A double-blind, multicentre, placebo-controlled study of tiludronate in Paget's disease of bone

William D Fraser, Trevor C Stamp, Robert A Creek, James P Sawyer, Christopher Picot

Summary
A multicentre, randomised, placebo-controlled, dose-ranging study was conducted to investigate the therapeutic activity and sustained efficacy of tiludronate (200 mg, 400 mg and 600 mg once daily) taken orally for 12 weeks in patients with Paget's disease. Serum alkaline phosphatase concentrations were compared with baseline at weeks 12 and 24; treatment success was defined as a 50% reduction compared with baseline. Changes in the hydroxyproline:creatinine ratio were also measured. Pain was assessed using the Huskisson Visual Analogue Scale and by questionnaire. Patients completing at least 11 weeks of treatment were followed-up 18 months later by postal questionnaire.

Significantly greater numbers of patients in the tiludronate groups successfully responded to treatment compared with the placebo group. A dose-response was observed; the percentage of patients responding to treatment being 31% (200 mg), 52% (400 mg) and 82% (600 mg) at week 12 and 45% (200 mg), 70% (400 mg) and 82% (600 mg) at week 24. Tiludronate treatment also significantly reduced hydroxyproline:creatinine ratios compared with placebo, again showing a dose response. Dose-related gastrointestinal symptoms were the commonest adverse events, occurring in 2.4%, 11.0%, 5.5% and 18.9% of patients receiving placebo and tiludronate 200, 400 and 600 mg daily, respectively. The response to oral tiludronate was sustained for more than 18 months in some patients and there was evidence of a reduction in the longer term complications of the disease. These results show that oral tiludronate is an effective, well-tolerated treatment for Paget's disease; the 400 mg once daily dose appears to offer the optimum balance of efficacy and tolerance.

Keywords: tiludronate, Paget's disease, dose-response relationships, osteitis deformans, bisphosphonates, alkaline phosphatase, hydroxyproline:creatinine ratio

This leads to excessive osteoblastic activity, with a variable mixture of immature 'woven' bone and irregular, mature lamellar bone being laid down. Increased bone resorption is associated with a marked increase in urinary hydroxyproline excretion whereas increased osteoblast activity is indicated by increases in serum alkaline phosphatase concentrations.

The bone changes associated with Paget's disease are asymptomatic in the majority of patients but complications such as fractures (11%) or deformity (17%) can occur. Pain, the main symptom of Paget's disease, can arise from the Pagetic bone itself or from complications, especially osteoarthritis, secondary to Paget's disease.

Two main classes of drug therapy are currently available for the treatment of Paget's disease. The calcitonins are moderately successful in suppressing disease activity, reducing urinary hydroxyproline and serum alkaline phosphatase concentrations by approximately 55% during treatment. However, relapse occurs rapidly when calcitonin is stopped and approximately 20% of patients become resistant to ongoing treatment. Another disadvantage of calcitonin treatment is the need for parenteral injection, although intranasal preparations are being developed. In addition, adverse gastrointestinal side effects and flushing can be troublesome.

A second class of drugs, the bisphosphonates, inhibit osteoclastic resorption and thus reduce bone turnover. They have demonstrated efficacy in Paget's disease with a dose-dependent response. Disodium etidronate is widely used for the treatment of Paget's disease although its value is compromised by dose-dependent inhibition of bone mineralisation. Tiludronate (chloro-4-phenyl thiomethylene bisphosphonate; SR 41319B) is a second generation bisphosphonate with experimental anti-osteoclastic activity. In an earlier double-blind clinical study, 800 mg/day of a capsule formulation proved the optimum dose in patients with Paget's disease. A subsequent tablet formulation of tiludronate, with two to three times greater bioavailability, has been shown, in an open study using 400 mg/day, to effect similar reductions in serum alkaline phosphatase and urinary hydroxyproline (Reginster et al, unpublished observations). We therefore sought to confirm the efficacy and optimum dose of tiludronate tablets in patients with Paget's disease of bone in a placebo-controlled, double-blind, randomised, multicentre dose-ranging study.

