CLINICAL STUDY

The effect of timing of teriparatide treatment on the circadian rhythm of bone turnover in postmenopausal osteoporosis

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Abstract

Background: We hypothesized that with the administration of teriparatide (TPTD) treatment at different times, we would be able to modify the physiological circadian rhythm of bone turnover.

Methods: The concentration of serum C-terminal telopeptide of collagen type I (βCTX), serum N-terminal propeptide of procollagen type I (P1NP), serum ionized calcium (iCa), and plasma PTH were measured every 3 h over a 24 h period in 14 postmenopausal osteoporotic women (aged 72.4 ± 9.3 years) treated with 20 μg TPTD for long term, given at different times of the day. General linear model-repeated measurements (GLM RM) were performed to analyze the circadian rhythms as well as intergroup comparisons.

Results: GLM-RM for both related groups showed a significant influence of time of day on all measured variables except P1NP. The analysis for each group separately provided a powerful model for βCTX (P<0.001, η²=0.496), serum iCa (P<0.001, η²=0.423), plasma PTH (P<0.001, η²=0.283), and serum P1NP (P<0.001, η²=0.248). While the evening TPTD treatment showed a marked circadian rhythm for serum βCTX, the morning TPTD treatment rather suggested circasemidian rhythm. The P1NP rhythm followed a much smaller amplitude of the rhythm than βCTX. Changes in serum iCa were positively related to changes in serum βCTX (P<0.001) and negatively related to changes in PTH (P<0.001).

Conclusion: Timing of TPTD administration may significantly change the 24 h variation in bone turnover markers as well as calcium-parathyroid axis in postmenopausal osteoporotic women.

European Journal of Endocrinology 164 643–648

Introduction

Circadian rhythmicity is an essential feature of bone and mineral homeostasis. The biochemical markers of bone resorption, such as collagen type I degradation products, e.g. C-terminal or N-terminal telopeptide of type I collagen, follow a circadian rhythm, decreasing during the day and peaking at night (1, 2). For biochemical markers of bone formation, such as plasma osteocalcin and serum N-terminal or C-terminal propeptide of type I procollagen, a similar pattern of circadian rhythm with much smaller amplitude than that seen in bone resorption markers has been found (3–5). While rhythmicity of serum osteocalcin has been linked to cortisol secretion (4), the physiological basis for the circadian rhythm of bone resorption is not clearly understood and seems to be unaffected by age, menopausal status, gender, bed rest (2), or cortisol secretion (6). However, fasting significantly diminishes the circadian rhythm of bone-resorptive activity (7) and feeding deepens the morning decrease in bone-resorptive activity and accentuates its nocturnal peak (2). PTH is one of the most important systemic regulators of bone and mineral homeostasis, which follows a circadian rhythm and responds to feeding, especially to calcium/phosphate intake. In healthy individuals, plasma concentrations of PTH show a circadian variability, with minimal concentrations around 1000 h and maximum levels at about 0300 h (8). Although the exact role for this rhythm has not been established yet, previous studies had showed an association between PTH rhythm and bone-resorptive activity (9). The PTH rhythm may set the amplitude of the circadian rhythm of bone resorption, especially by increasing the resorptive activity of existing osteoclasts during nocturnal fasting (10, 11). The nocturnal peak of PTH is abolished by fasting (12). The amplitude of the PTH circadian rhythm is smaller in osteoporotic women when compared with that seen in healthy women (3, 13, 14). Studies in which calcium was administered for a longer time period either in the morning or in the evening, as opposed to acute suppression of PTH in clamp studies (11), have shown that timing of intermittent calcium supplementation influences the circadian rhythm of bone resorption (10).
The aim of this study was to test our hypothesis that the effect of PTH treatment on bone remodeling and the calcium–PTH axis may be influenced by a different time of PTH administration. We investigated changes in bone turnover markers and the calcium–PTH axis over a 24 h period in postmenopausal osteoporotic women who were treated for a long term with teriparatide (TPTD), administered daily either in the morning or in the evening.

Subjects and methods

Subjects

All subjects were ambulatory women with established postmenopausal osteoporosis, 50–85 years of age, recruited from the Bone Center at the University Hospital, Prague, from July to December 2008. Patients were included if they were at least 5 years postmenopause, had a bone mineral density (BMD) T-score below −2.5 at the lumbar spine and/or the femoral neck, and had been treated with TPTD for at least 6 months, given either in the morning (approximately at 0800 h, before breakfast) or in the evening (approximately at 0200 h, after dinner). All patients had a total daily calcium intake of at least 1000 mg, either through diet or through diet enriched with calcium supplements, and were supplemented with vitamin D3 (800–1000 IU daily). Exclusion criteria were patients with abnormal serum or urinary calcium levels, with vitamin D insufficiency (serum 25-hydroxyvitamin D concentration < 50 nmol/l); diseases other than osteoporosis intervening with bone metabolism; and treatment with other drugs affecting bone metabolism such as corticosteroids, cyclosporine, fluoride, or thiazide diuretics. Out of 20 invited patients, 14 women were eligible for the study. The study was attempted with the understanding and written consent of each subject, with the approval of the ethics committee of the Faculty of Medicine, Charles University, Prague, and in compliance with national legislation and the code of ethical principles for medical research involving human subjects of the World Medical Association (Declaration of Helsinki).

