Progressive structural bone changes and their relationship with treatment in patients with psoriatic arthritis: a longitudinal HR-pQCT study

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Abstract

Background: Although the short-term effects of tumor necrosis factor alpha (TNF-α) and interleukin-17A (IL-17A) inhibition on the structural changes in psoriatic arthritis (PsA) using high-resolution peripheral quantitative computed tomography (HR-pQCT) have been reported, no studies have investigated the long-term structural changes in PsA patients receiving routine care. We reported longitudinal changes of erosions and enthesiophytes using HR-pQCT and their relationship with treatments in PsA patients over a 5-year period.

Methods: HR-pQCT examination at the second and third metacarpal heads (MCH2 and MCH3) was performed in 60 PsA patients at baseline and after 5 years. The size of each individual lesion was quantified. Erosion and enthesiophyte progression were defined as change exceeding the smallest detectable change (SDC).

Results: A total of 108 bone erosions and 99 enthesiophytes were detected at baseline. Three new bone erosions but no new enthesiophytes were evident at 5 years. A significant increase in mean (+SD) erosion (0.58 ± 1.50 mm³, \( P < 0.001 \)) and enthesiophyte (0.47 ± 0.76 mm³, \( P < 0.001 \)) volume was observed. Erosion and enthesiophyte progression were found in 37/111 (33.3%) and 50/99 (50.5%) lesions, respectively. During this 5-year period, 26 (43%) out of the 60 patients achieved sustained Disease Activity index for Psoriatic Arthritis (DAPSA) low disease activity (LDA) (SDL group, defined as achieving DAPSA-LDA at both baseline and 5 years). Fourteen (23%) out of 60 patients received a TNF inhibitor throughout the 5-year period (TNFi group). Fewer erosions progressed (12/51 [23.5%] vs 25/60 [41.7%], \( P = 0.047 \)) and the increased in enthesiophyte volume was significantly less (0.28 ± 0.67 vs 0.61 ± 0.80 mm³, \( P = 0.048 \)) in the SDL group than in the non-SDL group. However, no significant difference between the TNFi and non-TNFi groups was detected in terms of the change in volume or progression of bone erosion and enthesiophyte.

Conclusion: Damage accrual in terms of bone erosion and enthesiophyte was observed in PsA patients over a period of 5 years despite receiving routine clinical care. Nonetheless, sustained control of disease activity may be able to prevent these bony damages.

Keywords: Psoriatic arthritis, Progression, Erosion, Enthesiophyte, Low disease activity
**Background**

Psoriatic arthritis (PsA) is an inflammatory arthritis associated with psoriasis (PsO), involving not only the appendicular joints but also the entheses and the axial skeleton, resulting in bone damage [1, 2]. Chronic inflammation at these synovial and entheseal sites leads to erosions and enthesiophytes, respectively [1, 2]. By the time PsA patients first present to a rheumatologist, differences in bone erosion and enthesiophyte between PsA patients and healthy individuals are already evident [3–5]. There are age-related bone erosions and osteophytes in the hand joints of healthy individuals, suggesting that the threshold between “normal” and “pathological” is shifted with the increase of age [6]. However, whether there are progressive structural bone changes after the onset of PsA remains a topic of debate. Determining whether and how structural bone damages continue during the course of the PsA is important for understanding its pathophysiological feature and has implications for the treatment.

Recent recommendations emphasized that remission or low disease activity (LDA) as assessed by Disease Activity index for Psoriatic Arthritis (DAPSA) and minimal disease activity (MDA) are the principal targets for the treatment of PsA [7, 8]. Tight control of PsA disease activity through a treat-to-target approach significantly improves joint outcomes for newly diagnosed patients [9]. Tumor necrosis factor (TNF) and interleukin-17A (IL-17A) are the key cytokines involved in both the inflammation and structural changes seen in PsA patients [10]. Using high-resolution peripheral quantitative computed tomography (HR-pQCT), short-term inhibition of erosion and enthesiophyte progression has been demonstrated in the metacarpophalangeal joints of PsA patients after IL-17 inhibition by secukinumab [11]. On the other hand, while erosions showed an arrest of progression in PsA patients treated with either methotrexate (MTX) or TNF inhibitor (TNFi), new bone formation progressed in both groups after 1 year [12].

