**Title:** Development of an Ultrasound Imaging Atlas (AUTUSI atlas) for Grading Osteoarthritis in the First Metatarsophalangeal Joint

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**Conflicts of interests**

The authors declare no conflicts of interest in relation to this work.

**Abstract**

**Objective**

Ultrasound (US) imaging may play a fundamental role in the earlier detection and assessment of first metatarsophalangeal joint (MTPJ) osteoarthritis (OA) due to its ability to depict tissue-specific morphological changes before the point of irreversible structural damage. However, the role of US in supporting the diagnosis of OA in foot joints has not been clearly defined. The aims of the study were to develop a semiquantitative US atlas (the AUTUSI atlas) to grade the degree of osteoarthritic change in the first MTPJ and to evaluate the intra-examiner and inter-examiner reproducibility of using the atlas.

**Design**

Ultrasound images were obtained from 57 participants (30 participants with radiographically confirmed first MTPJ OA). The AUTUSI atlas supports the examination of grading joint effusion, synovial hypertrophy, synovitis, osteophytes, joint space narrowing, and cartilage thickness. Six examiners used the atlas to independently grade 24 ultrasound images across two sessions. Intra-examiner and inter-examiner reproducibility were determined using percentage agreement and Gwet’s AC2.

**Results**

Observations using the AUTUSI atlas demonstrated almost perfect-to-perfect inter-examiner agreement (percentage agreement ranged from 96% to 100%, and Gwet’s AC2 values ranged from 0.81 to 1.00) and moderate-to-perfect intra-examiner agreement (percentage agreement ranged from 67% to 100%, and Gwet’s AC2 values ranged from 0.54 to 1.00).

**Conclusion**

The AUTUSI atlas demonstrated excellent intra-examiner and inter-examiner reproducibility for evaluating first MTPJ joint effusion, synovial hypertrophy, synovitis, joint space narrowing, osteophytes, and cartilage thickness. The AUTUSI atlas affords an opportunity to detect prognostic markers of OA earlier in the disease cascade and has the potential to advance understanding of the pathological process of first MTPJ OA.

**Keywords**

Osteoarthritis

Metatarsophalangeal Joint

Ultrasound imaging

Foot

**Significance and innovations**

* The AUTUSI atlas recognises the heterogeneous involvement of multiple joint tissues and provides a reliable means of evaluating both structural and inflammatory features specific to first MTPJ OA
* The AUTUSI atlas offers the opportunity to identify prognostic inflammatory features earlier in the course of the disease before potentially irreversible damage or disability occurs.
* Application of the AUTUSI atlas may provide an opportunity to detect inflammatory features before the point of pain and before the point of irreversible structural damage. The ability to detect inflammatory features of OA before the point of irreversible structural change may provide a window of opportunity where individuals may be more responsive to interventions, thus broadening the scope and capabilities of targeted interventions to alter disease progression and symptomatic consequence. Consequently, the AUTUSI Atlas will expand people's eligibility for OA research and enable the study of targeted interventions at different stages of the disease process.

**Introduction**

Osteoarthritis (OA) is a prevalent, chronic, progressive, and disabling joint disease that imposes a remarkable global health burden, with notable implications for the individuals affected, healthcare systems, and wider socioeconomic costs (1, 2). Foot OA results in functional limitations and significant impairments in balance, strength, and locomotor ability, and negatively impacts work ability (3). However, the feet are often overlooked as a site of involvement relative to other joints commonly affected by OA (4). Within the foot, the most commonly reported affected foot site is the first metatarsophalangeal (MTPJ) (5).

The traditional view of OA as a degenerative disorder of articular cartilage resulting from normal bodily wear and tear is obsolete (6-8). The contemporary concept of OA emphasises the complex pathogenesis of the disease as a disorder of the joint as a whole organ, which involves not only hyaline cartilage but an additional and integrated role of bone and synovial tissue (6). Attention has now turned to the prognostic value and role of inflammatory markers (6, 9, 10), with several studies reporting an association between active synovitis and structural OA progression (11-14).

