ROLE OF CARNOSINE IN PREVENTION AND TREATMENT OF OSTEOPOROTIC LUMBAR VERTEBRAE OF OVARIECTOMIZED HAMSTERS

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INTRODUCTION

Estrogen plays an important role in the growth and maturation of bone as well as in the regulation of its turnover to maintain bone balance in adults. During bone growth, estrogen is needed for proper closure of epiphyseal plates both in females and males (Vaananen and Harkonen, 1996). In the adult, both bone formation and resorption occur side by side in a process called remodeling where bone formation equals the amount of that lost during resorption. With aging, resorption becomes more active than formation with a net loss of bone mass (Lindsay et al., 1980). Estrogen withdrawal as a result of menopause or induced by ovariectomy leads to osteoporosis as a result of a decrease in bone formation and an increase in bone resorption (Sun et al., 1997; Chen et al., 2000). However, the decrease in bone formation may precede the increase in bone resorption (Seifert-Klauss et al., 2002). The mechanisms which have been suggested for the development of osteoporosis following estrogen withdrawal include stimulation of the production of bone resorbing cytokines which stimulates osteoclast formation in bone marrow (Vaananen and Harkonen, 1996), reduction of osteoprotegerin in the osteoblasts resulting in stimulation of bone resorption (Liao et al., 2002) and reduction of the life-span of osteoblasts and increase of the life-span of osteoclasts (Chen et al., 2000).
Certain medications are in therapeutic use to prevent fractures in women with estrogen deficiency. These included estrogen replacement therapy, raloxifene, nasal calcitonin, bisphosphonates and parathyroid hormone (Watts, 1999). Although, theoretically, the most appropriate therapy for osteoporosis in younger women is estrogen replacement therapy (Castelo-Branco, 1998; Watts, 1999), yet the trial-based evidence of the ability of hormone replacement therapy (HRT) to prevent fractures is not strong. Moreover, the menstrual bleeding and increased risk of endometriosis, endometrial cancer and breast cancer are among the factors that lead to limitation of the use of HRT (Gibaldi, 1997; Colditz, 1998).

Carnosine (β-alanyl-L-histidine) is a dipeptide abundantly present in the skeletal muscles and in the nervous tissues of the vertebrates including humans (Boldyrev and Severin, 1990; Abe, 1991). Carnosine was proved to aid the recovery of fatigued muscle, to act as a membrane-stabilizing agent, to possess antioxidant activity and to have certain therapeutic properties (Perelman et al., 1989; MacFarlane et al., 1991; Bogardus et al., 1992). Many researches pointed to the wide safety of carnosine. No toxic effects of carnosine were observed even at concentrations of up to 500 mg / kg body weight (Ermakova et al., 1988). In addition, Soliman and Abdel Monem (2001) recorded carnosine LD 50 in mice as 18.5 gm / kg, a fact pointing to its safety.

A specific biological action of carnosine that may be of orthopedic importance is that it can uniquely chelate zinc ion forming β-alanyl-L-histidinato zinc (AHZ) that has more intensive effect than zinc sulfate on bone formation suggesting its role in treatment of osteoporosis (Kishi & Yamaguchi, 1994 - a & b; Yamaguchi, 1995; Yamaguchi & Kishi, 1995 - a). The dipeptide of AHZ may be useful in the penetration of zinc ions into marrow cells with stimulation of the osteoblastic cells (Hashizume & Yamaguchi, 1994; Yamaguchi et al., 1994; Yamaguchi and Matsui, 1997) and inhibition of the osteoclastic cell formation (Yamaguchi, 1995; Yamaguchi & Kishi, 1995 - a).

The aim of this work was to evaluate the degree of efficiency of the carnosine in prevention and treatment of osteoporosis through a histological and histomorphometric study of its effect on lumbar vertebrae in ovariectomized hamsters.

**MATERIALS AND METHODS**

Thirty-three adult female hamsters weighing 105 - 120 gms were used in this study. They were housed in cages, five animals each, under good hygienic
conditions and food and water were allowed ad libitum. The animals were divided into four groups as follows:

Group I (sham-operated group; control group; n = 9):
the animals were subjected to sham operation by mobilizing the ovaries of both sides without doing ovariectomy.