We report herein the results of a longitudinal HR-pQCT study of PsA. We measured the change in bone erosion and enthesiophyte volumes over a period of 5 years using HR-pQCT. The aims of this study are (1) to relate to treatments in PsA patients.

**Methods**

**Patients**

Sixty consecutive PsA patients were included in this study. Baseline distal radial densitometric and microstructural features in 53 out of the 60 PsA patients have been published [13]. All PsA patients fulfilled the Classification criteria for Psoriatic ARthritis (CASPAR) and were rheumatoid factor negative. Patients with HR-pQCT-detected joint destruction were excluded [14]. All patients had a comprehensive clinical and HR-pQCT assessment at baseline and after 5 years.

All patients in this cohort received routine clinical care at the Prince of Wales Hospital. The choice of treatment, which included conventional synthetic disease-modifying anti-rheumatic drugs (csDMARDs) and biologic DMARDs (bDMARDs), was at the discretion of the patient and his or her treating rheumatologists. Ethics committee approval was obtained from the Ethics Committee of The Chinese University of Hong Kong-New Territories East Cluster Hospitals (CRE-2016.366). All patients provided written informed consent.

**Clinical assessment**

Clinical and demographic parameters recorded include age, body weight, body height, and smoking habits. Clinical assessment included the number of swollen, tender, and deformed joints, Maastricht Ankylosing Spondylitis Enthesitis Score (MASES), and the presence of dactylitis. Disease activity was assessed using the Disease Activity index for PsA (DAPSA) [15], Psoriasis Area and Severity Index (PASI), erythrocyte sedimentation rate (ESR), and C-reactive protein (CRP). Physical function was assessed using the Health Assessment Questionnaire (HAQ) disability index. Use of non-steroidal anti-inflammatory drugs (NSAIDs), csDMARDs, and bDMARDs over the 5-year period was retrieved from the electronic management system.

Remission was defined as very low disease activity (VLODA) or DAPSA ≤4. LDA was defined as MDA or DAPSA ≤14. Sustained disease control was defined as achieving these targets at baseline and 5 years.

**High-resolution peripheral quantitative CT**

All patients underwent HR-pQCT examination (XtremeCT scanner, SCANCO Medical AG, Brüttisellen, Switzerland) at a nominal isotropic voxel size of 82 × 82 × 82 μm of the metacarpophalangeal (MCP) joint of the non-dominant forearm at baseline and 5 years. The non-dominant forearm was chosen to avoid the effects of mechanical stress, similar to other studies [16]. Assessment was confined to the second and third metacarpal head (MCH) regions to optimize time efficiency and minimize motion artifact. The second and third MCHs are the frequently affected MCHs in PsA when assessed by HR-pQCT. Erosive lesions were more commonly found at the MCP2 (82.1%) and MCP3 (85.7%) joints than the MCP4 joint (57.0%) [17]. Similarly, osteophytes were more common at the MCP2 (82%) and MCP3 (86%) joints than at the MCP4 joint (50%) [17].

The patients' forearm was immobilized in a carbon fiber cast fixed within the scanner gantry. A dorsopalmar...
projection image was obtained to define the tomographic scan region. The scan region started at the distal end of the MCH and spanned proximally 9.02 mm (110 slices) at baseline, extended between 80 slices distal to the edge of the MCH and 242 slices proximal to it at 5 years (322 slices).

**Co-localization of VOI: baseline-indexed image registration and slice matching**
A fully automated baseline-indexed 3D image registration and re-slicing (slice matching) [18] based on ITK-SNAP [19] were performed to acquire precisely matched baseline and follow-up volumes of interest (VOIs) at the second and third metacarpal heads (MCH2 and MCH3) (Additional file 1: Figure S1). Follow-up scans were coarsely aligned based on image center and principal axes and registered onto the baseline images using rigid transformation and mutual information image similarity metrics. Then, these scans were re-sliced into the same space of the baseline image using linear interpolation to account for translation and rotation.