The development of the radiographic La Trobe Foot Atlas (LFA) (15) led to significant improvements in estimating foot OA disease prevalence (16), as well as understanding different patterns of foot joint involvement (17). However, the current method of diagnosing OA by conventional radiography remains reactive and captures OA later in the disease process, once the condition is significantly advanced (15, 18). This necessitates a new model of diagnosis and care for OA that is proactive and preventative. More advanced modalities, including magnetic resonance imaging (MRI) and ultrasound (US) imaging have emerged as more accurate evaluators of both bone and soft tissue abnormalities in foot OA (19, 20). Magnetic resonance imaging has played a principal role in changing the understanding of OA pathologies when evaluating OA as a whole organ disease (10, 21). Munteanu et al. (19) recently developed an MRI atlas for the assessment of first MTPJ OA which demonstrated excellent intra-examiner and inter-examiner reproducibility (19, 22). However, shortcomings for MRI include high costs, prolonged duration of image acquisition, limited availability in community care, and it is contraindicated in certain conditions such as metal implants (23, 24). Notably, Oo et al.(25) demonstrated that the correlations between quantitative knee OA US features and corresponding MRI findings were very strong (ICC range = 0.85–0.98) (25).

Ultrasound imaging potentially affords inherent advantages for the diagnosis of first MTPJ OA. Ultrasound imaging can be performed chairside, has the ability to detect inflammatory joint pathology that is otherwise not detected by clinical examination, (26, 27) and can reliably quantify both bone and soft tissue abnormalities (28, 29). Ultrasound imaging may play a fundamental role in the earlier detection and assessment of foot OA (30, 31), thus enabling more targeted and timely interventions that may provide the capacity to alter disease progression. However, no foot-specific US grading system exists to classify the degree of pathological change in joint tissues. Therefore, the role of US in the diagnosis of OA in foot joints has not been clearly defined. The development of a US imaging atlas could provide capacity for earlier detection, standardisation of diagnosis, and a more sensitive method for classifying and grading the disease process. Therefore, the aims of the current study were to develop a semiquantitative US imaging atlas (image-based grading system) to grade the degree of osteoarthritic change in the first MTPJ and to evaluate the intra-examiner and inter-examiner reproducibility of using the atlas.

**Methods**

**Development of the AUTUSI atlas**

***Background work***

The AUTUSI atlas was developed using an evidence-based approach, using findings from a systematic review (32), a scoping review (33), a Delphi consensus study (34), and a US imaging reliability study (29) as the basis for development. The systematic review identified inconsistencies in the assessment of US imaging features, the definition of features, and the grading systems used to determine the degree of OA change in peripheral joints (32). Inconsistencies were identified by the scoping review against international guidelines and limited implementation of consensus-based recommendations for US imaging procedure guidance when evaluating the first MTPJ (33). To address these research gaps, an international multi-speciality Delphi study was conducted to gain consensus concerning which US imaging features should be assessed and graded, and what US imaging procedure should be performed when examining the first MTPJ for features of OA (34). The Delphi study identified the essential components that the US imaging acquisition procedure should encompass when examining the first MTPJ for features of OA (34). Our reliability study details the US imaging acquisition procedure and semiquantitative grading system for evaluating first MTPJ OA features in participants with radiologically confirmed OA (29). The US imaging acquisition procedure and grading system were reliable in evaluating joint effusion, synovial hypertrophy, synovitis, joint space narrowing, osteophytes and cartilage thickness in participants with radiologically confirmed OA. Two examiners demonstrated good to excellent inter-examiner reliability across two sessions (ICCs 0.6 to 1.0; ICCS 0.55 to 1.0) and intra-examiner reliability (ICCs 0.58 to 0.92) for all features graded on a semiquantitative scale (29). The semiquantitative grading of these US features was incorporated into the methodological development of the AUTUSI atlas. The Southern Health and Disability Ethics Committee, HDEC Ethics Reference: 2022 FULL 12721, granted ethical approval for the study.

