of functional tendon abnormalities and tendon pain.

The aims of the present study were to: I) investigate the prevalence of patellar tendon abnormalities in elite players of “pallapugno”, a game practiced in the north-west of Italy (similar to frisian handball, Basque pelota, and French game called “ballon au poing”) and that implies in throwers a repetitive overload of the non-dominant lower limb; II) compare the diagnostic accuracy of qualitative to the accuracy of quantitative assessments of tendon size and structure in detecting structural tendon abnormalities; III) establish whether structural patellar tendon abnormalities predict tendinopathy development.
Materials and methods

Subjects
Fourteen male throwers (age, mean ± SD: 21.8 ± 4.2 years) volunteered to participate in the study. They were free from neuromuscular or skeletal impairments and were asked to refrain from performing strenuous physical activity during 24 h before the experimental session. Side dominance was assessed with the “Waterloo Handedness and Footedness Questionnaires - Revised”\textsuperscript{14}. All subjects were right side dominant. Before participating to the study, the subjects received a detailed explanation of the protocol and gave written informed consent. The study conformed to the guidelines of the Declaration of Helsinki, met the ethical standards of the journal\textsuperscript{15} and was approved by the local ethics committee.

Procedures
Ultrasound B-mode images of the patellar tendon were acquired in each subject during a single experimental session. Both sides were investigated. The same experienced sonographer (MG) performed all assessments. Twenty scans were acquired in total for each subject while he was lying in a supine position and with the quadriceps muscle relaxed: 12 scans were acquired with a knee angle of 0° (fully extended knee), while 8 scans were acquired with a knee angle of 30° (Tab. I).

In addition, ultrasound analysis and qualitative assessments of tendon structure and neovascularization (see below) were repeated 6 months after the first investigation in players who became symptomatic for patellar tendinopathy.

Ultrasound equipment
All measurements were performed using a ClearVue 550 ultrasound device (Philips Medical Systems, Milan, Italy) equipped by a linear-array transducer with variable frequency band (4-12 MHz). Gain was set at 50% of the range, dynamic image compression was turned off, and time gain compensation was maintained in the same (neutral) position for all scans. Power Doppler imaging was performed with slow perfusion settings. All system-setting parameters were kept constant throughout the study and for each subject. Pictures were stored as DICOM files and transferred to a computer for processing.

<table>
<thead>
<tr>
<th>Images</th>
<th>Acquisition</th>
<th>Analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Knee angle: 0°</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 1</td>
<td>Transversal plane, proximal probe position, non-dominant (dominant) side</td>
<td>CSA and echo-intensity quantitative analyses</td>
</tr>
<tr>
<td>(Image 7)</td>
<td></td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>Image 2</td>
<td>Transversal plane, proximal probe position, non-dominant (dominant) side (including PD registration)</td>
<td>Neovascularization qualitative analysis</td>
</tr>
<tr>
<td>(Image 8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 3</td>
<td>Longitudinal plane, proximal probe position, non-dominant (dominant) side</td>
<td>Thickness and echo-intensity quantitative analyses</td>
</tr>
<tr>
<td>(Image 9)</td>
<td></td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>Image 4</td>
<td>Longitudinal plane, proximal probe position, non-dominant (dominant) side (including PD registration)</td>
<td>Neovascularization qualitative analysis</td>
</tr>
<tr>
<td>(Image 10)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 5</td>
<td>Transversal plane, central probe position, non-dominant (dominant) side</td>
<td>CSA and echo-intensity quantitative analyses</td>
</tr>
<tr>
<td>(Image 11)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 6</td>
<td>Longitudinal plane, central probe position, non-dominant (dominant) side</td>
<td>Thickness and echo-intensity quantitative analyses</td>
</tr>
<tr>
<td>(Image 12)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Knee angle: 30°</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 13</td>
<td>Transversal plane, proximal probe position, non-dominant (dominant) side</td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>(Image 17)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 14</td>
<td>Longitudinal plane, proximal probe position, non-dominant (dominant) side</td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>(Image 18)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 15</td>
<td>Transversal plane, central probe position, non-dominant (dominant) side</td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>(Image 19)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image 16</td>
<td>Longitudinal plane, central probe position, non-dominant (dominant) side</td>
<td>Structure qualitative analysis</td>
</tr>
<tr>
<td>(Image 20)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Qualitative assessments of tendon structure and neovascularization
The same experienced sonographer (MG) who performed the image acquisitions performed also the qualitative assessments. Tendon structure was evaluated both in the longitudinal and in the transversal scans by using the four-grade scale proposed by Sunding et al.\textsuperscript{16}: 0: normal structure (homogeneous echogenicity); 1: light structural changes (discrete hypo-echogenic areas); 2: moderate structural changes (some well-defined hypo-echogenic areas); 3: severe structural changes (extended hypo-echogenic areas). Neovascularization was evaluated both in the longitudi-