Slice-matched co-localized regions of interest (ROIs) were specified on the baseline and follow-up HR-pQCT examinations to assess longitudinal changes in erosions and enthesiophytes. The ROI criteria applied were to (1) include the metacarpal bone only, (2) exclude regions without bone, and (3) exclude the proximal or distal three slices to obviate boundary effect, if necessary.

**Image analysis**
The MCP joints were evaluated for the following criteria: number, size, and volume of erosions and enthesiophytes on the palmar, ulnar, dorsal, and radial quadrants of the MCH. Enthesiophytes were defined as new bone formation arising from the periosteal bone cortex at the insertion sites of the capsule, ligament, or tendons or at the location of functional enthesis [5, 20]. Erosions were defined as a clear break in the outer cortical margin evident on at least two consecutive slices and in two orthogonal planes [5, 17]. Two compartments were imaged corresponding to MCH2 and MCH3 for each patient. A semi-automated method based on ITK-SNAP [21] was used to calculate erosion and enthesiophyte volume on co-localized baseline and follow-up VOIs [4].

**Smallest detectable change**
To determine the smallest detectable change (SDC), 25 enthesiophytes and 30 erosions were randomly chosen and scored twice. SDC was then calculated for enthesiophyte and erosion volume using the formula

\[
SDC = 1.96 \times SD_{\Delta(CLEAN-SCORES)} / (\sqrt{2} \times \sqrt{k}),
\]

where \(SD_{\Delta(CLEAN-SCORES)}\) is the standard deviation of change in scores and \(k\) the number of readings [22].

**Definition of progression of bone erosion or enthesiophyte**
Erosion or enthesiophyte progression was defined as either (i) an increase in erosion or enthesiophyte volume exceeding SDC (SDC for erosion: 0.5 mm³, enthesiophyte: 0.3 mm³) or (ii) new development of an erosion or enthesiophyte. Erosion or enthesiophyte regression was defined as a decrease in erosion or enthesiophyte volume exceeding SDC. Individual erosions or enthesiophytes which did not fulfill the progression or regression criteria were classified as stable.

**Outcomes**
The purpose of this cohort study was (1) to determine the longitudinal change in erosion and enthesiophyte volume over a period of 5 years and (2) to evaluate the effect of (i) achieving sustained DAPSA-LDA (SDL) and (ii) long-term anti-TNF therapy on the progression of erosions and enthesiophytes.

**Statistical analysis**
After testing for Gaussian distribution, paired \(t\) test or Wilcoxon signed-rank test was applied to compare the structural bone changes at baseline and 5 years as appropriate. Comparisons between the two groups were performed using the chi-square test for categorical variables. Generalized mixed linear models were used to adjust for confounding factors including gender, age, body mass index (BMI), disease duration, disease activity (DAPSA) at baseline, and treatment over the 5-year period. Data were analyzed using the Statistical Package for the Social Sciences software for statistics (IBM SPSS V.22.0, IBM Corporation, Armonk, NY, USA), and a \(P\) value of less than 0.05 was considered statistically significant.

**Results**
**Demographic and clinical features**
At baseline, the mean age of the cohort was 51.9 ± 8.9 years, 27 (45.0%) were female, and 13 (21.7%) were ever-smokers (Table 1). The mean duration of psoriasis and PsA before study entry was 18.7 ± 16.7 years and 14.3 ± 6.9 years. Twenty-nine patients (48.3%) had radiologic evidence of juxta-articular new bone formation. Most patients had mild to moderate disease activity. Three (5.0%) patients received corticosteroids, 44 (73.3%) were on csDMARDs, and 16 (26.7%) received bDMARDs.

