Comparison of clinical, radionuclide, and radiographic features of osteoarthritis of the hands

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Abstract
Simultaneous clinical, scintigraphic, and macroradiographic assessments were carried out on 32 patients with hand osteoarthritis and the results at entry and one year reported. The presence and growth of osteophyte correlated with symptoms and a positive scan. The scan did not detect the radiographic features of juxta-articular radioactivities, subchondral sclerosis, or cartilage thinning. Osteophytes, particularly when fast growing, produce pain, a 'hot' scan, and may predict disintegration of joint architecture.

Conventional radiography is used to identify the features of osteoarthritis of the hands, but the disease is well established at this stage and it is not known which features represent disease and which repair. Long term studies of progression are hampered by difficulty in detecting early cases, which may be asymptomatic, the slow disease process, and the poor resolution of x ray features. Isotope bone scanning is a sensitive technique and can detect early changes in bone, blood flow and metabolism and in osteoarthritis gives a distinctive, localised, well defined hot spot in the typical distribution of thumb base and distal interphalangeal joints, but also in the wrist and metacarpophalangeal joints. A study comparing scintigraphy with conventional radiography suggested that the scan may detect early or active disease, which may either resolve or progress to irreversible damage. The study was limited because the highly sensitive but non-specific scan image was compared with a poor resolution standard anteroposterior hand radiograph in which the anatomical changes could only be graded on a four point severity scale.

The technique of microfocal radiography uses a small source of x rays (6-8 μm) and produces high magnification, high resolution macro-radiographs, from which accurate measurements of individual x ray features can be made at each joint in the hand. It also permits the detection of change over a reasonably short time span. We compared measurements from bone scan imaging, clinical observation, and microfocal radiography and applied statistical analysis in an effort to define the pathological processes taking place in osteoarthritis.

CLINICAL ASSESSMENT
Patients were examined at six monthly intervals by a single observer for 18 months, making four visits in all. Clinical assessment and microfocal radiography were carried out at each visit and on the same day, and bone scanning was added at the first and third visits. Assessment was made of tenderness over the distal interphalangeal, proximal interphalangeal, metacarpophalangeal, and carpometacarpal joints of the thumb and the radial and ulnar portions of the wrist by direct pressure and graded on a four point severity scale: 0=no pain; 1=tender, 2=tender and wince, 3=wince and withdraw. Heberden's and Bouchard's nodes were assessed by observation and palpation on the radial and ulnar sides of each finger and scored for number and for size on a five point scale: 0=absent, 1= palpable, 2=small, 3=medium, 4=large.

MICROFOCAL RADIOGRAPHY
Stereopair macroradiographs were prepared of the right and left hands and wrists of each patient, excluding the terminal interphalangeal joint of the thumb, which would have required an additional x ray plate. The hand was placed in a stereotactic unit, positioned close to the source and displaced by 6 mm between the first and second x ray exposure. The stereopair macroradiographs thus obtained were examined under a large format stereoscope (Ross Instruments, Salisbury, UK) for three dimensional evaluation of the joint structures. The right
sided, back illuminated image of the stereoscope contained a digitiser tablet linked to a MOF-videoplan (Carl Zeiss, Hertfordshire, UK). A cross wire cursor was used to outline the structure in the macroradiograph overlying the digitiser. The data initially recorded on microcomputer disks were transferred onto one large disk on an IBM PC/AT using a standard format for each patient/hand/visit and recorded so that comparisons could be made of data both within and between patients.

The following x ray features were measured:

(1) Joint space width: the interbone distance representing the space occupied by opposing articular cartilage. Where this was reduced, measurements were made across the narrowest part of the joint if it was uneven and across the midline when it was even.

(2) Subchondral sclerosis: the thickness in millimetres of the subchondral cortex was measured at three equidistant points along the margin of each joint and the readings averaged.

(3) Osteophytes: the number and area in square millimetres were measured at the articular margins and capsular insertions on the radial and ulnar sides of the proximal and distal ends of each joint.

(4) Juxta-articular radiolucencies: these radiological features of osteoarthritis have been referred to by Altman et al as erosions. The number and area in square millimetres were measured using a program developed for the evaluation of erosions in rheumatoid arthritis.

BONE SCAN
A four hour scintigraphic image was obtained of the hands after injection of 555 MBq of technetium-99m methylene diphenosphonate by acquiring counts for 15 minutes with a gamma-camera. The films were read at one sitting and were compared rather than read blind. The intensity of the image from each joint was assessed by one observer and scored: 1=suspicious, 2=positive, 3=strongly positive, and 4=intense. The wrist was considered as two units: radial and ulnar.

ANALYSIS
For each joint in the hand and the two wrist units the bone scan, x ray features, and clinical features were collated and the distribution of each throughout the hand was determined. Correlations between features were calculated using Kendall’s tau C, and changes between visits were tested for significance using Wilcoxon’s signed ranks test for imaging data and Wilcoxon’s signed rank test for clinical data. All statistical evaluations were carried out on an IBM PC/AT using SPSS/PC software, and significance was set at p=0.05.

