Strontium Ranelate Effect on the Repair of Bone Defects and Molecular Components of the Cortical Bone of Rats

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Introduction

Strontium ranelate (SR) is a drug developed for the treatment of osteoporosis. Its molecule consists of two stable strontium atoms (Sr²⁺), which are the active part of the compound and the ranelic acid organic molecule (1).

Among the drugs used in the treatment of osteoporosis, SR is the only one that demonstrates a dual mode of action on bone metabolism. It favors bone formation (osteogenesis) by stimulating division of osteoblast precursors and increasing synthesis of collagen and non-collagen proteins. Additionally, it simultaneously decreases bone resorption by inhibiting osteoclast differentiation and activity (2,3). Clinical use of SR for the treatment and prevention of osteoporosis has shown positive effects such as increase in bone mineral density and decrease in the risk of bone fracture (4).

The effect of SR on bone repair was investigated in animal models with tibial fracture and showed beneficial results such increases in bone callus and improvement in the biomechanical properties of this tissue (5,6). Use of strontium (Sr) incorporated with a bioactive material has been recently studied for the repair of bone defects in osteopenic animals and the results showed that local release of Sr atoms could improve bone repair (7). Although this drug was proven effective in bone repair, few data regarding their effects on the repair of bone defects are available. Furthermore, little is known about the molecular components of the bone formed during treatment.

Fourier transform infrared (FTIR) spectroscopy has been extensively used in the analysis of molecular components of biomaterials and tissues (8). Studies on bone metabolism and chemical and structural changes caused by bone aging and changes caused by diseases, such as osteoporosis and effects of therapies, were performed in mineralized tissues with FTIR (9-12).

The objectives of this study were to evaluate the effects of ovariectomy and systemic treatment with SR on the organic and inorganic components of cortical bone tissue and to assess whether SR treatment can promote repair of bone defects and improve microarchitecture of the newly formed bone.

Material and Methods

Experimental Animals

Adult rats (Rattus norvegicus, albinus, Wistar, n=27; 3 months old; average weight: 250 g) were used. The animals were ovariectomized (OVX; n=18) or subjected to sham surgery (SHAM; n=9). The rats were kept in rooms with controlled temperature (22 °C) rooms, with periodic change (12/12 h) of light, and specifically fed a pelleted diet containing calcium (Ca; 1%), phosphorus (P; 0.8%) and vitamin D3 (5,000 IU/kg). Water and pelleted food were available ad libitum. All experimental procedures
were approved by the Ethics Committee for Animal Use of the Institute of Biomedical Sciences, University São Paulo (USP), under protocol number 134/2012.

Production of Bone Lesion

Thirty days after ovariectomy and sham surgery, a bone defect was made in all animals. In all surgical procedures, general anesthesia was obtained by applying an intramuscular injection of 2% xylazine hydrochloride with 10% ketamine hydrochloride, in the proportion of 1.0/0.5 mL, with a 0.1 mL/100 g body weight dosage. After dissection of the subcutaneous tissues, a bone defect was made in the left femur (diaphysis region; distal portion). A spherical carbide drill with 2.5 mm external diameter driven by an electric motor (speed: 1500 rpm) under constant irrigation (0.9% NaCl sterile solution) was used. The drill was positioned perpendicular to the long axis of the femur and the cortical bone was cut in a circular shape. After the bone defect was made, the soft tissues were repositioned and sutured.

Pharmacological Treatment

Treatment started on the next day after the bone lesion. All ovariectomized rats were randomly divided into two groups: OVX (n=9) and OVX + SR (n=9). Animals in the OVX + SR group received SR (Protos®, Servier, Gidy, France) orally (625 mg/kg per day; aqueous suspension) by gavage (0.3 mL per animal). The drug was diluted in Milli-Q® Water (Millipore, Molsheim, France) and the suspension was administered immediately after it was prepared. A curved stainless-steel needle (length: 38 mm) for gavage of rats was used. An intramuscular injection of 2% xylazine hydrochloride, in the proportion of 10% ketamine hydrochloride, was made to induce general anesthesia. After procedures, the animals were euthanized four weeks after the bone defect surgery. After this procedure, the femur was removed from each animal and excess soft tissue was removed. A scraping with a scalpel blade was made on the cortical surface near the bone defect to obtain samples for spectroscopy. These samples were frozen and then lyophilized for 24 h. The bone specimens were fixed in formaldehyde solution (4%; phosphate buffer, 0.01 M). Ovariectomy was evaluated at necropsy by confirming uterine horn atrophy.