Group II (ovariectomy only; n = 9):
bilateral ovariectomy was performed for the animals of this group.

Group III (ovariectomy with early carnosine treatment; n = 9):
bilateral ovariectomy was performed for these animals followed by a daily dose of intramuscular injection of carnosine (10 mg / kg body weight), starting from the next day after the operation.

Group IV (ovariectomy with late carnosine treatment; n = 6):
the animals were subjected to bilateral ovariectomy. Four weeks after the operation, the animals started to receive a daily dose of intramuscular injection of carnosine (10 mg / kg body weight).

In each group, the animals were anaesthetized by intramuscular injection of 0.5 ml of sodium thiopental of concentration of 1 gm / 20 ml of distilled water and a midline abdominal incision was performed. Both ovaries were identified and mobilized with great care to avoid injuring the surrounding structures. Bilateral ovariectomy was done in groups II, III and IV. After the operation, the incision was closed by interrupted 0.5-silk sutures and the wound was sprayed by antibiotic powder.

The animals of each group were killed - three animals at a time - by intraperitoneal injection of 1 ml of sodium thiopental at the specified intervals of 4, 6 and 8 weeks, from the date of the operation, in groups I, II and III and at intervals of 6 and 8 weeks, from the date of the operation, in group IV. The lumbar vertebrae were removed from each animal and fixed in 10% formol saline. The specimens were decalcified using 2% formic acid, and then paraffin blocks were prepared. Longitudinal sections of 7 μm were cut stained with Hx & E and Masson's trichrome (Masson, 1924) stains.

Morphometric quantification:
The areas of bone trabeculae (Ab, in μm²) and their perimeters (Pb, in μm) were measured using the binary image of the image-analyzer computer assisted by
the software Leica Qwin 500 with a standard measuring frame (At) of 720240.8 μm². These data were measured in 10 fields of each specimen and the mean values were obtained. The following parameters were calculated using the following equations, according to Parfitt et al. (1983):

- Trabecular bone volume (TBV) % = (Ab / At) × 100.
- Mean trabecular plate thickness (MTPT; μm) = (2.000 / 1.199) × (Ab / Pb).
- Mean trabecular plate density (MTPD; per millimeter) = (1.199 / 2.000) × (Pb / At).

Statistical analysis (Table):

The Statistical Package for the Social Sciences (SPSS version 7.5) was used in data analysis. Data were expressed as mean ± SE. One-way analysis of variance (ANOVA) was used. The percentage of reduction compared to the control was calculated as follows:

\[ \% = \left( \frac{\text{treated} - \text{control}}{\text{control}} \right) \times 100. \]

RESULTS

Morphometric study (Table; Figs. 1, 2, 3):

In group II, where the animals underwent ovariectomy only, the measurements of each of TBV%, MTPD and MTPT showed reduction in their values compared with those of the control group I as well as with groups III and IV, along the whole period of the experiment. Comparing to the control group, the reduction in the values of TBV% and MTPT were statistically significant (P < 0.05), at the 4th week of the operation and statistically highly significant (P < 0.01) at the other period intervals. On the other hand, the reduction in MTPD became statistically significant (P < 0.05) from the 6th week onwards.

The measurements of TBV%, MTPD and MTPT of group III (ovariectomy + early carnosine treatment) were higher than those of group II but still showed reduction in their values compared with the control group. The reduction of the values of measurements of MTPD was statistically insignificant (P > 0.05) along the whole period of the experiment while the reduction of those of TBV% and MTPT were statistically insignificant at the 4th week from the date of the operation then statistically significant at the other intervals.

In group IV (ovariectomy + late carnosine treatment), the measurements of TBV% and MTPD were higher than those of group II but lower than those of groups
I and III. Comparing with the control group, the reduction in the values of the measurements of $\text{TBV}\%$ were statistically significant along the whole period of the experiment while the reduction of the measurements of MTPD were statistically significant at the 8th week of the operation. On the other hand, the values of measurements of MTPT were higher than those of groups II and III and lower than those of group I, where the reduction was statistically significant, along the whole period of the experiment.