![Longitudinal scan: proximal thickness](image1)

![Longitudinal scan: central thickness](image2)

![Longitudinal scan: proximal ROI](image3)

![Longitudinal scan: central ROI](image4)

![Transversal scan: proximal ROI](image5)

![Transversal scan: central ROI](image6)

Figure 1. Representative patellar tendon scans showing the targets for the thickness measurement (upper panels), longitudinal regions of interest (middle panels), and transversal regions of interest (lower panels) for the echo-intensity measurements: ROI: region of interest.
dinal and in the transversal scans by using the four-grade scale proposed by Sunding et al. 16: 0: no neovascularization; 1: mild neovascularization (a few solitary blood vessels); 2: moderate neovascularization (moderate quantity, mostly transversal blood vessels); 3: severe neovascularization (several, mostly horizontal blood vessels spread in the whole depth of the tendon).

We arbitrarily defined “subclinical tendinopathy” as the presence of either light structural changes in association with at least mild neovascularization or moderate/severe structural changes with/without neovascularization.

**Measurement of tendon thickness**
Tendon thickness was measured in the longitudinal scans acquired with 0-degree knee angle: as shown in the representative images in Figure 1 (upper panels), the target for the thickness measurement was the thickest part of the tendon both for the proximal probe position (i.e., just below the apex of the patella: left panel) and for the central probe position (i.e., midportion of the tendon: right panel).

**Measurement of tendon cross sectional area**
Tendon CSA was measured in the transversal scans acquired with 0-degree knee angle: as shown in the representative images in Figure 1 (lower panels), a region of interest was chosen in each scan to include as much of the tendon as possible. The area was determined for the selected region of interest by a custom software developed in MATLAB (The MathWorks, Inc., Natick, MA, USA).

**Measurement of tendon echo-intensity**
Tendon echo-intensity (for images acquired with 0-degree knee angle) was measured both in the longitudinal (Fig. 1: middle panels) and in the transversal scan (Fig. 1: lower panels) regions of interest: the mean echo intensity (8-bit resolution, resulting in a number between 0 and 255, where black=0, white=255) was determined by a custom software developed in MATLAB (The MathWorks, Inc., Natick, MA, USA).

**Statistical analysis**
Since the Shapiro-Wilk test for the normal distribution of the data failed, non-parametric tests were used. The Friedman’s ANOVA followed by Dunn’s post-hoc test and the Wilcoxon test were adopted for comparisons of size parameters (i.e., thickness and cross sectional area) and echo-intensity both between the two tendon portions investigated (proximal vs central) and between the two sides (left vs right).

K-means cluster analysis followed by F-ratio calculation were applied to the side-to-side differences in size and echo-intensity of the tendon proximal portion in order to discern between different groups of players.

Data are expressed as median (and range) and are represented as box-and-whisker plots. Threshold for statistical significance was set at P<0.05. Statistical tests were performed with the IBM SPSS Statistics (version 20 - IBM Corporation, Armonk, NY, USA) software package.

**Results**

*Qualitative assessments of tendon structure and neovascularization*
Figure 2 reports a representative example of one thrower player showing in both sides a normal structure of the patellar tendon without neovascularization, while Figure 3 shows for another representative thrower player moderate structural changes and mild neovascularization in the proximal portion of the left patellar tendon and normal structure in the homologous portion of the contralateral tendon.

Similar to these representative examples, the qualitative assessment of the longitudinal and transversal scans of all players showed normal structure without neovascularization in 7 out of 14 throwers and structural changes with or without neovascularization in the proximal portion of the non-dominant tendon of the other 7 throwers. The structure and neovascularization scores of the non-dominant tendon of all players are listed in Table II: in total, 5 out of 14 throwers (35% of cases: # 4-8-9-11-13 in Tab. II) were considered affected by “subclinical tendinopathy” of the non-dominant tendon because of the presence of light structural changes in association with at least mild neovascularization (two out of five players) or moderate/severe structural changes with or without neovascularization (three out of five players).

*Quantitative assessments of tendon size and echo-intensity*
Figure 4 reports the box-and-whisker plots of the tendon size parameters (thickness and CSA) and of the echo-intensity for both sides (left and right) and for both portions (proximal and central) investigated.