Fourier Transform Infrared (FTIR) Spectroscopy

Potassium bromide (KBr; 150 mg) pellets containing lyophilized cortical bone tissue (1 mg) were prepared. Spectra (transmission mode; 4 cm\(^{-1}\) resolution; 12 scans) in the range 4000-400 cm\(^{-1}\) were obtained at controlled temperature (18-20 °C). The Spectrum GX FT-IR (Perkin Elmer, Boston, MA, USA) spectrometer was used. The spectra were preprocessed with the Spectrum 5.2 (Perkin Elmer) software for baseline correction and normalization. Spectral analyzes were performed using the absorbance mode.

The mineral/matrix proportion was calculated as the ratio between the phosphate (1180-905 cm\(^{-1}\)) and amide I + II amide (1721-1520 cm\(^{-1}\)) band areas (13). The crystallinity index (CI) was calculated as the ratio between intensities of the 1057 and 1023 cm\(^{-1}\) sub-bands obtained by curve fitting in the 1200-900 cm\(^{-1}\) region (14). Maturation of collagen was estimated as the ratio between the areas of amide I sub-bands on 1660 and 1690 cm\(^{-1}\) (15).

Curves were adjusted using the Origin Pro 8.0 (Origin®, Northampton, MA, USA) software based on information about position provided by deconvolution and second derivative. These adjustments were made using the average spectrum and Gaussian function for calculation of CI (six points) and amide I (five points).

Energy-Dispersive X-ray Spectroscopy (EDS)

Presence or absence of Sr atoms incorporated into bone tissue in the OVX + SR group and the amount of calcium (Ca) and phosphorus (P) atoms present in bone tissue samples from all groups were observed by scanning electron microscopy (model EVO-MA40; Zeiss, Jena, Germany) with EDS. For analysis by EDS, lyophilized bone scrapings were placed under a double-sided carbon adhesive tape and observed without any coating.

X-ray Computed Microtomography (µCT)

Repair of bone defects in femurs was evaluated using the Skyscan (model 1172, Kontich, Belgium) microscanner and the NRecon, DataViewer, CT and CT-Analyzer-Vol software provided by the scanner manufacturer. The images were obtained under the following conditions: voltage: 60 kV; tube current: 165 µA; aluminum filter; thickness: 0.5 mm; rotation: 180°; angular increment: 0.6° and acquisition of the image with a CCD camera (10 Mp, 8 radiographic projections per rotation; resolution: 5 µm). The volume of newly formed bone within the defect was defined as volume of interest (VOI), and the volume of the medullary canal and the original bone tissue were excluded from the evaluation. Within VOI after segmentation, the following parameters were measured: bone volume (BV), relative bone volume (BV/TV), and number (Tb.N), thickness (Tb.Th) and separation (Tb.Sp) of trabeculae.

Statistical Analysis

Data were presented as mean and standard deviation. Statistical analyzes were performed using the R 3.1.0 (R Core Team, 2014; Vienna, Austria) software. Differences between groups were examined using analysis of variance...
(one-way ANOVA). When effects showed to be significant (p<0.05), multiple comparisons were performed by the Tukey’s test. Differences were considered significant for all tests with p<0.05.

Results

FTIR

Figure 1 shows the average spectra for each experimental group. Statistical analysis showed that the ratio between areas of the phosphate band and those of amide I and II showed no significant difference between groups (p=0.23) (Table 1). In the calculation of CI, the values for ratio between intensities of the 1057 and 1023 cm\(^{-1}\) bands were 1.024, 1.015, and 1.108 for the SHAM, OVX + SR and OVX groups, respectively (Fig. 2). Figure 3 shows a spectrum with its underlying bands solved by second derivative and curve-fitting techniques for the amide I band. Two of the underlying bands (1660 and 1690 cm\(^{-1}\)) are of particular interest because they correspond to the collagen crosslinking pyridinoline (PYR) and dihydroxylysinoornithine (DHLNL), respectively. The ratio PYR/DHLNL was 3.502, 3.355, and 2.514 in the SHAM, OVX and OVX + SR groups, respectively.