***Participants***

Ultrasound images were obtained from 57 participants who participated in an earlier reliability study (29). Participants over 20 years of age with a previous imaging diagnosis (confirmed by an imaging modality) or clinically diagnosed (diagnosed based on clinical assessments) first MTPJ OA were recruited from the general population in Auckland, New Zealand. Participants were recruited through professional interactive networks, social media (Twitter / X and Facebook) and local newspaper advertisements. Exclusion criteria were the possibility of pregnancy, the presence of any other inflammatory musculoskeletal condition, history of a first MTPJ surgery, or foot and/or ankle surgery in the last three months. Study participants were categorised into two groups (radiographically confirmed OA and non-OA group). The LFA was used to determine the presence of radiographic first MTPJ OA (15). Fifty-seven participants were screened in the first imaging session, of whom 30 had radiographically confirmed first MTPJ OA. All 30 participants with radiographic first MTPJ OA were invited to return for a repeat US imaging examination as part of the reliability study (29). Across both US imaging sessions, 87 sets of US images (one static image per feature) were acquired. Each participant was randomly assigned an alphanumeric code upon entry into the study. Written informed consent was obtained from all participants before the study. **Figure 1** provides an overview of participant engagement and how US images were acquired to develop the atlas.

***Participant and imaging data***

All imaging was performed at a private medical imaging facility and was conducted sequentially within a 60-minute session in three separate rooms. First, demographic data were obtained for each participant (age, gender, height, weight, body mass index, ethnicity, and first MTPJ affected). Second, the presence of radiographic first MTPJ OA was determined through radiographic screening and assessment. Third, participants underwent an US examination using the US imaging acquisition procedure and grading system (29).

***INSERT FIGURE 1 NEAR HERE***

***Radiographic assessment***

Weightbearing dorsal/plantar and lateral radiographs were obtained by an experienced radiographer. A radiologist (PC with more than 20 years’ experience) used the LFA to determine the presence of radiographic first MTPJ OA (15). The LFA considers OA to be present when a score of 2 or greater for osteophytes or joint space narrowing is documented from either the dorsal/plantar or lateral view (15). One radiologist (PC) assessed and reported on all radiographs.

***US imaging assessment***

Directly after the radiographic examination all 57 participants received an US examination. US features included in the examination were joint effusion, synovial hypertrophy, synovitis, osteophytes, joint space narrowing, and cartilage thickness. All US features were assessed in the dorsal view with the probe positioned longitudinally; a transverse orientation was also applied to examine joint effusion and synovial hypertrophy, to ensure the extent of the effusion and synovial thickening was fully investigated. A semiquantitative grading system was applied to all features (0=Absent, 1=Mild, 2=Moderate, 3=Severe). To mitigate problems with adequate discrimination between intermediary grades of cartilage thickness, a 0-2 semiquantitative grading system was applied **(Supplementary file 1)**. A sonographer (KF with more than 15 years of experience in musculoskeletal US imaging) applied the US imaging acquisition procedure and grading system to evaluate the first MTPJ for features of OA. A radiologist (PC) applied the grading system to the static images acquired by the sonographer. Both examiners were blinded to the radiographic results, clinical data, and each other’s grades. All ultrasound images were stored for the purpose of developing the AUTUSI atlas.

***Equipment***

A Philips Epiq Elite HW B.2 ultrasound machine, equipped with a multifrequency linear transducer (eL18–4 MHz), was used to acquire images of the first MTPJ. The US device did not change during the session or between sessions. Grey scale was used to examine all features (128Hz Gray Map 3, grey scale gain 30-40%, dynamic range 74-68%, Med, 2D Opt Res, SonoCT, XRES 4) and power Doppler was applied for the examination of synovitis (PRF 700Hz, Non-directional Flow Colour Map CPA 3, Colour gain 50-65%, Wall Filter 63Hz, Frequency 6.2MHz). The grey scale and power Doppler settings were adjusted to optimise image resolution and sensitivity to detect flow. At the beginning of each scanning session, focus was positioned at the level of the region of interest. Colour gain was adjusted below the degree that caused the appearance of noise artefacts.

***AUTUSI atlas image selection and consensus***

The first MTPJ OA AUTUSI atlas was developed by a podiatrist (PM), sonographer (KF) and radiologist (PC), who were all involved in the reliability study (29). **Figure 2** displays the image assessment and categorisation process used to develop the AUTUSI atlas. The three atlas developers (PM, KC, and PC) assessed all 87 sets of ultrasound images. Each set of ultrasound images included six longitudinal images illustrating joint effusion, synovial hypertrophy, synovitis, osteophytes, joint space narrowing, cartilage thickness, and two transverse images illustrating joint effusion and synovial hypertrophy. PM categorised all 696 static US images by feature and severity according to grading by the radiologist and sonographer from the authors’ reliability study (29). All 696 static images representing each feature with varying degrees of severity were reviewed to select the most representative examples of each grade of each feature. PM, KF and PC independently selected five images that best represented each feature definition and rating description. From here, the sonographer and radiologist independently selected the best image for each grade of each feature. If there was a disagreement, the image in question was replaced with another image considered a better representation. This process was repeated until consensus was achieved. This process led to the final presentation of the first MTPJ OA AUTUSI atlas.