There was an overall variation significance among the different animal groups, which became more significant with the increased time interval as indicated from the greater numerical value of F-ratio by the 8th week from the date of the operation.
Table: Measurements of the trabecular bone volume percent (TBV %), mean trabecular plate density (MTPD / mm³) and mean trabecular plate thickness (MTPT, μm) in the different experimental groups.

<table>
<thead>
<tr>
<th>weeks</th>
<th>Groups</th>
<th>TBV % Mean ± SE</th>
<th>MTPD / mm³ Mean ± SE</th>
<th>MTPT (μm) Mean ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 weeks</td>
<td>Control</td>
<td>42.44 ± 2.37</td>
<td>7.46 ± 0.88</td>
<td>57.43 ± 1.52</td>
</tr>
<tr>
<td></td>
<td>Group II PR</td>
<td>35.11 ± 2.83</td>
<td>7.16 ± 0.73</td>
<td>50.16 ± 2.21</td>
</tr>
<tr>
<td></td>
<td>Group III PR</td>
<td>42.09 ± 6.17</td>
<td>7.44 ± 0.68</td>
<td>55.52 ± 3.09</td>
</tr>
<tr>
<td>F - ratio</td>
<td>0.47</td>
<td>0.95</td>
<td>0.27</td>
<td></td>
</tr>
<tr>
<td>6 weeks</td>
<td>Control</td>
<td>42.44 ± 2.37</td>
<td>7.46 ± 0.88</td>
<td>57.43 ± 2.52</td>
</tr>
<tr>
<td></td>
<td>Group II PR</td>
<td>26.33 ± 2.70</td>
<td>6.24 ± 0.41</td>
<td>41.97 ± 2.95</td>
</tr>
<tr>
<td></td>
<td>Group III PR</td>
<td>32.14 ± 2.17</td>
<td>7.18 ± 0.50</td>
<td>44.83 ± 3.47</td>
</tr>
<tr>
<td></td>
<td>Group IV PR</td>
<td>29.83 ± 3.77</td>
<td>6.50 ± 0.85</td>
<td>45.82 ± 5.17</td>
</tr>
<tr>
<td>F - ratio</td>
<td>2.69</td>
<td>1.66</td>
<td>2.74</td>
<td></td>
</tr>
<tr>
<td>8 weeks</td>
<td>Control</td>
<td>42.44 ± 2.37</td>
<td>7.46 ± 0.88</td>
<td>57.43 ± 1.52</td>
</tr>
<tr>
<td></td>
<td>Group II PR</td>
<td>18.01 ± 4.19</td>
<td>4.82 ± 0.58</td>
<td>37.65 ± 4.12</td>
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<tr>
<td></td>
<td>Group III PR</td>
<td>29.36 ± 4.51</td>
<td>6.97 ± 1.64</td>
<td>42.27 ± 5.76</td>
</tr>
<tr>
<td></td>
<td>Group IV PR</td>
<td>28.18 ± 2.71</td>
<td>5.36 ± 0.41</td>
<td>52.46 ± 1.21</td>
</tr>
<tr>
<td>F - ratio</td>
<td>4.64 #</td>
<td>1.25</td>
<td>8.84 ##</td>
<td></td>
</tr>
</tbody>
</table>

PR: Percentage of reduction compared to the control.
SE: Standard error
*: Significant with respect to the control group (p < 0.05)
**: Highly significant with respect to the control group (p < 0.01)
#: Significant at p < 0.05 (ANOVA)
##: Significant at p < 0.01 (ANOVA).

Note: The F - ratio indicates the overall significance among the different animal groups. The greater the numerical value of the F - ratio the more significant the difference is.
Fig (1): A histogram showing:
A) The trabecular bone volume (TBV %).
B) Mean trabecular plate density (MTPD / mm³).
C) Mean trabecular plate thickness (MTPT, μm).
All measurements are demonstrated in the different experimental groups 4 weeks after the operation.
* Significant with respect to the control group (p < 0.05).
Fig (2) : A histogram showing:
A) The trabecular bone volume (TBV %).
B) Mean trabecular plate density (MTPD / mm³).
C) Mean trabecular plate thickness (MTPT, μm)
All measurements are demonstrated in the different experimental groups 6 weeks after the operation.
* Significant with respect to the control group (p < 0.05).
** Highly significant with respect to the control group (p < 0.01).
Fig (3): A histogram showing:
A) The trabecular bone volume (TBV %).
B) Mean trabecular plate density (MTPD / mm³).
C) Mean trabecular plate thickness (MTPT, μm).
All measurements are demonstrated in the different experimental groups 8 weeks after the operation.
* Significant with respect to the control group (p < 0.05).
** Highly significant with respect to the control group (p < 0.01).
Histological study:

Group I (sham-operated, control group):

Longitudinal sections of the lumbar vertebrae showed a thick shell of compact bone with prominent Haversian system (Fig. 4). The bone trabeculae were thick and arranged in vertical and transverse directions with good intertrabecular connectivity (Fig. 5). Well-aligned and intact cement lines could be seen in both compact and trabecular bones (Figs. 4 & 5).

Ovariectomized groups:

Four weeks post-operative:

* Group II (ovariectomy only):

The histological picture of the lumbar vertebrae of these animals showed a thin shell of compact bone containing few osteoporotic cavities (Fig. 6). A number of osteoclasts eroding the inner aspect of the compact bone could be seen (Fig. 7). The bone trabeculae were thin and fewer in number - compared with those of the control group - and arranged mainly in a vertical direction with many free ends and decreased intertrabecular connectivity (Fig. 8). Areas containing cartilaginous matrix and cartilage cells were demonstrated in many of bone trabeculae (Figs. 8, 9 - a& b). The bone marrow cavities were wide and contained a number of fat cells (Fig. 8).

* Group III (ovariectomy with early carnosine treatment):

The lumbar vertebrae of the animals of this group showed a thick shell of compact bone with intact cement lines (Fig. 10). The bone trabeculae were thick and arranged in vertical and transverse directions with good intertrabecular connectivity (Fig. 11). The bone trabeculae demonstrated few numbers of osteoporotic cavities (Fig. 11), many osteocytes and well-formed cement lines (Fig. 12).

Six weeks post-operative:

* Group II (ovariectomy only):

The histological picture of the lumbar vertebrae of these animals showed osteoporotic cavities within a thin shell of compact bone (Fig. 13). There were few thin bone trabeculae arranged mainly vertically with poor intertrabecular connectivity and wide bone marrow cavities (Fig. 13). Some of the trabeculae formed a thin shell around large mass of cartilage cells (Fig. 13).
* Group III (ovariectomy with early carnosine treatment) :

The lumbar vertebrae of this group showed thick bone trabeculae arranged both horizontally and vertically with good intertrabecular connectivity (Fig. 14). Cleavage of cement line (Fig. 14) and sites of osteoporotic cavities as well as few areas of cartilaginous matrix could be seen within some bone trabeculae (Figs. 14, 15). Proliferation of the cartilaginous plate was detected (Fig. 15) indicating active process of bone formation.

* Group IV (ovariectomy with late carnosine treatment) :

The histological picture of the lumbar vertebrae of this group showed many osteocytes within a relatively thin shell of compact bone, compared with that of the control group, (Fig. 16). The bone trabeculae were mainly arranged in vertical direction with decreased intertrabecular connectivity. The trabeculae demonstrated cartilaginous areas as well as few osteoporotic cavities (Fig. 17).

Eight weeks post-operative :

* Group II (ovariectomy only) :

The lumbar vertebrae of these animals still showed a thin shell of compact bone with some osteoporotic cavities. The bone trabeculae were mainly arranged vertically and showed large cartilaginous areas and sites of osteoporotic cavities. There was much reduction of the intertrabecular connectivity and increased trabecular free ends with widening of the bone marrow cavities (Fig. 18). Many osteoclastic cells were demonstrated, either imbedded inside the bone trabeculae or eroding their surfaces (Figs. 19 - a & b).