EDS

P and Ca elements showed to be the main constituents in the samples of the three groups. A significant difference

Table 1. Mean values of the variables, by group

<table>
<thead>
<tr>
<th>Variables</th>
<th>OVX</th>
<th>OVX+SR</th>
<th>SHAM</th>
<th>Total</th>
<th>p values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Phosphate/Amide I and II</td>
<td>4.05</td>
<td>0.27</td>
<td>4.09</td>
<td>0.08</td>
<td>3.82</td>
</tr>
<tr>
<td>P</td>
<td>21.2</td>
<td>8.92</td>
<td>20.2</td>
<td>5.82</td>
<td>26.88</td>
</tr>
<tr>
<td>Ca</td>
<td>81.4</td>
<td>15</td>
<td>47.42</td>
<td>11.01</td>
<td>81.02</td>
</tr>
<tr>
<td>Sr</td>
<td>0.18</td>
<td>0.34</td>
<td>1.36</td>
<td>0.79</td>
<td>0.17</td>
</tr>
</tbody>
</table>
was observed for the Ca and Sr elements in the OVX + SR group (p=0.009), with a decrease in the first and an increase in the second (Table 1).

**μCT**

Figure 4 shows two-dimensional micro-CT images of cross sections in the central region of bone defect in one animal of each group. In Table 2, the quantitative results obtained with micro-CT of bone defects were expressed as BV, BV/TV, Tb.Th, Tb.N and Tb.Sp. In the OVX group, all parameters showed lower values, except Tb.N. As the differences for BV, BV/TV and Tb.Th were significant (p<0.05), Tukey multiple comparisons were performed, and the value for p was set to identify pairs with significant differences.

Significant differences were observed between the SHAM and OVX groups when the BV and BV/TV variables were compared. In the SHAM group, BV were higher than those in the OVX group (p=0.014) and BV/TV were about 8% higher than in the OVX group (p=0.015). Comparison between the SHAM and OVX + SR groups showed borderline values for these variables. Regarding Tb.Th, a difference was observed between the OVX + SR and OVX groups. In the OVX + SR group, values for Tb.Th were about 1.8% higher than those in the OVX group (p=0.049). No statistically significant difference was observed between the SHAM and OVX + SR groups.

**Discussion**

The results of this study indicate that ovariectomy impaired bone microarchitecture and quality of bone tissue. This conclusion is consistent with results published in other studies in which this model was used (5,16,17). Ovariectomized animals exhibited the lowest averages in almost all morphological parameters (Table 2).

Bone microarchitecture, except Tb.N (p=0.092) and Tb.Sp (p=0.061), was markedly different among the experimental groups. As Tb.N remained constant, microarchitecture did not differ in terms of topology (Table 2). The OVX + SR group showed intermediate values for the BV and BV/TV variables.
but it had a significant increase in Tb.Th compared with the OVX group. Similar data were observed in another study in which the tibial fracture model was used. Four weeks after fracture, treatment with SR significantly increased the values for BV, TV, Tb.Th, Tb.N and BV/TV compared with the non-treated OVX group (5). Strontium is able to prevent changes in estrogen deficiency-induced bone turnover (18). Thus, inhibition of excessive bone resorption and the promotion of bone formation by SR in the initial period of bone repair may have contributed to these results.

SR therapy was well tolerated and safe and no adverse effect was observed in animals that received medication. This was also reported in a previous publication, in which the authors treated rats with 125, 250, or 625 mg/kg per day of SR and they did not observe any adverse effect on body weight gain (17).

The animals were fed a diet with normal Ca concentration (1%) and this is required for a proper therapeutic interpretation, as Sr serum levels, and thus the Sr bone levels, are influenced by Ca dietary levels (16). A SR dose equal to 625 mg/kg per day was used. Such dose in animals fed a normal Ca diet can lead to a Sr serum concentration similar to that observed in patients treated with a therapeutic dose of 2 g per day (5). The drug was administered orally in an aqueous suspension and the animals were not deprived of food. Since rats have nocturnal habits and eat mostly at night, gavage was carried out in the morning to reduce the possibility of the ration to hinder Sr absorption.

In the EDS analysis, Sr incorporation in bone samples was observed in the OVX + SR group. Such incorporation of Sr was reported to occur mainly in newly formed bone during treatment with SR (19). Regarding Ca concentration in the mineralized matrix, the OVX + SR group showed the lowest mean values among the three groups. This may have occurred because Sr uptake by bone mineral crystals may occur by ion exchange on the surface or substitution of Ca ions by Sr ions in the bone (20,21).