Paget's disease is common in the UK, affecting approximately 4.6% of the population over 55 years of age, and represents a significant cause of morbidity. Paget's disease of bone is marked by dramatic increases in the number of osteoclasts, resulting in increased, chaotic, bone resorption.
Tiludronate in Paget's disease of bone

Patients and methods

PATIENT POPULATION
A total of 112 patients, of either sex (age ≥18 years) with diagnosis of Paget's disease confirmed by scintigraphy (63%) and/or radiography (92%) entered the study. Serum alkaline phosphatase concentration had to be at least twice the upper limit of normal at the local laboratory. Patients were ineligible for study entry if they fulfilled any of the following criteria: prior treatment with bisphosphonates other than etidronate within the past two years; treatment with etidronate, mithramycin or calcitonin within six months; prior treatment at any time if the current serum alkaline phosphatase concentration was less than 30% above the lowest concentration then achieved; recent fracture or confinement to bed; current peptic ulceration; clinically significant hepatic, renal or haematological disorder; pre-menopausal women; malignant neoplasia within the previous five years or breast malignancy within the previous 10 years; recent change in dose of hormone replacement therapy, vitamin D or corticosteroids. Withdrawal from the trial was permissible at any time.

STUDY DESIGN
The double-blind, randomised study consisted of four parallel treatment groups receiving a single dose of placebo, or oral tiludronate (200 mg, 400 mg or 600 mg/day) for 12 weeks. Patients were recruited from 16 hospitals throughout the UK. A follow-up of patients who had completed at least 11 weeks of treatment was carried out 18 months after the completion of the last patient in the trial using a postal questionnaire. This assessed the need for, and time to, first retreatment for Paget's disease. The number of patients suffering from complications of Paget's disease (defined as any orthopaedic (including fractures), neurological (including pain), cardiovascular or sarcomatous event) were assessed. Local laboratory serum alkaline phosphatase concentrations from routine out-patient visits since the end of the main trial were also recorded.

All patients gave written, informed consent and each centre obtained written approval from local ethics committees. The study was conducted in accordance with the Declaration of Helsinki as revised in 1989.

Patients were allocated sequential study numbers and randomly assigned to one of the four treatment groups. Patients attended a total of six visits: at baseline (week 0), after two, four, eight and 12 weeks treatment, and 24 weeks after baseline. Medication was taken in the morning with water, at least two hours before or after taking food.

ASSESSMENTS
Blood samples were taken at every visit and analysed independently at a single central laboratory, using automated techniques for all routine haematology and biochemistry (including serum calcium and phosphate) tests. Serum alkaline phosphatase concentrations were the primary endpoint for efficacy. Fasting urine samples were collected over a period of at least two hours after an overnight void. Hydroxyproline concentrations were measured by a cation exchange resin system and quantitative spectrophotometric colorimetry.

Medical histories and a full clinical examination were completed at the first visit. A 12-lead electroencephalogram was performed at weeks 0 and 12. Compliance was assessed by tablet counts and concomitant medication and adverse events were recorded in detail throughout the study.

Pain was a secondary efficacy criterion. A visual analogue scale was used for patients to assess their Pagetic pain. The scale ranged from 'no pain' to 'agonising pain'. In addition, at each visit, symptomatic bone pain was assessed by the physician as 'none', 'some' or 'constant', with its severity at rest and on movement assessed as 'none', 'mild', 'moderate', or 'severe'.

STATISTICAL ANALYSIS
The study was intended to have an 80% power of detecting a true difference of 40% in success rates between any two treatment groups (α=0.05, two-sided), with a total of at least 100 patients. Data were analysed on an 'intention to treat' basis. A successful response was defined as at least a 50% reduction in serum alkaline phosphatase concentration after 12 weeks treatment; patients whose serum alkaline phosphatase concentrations decreased by 25% or less compared with week 0 were defined as resistant.

Between-group comparisons of the proportion of patients with a successful response and those resistant to treatment were made at weeks 12 and 24. Two by two contingency tables of each pair-wise comparison were analysed by Fisher's exact test and the p-values were subsequently corrected by the Sidak method for multiple testing.

Serum alkaline phosphatase concentrations and hydroxyproline:creatinine ratios were compared at weeks 12 and 24 by analysis of variance (ANOVA). Data were log transformed and presented as adjusted geometric mean ratios to allow for centre and baseline variability. Multiple comparisons of the least square means using the Sidak t-test were presented as Sidak-corrected p-values and simultaneous 95% confidence intervals. The associations between the gastrointestinal adverse events and concomitant use of nonsteroidal anti-inflammatory or a history of gastric disorder were analysed using the Breslow-Day test.

