Enthesitis of the hands in psoriatic arthritis: an ultrasonographic perspective

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Abstract
Psoriatic arthritis is a systemic inflammatory disease in which enthesitis and dactylitis are two of the main hallmarks of the disease. In the last years, ultrasonography is increasingly playing a key role in the diagnosis of psoriatic arthritis and ultrasonography of the entheses, particularly of the lower limbs, is commonly used to assess patients with that disease. New advancements in ultrasound equipment using high frequencies probes allowed us also to identify and characterize the involvement of the entheses of the hand in psoriatic arthritis, confirming the results of the experimental models of the disease and the theory of the sinovio-entheseal complex, even in small joints.

Keywords: ultrasonography; psoriatic arthritis; enthesitis; seronegative arthritis; synovio-entheseal complex

Introduction
Psoriatic Arthritis (PsA), usually included in the Spondyloarthritis (SpA) group, can affect different articular structures, from bone to soft tissues (e.g. tendons, entheses, and bursae) and enthesitis seems to be the earliest lesion in animal models of SpA [1,2]. From a clinical point of view, in 1971 Ball et al firstly described enthesitis in SpA [3]. Two decades later McGonagle et al demonstrated the concept of the involvement of the synovio-entheseal complex (SEC) and enthesis-related inflammation as an early event in PsA and SpA and its diagnostic usefulness to differentiate PsA from Rheumatoid Arthritis (RA) [4,5]. The European League Against Rheumatism (EULAR) recommends the use of imaging in diagnosis and management of SpA and, in the last years, ultrasound (US) of the entheses, particularly of the lower limbs, has been increasingly applied in diagnostic studies on PsA [6,7]. Differential diagnosis in seronegative polyarthritis of the small joints is a challenge for clinicians and with potential diagnostic errors often occur. Recently, several studies have highlighted the role of hands US to assist the differential diagnosis between PsA and RA, demonstrating more severe extra-synovial involvement which includes the SEC in the former group [8–10]. In 2001, Benjamin et al proposed to distinguish entheses in classical and functional: in the latter fibrocartilage are in contact with bone but are not attached [11]. Functional entheses plays a key role in resisting shear and compressive mechanical stress of the entheseal organ [4,12].

The aim of this pictorial essay is to describe and show the most common US findings of functional and classical enthesitis of the hands in patients with PsA.
Functional entheses

The digits have several functional entheses associated with the presence of fibrocartilage to reduce mechanical stress. Here we report examples of hands functional enthesitis during PsA and the description of their US elementary lesions.

Extensor tendon on metacarpophalangeal joint

Despite the crucial role of tendon involvement in chronic inflammatory arthritis, most of US research has been focused on synovitis, some on tenosynovitis, and very few on the peritenonitis (PTI) [13]. Gutierrez et al described the presence of US detectable tendon thickening, echostructural hypoechoogenicity, and hypoechoic swelling of the tissue surrounding the tendon, with or without peritendineal power Doppler (PD) signal at the level of tendons without synovial sheath [14]. One of the images they presented, showed the peritendinitis of the extensor digitorium tendons at the level of the metacarpophalangeal (MCP) joints, defined as hypoechoicgicity at the level of peritenon with associated oedema of the surrounding soft tissues and PD signal [14]. The same group published in 2011 preliminary data on high frequency US of peritenon extensor tendon inflammation in PsA and RA patients. PTI, defined as a hypoechoic swelling of the soft tissue surrounding the extensor digitorum tendon (with or without peri-tendinous PD signal), was found in 65.8% of the clinically involved MCP joints of the PsA group and in none of the RA patients examined [9] (fig 1). At the level of the MCP joints the PTI pattern could be intended as a part of an inflammatory involvement of the SEC, consisting of a functional enthesis formed by extensor digitorum tendon, sesamoid fibrocartilage in the tendon as it crosses the MCP joints, superficial and deep peritendinous tissues, and joint synovial capsule [11] (fig 2). These data suggested the relevant potential role of US in the differential diagnosis between RA and PsA at MCP joints level [9], especially in early disease [8]. Recent preliminary works underline that PTI is as frequent as intra-articular synovitis, that it can be a cause of MCPs swelling and that PD positivity at PTI is able to differentiate PsA development in early arthritis settings [15,16].