Study design

All subjects continued their TPTD and calcium and vitamin D supplements on the day before the study and fasted overnight before the start of the study. Standardized diet was served from 0800 to 0830 h, 1130 to 1200 h, and 1730 to 1800 h. The intake of fluid was defined. The period of sleep and wakefulness was unbroken. Blood samples were obtained in the 3 h intervals via indwelling venous cannulae immediately before the administration of TPTD and meal intake (0800 h) in the fasted state and over the next 24 h. TPTD was administered either in the morning (0800 h) or in the evening (2000 h). The plasma and serum specimens were stored at −70 °C. Measurement of the serum ionized calcium (iCa) was performed on the same day after anaerobic collection of all specimens, which were anaerobically stored at 4 °C to avoid pH shifts.

Biochemical analysis

All samples, except the serum iCa, were simultaneously assayed. Serum iCa was measured using the ion-selective electrode with an AVL 9180 (Roche Diagnostics GmbH). The within-run imprecision was below 2% and between-run imprecision was below 4%. The plasma concentrations of the immunoreactive intact PTH were determined using an electrochemiluminescence-based immunoanalysis (the Elecsys 1010 Analyzer, Roche Diagnostics GmbH). The within-run imprecision was below 6%.

The serum concentrations of βCTX were measured using the electrochemiluminescence-based immunoanalysis (the Elecsys 1010 Analyzer, Roche Diagnostics). The within-run imprecision for the βCTX was below 7%. The serum concentrations of intact N-terminal propeptide of type I procollagen (PINP) were measured by RIA (Procollagen Intact PINP, Orion Diagnostica, Espoo, Finland). The within-run imprecision was below 5%.

Serum concentrations of 25-hydroxyvitamin D (25-OHD) were determined by using enzymoimmunoassay (OCTEA-25-Hydroxy Vitamin-D, Immunodiagnostic Systems Limited, Bolton, UK). Concentrations of 25-OHD above 50 nmol/l were considered normal.

Statistical analysis

The general linear model-repeated measured variant (GLM-RM) was used to assess the 24 h cyclic changes in the measured variables. The circadian rhythm was considered proven if the model revealed either the existence of a characteristic curve describing the daily rhythm similar for all patients (influence by time) or the existence of two curves for each group significantly demonstrating the similarity of the rhythm within each group, yet differences between the groups (influence by group interaction). In addition to testing the significance of levels observed depending on the selected variables (group, time, and group by time interaction), we used GLM RM to estimate the proportion of explained variability and thus effect size of this dependence (R^2). Cohen’s convention indicates a large size effect threshold R^2 > 0.137. GLM-RM also served for intergroup comparisons. With Pearson’s correlation coefficient, calculated for the entire sample and for each group separately, we explored the relationship between changes in the levels of serum βCTX, iCa, and PTH during adjacent sampling (difference of 3 h each). The cosine model Y = mesor + AMP × COS (2π/24 t + phase) has been used as an adjunct to the analysis of circadian rhythms. A description of the method and
Table 1 Patients’ characteristics at baseline. Values are mean ± S.D. Differences between groups were analyzed by unpaired t-test.

<table>
<thead>
<tr>
<th></th>
<th>Morning TPTD</th>
<th>Evening TPTD</th>
<th>P</th>
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</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>7</td>
<td>7</td>
<td>NS</td>
</tr>
<tr>
<td>Age (years)</td>
<td>69.7 ± 9.2</td>
<td>75.1 ± 9.1</td>
<td>NS</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>66.6 ± 7.0</td>
<td>53.9 ± 4.1</td>
<td>0.004</td>
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<tr>
<td>Height (m)</td>
<td>1.65 ± 0.06</td>
<td>1.53 ± 0.06</td>
<td>0.006</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>25.6 ± 2.6</td>
<td>23.0 ± 1.1</td>
<td>NS</td>
</tr>
<tr>
<td>Lumbar spine BMD (T-score)</td>
<td>−3.2 ± 0.7</td>
<td>−3.1 ± 0.8</td>
<td>NS</td>
</tr>
<tr>
<td>Total hip BMD (T-score)</td>
<td>−2.1 ± 0.9</td>
<td>−2.4 ± 0.7</td>
<td>NS</td>
</tr>
<tr>
<td>Duration of TPTD treatment</td>
<td>9.4 ± 1.8</td>
<td>11.9 ± 2.3</td>
<td>NS</td>
</tr>
<tr>
<td>(months)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S-iCa (mmol/l)</td>
<td>1.29 ± 0.05</td>
<td>1.31 ± 0.05</td>
<td>NS</td>
</tr>
<tr>
<td>P-PTH (ng/l)</td>
<td>28.9 ± 7.4</td>
<td>25.8 ± 11.3</td>
<td>NS</td>
</tr>
<tr>
<td>S-βCTX (ng/l)</td>
<td>662.5 ± 168</td>
<td>940.1 ± 461</td>
<td>NS</td>
</tr>
<tr>
<td>S-PINP (µg/l)</td>
<td>118.3 ± 39</td>
<td>169.9 ± 92</td>
<td>NS</td>
</tr>
</tbody>
</table>