As expected, the thickness of the proximal portion of both sides of the tendon was higher compared to that of the central portion (left side: P<0.001; right side: P<0.01), while no differences were observed in tendon CSA and echo-intensity between the proximal and central portion of both sides (P values > 0.05). Further, no differences in thickness, CSA, and echo-intensity of the tendon proximal portion were observed between the two sides (P values > 0.05).

Analysis of individual data and K-means cluster analysis unraveled a remarkable interindividual variability in the side-to-side differences in size and echo-intensity of the tendon proximal portion. In fact, 6 out of 14 players (43% of cases) presenting comparable values of thickness, CSA, and echo-intensity between the proximal portion of the two sides were classified in cluster 1 (Tab. III: P>0.05 for all side-to-side comparisons), while 8 out of 14 players (57% of cases)
presenting side-to-side differences in CSA (P=0.01) and echo-intensity (P=0.01) were included in cluster 2 (Tab. III: P<0.05 for side-to-side comparisons in CSA and echo-intensity). In this subgroup of 8 players, CSA was significantly higher and echo-intensity was significantly lower in the non-dominant tendon compared to the dominant one. The following significances (relative to the F-ratio for each variable) were obtained for the differences between the two clusters: thickness: P=0.165; CSA: P=0.005; longitudinal echo-

Figure 2. Representative patellar tendon scans of one thrower player showing in both sides a normal structure of the tendon without neovascularization.
intensity: $P=0.001$; transversal echo-intensity: $P=0.003$.
Briefly, CSA and echo-intensity were the quantitative variables most discriminant between the two clusters and resulted significantly different between the dominant and the non-dominant tendon in a large subgroup (57% of cases) of asymptomatic thrower players.

Figure 3. Representative patellar tendon scans of one thrower player showing moderate structural changes and mild neo-vascularization in the proximal portion of the left patellar tendon and normal structure in the homologous portion of the right tendon.
Correspondence between qualitative and quantitative assessments

The 5 players identified by the qualitative analysis as affected by subclinical tendinopathy (#4-8-9-11-13 in Tab. II) were a subgroup of the 8 players classified in cluster 2.

Briefly, the qualitative assessments of tendon structure and neovascularization were in good agreement with the results of the cluster analysis applied to quantitative data of CSA and echo-intensity.

Do patellar tendon abnormalities predict tendinopathy?

Two players (#4 and 9 in Tab. II) identified by qualitative analysis as affected by subclinical tendinopathy and classified in cluster 2 (2 out of 8 cases: 25%) developed a clinical tendinopathy (that was confirmed by ultrasonography through qualitative assessments of tendon structure and neovascularization) after a follow-up of six months, while no players classified in cluster 1 (0 out of 6 cases: 0%) developed a tendinopathy (Tab. II, last column). Therefore, patellar tendon abnormalities at baseline seems to increase (although not significantly) the risk of development of subsequent patellar tendinopathy (relative risk=3.89, 95% CI 0.22 - 68.67, P=0.35).

Table II. Results of the qualitative assessments of tendon structure and neovascularization (second and third column: 5 players affected by "subclinical tendinopathy" of the non-dominant patellar tendon are highlighted in bold) and of the cluster analysis applied to quantitative data of cross-sectional area and echo-intensity (fourth column). The last column reports that a clinical tendinopathy of the non-dominant patellar tendon was found in two players (#4 and 9) after a follow-up of six months.

<table>
<thead>
<tr>
<th>Player</th>
<th>Structure score</th>
<th>Neovascularisation score</th>
<th>Cluster</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>+</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>7</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>8</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>9</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>+</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>11</td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>12</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>13</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>14</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure 4. Patellar tendon thickness, cross sectional area, longitudinal echo-intensity, and transversal echo-intensity for both sides (left and right) and for both portions (proximal and central) investigated in the whole group of 14 thrower players.
In the present study, patellar tendon ultrasound images were acquired from both sides of 14 asymptomatic thrower players and qualitative assessments (of tendon structure and neovascularization) and quantitative measurements (of tendon thickness, CSA, and echo-intensity) were performed. Referring to the aims listed in the Introduction, the main results can be summarized as follows: I) qualitative assessments showed a subclinical tendinopathy of the non-dominant tendon in 5 out of 14 players (35% of cases); II) quantitative assessments showed abnormalities of the non-dominant tendon in 8 out of 14 players (57% of cases); III) qualitative assessments of tendon structure and neovascularization were in good agreement with the results of the cluster analysis applied to quantitative data of CSA and echo-intensity; IV) CSA and echo-intensity were the discriminative tendon parameters most discriminant between asymptomatic players without structural tendon abnormalities and those with tendon abnormalities; V) patellar tendon abnormalities at baseline seem to increase the risk of development of subsequent patellar tendinopathy.