Results

CLINICAL ASSESSMENT
There was a significant increase in the number of interphalangeal nodes over the 18 months from a mean (SD) for each patient of 3.4 (4.5) to 7.5 (5.4) (p=0.0014). The mean (SD) size for each patient increased from 9.6 (12) to 21 (17.4) (p=0.002). The mean (SD) number of tender joints per patient at entry was 7.7 (6.9). The mean severity for the tender joints was 9.4 (8.4). Neither changed significantly.

MICROFOCAL RADIOGRAPHY
A detailed description of the macroradiographic features and progression of hand osteoarthritis has been reported elsewhere, and the principal findings are summarised below.

Subchondral sclerosis At baseline all patients with osteoarthritis had significantly increased subchondral cortical thickness in the right (p=0.003) and left (p=0.002) wrist and hand. By 18 months the subchondral cortical thickness had increased significantly in the patients with osteoarthritis as a group (p=0.006). Three groups were identified on the basis of their radiographic progression: 19 (60%) were considered to have progressed because each patient showed an increase in subchondral sclerosis greater than the coefficient of variation of 4%. Two patients (6%) showed no change in subchondral cortical thickness, whereas the remaining 11 patients (34%) showed a significant decrease (p<0.005).

Osteophytes were noted at baseline visit in the right and left hands of all patients compared with controls. During the 18 month period osteophytes in both hands increased significantly in both number (p<0.005) and area (p<0.005).

Juxta-articular radiolucencies were seen in the bones of the wrist and finger joints of all patients with osteoarthritis compared with controls. During the study period these radiolucencies showed no significant change in number but increased significantly in size (p=0.002).

Joint space width Two main groups were identified from the baseline measurements: 18 (56%) patients with osteoarthritis had a joint space slightly but not significantly wider than the non-arthritic subjects in the right and left wrists. The remaining 14 (44%) patients had a joint space significantly narrower than the non-arthritic controls. These two groups were further subdivided into four groups (A, B, C, and D) on the basis of radiographic progression: A, 7 (22%) were normal at entry and remained unchanged; B, 11 (34%) normal at entry and narrowed; C, 10 (31%) narrow at entry but did not change; D, 4 (13%) narrow at entry and narrowed further.

BONE SCAN
All patients had one or more joints which scored one or more on the scan. At entry the mean (SD) number for each patient was 14.5 (5.8) and the mean (SD) intensity score was 34.3 (19.4). There was no significant change in these means at 12 months.

Comparison of the distribution of joints with a positive scan in the hands at entry with that at 12 months showed a similar pattern in both hands with the following joints being most commonly affected: thumb metacarpophalangeal joint (62%), index distal interphalangeal...
Comparison of assessments of osteoarthritis of the hands

Mean (standard deviation) of the changes in osteophyte number and area during 12 months grouped by change in the bone scan score. The significance of differences was calculated between the group with raised or increasing isotope uptake and the group with decreasing or no radionuclide uptake.

<table>
<thead>
<tr>
<th>Group</th>
<th>Change in number</th>
<th>Change in area (mm²)</th>
<th>Combined group change in number</th>
<th>Combined group change in area (mm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decrease in score</td>
<td>216</td>
<td>0.06 (0.08)</td>
<td>0.15 (0.74)*</td>
<td>0.007 (1.63)</td>
</tr>
<tr>
<td>Unchanged score: 0</td>
<td>349</td>
<td>0.20 (0.67)</td>
<td>0.09 (1.25)</td>
<td>0.08 (1.32)</td>
</tr>
<tr>
<td>Unchanged score: 1-4</td>
<td>188</td>
<td>0.14 (1.01)</td>
<td>0.22 (1.03)*</td>
<td>0.45 (3.59)</td>
</tr>
<tr>
<td>Increase in score</td>
<td>109</td>
<td>0.35 (1.06)</td>
<td>0.62 (2.25)</td>
<td>0.51 (2.91)*</td>
</tr>
</tbody>
</table>

*Not significant; p=0.005 (Mann-Whitney U test).

joint (59%), thumb distal interphalangeal joint (59%), carpometacarpal joint (59%), and middle distal interphalangeal joint (59%). There were slightly fewer joints with a positive scan in the left hand (216 compared with 238). Also, the severity score was lower in the left hand (517 compared with 554). The wrist showed considerable scintigraphic activity, with 16 (50%) patients having a positive scan in the ulnar compartment and 10 (31%) in the radial compartment.

Statistical analysis showed the following significant correlations at the first visit: scan and area of osteophytes (p=0.003 for the right hand and p=0.0013 for the left hand); scan and joint tenderness (p=0.0037 for the right hand and p=0.0004 for the left hand); number of radiological osteophytes and nodes (p=0.0004 for the right hand and 0.0015 for the left). Correlation between joint tenderness and nodes was highly significant in the left hand (p=0.0005) but just failed to reach significance in the right (p=0.059).