* Group III (ovariectomy with early carnosine treatment) :

By the end of the 8th week following the date of the operation, the compact bone shell of the lumbar vertebrae of these animals was still thick; having similar thickness to that of the control group with large number of osteocytes (Fig. 20). The bone trabeculae were arranged both vertically and horizontally with intertrabecular connectivity and few trabecular free ends. Small cartilaginous areas as well as few osteoporotic cavities could be seen within the trabecular bone (Fig. 21). Active process of bone formation was indicated by the appearance of osteocytes within the areas of the cartilaginous matrix occupying the bone trabeculae (Fig. 22).
**Group IV (ovariectomy with late carnosine treatment):**

The histological picture of the lumbar vertebrae of these animals showed bone trabeculae arranged mainly vertically and decreased intertrabecular connectivity with few trabecular free ends. Small areas of cartilaginous matrix and several osteoporotic cavities were also demonstrated within the bone trabeculae (Fig. 23). Similar to those of group III, osteocytes were seen within the areas of cartilaginous matrix occupying the bone trabeculae.
Fig. (6): A photomicrograph of a longitudinal section in a lumbar vertebra of group II (underwent ovariectomy only), 4 weeks after the operation, showing thin shell of compact bone containing osteoporotic cavities (o). Note the periostial-covering layer (arrow).

(Hx. & E.; x 100)

Fig. (7): A photomicrograph of a longitudinal section in a lumbar vertebra of group II (underwent ovariectomy only), 4 weeks after the operation, showing an osteoclast (OC) eroding the inner aspect of the compact bone (CB). Note the cartilage cells (CC) within a bone trabecula on the left side of the picture.

(Hx. & E.; x 400)
Fig. (8): A photomicrograph of a longitudinal section in a lumbar vertebra of group II (underwent ovariectomy only). 4 weeks after the operation, showing thin bone trabeculae which are few in number and arranged mainly in a vertical direction with many trabecular free ends. Decreased trabecular connectivity and areas containing cartilaginous matrix (arrows) within the bone trabeculae are clearly seen. Note the wide bone marrow cavities containing a number of fat cells (F) and the cartilaginous plate (PT).

(Hx. & E.; x 40)
Figs. (9 a & b) : Photomicrographs of different fields of a longitudinal section in a lumbar vertebra of group II (underwent ovariectomy only), 4 weeks after the operation, showing:

a) Cartilage matrix (CM) within a bone trabecula (TB). (Hx. & E.; x 400)

b) Cartilage cells (CC) within a bone trabecula (TB). (Hx. & E.; x 400)
Fig. (10): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early carnosine treatment), 4 weeks after the operation, showing a thick shell of compact bone (C) with intact cement line (arrow). Note the thick covering layer of periosteum (P) with muscle fibers (M) attached to it.

(Hx. & E.; x 100)

Fig. (11): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early carnosine treatment), 4 weeks after the operation, showing thick bone trabeculae arranged in vertical and transverse directions with good intertrabecular connectivity. Few osteoporotic cavities (o) can be seen within the bone trabeculae.

(Hx. & E.; x 40)
Fig. (12) : A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early camosine treatment), 4 weeks after the operation, showing one of the bone trabeculae with many osteocytes (os) and intact cement lines (crossed arrows).

(Hx. & E.: x 100)

Fig. (13) : A photomicrograph of a longitudinal section in a lumbar vertebra of group II (underwent ovariectomy only), 6 weeks after the operation, showing an osteoporotic cavity (o) within a thin shell of compact bone (CB). Few thin bone trabeculae arranged mainly in vertical direction with poor intra trabecular connectivity and wide bone marrow cavities can be seen. Note the thin layer of trabecular bone (TB) surrounding a large mass of cartilage cells (CC).

(Hx. & E.; x 40)
Fig. (14): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early carnosine treatment), 6 weeks after the operation, showing thick bone trabeculae arranged in transverse and vertical directions with good intertrabecular connectivity. Osteoporotic cavities (crossed arrows), cleavage of cement lines (arrows) and few areas of cartilaginous matrix (CM) can be clearly seen within the bone trabeculae.
(Hx. & E.; x 40)

Fig. (15): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early carnosine treatment), 6 weeks after the operation, showing proliferation of the cartilaginous plate (PT). Note the osteoporotic cavity (o) and areas of cartilaginous matrix (CM) within a bone trabecula.
(Hx. & E.; x 100)
Fig. (16) : A photomicrograph of a longitudinal section in a lumbar vertebra of group IV (underwent ovarietomy with late camosine treatment), 6 weeks after the operation, showing large number of osteocytes within a relatively thin shell of compact bone.

