Efficiency of Ibandronate in Monotherapy and in Combination with Alfacalcidol in Women with Postmenopausal Osteoporosis

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Abstract

The aim of this study was to carry out a comparative analysis of the efficiency of ibandronate monotherapy and combined therapies with ibandronate and alfacalcidol in women with postmenopausal osteoporosis.

Materials and Methods: A total of 53 women (mean age 60.7 years) with postmenopausal osteoporosis (PMO) were randomized to monotherapy with ibandronate 150 mg/month (Group Ib) (n=25) and therapy with ibandronate 150 mg/month plus alpha-D3 (AD3) 1μg (alfacalcidol) daily (Group Ib+AD3) (n=28). All women received calcium and vitamin D3 supplements. Patients were recruited in one center and were followed up for 6 months on a monthly basis. To assess the efficacy of therapy, BMD was measured at LS (L1–L4) and PF at the beginning and end of therapy by DEXA. Biochemical markers of bone turnover were also assessed.

Results: Statistically significant increases in BMD compared with baseline values and the control group were observed in both ibandronate treatment groups. Growth of BMD was significantly higher in Group Ib+AD3 compared to Group Ib. An assessment of CTX dynamics showed a notable decrease in CTX level in patients of both groups compared with levels before treatment. Generally, PTH level decreased insignificantly, but a more pronounced reduction was seen in Group Ib+AD3. TP1NP level significantly increased in Group Ib and was more pronounced in Group Ib+AD3.

Conclusion: Combined therapy with ibandronate sodium and the D-hormone analog alfacalcidol augments the effectiveness of treatment observed in ibandronate sodium monotherapy in PMO women. (Int J Biomed. 2016;6(3):190-194.)

Key Words: menopause • osteoporosis • bone mineral density • ibandronate treatment
powerful antiresorptive potential.\textsuperscript{[12-14]}

The clinical efficacy of ibandronate was confirmed by results of multicenter studies. Thus, the results of a multicenter, randomized, double-blind comparative study MOBILE (Monthly Oral Ibandronate In Ladi Es) of therapy with ibandronate demonstrate a reduction in the incidence of vertebral fractures (by 4.9% and 6.6%, respectively) and fractures of the thigh bone (by 3.2% and 6.2% respectively) after 1 and 2 years.\textsuperscript{[15-16]} The BONE study (Oral Ibandronate Osteoporosis Vertebral Fracture Trialin North America and Europe) found that women taking ibandronate on a daily or monthly basis have reduced risk of vertebral fractures and non-vertebral fractures in two populations – North America (60 and 54%, respectively) and Europe (50 and 48%, respectively).\textsuperscript{[17]}

The aim of this study was to carry out a comparative analysis of the efficiency of ibandronate monotherapy and combined therapies with ibandronate and alfacalcidol in PMO women.

**Materials and Methods**

The study included 53 PMO women aged between 51 and 75 years (mean age 60.7±6.13 years, median 61.0 years; interquartile range [IQR] 55.0–64.0 years) with PMO and menopause duration of at least 1 year. Osteoporosis was diagnosed according to WHO criteria (1994) using T score standard deviations (SDs) from the normative values of peak bone mass in healthy women. A value of not more than -1 SD was regarded as normal, a value of -1t0-2.5 SD as osteopenia, and ≤-2.5 SD as osteoporosis. Women were randomized to monotherapy with ibandronate 150 mg/month (Group Ib) (n=25) and therapy with ibandronate 150 mg/month plus alphas-D3(AD3)1μg (alfacalcidol) daily (Group Ib+AD3) (n=28). Both groups were homogeneous, particularly for BMD. All patients were informed about the disease and its complications and gave informed consent to participate in the study. All women received calcium supplements at a dose of 1,000mg and vitamin D3 800 IU/day. A control group of 16 women received only calcium 1,000 mg and vitamin D3 800 IU/day.

Physical load enhancement in the form of daily 30-min walking was recommended to all patients. Patients were recruited in one center and were followed up for 6 months on a monthly basis. Women with diseases affecting bone metabolism such as hyperparathyroidism, thyrotoxicosis, Cushing’s syndrome and disease, hypogonadism, malabsorption syndrome, kidney and liver disease, and malignancies were excluded, as were those taking medications that affect calcium metabolism during 12 months before the study.

**BMD Measurements**

The primary objective of the study was to assess the evolution of BMD on the background of the therapy, and the second objective was to follow the evolution of BTMs. To assess the efficacy of therapy, BMD was measured at LS (L1–L4) and PF at the beginning and end of therapy by dual energy x-ray absorptiometry (DEXA) using a bone densitometer (Prodigy, CE Lunar Corporation, USA) in private clinic “Doctor Summit” (Summit Trading Company, Ltd., Tashkent, Uzbekistan). In compliance with WHO criteria, BMD was expressed as g/cm² and T-score. Efficacy analyses were conducted according to the intention-to-treat (ITT) principle. The ITT population included all randomized patients who had taken at least one dose of treatment and who had a baseline and one post-baseline evaluation.