In 13 (41%) patients there was a change in the number of joints with a positive scan between the first and second visit. The table shows the mean change in osteophyte number and area during the one year period grouped by change in bone scan. At joints where there was either a decreased or negative score only a very small increase in osteophyte was detected. At joints where there was either a positive but unchanged or an increased score there was a significant increase in osteophyte size but not in number. No correlation was found between the bone scan and the extent or the change in the dimensions of subchondral sclerosis, joint space width, and juxta-articular radiolucenties.

Discussion

Osteoarthritis is not a disease in the conventional sense but rather a process of joint disintegration with many causes, characterised by cartilage loss and reactive new bone formation at joint margins with associated variable degrees of synovitis and fibrous thickening of the joint capsule. In the large weightbearing joints, such as the hip and knee, abnormal development and trauma are implicated as causes but the cause of osteoarthritis of the hands is unknown and may be multifactorial with an underlying predisposition and genetic factors having an important role. Radiologically, osteoarthritis, is common so that it becomes difficult to determine what is the result of disease and what reflects normal aging, particularly when despite radiological change there is no pain or interference with joint function. Radiographs reflect the morphology of mineralised bone and show the result of repair and destruction. A scan is a study of function and depends on osteoblastic and osteoclastic activity and bone vascularity and reflects the dynamic response of bone to a variety of insults, such as trauma, inflammation, and neoplasia.

A comparison of diagnostic modalities reported that scintigraphy showed more extensive abnormality than either radiography or arthrography in osteoarthritis of the knee. In their paper comparing bone scanning with conventional radiography Hutton et al studied 33 patients with nodal osteoarthritis of the hands and showed an overall similar pattern of involvement between scanning and radiography but a marked dissimilarity in some patients between the severity on radiography compared with scanning. Some joints were found to be abnormal on one or other investigation only. They also noted that a joint with a positive scan was likely to show subsequent radiological deterioration, but they were unable to locate the site of deterioration or measure this because of the poor resolution of conventional hand radiographs.

The site of uptake of radiopharmaceuticals in bone has been studied in experimentally induced osteoarthritis in rabbits. In early disease uptake is concentrated in developing osteophytes and, later, in the subchondral bone. In human femoral heads with end stage osteoarthritis, removed at operation after injection of bone seeking isotopes, autoradiography showed uptake primarily in the subchondral bone and in the walls of cysts.

Increased thickness of subchondral bone is a major feature of osteoarthritis and was present in all our patients. It increased radiologically in 19 (60%) but we found no correlation between this feature and the scan, despite the animal models. There may be two stages of sclerosis: an early stage, which may be the response to initiating factors such as subchondral microstress fractures, and a later one, which represents the response to altered weight bearing consequent on cartilage loss and disordered alignment. In the hand neither process seems scintigraphically active, but this may be because of the difference in size of the joints—those of the hand being much smaller than the hip in man and the knee in the rabbit—or the degree of loading to which the hand joints are subjected.
Juxta-articular radiolucencies are not reported to be a characteristic feature of the radiography of osteoarthritis and their exact nature is unclear, but they may be an important early feature. The similarity of their distribution to that of rheumatoid erosions suggests that they may have an inflammatory origin. We found no correlation between juxta-articular radiolucencies, tenderness, and a positive scan, however, which makes this unlikely, and their exact origin and significance remain obscure despite the fact that they were present in all patients and increased significantly in size.

A cardinal feature of osteoarthritis is loss of articular cartilage. In the early stages disruption of the collagen matrix results in swelling as water content is increased, and this is followed by fissuring and loss of cartilage substance. In just over half of our patients we found a slightly increased joint space, suggesting that microfocal radiography could detect cartilage swelling in early disease. In the others there was already established thinning. In the subgroups described, in whom progressive cartilage loss was shown on serial radiographs, no correlation was found with the scan, so this key process in osteoarthritis seems to be scintigraphically inert, at least in the wrist and hand.

When the data were analysed according to the presence and growth of osteophytes, highly significant correlations were found with the scan and to a lesser extent with joint tenderness, though this finding must be interpreted with caution as the patients were allowed to take analgesic and anti-inflammatory drugs as required. Osteophyte formation seemed to be a key process in causing clinical symptoms. If the work of Hutton et al is correct osteophytes would also seem to be the marker for subsequent disintegration of the joint architecture as this was predicted by a positive scan and hence, as our study has shown, by the presence of actively remodelling osteophyte. This is a surprising finding in view of the previous suggestion that osteophytes may be an independent phenomenon related to age. Possibly, the rate of remodelling of osteophyte is the key: the slower the change in size, the less likely that the joint is seriously compromised. Previously scintigraphy was considered a sensitive but non-specific tool in osteoarthritis, but this study has shown that it detects very specific pathological processes of clinical significance. With the development of chondroprotective agents more studies are likely to be carried out in patients with osteoarthritis. In a disease with such a long, intermittent, and poorly described natural history, scintigraphic grading of disease activity may prove a useful way of standardising treatment groups. The role of osteophytes in the causation of symptoms in osteoarthritis in large weightbearing joints also merits further study.

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