Digital pulleys

The pulley system is essential for the accurate tracking of the flexor tendon and for maintaining the correct position of the tendon and bone across the joint and it provides a fulcrum to elicit flexion and extension. Considering the presence of fibrocartilage and its mechanical role, the pulley system is a typical example of functional enthesis. Currently, the most frequent US evaluation of the pulley system described in the literature is focused at the A1 pulley, located on the palmar aspect of the hand, at the level of the MCP joint [17]. For the thickness of that pulley, a cut-off value of 0.62 mm was proposed to discriminate between healthy and diseased structures [18]. Moreover, a positive correlation was found between US and intra-operative measurements of pulley thickness thus confirming the validity of US in the assessment of those structures [18]. In addition, non specific studies focused on the attachment of the pulley on the volar ridges of the proximal and middle phalanges. In inflammatory arthritis, pulleys can be affected during tenosynovitis as their loosening can be seen as secondary to the inflammatory process and this event could be a marker for dis-
ease severity, affecting the function and other outcomes of the patients. Thus this feature could potentially be a marker of disease severity, affecting the hand function. Recently, Tan et al demonstrated by magnetic resonance imaging (MRI) inflammatory changes at digital pulleys and tendons in dactylitis in PsA patients, suggesting a form of functional enthesitis that could be related to the dactylitis process [19]. Moreover, Tinazzi et al showed pulley thickening in subjects affected by PsA compared with RA and healthy controls [20]. As depicted in figure 3 A1, A2 and A4 pulleys in PsA are easily detectable and measurable in longitudinal and transverse scan, but additional studies on early PsA patients are needed to ascertain whether these pulley changes may trigger PsA tenosynovitis or whether they represent a reactive process.

**Fibrous skeleton**

Dactylitis (also named sausage digit) is the clinical hallmark of PsA and it is an important predictor of structural damage [21]. In imaging studies (US and MRI), soft tissue oedema and flexor tenosynovitis are the two most important elementary lesions described in psoriatic dactylitis [10,19,22]. Overall, oedema of the soft tissues can be interpreted as an inflammation of the “fibrous skeleton” of the digit, which is made up of ligaments, fibrous capsular bands, palmar fasciae, and fibrous sheaths that attach to the bone or dermis [10]. However, this theory linking soft tissue oedema and functional enthesitis of the fibrous skeleton needs to be confirmed by pathology. Indeed, to date only a single case report focused on the histologic features of dactylitis: enthesitis and inflammation of tenosynovium were found to be prominent in that case and this inflammation reflected both a chronic T cell-rich inflammatory infiltrate and a stromal reaction [23]. In US studies, soft tissue oedema of the digit was usually defined as a thickening of the soft tissue around the flexor tendon with an intense PD signal [8,10,24] and it was also called by some authors as ‘pseudotenosynovitis’ [24,25] (fig 4). Such soft tissues thickening was also detected in fingers without clinical dactylitis, suggesting that this lesion could have a central role in the genesis of the psoriatic sausage digit. Moreover, Zabotti et al recently demonstrated that US detection of soft tissue oedema around the flexor tendon was highly specific for early PsA, if compared to early RA [8].

### Classical entheses

The sites where tendons, ligaments and joint capsules attach to bone (i.e. classical enthesis) are the typical sites of inflammation in PsA. Currently, most of the US studies focused on the assessment of classical entheses at the level of the lower limbs in PsA for diagnosis and prognosis; however, other areas can be examined for this purpose, particularly the hand. An analysis of hands entesitis by US is reported as follows.