(Hx. & E.; x 100)

Fig. (17) : A photomicrograph of a longitudinal section in a lumbar vertebra of group IV (underwent ovarietomy with late camosine treatment), 6 weeks after the operation, showing bone trabeculae arranged mainly in a vertical direction and containing cartilaginous areas (arrows) and few osteoporotic cavities (crossed arrows). Note the decreased intertrabecular connectivity.

(Hx. & E.; x 40)
Fig. (18) : A photomicrograph of a longitudinal section in a lumbar vertebra of group II (underwent ovarioectomy only), 8 weeks after the operation. The bone trabeculae are arranged mainly in a vertical direction and showed large cartilaginous areas (arrows) and sites of osteoporotic cavities (o). Much reduction of the intertrabecular connectivity, increased trabecular free ends and widening of the bone marrow cavities can be seen clearly.

(Masson's trichrome; x 40)
Figs. (19 a & b): Photomicrographs of different fields of a longitudinal section of a lumbar vertebra of group II (underwent ovariectomy only), 8 weeks after the operation, showing:

a) Osteoclast cells (oc) embedded inside a bone trabecula (TB). Note the osteoblasts (OB) on the surface of the bone trabecula and the part of the cartilaginous plate (PT).

(Hx. & E.; x 400)

b) Osteoclast cells (oc) eroding a cartilaginous surface of a bone trabecula.

(Hx. & E.; x 400)
Fig. (20): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early camosine treatment), 8 weeks after the operation, showing a thick shell of compact bone with large number of osteocytes (arrows).

(Hx. & E.; x 100)

Fig. (21): A photomicrograph of a longitudinal section in a lumbar vertebra of group III (underwent ovariectomy with early camosine treatment), 8 weeks after the operation, showing bone trabeculae arranged in both vertical and horizontal directions with intertrabecular connectivity and few trabecular free ends. Small cartilaginous areas (arrows) as well as osteoporotic cavities (crossed arrows) can be seen within the bone trabeculae.

(Hx. & E.; x 40)
Fig. (22): A higher magnification of a different field of the previous section showing osteocytes (arrows) within a cartilaginous area (CM) occupying one of the bone trabeculae.

(Hx. & E.; x 400)

Fig. (23): A photomicrograph of a longitudinal section in a lumbar vertebra of group IV (underwent ovariectomy with late carnosine treatment), 8 weeks after the operation, showing bone trabeculae arranged mainly vertically. Small cartilaginous areas (crossed arrows) and several osteoporotic cavities (arrows) can be seen within the bone trabeculae. Note the decreased intertrabecular connectivity and the few trabecular free ends.

(Hx. & E.; x 40)

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DISCUSSION

Bone loss already occurs in young adult women or men. However, the rate of loss is slow because the remodeling rate is low in this young age while the loss accelerates in women at menopause because the remodeling intensity increases and the number of basic multicellular units becomes more negative with reduction of the mineral content of bone tissue (Riggs, 2002).

In the present work, bilateral ovariectomy flared up osteoporotic changes which, to a great extent, imitated the senile and postmenopausal osteoporotic alteration. The operation led to cancellous as well as compact bone loss of the lumbar vertebrae and its severity increased with time lapse. The compact bone showed enhanced endocortical resorption as well as increased intracortical porosity. In agreement, Vaananen and Harkonen (1996) studying the effect of estrogen deficiency on young skeletons as well as on those following menopause, obtained the same findings. Furthermore, Vaananen and Harkonen (1996) emphasized that the endocortical resorption represented the first response of estrogen withdrawal and preceded the intracortical porosity. In the untreated ovariectomized animals (group II) of the current work, the presence of a few number of intracortical osteoporotic cavities with the early appearance of the osteoclasts eroding the inner surface of the compact bone resulting in its considerable thinning, 4 weeks post-operatively, confirms this view of Vaananen and Harkonen (1996). Concerning the cancellous bone, the vertically directed thin bone trabeculae containing many osteoporotic cavities with loss of the inter trabecular connectivity, the increased trabecular free ends and the widening of the bone marrow cavities were the common histological findings and were equivalent to those obtained by Chow et al. (1992), Vaananen and Harkonen (1996), Kinney et al. (1998) and Chen et al. (2000). These previous researches pointed out that the decrease in the amount of the trabecular bone and their interconnection was the most serious manifestation of estrogen loss. They, further, attributed the decreased bone mass and reduced bone strength to this disturbed bone architecture because of penetrative bone resorption and microfractures.