Compared with baseline, disease activity at 5 years remained relatively stable as only seven patients were started on bDMARDs. Fourteen (23%) patients received TNFi with \(n = 6\) or without \(n = 8\) concomitant methotrexate (MTX) throughout the 5-year period. Seven patients received TNFi less than 5 years, two patients received ustekinumab and secukinumab for less than 2
### Table 1 Changes in demographic and clinical characteristics over 5-year

|                          | Baseline (n = 60) | 5 years (n = 60) | Change (n = 60) | P value     |
|--------------------------|------------------|-----------------|----------------|------------|
| **Demographic characteristics** |                  |                 |                |            |
| Female, N (%)            | 27 (45.0)        |                 |                |            |
| Smokers, N (%)           | 13 (21.7)        |                 |                |            |
| Age, year                | 51.9 ± 8.9       |                 |                |            |
| BMI                      | 25.4 ± 4.3       | 25.8 ± 4.5      | 0.46 ± 1.9     | 0.060      |
| Height (cm)              | 163.6 ± 8.5      | 163.0 ± 8.4     | −0.57 ± 1.3    | 0.002      |
| Weight (kg)              | 68.2 ± 14.3      | 68.8 ± 14.0     | 0.62 ± 5.1     | 0.345      |
| **Disease-specific characteristics** |                  |                 |                |            |
| Duration of psoriasis (year) | 18.7 ± 16.7     |                 |                |            |
| Duration of PsA (year)   | 14.3 ± 6.9       |                 |                |            |
| Juxta-articular new bone formation | 29 (48.3) | | | |
| Polyarticular            | 21 (35.0)        |                 |                |            |
| Oligoarticular           | 17 (28.3)        |                 |                |            |
| Presence of enthesitis   | 13 (21.7)        |                 |                |            |
| Presence of dactylitis   | 8 (13.3)         |                 |                |            |
| Tender joint at MCP2     | 8 (13.3)         |                 |                |            |
| Tender joint at MCP3     | 4 (6.6)          |                 |                |            |
| Swollen joint at MCP2    | 3 (5.0)          |                 |                |            |
| Swollen joint at MCP3    | 2 (3.3)          |                 |                |            |
| CRP (mg/L)               | 4.4 ± 4.9        | 4.0 ± 5.2       | −0.4 ± 5.7     | 0.629      |
| ESR (mm/h)               | 190 ± 15.9       | 275.1 ± 17.4    | 87.1 ± 13.3    | 0.000      |
| DAPSA                    | 11.4 ± 9.3       | 9.7 ± 5.9       | −1.7 ± 9.0     | 0.150      |
| PASI                     | 6.3 ± 8.6        | 4.7 ± 6.4       | −1.6 ± 7.2     | 0.088      |
| HAQ                      | 0.3 ± 0.4        | 0.4 ± 0.5       | 0.1 ± 0.4      | 0.067      |
| MASES                    | 0.9 ± 2.4        | 0.5 ± 1.4       | −0.5 ± 1.8     | 0.035      |
| VAS pain (mm)            | 31.1 ± 26.1      | 31.1 ± 24.5     | 0 ± 26.4       | 1.000      |
| VAS PhyGA (mm)           | 23.8 ± 18.9      | 18.5 ± 18.2     | −5.4 ± 21.8    | 0.060      |
| VAS PatGA (mm)           | 41.0 ± 28.3      | 37.0 ± 22.9     | −4.0 ± 26.0    | 0.243      |
| Tender joint count       | 2.9 ± 5.7        | 1.8 ± 2.8       | −1.2 ± 5.4     | 0.103      |
| Swollen joint count      | 0.8 ± 1.5        | 0.7 ± 1.6       | −0.1 ± 1.7     | 0.597      |
| Deformed joint count     | 3.8 ± 6.4        | 7.0 ± 7.4       | 3.2 ± 5.5      | 0.000      |
| **Current/ever treatment** |                  |                 |                |            |
| NSAIDs, N (%)            | 25 (41.7)        | 28 (46.7)       |                | 0.080      |
| Corticosteroids, N (%)   | 3 (5.0)          | 4 (6.7)         |                | 1.000      |
| csDMARDs, N (%)          | 44 (73.3)        | 49 (81.7)       |                | 0.274      |
| bDMARDs, N (%)           | 16 (26.7)        | 23 (35.3)       |                | 0.172      |
| Any DMARDs, N (%)        | 46 (76.7)        | 53 (88.3)       |                | 0.093      |
| csDMARDs + bDMARDs, N (%)| 16 (26.7)        | 19 (31.7)       |                | 0.547      |
| **Clinical response**    |                  |                 |                |            |
| VLDA, N (%)              | 4 (6.7)          | 8 (13.3)        |                | 0.083      |
| MDA, N (%)               | 15 (25.0)        | 22 (36.7)       |                | 0.166      |
| REM (DAPSA ≤4), N (%)    | 14 (23.3)        | 13 (21.7)       |                | 0.044      |
| LDA (DAPSA ≤14), N (%)   | 26 (43.3)        | 29 (48.3)       |                |            |
| MDA (DAPSA ≤28), N (%)   | 17 (28.3)        | 18 (30.0)       |                |            |
years, including one patient who switched from ustekinumab to secukinumab. No patient received bone-active drugs including denosumab.