Data from the follow-up questionnaire were analysed using the Chi-squared statistic unless all the questions were not answered in which case a Fisher's exact test was used. All the serum alkaline phosphatase samples collected during the follow-up period were analysed locally. To allow comparison, these data were converted to a percentage of the mid point of the appropriate reference range.
Results

Patient characteristics and randomisation at entry are given in table 1. The most prevalent lesion sites at recruitment were the pelvis (75% of patients) and the lumbar spine (39%); 30% of patients had disease at one site only and 50% of patients had received no previous therapy for Paget’s disease. Seven patients were withdrawn for adverse events (one on placebo, three on 200 mg, one on 400 mg and two on 600 mg), two died (both on 200 mg), five were withdrawn for protocol violation (three on placebo, one each on 400 mg and 600 mg), and four were lost to follow-up (two on placebo, one each on 200 mg and 600 mg). Three patients, all on placebo, received alternative treatment for Paget’s disease during the initial 12 week follow-up period. Patient characteristics at entry were similar in each group (table 1), although known duration of disease was longer in the 400-mg dose group. The mean compliance calculated from tablet counts was >96% in each treatment group.

Efficacy

No changes in efficacy parameters were observed in patients treated with placebo. The percentage of patients achieving a ≥50% reduction in serum alkaline phosphatase concentrations on each treatment is given in table 2. All tiludronate doses were significantly more effective than placebo at weeks 12 and 24 (p<0.0001 except for the 200 mg dose at week 12 (p<0.05)). The tiludronate response was dose-related and increased over time (figure 1). Serum alkaline phosphatase levels fell significantly more with 600 mg than with 200 mg at week 12 (p<0.001; 95% confidence intervals CI 0.524–0.837) and at week 24 (p<0.001; 95% CI 0.463–0.832), and more than 400 mg at week 12 (p=0.043; 95% CI 0.630–0.995). The reduction in serum alkaline phosphatase was evident from two weeks and continued in the 12 weeks after treatment ended. There was no evidence to suggest that efficacy differed in patients who had received previous bisphosphonate treatment (p>0.20; type III sum of squares). Furthermore, any differences in efficacy between treatment groups were found to be unrelated to the duration of Paget’s disease (p>0.23; type III sum of squares).

Normal serum alkaline phosphatase levels (<115 IU/l) were achieved in a significantly higher percentage of 200 mg (31%), 400 mg (39%) and 600 mg (44%) recipients than placebo recipients (0%) at 24 weeks (p<0.05). Resistance to treatment by week 24 was 91% for placebo, 8% for tiludronate 200 mg and 4% for both tiludronate 400 mg and 600 mg groups (p<0.05 for all group comparisons to placebo).

Changes in the geometric mean hydroxyproline:creatinine ratio over time for each treatment are illustrated in figure 2.

Table 1 Patient characteristics at entry

<table>
<thead>
<tr>
<th></th>
<th>Placebo (n=26)</th>
<th>200 mg (n=29)</th>
<th>400 mg (n=29)</th>
<th>600 mg (n=28)</th>
<th>Total (n=112)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (M/F)</td>
<td>12/14</td>
<td>12/17</td>
<td>18/11</td>
<td>18/10</td>
<td>60/52</td>
</tr>
<tr>
<td>Race (Caucasian/Negroid)</td>
<td>26/0</td>
<td>29/0</td>
<td>29/0</td>
<td>27/1</td>
<td>111/1</td>
</tr>
<tr>
<td>Mean age (years) (SD)</td>
<td>69.9 (7.3)</td>
<td>70 (7.8)</td>
<td>72 (7.6)</td>
<td>68 (7.6)</td>
<td>70 (8.1)</td>
</tr>
<tr>
<td>Mean time since diagnosis of Paget's disease (years) (SD)</td>
<td>8.5 (7.0)</td>
<td>7.1 (9.0)</td>
<td>11.1 (9.6)</td>
<td>5.3 (8.0)</td>
<td>8.1 (8.6)*</td>
</tr>
<tr>
<td>Alkaline phosphatase** (IU/l)</td>
<td>388 (139–1611)</td>
<td>397 (174–1795)</td>
<td>435 (197–1614)</td>
<td>436 (131–1268)</td>
<td>414 (131–1795)</td>
</tr>
<tr>
<td>Hydroxyproline: creatinine ratio†</td>
<td>0.05 (0.01–0.21)</td>
<td>0.06 (0.01–0.38)</td>
<td>0.07 (0.02–0.55)</td>
<td>0.05 (0.01–0.40)</td>
<td>0.06 (0.01–0.35)</td>
</tr>
<tr>
<td>No of patients with Pagetic bone pain (%)</td>
<td>17 (65)</td>
<td>18 (62)</td>
<td>20 (68)</td>
<td>16 (57)</td>
<td>70 (63)</td>
</tr>
</tbody>
</table>