With time lapse after ovariectomy, it was obvious that cancellous bone was much more affected than the compact one. This finding is consistent with the statement of World Health organization (1994) reporting that bone loss in postmenopausal osteoporosis was much slower in the cortical than the trabecular bone. Riggs et al. (1998) also assumed that there was a profound and accelerated bone loss mainly in cancellous bone during the first postmenopausal decade. Further, Riggs (2002)
correlated this bone loss to the abrupt loss of action of estrogen and the loss of its direct restraining effect on bone cell function.

Many mechanisms have been suggested for the development of osteoporosis following estrogen withdrawal. Vaananen and Harkonen (1996) attributed it mainly to increased osteoclast formation, whereas, Turner et al. (1987) and Wronski et al. (1988) postulated that the bone loss was the sequence of an increase in bone resorption associated with a slight increase in bone formation. Further, Sun et al. (1997), Chen et al. (2000) and Seifert-Klauss et al. (2002) ascribed the osteoporotic changes following estrogen withdrawal to inhibition of osteoblastic formation and stimulation of osteoclastic activity. Riggs (2002) pointed out that estrogen deficiency reduced osteoblast life-span and increased osteoclast life-span as well as reduced the mineral content of bone tissue. An additional statement by Seifert-Klauss et al. (2002) postulated that decreased bone formation may precede increased bone resorption. Comparable findings were previously presented by Gibaldi (1997) who deduced that estrogen withdrawal following ovariectomy induced a four-fold increase in the osteoclasts which erode the bone trabeculae leading to their perforation and eventually to their fading. This postulation perfectly explains the present increased number of osteoclasts eroding both the compact and trabecular bones as well as the marked reduction in the bone trabecular interconnection, density and thickness observed in the ovariectomized animals (group II). The increased activity of the osteoclasts as result of estrogen withdrawal was explained by Vaananen and Harkonen (1996) who suggested that estrogen withdrawal stimulates the production of bone resorbing cytokines which regulate osteoclast formation in the bone marrow microenvironment. However, Liao et al. (2002) attributed this to the reduction of the osteoprotegerin in the osteoblasts which has an inhibitory effect on bone resorption. In these ovariectomized animal specimens, the presence of large areas of cartilaginous matrix occupying the bone trabeculae with absence of signs of active process of bone formation are in favor of the assumption that estrogen withdrawal leads, as well, to inhibition of osteoblastic activity and bone formation (Sun et al., 1997; Chen et al., 2000; Seifert-Klauss et al., 2002). Whether the effect of estrogen on osteoblasts is by direct mechanisms is still uncertain (Vaananen and Harkonen, 1996).

In group III (early carnosine-treated animals), the large number of osteocytes within a thick shell of compact bone as well as the presence of thick bone trabeculae arranged both vertically and horizontally with good intertrabecular connections denoted the role of early treatment with carnosine following ovariectomy, in
conserving the normal bone architecture compared with those without treatment (group II). Equivalent observations were presented by Yamaguchi and Kishi (1993), Kishi and Yamaguchi (1994 - b) and Kishi et al. (1994). Yamaguchi and Kishi (1993) pointed out that prolonged administration of tested doses of 10, 30, 100 mg / kg / d carnosine could completely prevent bone loss in the ovariectomized rats. Moreover, the histomorphological study of Kishi et al. (1994) supported this view. In the current work, the presence of sites of cleavage of cement lines and osteoporotic cavities in the trabecular bone as well as reduction in TBV % demonstrated in carnosine-treated groups might contradict the view of Yamaguchi and Kishi (1993) of a complete preventive effect of carnosine on bone loss. However, the short-term administration of carnosine (8 weeks) and the use of its lowest tested dose (10 mg / kg / d) can explain this contradiction. In agreement, Kishi and Yamaguchi (1994 - b) deduced that the osteoporotic preventive effect of carnosine was concentration dependent.