The effects of ovariectomy and treatment with SR on cortical bone tissue molecular components were also assessed in this study. Geometry and material properties determine the mechanical strength of bone tissue. Such properties include mineral content, matrix mineral composition, cellular activity, and crystal size and distribution. Most of these material properties, except cellular activity, can be determined by infrared spectroscopy (22).

The study of bone chemistry is hampered by differences between responses of bone trabecular and cortical regions to diseases and medications (23). In the present study with FTIR, were used KBr pellets made with scrapings from bone tissue cortical surface. The Ci, as measured by the technique of curve fitting in the phosphate band region, increased about 10% in the OVX group. The OVX + SR group showed a value close to that of the SHAM group, indicating that the treatment maintained bone crystallinity near to that of the control group and this effect is beneficial to the tissue. Studies show that an increase of this index is associated with osteoporotic bone fragility (9,11). Increase in bone crystallinity increases its risk of fracture (13).

The ratio between inorganic and organic components showed no significant difference among groups. Huang et al. (23) used monkeys in their study and observed an increase in the ratio between the phosphate and amide I formed in cortical bone after ovariectomy, whereas trabecular bone did not show any change in phosphate content.

Regarding analysis of collagen maturation, the areas of amide I sub-bands at 1660 and 1690 cm\(^{-1}\) were calculated. The percent ratio of relative areas of these sub-bands is related to enzymatic mature (PYR) and immature (DHLNL) collagen cross-links, which are abundant in mineralized tissues (15). The results showed that treatment with SR caused a decrease in this ratio, as compared with other groups. Abnormalities in enzymatic and non-enzymatic collagen cross-links affect the mineralization process and may lead to microdamage accumulation (24). Gourion-Arisquaud et al. (13) used FTIR to analyze biopsy of the iliac crest in women with and without fracture and found that the value for PYD/DHLNL was significantly higher in patients with fracture. Farlay et al. (25) used mice with a characterized decrease in collagen cross-linking after inhibition of lysyl oxidase. They used FTIR to determine the ratio 1660 and 1690 cm\(^{-1}\) in bone tissue and compared it with collagen cross-links as determined by liquid chromatography. They concluded that this ratio is unrelated to collagen cross-linking but it increases with bone mineral age, suggesting that a modification of this ratio could be mainly due to a modification of the collagen secondary structure related to the mineralization process. However, the authors did not use second derivative spectroscopy or deconvolution methods to determine the number and position of the underlying peaks of the amide I band.

In conclusion, the present study demonstrated that systemic treatment with SR promoted bone repair and improved microarchitecture of the newly formed bone within the defect. However, the results of the cortical surface analyzes near the bone defect with FTIR data suggest that Sr decreased cross-linking of mature collagen.

**Resumo**

Este estudo foi conduzido para avaliar os efeitos do tratamento com ranelato de estrôncio (RE) na reparação de defeitos ósseos e componentes moleculares de osso nos fêmures. Ratas adultas (n = 27) foram submetidas a ovariectomia (OVX) ou cirurgia Sham. Trinta dias após a cirurgia, um defeito foi feito no fêmur e os animais foram então divididos em três...
groups: OVX, SHAM e OVX+RE. A eutanásia foi realizada quatro semanas após a cirurgia de preparo do defeito ósseo. A reparação do defeito ósseo foi avaliada por microtomografia computorizada (µCT) e a composição química do osso cortical foi analisada por espectroscopia de infravermelho de transformada de Fourier (FTIR) e espectroscopia por energia dispersiva de raios X (EDS). A espessura do osso trabecular (Tb.Th) recém formado no grupo OVX+SR foi significativamente maior que a do grupo OVX. A maturidade do colágeno no grupo OVX+SR foi menor do que nos outros dois grupos. Neste grupo, observou-se também um aumento significativo na quantidade de estrôncio (Sr) e uma diminuição na quantidade de cálcio (Ca) no tecido ósseo. O tratamento sistêmico com RE melhorou a microarquitetura do osso recém formado dentro do defeito, mas diminuiu a reticulação do colágeno maduro no osso cortical.

References


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Strontium ranelate effects in cortical bone