*Significantly different from placebo (p<0.05). †Significantly different from placebo (p<0.001). **Significantly different from placebo and tiludronate 200 mg (p<0.001). ††Significantly different from placebo, tiludronate 200 mg (p<0.0001) and 400 mg (p<0.005)
xyproline:creatinine ratios were substantially decreased by tiludronate, but not by placebo, with a dose-response effect. The most rapid reduction of hydroxyproline occurred in the first four weeks of treatment. Tiludronate 400 mg and 600 mg decreased hydroxyproline: creatinine ratios to a significantly greater extent than placebo at week 12 (p<0.001) and week 24 (p<0.05). Although tiludronate 200 mg decreased the hydroxyproline:creatinine ratio by more than 34% compared with placebo, the reduction was not statistically significant; 600 mg was significantly more effective than 200 mg at weeks 12 and 24 (p<0.05), but other between-treatment differences were not statistically significant.

Only 70 out of 112 patients (63%) complained of Pagetic bone pain on entry. The proportion of patients complaining of bone pain at weeks 12 and 24 was greater with placebo (57% and 67%, respectively) and lowest with tiludronate 600 mg (36% and 37%, respectively). However, there were no significant differences between any of the treatment groups at week 12 or 24. The number of patients reporting Pagetic bone pain at entry were 17, 18, 19 and 16 for placebo, 200 mg, 400 mg and 600 mg, respectively. By week 24 the numbers were 16, 14, 13 and 10, respectively, although patients reporting pain at week 24 did not necessarily report pain at entry. Median Visual Analogue Scale scores for pain were lower in the tiludronate 600 mg group at weeks 12 and 24 (0 and 0 mm, respectively) than on placebo (15 and 27 mm, respectively) but there were no significant differences between any of the treatment groups.

SAFETY
The few changes in haematological or biochemical safety measurements that were of clinical significance are described below. One 62-year-old man (600 mg dose) developed epigastric pain, nausea and an elevated serum creatinine concentration (182 μmol/l; reference range 45–125) at week 24. During treatment, creatinine concentrations had remained stable and within the reference range. No significant changes in liver function tests were observed during the trial.

One 79-year-old man (600 mg dose) was found to have raised liver enzymes during the study. On weeks 8 and 12 his alanine transaminase concentrations were above the reference range (47 IU/l and 43 IU/l, respectively; reference range 5–40 IU/l), and throughout the entire duration of the study his y-glutamyl transpeptidase concentrations were also above the reference range (week 0: 126 IU/l, week 2: 99 IU/l, week 4: 90 IU/l, week 8: 86 IU/l, week 12: 85 IU/l and week 24: 67 IU/l; reference range 0–60 IU/l). With the exception of the y-glutamyl concentration at week 0, none of these raised enzyme concentrations were considered to be clinically significant.

A clinically significant eosinophilia was reported in a 69-year-old man (placebo) at week 8 of the study (1.6 x 10⁷/l; reference range <0.5 x 10⁷/l), but this had returned to within the reference range at subsequent visits.

There was a tendency for adjusted calcium concentrations to move from normal to low normal in tiludronate recipients. However, these abnormalities were not considered to be of clinical significance by the investigators.

Adverse events were classified according to the WHO scheme. A total of 164 adverse events were reported by 79 patients; those occurring in seven or more patients are shown in table 3. The incidence of adverse events was higher on tiludronate (65/86 patients (76%)) than on placebo (14/26 patients (54%)) and was dose-related. The majority (54%) of adverse events were mild and 56% of adverse events were not thought to be related to the study medication. Previous treatment with bisphosphonates did not alter the incidence of adverse events.