***INSERT FIGURE 2 NEAR HERE***

***Preparation of ultrasound images***

An initial AUTUSI atlas was developed consisting of six features with an associated semiquantitative grading system. The AUTUSI atlas consisted of 31 static ultrasound images. To assist in the differentiation between grades and features, PM designed line drawings to align with each ultrasound image. The AUTUSI atlas was distributed to KF and PC for approval. The AUTUSI atlas was independently evaluated and confirmed appropriate by co-authors (MC, CB, RE, and KR). Agreement was reached regarding image selection and format design. **Figure 3** shows the AUTUSI atlas images for the assessment of synovitis. Power Doppler signals must be detected within synovial hypertrophy to be considered as a sign of synovitis. The scoring of power Doppler signals was performed according to a semiquantitative scale (0 = No flow in the synovium, 1 = Mild, single vessel signals (one or more), 2 = Moderate, confluent vessel signals in less than half of the area of the synovium, 3 = Severe, vessel signals in more than half of the area of the synovium). The full AUTUSI atlas is presented in **Supplementary file 2.**

***INSERT FIGURE 3 NEAR HERE***

**Reproducibility of the AUTUSI atlas**

***Image selection for grading***

To assess the reproducibility of the AUTUSI atlas, a random sample of 24 images were selected from the total sample of 696 available images for grading. Ultrasound images were grouped by participant code according to feature and grade using Microsoft® Excel® (version 2307, Microsoft Corp., Redmond, Washington, USA). Participant codes representing images included in the AUTUSI atlas were deleted from the spreadsheet to ensure atlas examiners would not grade images that had been previously selected as atlas images. Ultrasound images were randomly selected by row number using a random number generator. Twenty-four US images (three images per feature) were randomly selected from the remaining 665 images. Selected images were compiled into a document and sent with the AUTUSI atlas to examiners.

***Atlas examiners***

Six examiners with experience in US imaging from different countries (New Zealand [n=4], United Kingdom [n=1] and Canada [n=1]) and different health professions (radiologists [n=2], podiatrist [n=2], sonographer [n=1], and physical therapist [n=1]) independently rated the randomly selected ultrasound images using the AUTUSI atlas. All examiners were provided with written instructions detailing the AUTUSI atlas, the US imaging grading sheet, and the tasks required of them for grading sessions one and two. An additional video outlining the process was also provided. This step included three examiners (PM, KF, and PC) who developed the atlas and three examiners who were not involved in the US imaging acquisition or the atlas development – a radiologist, a physical therapist, and a podiatrist. To determine the inter-examiner reproducibility of the AUTUSI atlas, all six examiners (PM, KF, PC, RG, JW, and CD) independently rated the random sample of 24 US images (3 images per feature of varying severity) using the atlas. To determine intra-examiner reproducibility, this process was repeated 3-4 weeks later (without reference to previous ratings).

***Statistical Analysis***

Gwet’s AC2 values were calculated using the irrCAC package in R, version 1.0, to determine inter-examiner and intra-examiner reliability. Quadratic weighting was used for all calculations due to the ordinal nature of scoring across the US features. Gwet's AC2 values were interpreted using the following cutoffs: <0 poor; 0.01–0.20 slight; 0.21–0.40 fair; 0.41–0.60 moderate; 0.61–0.80 substantial; 0.81–1.00 almost perfect or 1.00 perfect (35). Gwet's AC1 values were calculated because this approach has been shown to provide a more robust measure of reliability than the traditionally used kappa statistic (36).

**Results**

**Table 1.** details the demographic characteristics of the 57 participants (25 female, 5 male) included in the AUTUSI atlas development study.

***INSERT TABLE 1 NEAR HERE***

**Inter-examiner reproducibility**

**Table 2.** displays the reproducibility statistics for inter-examiner analyses. Percentage exact agreement ranged from 96% to 100%. Gwet’s AC2 values ranged from 0.81 to 1.00, indicating almost perfect-to-perfect levels of agreement. Inter-examiner reproducibility was similar between session one and session two grading. Although acceptable, the grading of synovitis was not as consistent as other US features.