**Biochemical Measurements**

To assess the metabolic activity of bone remodeling, biochemical markers such as CTX (b-CrossLaps test) and TP1NP were measured by electrochemiluminescence assays (Elecsys biochemical analyzer, Elecsys b-CrossLaps) at baseline and after 6 months of therapy. An absorptiometric method was used to determine serum total calcium and nonorganic phosphorus, and a kinetic method was used to determine alkaline phosphatase activity by the amount of liberated 4-nitrophenol. Commercially available kits (CIS Bio International, France) were used to measure levels of parathyroid hormone (PTH).

The study was approved by the local Ethics Committee. Written informed consent was obtained from each patient.

Statistical analysis was performed using the statistical software «Statistica» (v6.0, StatSoft, USA). Baseline characteristics were summarized as frequencies and percentages for categorical variables and as mean ± SD for continuous variables. The data for variation indices with nonparametric distribution are presented with medians and interquartile range [IQR]. The Mann-Whitney (U Test) was used to compare the differences between groups. Group comparisons with respect to categorical variables are performed using chi-square test with Yates correction. A probability value of \( P<0.05 \) was considered statistically significant.

**Results**

The studied groups were comparable in terms of medical history (Table 1). High BMI was observed in all groups, which is likely associated with a sedentary lifestyle and poor diet (prevalence of pastry, fried food with use of animal fats) among postmenopausal women.

**Table 1.**

**Baseline characteristics of patients**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Control group (n=16)</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Group Ib</td>
</tr>
<tr>
<td>Age,y</td>
<td>62.9±6.4</td>
<td>65.7±3.8</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>31.2±4.8</td>
<td>29.1±4.4</td>
</tr>
<tr>
<td>Menopause age, y</td>
<td>50.5±3.6</td>
<td>48.9±4.7</td>
</tr>
<tr>
<td>Menopause duration, y</td>
<td>12.4±4.3</td>
<td>16.8±5.9</td>
</tr>
<tr>
<td>Fractures in anamnesis, n (%)</td>
<td>1(6.3)</td>
<td>1(4.0)</td>
</tr>
<tr>
<td>Hip</td>
<td></td>
<td>1(6.3)</td>
</tr>
<tr>
<td>Wrist</td>
<td></td>
<td>1(4.0)</td>
</tr>
<tr>
<td>Spine</td>
<td></td>
<td>1(4.0)</td>
</tr>
<tr>
<td>Humerus</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fractures in relatives, n (%)</td>
<td>2(12.5)</td>
<td>2(8.0)</td>
</tr>
<tr>
<td>Menopause before 45 yrs, n (%)</td>
<td>1(6.3)</td>
<td>3(12.0)</td>
</tr>
</tbody>
</table>
Patients who had previous fractures were 2 women in Group Ib (fractures of wrist and humerus), 3 women in Group Ib+AD3 (hip, vertebra, and humerus), and 2 women in the control group (hip and wrist). Patients who had a family history of fractures were 2 (8.0%) women in Group Ib, 3 (10.7%) women in Group Ib+AD3, and 2 (12.5%) women in the control group. Onset of menopause before the age of 45 years had occurred in 3 (12.0%), 5 (17.9%) and 1 (6.3%) women of Group Ib, Group Ib+AD3 and the control group, respectively.

Basal LS-BMD and PF-BMD equaled to $0.746\pm0.05$ g/cm$^2$ (median of 0.741 g/cm$^2$; IQR 0.709-0.781) and $0.747\pm0.05$ g/cm$^2$ (median of 0.746 g/cm$^2$; IQR 0.728-0.774), respectively.

**Bone Mineral Density**

An insignificant increase in BMD was noted in the group of women who used calcium supplements and vitamin D3. In this group, LS-BMD decreased in 8 (50%) women and increased only in 4 (25%) women. The average growth of BMD amounted to 0.24% (Fig.1). In the control group, PF-BMD did not change in 8 (50.0%) patients and an insignificant increase was seen in 7 (43.8%). The average growth amounted to 0.06% (Fig.2).

In Group Ib, a significant increase in LS-BMD and PF-BMD was found. However, we observed a decrease in LS- and PF-BMD in 3 (12.0%) and 4 (16.0%), respectively. Growth of BMD was significantly higher in Group Ib+AD3 compared to Group Ib and the control group; a slight decrease in LS-BMD was seen in 2 patients of Group Ib+AD3.

**Biochemical Markers of Bone Turnover**

An assessment of CTX dynamics showed a notable decrease in CTX level in patients of both groups compared with levels before treatment (Fig.3). Generally, PTH level decreased insignificantly, but a more pronounced reduction was seen in Group Ib+AD3. TP1NP level significantly increased in Group Ib (30.2%) and was more pronounced in Group Ib+AD3 (52.2%).