Twenty-six (43%) patients achieved sustained DAPSA-LDA (SDL), defined as achieving DAPSA-LDA at baseline and at 5 years) while 9 (15%) patients achieved sustained minimal disease activity (sMDA), defined as achieving MDA at baseline and at 5 years.

Change in bone erosion and enthesiophyte after 5 years

There were 108 erosions and 99 enthesiophytes at baseline. After 5 years, three new erosions but no new enthesiophytes were evident (Table 2). There was a significant increase in the volume of individual erosions (0.6 ± 1.5 mm³, \( P < 0.001 \)) (Additional file 1: Figure S2A) and enthesiophytes (0.5 ± 0.8 mm³, \( P < 0.001 \)) (Additional file 1: Figure S2B).

Probability plots showed a clear shift to progression of erosion volume (>0.5 mm³ increase in volume: \( N = 37/111 \) (33.3%) (Fig. 1a). The mean ± SD increase in volume of this subgroup was 1.0 ± 0.6 mm³. Enthesiophyte volume was stable in 38/99 (38.4%), with a change in size of 0.1 ± 0.1 mm³. Only a few enthesiophytes showed evidence of regression (>0.3 mm³ decrease in volume: \( N = 11/99 \) (11.1%), with a change in size of −0.7 ± 0.4 mm³.

Erosions or enthesiophytes that showed progression have a significantly larger baseline volume compared to those lesions which showed no progression (either stable or regression) (Additional file 1: Figure S3).

Effects of achieving sustained DAPSA-LDA

The change in erosion volume was similar in patients who achieved sustained DAPSA-LDA (SDL group, \( n = 26 \)) and those who did not achieve sustained DAPSA-LDA (non-SDL group, \( n = 34 \)) (Table 3, Additional file 1:

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**Table 1** Changes in demographic and clinical characteristics over 5-year (Continued)

| Baseline (n = 60) | 5 years (n = 60) | Change (n = 60) | \( P \) value |
|------------------|----------------|----------------|-------------|
|                  |                 |                |             |
| HDA (DAPSA >28), N (%) | 3 (5.0) | 0 (0) |            |

Results are mean ± SD or number (percentage) unless otherwise indicated. Significant results are highlighted in italics.

BMI body mass index, MCP metacarpophalangeal joint, CRP C-reactive protein, ESR erythrocyte sedimentation rate, DAPSA Disease Activity in Psoriatic Arthritis, PASI Psoriasis Area and Severity Index, HAQ Health Assessment Questionnaire, VAS Visual Analogue Scale, PhyGA Physician Global Assessment, PatGA Patient Global Assessment, NSAIDs non-steroidal anti-inflammatory drugs, csDMARDs conventional synthetic disease-modifying anti-rheumatic drugs, bDMARDs biologic disease-modifying anti-rheumatic drugs, VLDAs very low disease activity, MDA minimal disease activity, REM remission, LDA low disease activity, MDA moderate disease activity, HDA high disease activity.