***INSERT TABLE 2 NEAR HERE***

**Intra-examiner reproducibility**

**Table 3.** displays the reproducibility statistics for intra-examiner analyses. Percentage exact agreement ranged from 67% to 100%. Gwet’s AC2 values ranged from 0.54 to 1.00, indicating moderate-to-perfect levels of agreement. Intra-examiner reproducibility was similar for the examiners 1, 2, 3, and 5. Although acceptable, the grading of synovitis was not as consistent as other US features for two examiners.

***INSERT TABLE 3 NEAR HERE***

**Discussion**

This study developed and assessed the reproducibility of the AUTUSI atlas to evaluate the presence and severity of joint effusion, synovial hypertrophy, synovitis, osteophytes, joint space narrowing, and cartilage thickness in the first MTPJ. The AUTUSI atlas demonstrated excellent intra-examiner and inter-examiner reproducibility for evaluating first MTPJ joint effusion, synovial hypertrophy, joint space narrowing, osteophytes, and cartilage thickness. Although acceptable, the grading of synovitis was not as consistent as other US features.

The uncertainty to differentiate between images representing different grades of synovitis may be partly attributable to images depicting a similar volume of power Doppler signal. As outlined in the synovitis description of the AUTUSI atlas, power Doppler signals must be detected within the synovium to be considered as a sign of synovitis (0 = No flow in the synovium, 1 = Mild, single vessel signals [one or more], 2 = Moderate, confluent vessel signals in less than half of the area of the synovium, 3 = Severe, vessel signals in more than half of the area of the synovium). The randomly selected synovitis images chosen for the US imaging grading sheet represent different grades of severity. However, when assessing the power Doppler signal across the entire image and not just the Doppler signal contained to the synovium, the volume may have appeared similar. Therefore, there may have been cases where the Doppler signal was graded outside the synovium, consequently increasing the severity grade. This finding may also reflect the use of previous US imaging grading systems originally constructed from rheumatoid arthritis (RA) populations, where the degree of inflammation is distinctly much higher (37, 38). Additionally, the marked variations across interpretations of how the synovial pathology has been defined and categorised in previous investigations may also be reflected in the scoring variation (32). Variation may also be explained by the number of choices with a semiquantitative grading scale; synovitis grading variation may be attributed to one examiner overscoring and one underscoring when the image was doubtful. Therefore, additional training of all examiners prior to use of the AUTUSI atlas may improve performance and ensure understanding of grading descriptions.

To elucidate the role of inflammation in foot OA, grey scale inflammatory abnormalities for synovial hypertrophy and joint effusion were graded separately. Synovitis was also examined as a separate entity by power Doppler signal (flow signal detected within synovial hypertrophy was considered a sign of synovitis) (39, 40). Differentiating between inflammatory features addresses past inconsistencies and interpretation of the different entities of synovial pathology. Previous discrepancies in how synovitis, synovial hypertrophy and joint effusion have been defined and categorised make it unclear if synovial pathology is best represented as separate entities (joint effusion and synovial hypertrophy) or combined as proxy measures for synovitis (32). Due to the marked variation in prevalence between grey scale and Doppler-detected inflammatory features demonstrated in hand OA (40). Including grey scale features indicative of inflammation solely may result in OA being underestimated. The distinction between inflammatory pathology is further enhanced by our semiquantitative grading system specific to OA. To date, US imaging grading systems applied to OA have been largely extrapolated from those originally designed and validated to quantify inflammatory change in RA (32). The distinct difference of inflammation experienced in OA compared to RA (12, 37), reinforced the need for an OA-specific grading that truly depicts disease progression, as in this work.

The AUTUSI atlas will allow for quantification of the presence and severity of pathologic change occurring in joint tissues in first MTPJ OA. The AUTUSI atlas has the potential to advance understanding of the pathological process of first MTPJ OA, provide capacity for earlier detection, support standardisation of diagnosis, and subsequently broaden the scope and capabilities of targeted interventions. From this work, it is evident that there is a role for using US imaging in detecting different stages of first MTPJ OA. The AUTUSI atlas may be the catalyst in developing a classification criteria specific to the first MTPJ OA, which could be utilised both clinically and in future research. The AUTUSI atlas will enhance the capability to detect and characterise first MTPJ OA across various stages of disease progression rather than being restricted solely to the point of irreversible end-stage disease.