Laboratory investigations demonstrated normal parameters for calcium–phosphorus metabolism (Ca, P and alkaline phosphatase) in all groups without significant intergroup differences. Levels of calcium and phosphorus as well as alkaline phosphatase activity were in the reference limits, not changing significantly in both groups.

**Safety**

A flu-like syndrome was observed in 9 women (4 patients in Group Ib and 5 patients in Group Ib+AD3) during study. The intensity of the syndrome was light to moderate, resolved spontaneously or after administration of antipyretics, and did not require discontinuation of therapy.

**Discussion**

Currently, ibandronate is one of the most in-demand drugs with proven ability to reduce the risk of vertebral
fractures and a dose-dependent activity in relation to peripheral fractures. Ibandronate has been studied in clinical trials involving nearly 9,000 patients. The results of evaluation of ibandronate efficacy in osteoporosis were reported in large randomized studies: BONE,[17] MOPS,[18] MOBILE.[15]

In BONE study, both daily and monthly administration of ibandronate led to a significant increase of vertebral BMD (to 5.4 and 4.4% - North American population, and to 7.1 and 6.3% - European population) compared to its baseline values. Growth of PF-BMD by 2.6% and 3.7% was seen with daily intake and by 2.5% and 3.1% with monthly intake for North American and European populations of patients.[17] During a 2-year (MOBILE study), LS-BMD of patients receiving ibandronate 150 mg once monthly increased by 7.6% (from baseline), and it increased by 6.4% from baseline in patients receiving 100 mg per month. After 3 years of treatment, patients in both groups showed an increase in PF-BMD as compared with the original mineral density (3.4% - 100 mg and 4.1% - 150 mg).[15]

In our study, within 6 months PMO women received ibandronate 150 mg once monthly, as well as the same dose of ibandronate plus Alpha-D3 1mg/day. Significant improvements of LS-BMD by 4.9% and PF-BMD by 3.7% were observed on the background of ibandronate monotherapy. In combination therapy of ibandronate with Alpha-D3, an increase in LS-BMD and PF-BMD comprised 8.5% and 8.3%, respectively. A decrease in LS-BMD and PF-BMD was seen in 3 and 4 women, respectively, in Group Ib. At the same time, only 2 patients treated with Ib+AD3 demonstrated a slight decrease in LS-BMD.

Two opposed processes, bone formation and bone resorption, characterize bone metabolism. The type 1 collagen degradation product CTX is a parameter characterizing the degree of bone tissue resorption, while TP1NP is a marker of bone matrix formation. Increases in CTX are believed to be the primary and most sensitive parameter of the shift in balance from bone remodeling towards bone resorption.[19]

Study of markers of bone metabolism showed that CTX level decreased significantly in the treatment with ibandronate, and an even greater reduction resulted from the combination therapy. Effects of ibandronate were also associated with a significant increase in TP1NP level, more pronounced in Group Ib+AD3.

Bisphosphonates reduce the number and activity of osteoclasts, induce their apoptosis and inhibit enzymes of the mevalonate pathway, which ultimately results in reduced bone resorption. However, D-hormone deficiency may prevent the bisphosphonates from exerting their antiresorptive effects. D-hormone deficiency may arise from impaired vitamin D conversion into D-hormone and from primary (nutritional) shortage of calcium and vitamin D. As compared with bisphosphonates, active metabolites of vitamin D posses more diverse actions, which have a complementary effect: they reduce osteoclast genesis, stimulate differentiation of preosteoblasts and bone formation (anabolic effect), and also have metabolic effects that appear as reduced levels of PTH and pro-inflammatory cytokines.[20] The addition of alfacalcidol to the bisphosphonates may not only contribute to the effectiveness of therapy, but can also solve such a clinical problem as resistance to bisphosphonates, which sometimes is as high as 83%.[21]

PTH is a key regulator of calcium–phosphorus metabolism. It is synthesized by the parathyroid glands in response to a reduction in extracellular calcium concentrations, and it activates osteoclasts to increase bone resorption (demineralization, bone destruction); as a result, calcium and phosphorus are released into the circulation. Increases in PTH levels, particularly in elderly individuals with vitamin D3 deficiency, can result in osteomalacia, increased bone remodeling, bone mass reduction, and fractures.[22] Reduction of PTH levels in our study was observed in both treatment groups with more significant reduction in combined therapy.

Conclusion

Thus, combined therapy with ibandronate sodium and the D-hormone analog alfacalcidol augments the effectiveness of treatment observed in ibandronate sodium monotherapy in PMO women.

Sources of Funding

This study was conducted as an investigator driven study. However, F. Hoffmann-La Roche Ltd has funded the Republican Specialized Scientific-Practical Medical Center of Endocrinology to perform the study. F. Hoffmann-La Roche Ltd was not involved in the acquisition of the data, in the statistical analysis, or in the drafting and revision of the article.

Competing interests

The authors declare that they have no competing interests.

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