Two deaths were reported during the study. One was due to pulmonary embolus and the

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**Figure 2** Effect of tiludronate (200, 400 or 600 mg daily) on mean serum hydroxyproline:creatinine ratio

**Table 3** Number of patients reporting an adverse event (%)

<table>
<thead>
<tr>
<th></th>
<th>Placebo (n=26)</th>
<th>Tiludronate 200 mg (n=29)</th>
<th>Tiludronate 400 mg (n=29)</th>
<th>Tiludronate 600 mg (n=28)</th>
<th>p-value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea</td>
<td>2 (7.7)</td>
<td>5 (17.2)</td>
<td>3 (10.3)</td>
<td>7 (25.0)</td>
<td>0.351</td>
</tr>
<tr>
<td>Diarrhoea*</td>
<td>0 (0.0)</td>
<td>3 (10.3)</td>
<td>4 (13.8)</td>
<td>7 (25.0)</td>
<td>0.037</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>2 (7.7)</td>
<td>5 (17.2)</td>
<td>1 (3.4)</td>
<td>2 (7.1)</td>
<td>1.00</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>0 (0.0)</td>
<td>3 (10.3)</td>
<td>1 (3.4)</td>
<td>5 (17.9)</td>
<td>0.114</td>
</tr>
<tr>
<td>Skeletal pain</td>
<td>3 (11.5)</td>
<td>1 (3.4)</td>
<td>3 (10.7)</td>
<td>3 (21.4)</td>
<td>0.385</td>
</tr>
<tr>
<td>Vomiting</td>
<td>0 (0.0)</td>
<td>1 (3.4)</td>
<td>0 (0.0)</td>
<td>6 (21.4)</td>
<td>0.198</td>
</tr>
</tbody>
</table>

*Significantly higher incidence on tiludronate than on placebo. There were no other significant differences between placebo and tiludronate. †Fisher’s exact test: 2-tailed, placebo vs all tiludronate
other followed a myocardial infarction; neither event was attributed to study medication. Of the seven patients withdrawn because of adverse events, four were considered to be related to study medication: two from the 600 mg dose group (both for gastrointestinal symptoms), one from the 200 mg dose group (also gastrointestinal) and one on placebo (asthenia/dizziness). The most common adverse events were gastrointestinal (38%) with diarrhoea occurring significantly more often on tiludronate than on placebo (p = 0.0372), but none were considered serious. The number of gastrointestinal events was dose-related (placebo: four, 200-mg group: 18, 400-mg group: nine, 600-mg group: 31). The higher incidence of gastrointestinal adverse events on tiludronate was not related to either concomitant use of nonsteroidal anti-inflammatory agents (p=0.107) or a previous history of gastrointestinal disorders at entry (p=0.969).

There was one traumatic fracture which occurred during the study after 21 days treatment with 400 mg daily involving a non-Pagetic rib.

FOLLOW-UP
Eighteen months after the last patient had completed the trial, patients who had completed at least 11 weeks of treatment were followed up to assess the sustained clinical efficacy of tiludronate. Efficacy was assessed in terms of time interval between completion of the initial study and further treatment, number of patients requiring retreatment with a specific anti-Paget agent, number of patients suffering from complications of Paget’s disease, and serum alkaline phosphatase concentrations.

Of the original 112 patients entered into the study, 98 were eligible for follow-up. Of the remaining 14 patients, five were withdrawn for protocol violations (three on placebo, one each on 400 mg and 600 mg), one had insufficient treatment duration (placebo), five had adverse events (two each on 200 mg and 600 mg, one on 400 mg), one dropped out (200 mg) and two died (both on 200 mg). Completed questionnaires were returned on 85 patients (19 for placebo, 22 for each of the tiludronate groups).

Thirty-six patients had required retreatment for Paget’s disease since the end of the original study (table 4). The reasons for retreatment stated by the investigators were pain (three), deterioration of Paget’s disease (26) and serum alkaline phosphatase raised 30% above the end of treatment concentration (seven). There was a distinct dose-dependent effect on the number of patients deemed as needing retreatment between treatments, with doses of 400 mg (p=0.019) and 600 mg (p=0.008) tiludronate being significantly different from placebo. The period of time until first retreatment or last review (if without further treatment) showed a statistically significant increase in ‘treatment-free time’ for all the treatment groups compared with placebo (p=0.043, 0.028, and 0.001 for 200 mg, 400 mg, and 600 mg tiludronate, respectively). However, there was no evidence of differences in ‘treatment-free time’ between the tiludronate groups.