**Table 2** Number and volume of bone erosion and enthesiophyte

| Number                      | Baseline (n = 60) | 5 years (n = 60) | Change (n = 60) | \( P \) value |
|-----------------------------|------------------|----------------|----------------|-------------|
| Erosions total, N            | 108              | 111            | 3              |             |
| Metacarpal head 2, N (%)     | 49 (45.4)        | 50 (45.1)      | 1              |             |
| Metacarpal head 3, N (%)     | 59 (54.6)        | 61 (54.9)      | 2              |             |
| Enthesiophytes total, N      | 99               | 99             | 0              |             |
| Metacarpal head 2, N (%)     | 58 (58.6)        | 58 (58.6)      | 0              |             |
| Metacarpal head 3, N (%)     | 41 (41.4)        | 41 (41.4)      | 0              |             |

Volume of each individual lesion

| Erosions                  | Baseline (n = 60) | 5 years (n = 60) | Change (n = 60) | \( P \) value |
|---------------------------|------------------|----------------|----------------|-------------|
| Metacarpal head 2 (mm³)   | 3.6 ± 4.6        | 4.0 ± 4.8      | 0.4 ± 1.5      | 0.050       |
| Metacarpal head 3 (mm³)   | 4.8 ± 3.7        | 5.5 ± 4.3      | 0.7 ± 1.5      | 0.001       |
| Metacarpal heads 2 and 3 (mm³) | 4.2 ± 4.2 | 4.8 ± 4.6 | 0.6 ± 1.5 | <0.001      |

| Enthesiophytes            | Baseline (n = 60) | 5 years (n = 60) | Change (n = 60) | \( P \) value |
|---------------------------|------------------|----------------|----------------|-------------|
| Metacarpal head 2 (mm³)   | 3.3 ± 2.2        | 3.7 ± 2.4      | 0.4 ± 0.7      | <0.001      |
| Metacarpal head 3 (mm³)   | 3.5 ± 2.5        | 4.0 ± 2.8      | 0.5 ± 0.9      | <0.001      |
| Metacarpal heads 2 and 3 (mm³) | 3.4 ± 2.3 | 3.9 ± 2.6 | 0.5 ± 0.8 | <0.001      |

Results are mean ± SD or number (percentage). Significant results are highlighted in italics.
However, a significantly lower proportion of erosions progressed in the SDL group compared with the non-SDL group (12/51 [23.5%] vs 25/60 [41.7%], $P = 0.047$) (Table 3).

Similarly, increase in enthesiophyte volume in the SDL group was significantly lower than that in the non-SDL group (mean ± SD change: 0.3 ± 0.7 vs 0.6 ± 0.8, $P = 0.048$) (Table 3, Additional file 1: Figure S4-B). Although a numerically lower proportion of enthesiophytes progressed in the SDL group compared to the non-SDL group (17/40 [42.5%] vs 33/59 [55.9%], $P = 0.153$), the difference did not reach statistical significance (Table 3).

Using the generalized mixed linear model, only sustained DAPSA-LDA is negatively associated with the change in total enthesiophyte volume after adjusting for the use of TNFi and other baseline parameters (Additional file 1: Table S1).

Effects of long-term anti-TNF therapy
The change in size of erosions and enthesiophyses was similar in the long-term TNFi ($n = 14$) and non-TNFi ($n = 46$) patient groups (Table 3, Additional file 1: Figure S5). A similar proportion of erosions and enthesiophyses progressed in the TNFi and non-TNFi groups (Table 3). These results were unchanged after excluding the seven patients who received <5 years of TNFi and other bDMARDs in the non-TNFi group (results not shown).
Discussion

To the best of our knowledge, no previous longitudinal HR-pQCT studies have reported the long-term change in bone erosion and enthesiophyte in PsA patients receiving routine care. We observed progressive increase in bone erosion and enthesiophyte volume in PsA patients, which might be prevented by sustained control of disease activity. Unfortunately, we did not have the longitudinal data of a healthy control group for comparison. One cross-sectional study revealed a 4% increase in erosion and osteophyte count, as well as erosion volume per year in the healthy subjects [6], which was much greater than the rate of change observed in the present study. Thus, how the longitudinal structural bone changes in PsA differ from the usual age-related trajectory in heathy subjects [6] needs further investigation.

Radiography is currently the main modality used to monitor structural damage in peripheral spondyloarthritids (SpA) including PsA [23], though commonly used scoring systems do not include bony proliferation as a criterion [24–27]. Although 91 to 98% of PsA patients across all trial groups met the radiographic criteria for non-progression in recent studies [28], our data demonstrated that structural bone damage accrual continued despite conventional treatment in terms of bone erosion and enthesiophyte. More effective treatments and strategies will need to be developed to prevent progression of bony damage.

