Quantitative sacroiliac scintiscanning: a sensitive and objective method for assessing efficacy of nonsteroidal, anti-inflammatory drugs in patients with sacroiliitis

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Summary Serial computer assisted quantitative sacroiliac scintiscanning (SI joint/sacrum ratios) 3 hours after low dosage (5 mCi) 99mTc methylene diphosphonate has been used as an objective index of sacroiliitis in a single blind 14-day cross-over comparison of azapropazone 600 mg b.d. and naproxen 500 mg b.d. in 18 patients with active sacroiliitis. Clinical assessments included visual analogue scales for measurement of pain and early morning stiffness, chest expansion, a modified Schober test, and goniometric measurement of thoracolumbar spinal flexion by means of an inclinometer. Statistically significant decreases in pain (p<0.001) and early morning stiffness (p<0.001) followed treatment with each NSAID, but there was no significant difference in the fall in these parameters, although 15 out of 18 patients expressed a preference for naproxen. Chest expansion and thoracolumbar flexion were not significantly affected by either drug. Serial quantitative scintigraphy showed a mean fall in joint sacrum ratios following each treatment which was statistically significant (p<0.02) only after naproxen. Serial quantitative scintigraphy can be used as an objective method of assessing sacroiliitis and was sufficiently sensitive to reflect the patients' subjective preference in a short-term comparison of 2 NSAID.

The sensitivity of sacroiliac (SI) scintigraphy as an objective measure of sacroiliitis has been greatly increased by computer quantification of the uptake of tracers of bone seeking isotopes over the SI joints compared with that over the sacrum. The technique has been shown to be particularly sensitive in early sacroiliitis but uptake is more variable in patients with advanced disease and established radiological changes.

Treatment with nonsteroidal anti-inflammatory drugs (NSAID) can lower the joint/sacrum scan ratio (SI/S ratio) and this may be partly responsible for the considerable overlap of SI/S ratios in normal subjects and patients with sacroiliitis found by some investigators.

Following early suggestions that serial quantitative SI scintigraphy (SQSS) might be useful in assessing the short-term efficacy of NSAID in the treatment of inflammatory sacroiliitis we have used it as a method of objective assessment in a short-term cross-over trial of 2 NSAID in a group of patients with active sacroiliitis. The results have been compared with those of conventional methods of clinical assessment.

Patients and methods

Trial design

The study was a single-blind, cross-over comparison of the effect of 14 days' treatment with azapropazone 600 mg b.d. and naproxen 500 mg b.d. in patients with active sacroiliitis. Eighteen patients (16 male, 2 female) with clinically active sacroiliitis who all satisfied the New York criteria for definite ankylosing spondylitis were randomly allocated into 2 groups, A and B. After a 7-day drug-free washout period group A were treated with azapropazone 600 mg b.d. and group B with naproxen 500 mg b.d. for 14 days. After a further 7-day drug-free washout period group A patients received naproxen 500 mg b.d. and group B patients azapropazone 600 mg b.d. for 14 days. No
other NSAID were allowed during the course of the study, but patients were given a supply of paracetamol as a rescue analgesic. Excluded from the trial were pregnant women or women likely to become pregnant, patients with peptic ulcers, bowel, renal, or hepatic disease, patients with blood dyscrasia and those receiving anticoagulant therapy, as well as all patients over the age of 65 years.

ASSESSMENTS

Clinical assessments and QSS were undertaken at the beginning and end of each treatment period on days 7, 21, 28, and 42. Global pain and early morning stiffness were assessed on vertical 10 cm visual analogue scales.\textsuperscript{10} Chest expansion (cm) was measured at the 4th intercostal space.\textsuperscript{11} Flexion of the thoracolumbar spine was measured by McCrea and Wright’s modification of the Schober test\textsuperscript{12} and by goniometric measurement with the improved inclinometer.\textsuperscript{13}

Radiographs of the sacroiliac joints were graded for sacroiliitis prior to the start of the trial on a 4-point scale: (1) normal, (2) suspicious, (3) definite, (4) sclerosis/fusion.

At the end of the 6-week trial patients were asked to express any preference for either treatment and a record was made of any adverse side effects.

QUANTITATIVE SACROILIAC SCINTIGRAPHY

5 mCi (200 MBq) of \textsuperscript{99m}Tc-methylene diphosphate were injected intravenously and the patient imaged 3 hours after the dose with a small-field Ohio Nuclear series Sigma 100 gamma camera fitted with specially designed extra high sensitivity collimator (giving approximately 4 times the sensitivity of the standard collimator). A total of 500 000 counts were accumulated with the gamma camera centred on the body of the 1st sacral vertebra.

Computer drawn isocount contours of 65% outlined the whole of each sacroiliac joint area. The sacral reference area comprised the bodies of the first 3 sacral vertebrae. Within each area of interest the total number of counts and area were calculated by the computer, the result being expressed in a ratio (the SI/S ratio) derived from the expression

\[
\text{SI/S ratio} = \frac{\text{mean SI counts/pixel}}{\text{mean sacral counts/pixel}}
\]

As there was no significant difference between left and right SI/S ratios in any of the subjects, a single SI/S ratio for each scan could be derived from the mean of the left and right SI/S ratios. By calculating the SI/S ratio in this manner errors arising from variability of tracer uptake throughout the SI joint are eliminated.

Statistical analyses were performed by Student’s paired \(t\) test.

Results

Groups A and B were well matched for age, sex, and disease activity (Table 1).

Global pain and early morning stiffness decreased significantly following treatment with both azapropazone and naproxen (Fig. 1), but there was no significant difference in the fall in these parameters following either drug. Fifteen out of 18 patients, however, expressed a definite preference for naproxen. Chest expansion and thoracolumbar flexion were not significantly affected by either drug (Table 2).

Serial measurements of QSS showed a mean fall in the SI/S ratio following treatment with both azapropazone and naproxen, by the fall was statistically significant only after treatment with naproxen (Fig. 2). This was true whether the drug was given first or

<table>
<thead>
<tr>
<th>Age, sex, and disease activity in the patients studied</th>
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<tbody>
<tr>
<td>Group A</td>
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<tr>
<td>Number</td>
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<tr>
<td>Sex</td>
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<tr>
<td>Age, mean (range) years</td>
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<tr>
<td>Pain, VAS mean (± SD), cm</td>
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<td>EMS Chest expansion, mean (± SD), cm</td>
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<td>Schober test, mean (± SD), cm</td>
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<td>Inclinometer, mean (± SD), degrees</td>
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<td>Radiological sacroiliitis (grades)</td>
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<td>4</td>
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<td>SI/S ratio, mean (± SD)</td>
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VAS = visual analogue scale. EMS = early morning stiffness.


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References