The high prevalence of subclinical tendinopathy and structural tendon abnormalities in the non-dominant side of the investigated group of thrower players can be related to the repetitive overload of the non-dominant lower limb: in fact, the throwing performance of these players is similar to that of javelin throwers who commonly present both subclinical and clinical patellar tendinopathy.

Patellar tendinopathy is commonly evaluated through the ultrasonographic assessment of localized tendon thickening, presence of hypoechoic areas, and altered vascularity. The two approaches (qualitative and quantitative) we adopted to evaluate the patellar tendons of throwers enabled to assess all these ultrasonographic features: tendon size, tendon echogenicity, and tendon vascularity. The main findings observed in players classified as affected by subclinical tendinopathy (and included in cluster 2) were presence of hypo-echogenic areas and decreased echo-intensity of the non-dominant tendon compared to the dominant one, presence of vessels on Power Doppler analysis, increased thickness and CSA of the non-dominant tendon compared to the dominant one. All these findings represent valid signs of some of the structural changes that occur during the tendinopathic process: increase in tendon thickness, increase in the number of vessels, disorganization of collagen fibers, increase in the hydrated components of the extracellular matrix, and breakdown of tissue organization. Other histopathologic findings (that cannot be assessed through ultrasonography) include increase in the number of sensory nerves, proliferation of type III collagen fibers, hypocellularity, increased number of inflammatory cells. All these features underlie tendon pathomechanics (i.e., reduced load-bearing capacity) and ultimately result in tendon pain (and increased risk of tendon rupture).
In the present study, the percentage of structurally abnormal asymptomatic patellar tendons that became symptomatic was 25%. This percentage and the relative risk of 3.89 are in agreement with the results of previous studies indicating that the ultrasonographic structural changes in asymptomatic subjects may represent markers of an early presymptomatic pathology. However, a frequently cited criticism of ultrasound findings is their poor reliability. For instance, ultrasonography is perceived to have a high risk of error when evaluating tendon thickness and CSA. Moreover, the qualitative assessment of Achilles and patellar tendon echogenicity showed poor to moderate inter-observer agreement. On the contrary, quantitative analyses of tendon thickness and CSA showed excellent inter-rater and intra-rater reliability. To our knowledge, this is the first study investigating the longitudinal and transversal echo-intensity in patellar tendons of asymptomatic humans. Previous experimental works assessing the echo-intensity of Achilles tendons in rats and of infraspinatus tendons in sheep found that the echo intensity of the injured tendons was significantly lower compared to intact tendons. Consistently, we observed in the 8 players classified in cluster 2 that both the longitudinal and the transversal echo-intensity were significantly lower in the proximal portion of the left (non-dominant) tendon compared to the right (dominant) tendon. Further, we showed that the quantitative variables most discriminant between structurally abnormal and normal patellar tendons were echo-intensity and CSA. Therefore, we recommend the systematic use of these quantitative ultrasound features in cross-sectional studies aimed to compare the structural tendon adaptations between different populations as well as in longitudinal studies aimed to establish the time course of the tendinopathic process. It may be hypothesized that the decrease in tendon echo-intensity (that is possibly related to the increase in the hydrated components of the extracellular matrix and to the local inflammatory response) precede the increase in tendon CSA (that is possibly related to disorganization of collagen fibers and proliferation of type III collagen fibers). However, further studies are required to confirm this hypothesis.

There are several limitations to this study. First, the small sample size and the peculiarity of the investigated population of elite throwers make generalization to other (tendons of other) populations of athletes difficult. Second, the short duration of the follow-up (six months) could have implied an underestimation of the risk of development of patellar tendinopathy. Third, the causal association between (ultrasound markers of) tendon histopathology and development of tendinopathy was not demonstrated.

In conclusion, this study showed a high prevalence of abnormalities of the non-dominant patellar tendon in the recruited group of elite players (35% and 57% of cases according to the qualitative and quantitative assessments, respectively). Qualitative assessments of tendon structure and neovascularization were in good agreement with the results of the cluster analysis applied to quantitative data of CSA and echo-intensity. Finally, we found that patellar tendon abnormalities at baseline seem to increase the risk of development of subsequent patellar tendinopathy.

Conflict of interest

None.

References