More importantly, it provides preliminary evidence that long-term control of disease activity (achieving sustained DAPSA-LDA) using various combinations of NSAIDs, csDMARDs, and bDMARDs may be useful in preventing the progression of enthesiophytes and bone erosions. Our findings concurred with previous studies reporting significantly greater radiographic progression in subjects with a moderate or poor outcome utilizing different composite disease activity measures including DAPSA over a period of 6 months [29]. Similar to other studies in newly diagnosed PsA patients [30], the proportion of PsA patients who achieved DAPSA-LDA was much higher than MDA after 5-year routine care. Higher DAPSA scores at 6 months were significantly and independently associated with probability of radiographic structural progression over a period of 1 year after adjusting for baseline radiographic score and treatment received [15]. Our findings add significant new information about the importance of sustained control of PsA disease activity, not only for improvement in pain and function but also for prevention of anabolic and catabolic structural joint damage.

Our data failed to detect a significant difference between patients with and without TNFi treatment throughout 5 years in terms of bone erosion and enthesiophyte, which confirmed previous findings that TNFi cannot inhibit enthesiophyte formation in PsA [12]. Inflammatory cytokines including TNF-α, IL-17, IL-23, and IL-22 can increase osteogenic differentiation [31]. Nonetheless, the overall impact of these cytokines on pathologic bone formation is not well understood [31]. For example, IL-22 enhances while a combination of TNF and interferon-γ (IFN-γ) with IL-22 suppresses osteogenic differentiation in human mesenchymal stem cells [32]. One can appreciate that IL-17 and TNF-α can inhibit or promote bone formation in a context-dependent manner. Therefore, it is not surprising that TNFi does not inhibit enthesiophyte formation in PsA as seen in the current study. Indeed, prospective studies have shown progression of spinal syndesmophyte formation over 2 years in patients with ankylosing spondylitis (AS) despite ongoing TNFi therapy [33, 34]. That said, recent retrospective analyses have suggested that long-term TNFi therapy can retard radiographic progression [35–37], while a prospective cohort of axial SpA patients also demonstrated a reduced likelihood of new
syndesmophyte formation with TNFi use [38]. Further studies will elucidate whether longer term use (>5 years) or earlier use (disease duration <10 years) of TNFi will be effective in preventing enthesiophyte formation in PsA, similar to what has been shown in the axial skeleton.

Receptor activator of nuclear factor-kappa B (RANK) ligand (RANKL), in the presence of macrophage colony-stimulating factor, is a major factor involved in osteoclastogenesis. TNF-α can amplify osteoclastogenesis in the presence of RANKL as well as having a direct effect on osteoclast formation when RANKL is not present [31]. Given the known capacity of TNF-α to promote osteoclastogenesis [39], one would expect suppression of this cytokine may reduce structural bone damage. A previous study reported no change in the width of erosion in both MTX- and TNFi-treated groups after 1 year, while the depth of the erosion even decreased slightly in the latter group [12]. In the current study, the increase in erosion size was numerically less in the TNFi group compared with the non-TNFi group, although statistically insignificant, probably due to the small sample size and long disease duration. Structural bone changes are generally more active in early disease [40]; therefore, the effects of anti-TNF in preventing erosion progression may be less obvious in our group of patients with long-standing disease compared to the cohort with shorter disease duration [12]. Retrospective studies have demonstrated that patients with erosive PsA receiving TNFi had a better radiographic outcome compared to those treated with MTX [41]. A recent randomized trial showed that etanercept-containing arms showed less radiographic progression compared with MTX monotherapy at week 48 [42]. Future studies with a larger sample size will be required to confirm whether TNFi is superior to MTX in preventing long-term erosion progression in PsA using HR-pQCT. Six-month treatment of secukinumab prevented erosion and enthesiophyte progression on HR-pQCT and reduced disease activity evident on magnetic resonance imaging in PsA patients [11]. We were not able to assess the effects of secukinumab as only two patients received this treatment, and the majority of our patients were treated with csDMARDs and NSAIDs.