**Insertion of extensor tendon on distal interphalangeal joint and nail area**

Nail disease is often present in PsA patients and can be an early predictor of PsA [26,27]. There is a well known link between extensor tendon enthesopathy, distal interphalangeal (DIP) joint involvement, and nail pathology. As Tan et al elegantly showed by histology, the supporting fascia of the nail root is a continuation of the enthesis of the extensor tendon [28]. This could explain why PsA is associated with a local inflammation that involves both nail root and adjacent bone [28,29]. The US and clinical
study of Aydin et al confirms the importance of extensor tendon distal enthesis in the pathogenesis of nail disease with or without clinical arthritis [29], as previously suggested also by Ash et al [26]. Several studies showed that US is very useful to study enthesis at the level of the distal attachment of the extensor tendon, as well as DIP joint synovitis and nail involvement [28,29]. The extensor tendon distal enthesis involvement can be seen by US as an abnormal hypoechoic tendon with loss of normal fibrillar architecture and/or thickened tendon at its distal phalanx attachment which are seen in two perpendicular planes and that may exhibit PD signals near to the bony cortex and/or bony changes, including enthesophytes, erosion or irregularities [30] (fig 5). The latest Outcome Measures in Rheumatology (OMERACT) definitions indicate for those abnormalities a location ≤2mm from the bony cortex as a landmark of enthesis in SpA [30]. However, they do not specify to which enthesis this is applicable and if this can be considered as a characteristic abnormality also of small structures such as the distal enthesis of the hand extensor tendons. Furthermore, it is crucial also to distinguish findings related to inflammation from those due to structural damage as well as to use US in the differential diagnosis between a DIP involvement due to osteoarthritis or PsA. In this context, a clear and solid evidence from US studies is needed. In terms of nail imaging, US is able to show abnormalities occurring at the level of different components of the nail unit [14]. The possibility to image by US the pathological changes of the nail, such as the loss of the normal trilaminar structure of the plate and the enhanced flow by PD [14,31,32], in such sites could be a matter of speculation regarding the physiopathology of psoriatic disease, according with the nail/enthesis hypothesis [33] (fig 6). Another crucial aspect to consider in this context is the significance of possible nail pathology detected by US in patients without clinical involvement [34]. Finally, new advances in US equipment using high frequencies probes

Fig 5. Longitudinal scan on the DIP joint. A. Thickening of the extensor tendon (et) with hypoechogenicity of tendon and enthesis (white arrowheads) and insertional enthesophyte. B Soft tissue oedema (white star) and synovial effusion (white point) of the DIP joint.

Fig 6. A Longitudinal scan of the nail: normal aspect. The nail plate (*) is as a flawless trilaminar structure with sharp borders. Under the nail plate the nail bed (^) appears as a hypo-isoechoic area under the nail plate and ending in the germinative matrix (#). The insertion and enthesis of the extensor tendon is next to the germinal matrix (°). A picture of a psoriatic nail is in B. The nail plate is altered, with loss of the trilaminar structure and thickening of the plate. The nail bed is thickened, with loss of homogeneity. In C the PD signal is depicted. Small vessels appear with a weak signal characterizing all the bed and some spots are also evident along the venule just next to the tendon. In D the signal is clearly increased and a single spot is also evident in the proximity of the fibrocartilage of the enthesis of the extensor tendon (§)
(e.g. \(\geq 22\)MHz) might allow a better explanation of the link between the nail unit and the DIP joint, possibly revealing more specific alterations for psoriatic disease.

**Insertion of central slip of extensor tendon on proximal interphalangeal joint**

At the site of proximal interphalangeal (PIP) joint, the enthesis of the central slip of the extensor tendon and the synovial fold and recess are intimately related. Central slip enthesis (CSE), firstly described by Filip-pou et al, is defined as grey scale evidence of hypoechoic and thickened enthesis insertion, when compared to the proximal part of the tendon and to the controlateral joint [35]. The detection of CSE has been described to be usually in association with articular synovitis in PsA patients [8,35], manifesting a typical example of hand’s SEC inflammatory involvement (fig 7).

**Conclusions**

Integrating US with clinical evaluation to improve diagnostic accuracy can nowadays be considered an achievable target. In this perspective the identification of highly specific US elementary lesions (i.e. enthesitis) could be useful to assist the rheumatologist in the diagnosis and therapeutic strategies of arthritis. In the light of the crucial role of functional and classical entheses at hand and厚ened enthesis insertion, when compared to the proximal part of the tendon and to the controlateral joint [35]. The detection of CSE has been described to be usually in association with articular synovitis in PsA patients [8,35], manifesting a typical example of hand’s SEC inflammatory involvement (fig 7).

**Conflict of interest:** none

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