BMI, body mass index; BMD, bone mineral density; S-iCa, serum ionized calcium; P-PTH, plasma intact PTH; S-βCTX, serum C-terminal telopeptide of collagen type I; S-PINP, serum propeptide of procollagen type I.

*Results*

The patients’ characteristics and the baseline values of the measured biochemical parameters are given in Table 1. Analysis of the variations of serum βCTX indicated a significant influence of time of day on the measured value (for time $P<0.001$, $\eta^2=0.492$, large size effect by Cohen’s convention). These results suggest that there is a characteristic curve of βCTX levels, which is inherent in all patient groups studied and which explains 49.2% of interindividual variability in measured levels of βCTX. Further analysis showed a more powerful model, which characterizes the two curves, for each group separately (time versus group interaction, $P<0.001$, $\eta^2=0.496$). In the evening TPTD group, serum βCTX levels indicated a circadian rhythm, whereas the morning TPTD treatment group suggested a circasevian rather than a circadian rhythm (Fig. 1A and B). The difference in 24 h mean βCTX concentrations between treatment groups did not approach significance ($P=0.098$); however, the power of the test was low (observed power=0.379). Analysis of the 12 h response curves showed a significantly greater daily mean value in serum βCTX levels (between 2000 and 0800 h) for the evening TPTD group compared with the morning TPTD group ($P=0.033, \eta^2=0.326$; Fig. 1A and B).

For serum P1NP, GLM-RM did not prove the influence of time of day on the measured value for the related groups ($P=0.298$). The analysis based on the two curves for each group separately demonstrated significantly different P1NP rhythms (time by group interaction, $P<0.001$, $\eta^2=0.248$), however, with much smaller amplitude than the bone resorption marker (Fig. 2A and B; Table 2). Probably, due to the wide inter-individual variability, the differences in 24 h mean values between morning and evening TPTD administration could not be detected.

Analysis of variations of iCa indicated a significant time effect ($P=0.021$), grouped by time interaction ($P<0.001$) and a large size effect by Cohen’s convention ($\eta^2=0.180$ and $\eta^2=0.423$, time and group by time interaction respectively). Curves from both groups were running against each other (i.e. high levels for the morning group occur at times of low levels for the evening group and vice versa (Fig. 3A and B)) and the common curve explained 18.0% of inter-individual variability in measured levels of serum iCa. The analysis after distribution explained 42.3% of the variability. The difference in 24 h mean iCa concentrations between treatment groups did not approach significance ($P=0.973$); however, the power of the tests is extremely low (0.050). Also, analysis of the 12 h response curves showed no significant difference between treatment groups, probably due to high

![Figure 1](https://via-free-access.bioscientifica.com/)

**Figure 1** (A and B) Changes in serum C-terminal telopeptide of collagen type I (βCTX) either after morning (A) or after evening (B) teriparatide (TPTD) treatment. Each hairline represents one individual. The bold line represents the mean ± S.E.M. Dotted lines represent normal reference range. TPTD administration is indicated by bold arrow. Meals were taken at the times indicated by triangles (0800–0830 h; 1130–1200 h; and 1730–1800 h).
of the cosinor model, in sharp contrast with most of our measured data, is a requirement that the daily maximal and daily minimal values are occurred in the opposition, i.e. 12 h after each other. Therefore, the estimates of the phase, estimated time for maximum, and estimated time for the minimum cannot be applied to our data, contrary to the estimates of the MESOR and amplitude has no effect. The results of the cosinor model are shown in Table 2. Out of the variables, only the amplitude of S-βCTX after evening administration of TPTD was significantly greater (P < 0.05).

**Discussion**

It is well known that treatment with exogenous PTH exerts both anabolic and catabolic effects on bone, depending on the concentration and duration of exposure (15); however, because of the physiological circadian rhythms of bone turnover and PTH, there is a possibility that the effect of TPTD treatment on bone turnover may also be influenced by the time of TPTD administration during the day. Despite the great interest in PTH as a therapeutic agent, the effect of time-dependent PTH treatment on bone turnover has not been studied previously.