This is the first long-term follow-up study in a PsA cohort to study the progression of catabolic and anabolic bone changes using HR-pQCT. We perform baseline-indexed image registration [43] and slice matching to acquire precisely matched baseline and follow-up VOI [44] to ensure reliability. In addition, we assessed the change in enthesiophyte volume instead of measuring maximal height [12] or using semi-quantitative grading methods [11, 17], which should provide a more objective assessment.

One limitation of this study is that validated disease activity assessments, e.g., DAPSA, were not routinely performed except at baseline and 5 years. Future longitudinal studies with serial disease activity assessments are needed to provide further information on the role of cumulative inflammatory burden and the development of bone damage in PsA. We also did not investigate minimal clinical meaningful volume change of bone erosion or enthesiophyte, especially for hand function. We acknowledged that the number of TNF-treated patients might not be sufficient to reach a solid conclusion. Thus, a well-designed long-term cohort study with a large sample size should be used to further address this question. Moreover, we did not repeat x-rays of all the hands and feet at 5 years, which might be useful to correlate radiologic new bone formation with HR-pQCT findings in future studies. Last but not least, we did not detect any differences in the risk of bone damage progression in patients with and without tender or swollen joints at baseline. Imaging modalities including magnetic resonance imaging (MRI) or ultrasonography would have been useful to objectively document and quantify the burden of inflammation. A multi-modality imaging study would be important to address whether synovitis confirmed by MRI or ultrasound at baseline may predict the development of bone erosion and enthesiophyte on HR-pQCT in the future.

Conclusions
Damage accrual in terms of bone erosion and enthesiophyte was observed in PsA patients over a period of 5 years despite receiving routine clinical care. Nonetheless, sustained control of disease activity may be able to prevent these bony damages. Future larger long-term HR-pQCT studies are needed to validate these initial observations.

Supplementary information
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Additional file 1: Figure S1. Representative image of baseline indexed image registration and slice matching. Figure S2. Examples of bone erosion and enthesiophyte in psoriatic arthritis (PsA) patients at baseline and 5-year follow-up. Figure S3. Baseline volume and dynamics of bone erosion and enthesiophyte, Figure S4. Mean within-subject volume changes in bone erosion and enthesiophyte between patients who achieved and did not achieve sustained DAPSA-LDA. Figure S5. Mean within-subject volume changes in bone erosion and enthesiophyte between patients with and without TNFi throughout 5-year. Table S1. Generalized mixed linear model with change in bone erosion and enthesiophyte volume as dependent variable.

Abbreviations
bDMARDS: Biologic disease-modifying anti-rheumatic drugs; BMI: Body mass index; CASPAR: ClASsification criteria for Psoriatic ARthritis; CRP: C-reactive protein; csDMARDs: Conventional synthetic disease-modifying anti-rheumatic drugs; DAPSA: Disease Activity index for PsA; ESR: Erythrocyte sedimentation...
rate; HAQ: Health Assessment Questionnaire; HR-pQCT: High-resolution peripheral quantitative CT; IL: Interleukin; LDA: Low disease activity; MASES: Maastricht Ankylosing Spondylitis Enthesitis Score; MCH: Metacarpal heads; MDA: Minimal disease activity; NSAIDs: Non-steroidal anti-inflammatory drugs; PASI: Psoriasis Area and Severity Index; PsA: Psoriatic arthritis; PsO: Psoriasis; SDC: Smallest detectable change; TNF: Tumor necrosis factor; VLDA: Very low disease activity

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Authors’ contributions
LST had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. DZW, JFG, SHML, PCHW, LS, EKL, IT, TKL, VWH, LQ, and LST collected the data. DZW, JFG, SHML, LS, VWH, LQ, and LST analyzed and interpreted the data. DZW, PCHW, JFG, LQ, and LST prepared and revised the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
The study protocol (CRE-2016.366) was approved by the Ethics Committee of The Chinese University of Hong Kong-New Territories East Cluster Hospitals. Written informed consent was obtained from all subjects prior to participate in the study.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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