Fourteen patients experienced a complication of Paget’s disease (including pain) during the follow-up period (table 4). There was a statistically significant reduction in the 400 mg tiludronate group compared with placebo (p=0.029) and comparisons between 600 mg tiludronate and placebo approached significance (p=0.078). Analysis of the decrease in mean serum alkaline phosphatase concentrations against time showed a dose-related effect. For all tiludronate groups the reduced concentrations following treatment were maintained for a plateau phase which lasted for several months. The subsequent slow increase in serum alkaline phosphatase concentrations reflected a gradual increase in disease activity.

The dose–response relationship observed during the six-month trial period persisted throughout the follow-up period, as illustrated in figure 3.

Discussion
Bisphosphonates have a low oral bioavailability17 which may be markedly altered by formulation changes, with consequent effects upon efficacy and tolerability. In this study, for example, a 400-mg tablet showed a similar effect to that previously demonstrated by an aqueous capsule,14 the difference being ascribed to the addition of sodium lauryl sulphate to the formulation.

The results showed that 400 mg tiludronate daily for 12 weeks was associated with a reduction of mean serum alkaline phosphatase concentrations by 67% at the end of 24 weeks. In this dosing group, 70% of the patients showed a 50% fall in serum alkaline phosphatase concentrations and 96% showed a 25% fall (used as an efficacy criterion in some studies18). The proportion of patients in the 400-mg group whose serum alkaline phosphatase concentration was normalised was 39%.

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Results of the follow-up survey, 18 months after completion of study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tildudronate</td>
<td>Placebo (n=19)</td>
</tr>
<tr>
<td>Number of patients requiring retreatment</td>
<td>13</td>
</tr>
<tr>
<td>Time until lower quartile (25%) retreatment (days)*</td>
<td>153</td>
</tr>
<tr>
<td>Complications of Paget’s disease</td>
<td>6</td>
</tr>
</tbody>
</table>

*At the time of survey the 400 mg and 600 mg tiludronate groups had not reached median (50%) retreatment time, therefore the lower quartile (25%) was compared.
Figure 3 Change in mean percentage of mid-normal range of serum alkaline phosphatase over time. Data points for the post-study period (after week 24) are plotted as rolling three-month means where n > 3.

This level of response is similar to that reported for higher doses of etidronate (20 mg/kg) and for other bisphosphonates, as reviewed by Kanis,1 and is higher than previously observed with injected calcitonin (25%). The results for 600 mg tiludronate were slightly higher than for 400 mg with 82% of patients achieving a 50% reduction in concentrations (mean reduction 74%) and 44% of patients achieving normalisation. Tiludronate maintained its suppression of serum alkaline phosphatase 12 weeks after withdrawal of treatment and its efficacy was unaltered by prior bisphosphonate therapy.

Tiludronate produced rapid, marked reduction of hydroxyproline:creatinine ratios, indicating that its chief action is to inhibit osteoclast activity. The percentage reduction in this ratio was approximately equal for 400 mg and 600 mg tiludronate (66% and 67%, respectively), although the 400 mg group had a higher mean hydroxyproline:creatinine ratio at baseline (0.073 compared with 0.052 for the 600 mg group).

Tiludronate was associated with few serious adverse events of clinical significance. However, as with other bisphosphonates, a dose-related increase in gastrointestinal symptoms was observed. These symptoms often resolved spontaneously, but occasionally required treatment to be discontinued. While 75% (21/28) of patients receiving 600 mg of tiludronate had gastrointestinal symptoms only 31% (9/29) of those on 400 mg reported similar complaints.

Although pain was a secondary endpoint for efficacy in this study, only 63% of patients complained of Pagetic pain at entry. The percentage of patients reporting Pagetic bone pain after 24 weeks exhibited an apparent dose response, being 67%, 53%, 46%, and 37%, on placebo, 200 mg, 400 mg, and 600 mg, respectively; however the difference from placebo was not significant.

The efficacy of a treatment for Paget's disease may be assessed either by direct clinical endpoints or biochemically. Biochemical as-

Summary/learning points
- Oral tiludronate is an effective and well-tolerated treatment of Paget's disease of bone.
- The reduction in alkaline phosphatase observed with tiludronate demonstrates a dose response.
- Adverse events are dose-related and the only adverse event that had a statistically significantly higher incidence with tiludronate compared to placebo was diarrhoea (p=0.037).
- Tiludronate in a dose of 400 mg once daily for three months offers the optimum balance of efficacy and tolerance.
12 weeks offers an optimum balance of efficacy and safety.

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