The AUTUSI atlas is the first step in a longitudinal analysis to determine the predictive value of US features in participants without radiographically confirmed OA. Incorporating the AUTUSI atlas as an outcome measure in population-based cohort studies could provide valuable insight into early detection. Furthermore, it provides the capacity to determine which US imaging features may progress to radiographic change and increased symptomatic consequences. Finally, further work is needed to determine which AUTUSI atlas features would be most accurate in the diagnosis of first MTPJ OA compared to radiographic grading, the relationship between the atlas and symptom severity, and how atlas features respond to clinical interventions.

A particular strength of the present study is the inclusion of six examiners from different countries and health professions, and yet all experienced users of US imaging. Additionally, three reviewers were blinded to the image acquisition and atlas development process. The justification for including six examiners from different professional backgrounds ensures the atlas is reproducible across a range of health professions that are likely to implement the atlas, such as podiatrists, physical therapists, radiologists, sonographers and researchers. In comparison, an MRI atlas and the LFA included the same two examiners for atlas development and reproducibility (15, 19). A further strength of our study is the accompaniment of supporting images in our atlas which clearly depicts the US imaging (patient and probe positioning) acquisition procedure. Ultrasound imaging has previously been demonstrated as being operator-dependent (41, 42). However, substantial developments in US software and transducers have advanced the resolution to better identify superficial structures, such as the first MTPJ, allowing more detailed assessment of associated pathology (28, 43, 44). To help mitigate operator dependence the authors wanted to ensure that a standardised procedure could be performed for all atlas users. Supporting images would ensure correct interpretation and allow subsequent studies to replicate our procedure.

This study must be viewed in the context of its limitations. First, despite examining 24 ultrasound images with the atlas, the reviewers only evaluated three images per feature. Therefore, future research is needed to determine the reproducibility of the atlas when grading a larger sample of images. Second, there may have been examiner bias because the same radiologist reported on radiographic screening and graded the acquired ultrasound images. To minimise the risk of bias, we randomly assigned an alphanumerical code to all participants upon entry into the study. Finally, the prevalence and burden of OA are not uniform across demographic groups. Minority populations, especially African American, Hispanic, Māori and Pacific people experience poorer health outcomes (such as pain and disability) (45, 46). Despite efforts to proactively recruit an ethnically diverse population representative of the broader New Zealand population, no Pacific peoples were included. Pacific peoples suffer from significant and longstanding health inequalities and poorer health outcomes compared to other New Zealanders. Therefore, the inclusion of ethnicity related to OA data is needed to better understand what factors contribute to these inequalities and to provide the capacity to measure progress.

**Conclusion**

The AUTUSI atlas demonstrated excellent intra-examiner and inter-examiner reproducibility for evaluating first MTPJ joint effusion, synovial hypertrophy, synovitis, joint space narrowing, osteophytes, and cartilage thickness. The AUTUSI atlas affords an opportunity to detect prognostic markers of OA earlier in the disease cascade, which would provide a window of opportunity and broaden the scope and capabilities of targeted interventions to alter disease progression. The AUTUSI atlas has the potential to advance understanding of the pathological process of first MTPJ OA.

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**Authors’ contributions**

PM, CB, RE, KR and MC were responsible for the conception and design of the research. PM was responsible for recruiting participants. PM, KF, and PC were responsible for conducting the research, image acquisition, and initial atlas generation. PM, KF, PC, JW, RG and CD were responsible for interpreting and applying the AUTUSI atlas. PM, CB, RE, KR and MC were responsible for analysing data and interpreting the results. All authors were responsible for the preparation and review of the manuscript prior to submission for publication. All authors read and approved the final manuscript.

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**Figure legend**

**Figure 1: Atlas development participant journey**

OA, Osteoarthritis; US, Ultrasound

**Figure 2: Image assessment and categorisation process used to develop the AUTUSI atlas**

US, Ultrasound

**Figure 3:** **Atlas images for synovitis of the first MTPJ**

1st Met, First metatarsal; 1st PP, First proximal phalanx; 1st MTPJ, First metatarsophalangeal joint; EHL, Extensor hallucis longus.