In this study, we found that time-dependent TPTD treatment is able to significantly change the 24 h variation in bone turnover markers, as well as serum calcium and endogenous PTH secretion, in women with established postmenopausal osteoporosis. Whereas the evening TPTD treatment (2000 h) followed the circadian rhythm of bone resorption marker βCTX with a nadir between 1100 and 1400 h and a peak between 2300 and 0200 h, the morning TPTD treatment (0800 h) resulted in an increase in βCTX between 0800 and 1400 h and a subsequent decrease until 2000 h, which was followed by a significant increase in bone resorption marker, suggesting rather
circasemidian than circadian rhythm. The concentrations of serum iCa transiently but significantly increased 3–6 h after TPTD administration in both treatment groups (Fig. 3A). Parallel movement of serum iCa and jCTX, which were significantly related to each other, indicates that changes in serum iCa reflect changes in osteoclast-resorptive activity. The PTH serum concentrations significantly decreased 3–6 h after TPTD administration, and the profiles of serum iCa and endogenous PTH were significantly inversely related and mirrored with each other, indicating that the parathyroid function is preserved during TPTD treatment. These results are in accordance with the previous studies in which an inverse correlation between circadian changes in serum iCa and endogenous PTH has been demonstrated (16, 17).

Our results indicate that the sensitivity of osteoclasts to TPTD treatment varies by the time of TPTD dosing. The percentage increase in serum jCTX during the first 6 h after TPTD administration was significantly more pronounced after the evening TPTD administration in comparison with its morning administration (P = 0.009; 22 ± 56 and 94 ± 38%; morning and evening TPTD treatment respectively). The mechanism(s) of this phenomenon is presently uncertain; however, the results imply a homeostatic mechanism that limits response of bone resorption to TPTD treatment in the morning and/or augments its response in the evening. Although we did not investigate the mechanisms underlying this phenomenon, we suggest at least two factors that may modulate the bone remodeling balance during the TPTD treatment. First, at a cellular level, the increase in bone-resorptive activity by PTH is believed to be an indirect effect of PTH via the production of local factors, such as RANKL from osteoblasts and/or a transient decrease in the production of osteoprotegerin (OPG), which acts as a decoy receptor for RANKL (18, 19). A recent study by Joseph et al. (9) demonstrated a circadian rhythm for circulating OPG levels with a daytime increase and nocturnal decrease in healthy subjects. Therefore, the endogenous circadian rhythm in OPG secretion may at least partly modulate the effect of TPTD treatment on osteoclast’s resorptive activity. However, further studies are needed to test this possibility in postmenopausal osteoporotic women. Secondly, the effect of PTH injection on osteoclasts may be attenuated due to the known suppressive effect of food intake on bone-resorptive activity in the morning (20). Interestingly, osteoclasts are less affected by feeding in the late afternoon and evening in non-fasting individuals (21). Further investigation is required for better understanding of these complex regulations and to optimize bone remodeling balance during TPTD treatment in postmenopausal women with osteoporosis.

In this study, serum PINP significantly varied according to the time of TPTD administration (P = 0.001) but to a much smaller extent than the bone resorption marker. Serum PINP showed a mild but significant decrease during 6 h after the morning (P = 0.033) or evening (P < 0.05) TPTD administration, with a subsequent mild increase in both treatment groups. The delayed response of serum PINP in our study may suggest that collagen synthesis requires more time than collagen degradation to show acute changes after TPTD administration. The inter-individual mean levels in PINP were highly variable and the overall mean 24 h level of PINP was not significantly different between the groups.

Several limitations of this study must be taken into account. First, due to the cross-sectional design, we could not evaluate the changes in 24 h variation in the measured parameters before and after TPTD treatment. Secondly, due to the small number of subjects and a high inter-individual variability, some differences did not reach statistical significance. However, our results clearly showed that the time of TPTD administration has a large effect on the measured variables. Thirdly, a more frequent sampling would be
required to study the 24 h variation of serum iCa and plasma endogenous PTH (9).
In conclusion, this is the first study to show that circadian variation in bone-resorptive activity significantly differs according to the dosing time of the TPTD treatment. Our data support our hypothesis that the response of bone cells to TPTD treatment differs by its dosing time. Whether the timing of TPTD administration will have an impact on the efficacy and safety of long-term TPTD treatment will need to be further analyzed in prospective studies. With respect to long-term efficacy, several parameters such as BMD need to be evaluated.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Funding
This work was supported by IGA Ministry of Health of Czech Republic NS 10564-3.

Acknowledgement
We thank Oldřiska Lukaskova for excellent technical assistance.

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Received 28 January 2011
Accepted 2 February 2011