Yamaguchi et al. (1994), Yamaguchi and Hashizume (1994 - a & b) and Sugiyama et al. (2000) found that beta-alanyl-L-histidine, which chelates zinc ions in various essential traces forming beta-alanyl-L-histidinato-zinc (AHZ), prevents bone loss following estrogen withdrawal through stimulation of bone formation and osteoblastic proliferation. Segawa et al. (1993), Yamaguchi and Kishi (1993), and Yamaguchi and Kishi (1994 - a & b) concluded that AHZ prevented deterioration of bone metabolism following ovariectomy by inhibiting the reduction of calcium and inorganic phosphorus concentrations in the serum as well as inhibiting the reduction of the alkaline phosphatase activity and calcium content of bone induced by various bone resorption factors. Further, Segawa et al. (1992) emphasized the ability of AHZ to prevent the development of deteriorating bone metabolism even in rats fed on low calcium and vitamin D-deficient diets. In the current work, the presence of large number of osteocytes within both compact and trabecular bones and signs of an active process of bone formation; proliferation of the cartilaginous plate and appearance of osteocytes within the cartilaginous areas of the bone trabeculae in carnosine treated groups are consistent with the postulation of the stimulatory effect of carnosine on bone formation.

The mechanism by which carnosine stimulates bone formation was proved by the finding of increased osteoblastic cell number, DNA content and protein concentration in the cells (Yamaguchi and Matsui, 1996). Furthermore, Soliman and Ali (2003) demonstrated that carnosine causes increase of the concentration of liver DNA and RNA as well as selective increase of serum proteins. This may add further
support to the possible ability of carnosine to stimulate bone protein building. The intermediary metabolism involved in this carnosine stimulatory action is through its direct effect on many cellular enzymes e.g. cellular alkaline phosphatase and protein kinase (Hashizume and Yamaguchi, 1994; Yamaguchi and Ehara, 1996; Yamaguchi and Matsui, 1997) and not through its inhibition to the effect of bone resorbing factors on osteoblastic cells (Yamaguchi and Hashizume, 1994 - a). In agreement, Yamaguchi and Hashizume (1994 - b) detected a marked increase in the concentrations of many cellular proteins, such as osteocalcin, insulin-like growth factor-I and transforming growth factor-beta, secreted from osteoblastic cells in the culture medium by the presence of AHZ. Furthermore, Hashizume and Yamaguchi (1994) and Yamaguchi et al. (1994) demonstrated that the other zinc-chelating compounds do not have an effect on cellular protein content. They attributed the potency of AHZ on this aspect to its dipeptide part (AH: carnosine) that may be useful in the penetration of zinc ions into marrow cells stimulating their protein synthesis.

In the current histological study, failure to detect any osteoclasts on the surfaces or within the compact or the trabecular bones in the two groups treated with carnosine (III & IV) versus to detection of many of them eroding the compact and trabecular bone in the untreated group (II), can lead to the assumption that carnosine may inhibit the osteoclastic bone resorption following estrogen withdrawal. Such assumption is in a full agreement with Kishi and Yamaguchi (1994 - a), Yamaguchi (1995) and Yamaguchi and Kishi (1996) who ascribed this phenomenon to the inhibitory effect of carnosine Zn compound (AHZ) on osteoclast-like cell formation, in mouse marrow culture in vitro, at the earlier stage of differentiation of marrow cells without affection of the osteoclastic number. In addition, Yamaguchi and Kishi (1995 - b) indicated that the transforming growth factor-beta has a stimulatory and an inhibitory effect on osteoclast-like cell formation in mouse marrow culture and that AHZ inhibited its stimulatory effect. Furthermore, Yamaguchi (1995) and Yamaguchi and Kishi (1995 - a) deduced that the preventive effect of AHZ on bone resorption in vivo is via its inhibitory action on the process of parathyroid hormone-induced protein kinase C activation, which increases the released cytoplasmic Ca^{2+} from its osteoclastic cell stores. The more intensive effect of AHZ than other zinc-chelating compounds was proved by Kishi and Yamaguchi (1994 - a) and Yamaguchi and Kishi (1995 - a and 1996) who attributed it to the beneficial effect of the dipeptide component of AHZ in the penetration of zinc ion into marrow cells influencing the process of protein kinase C activation.

From the present histomorphometric study, it was found that comparing with early carnosine treated group (III), the late carnosine treated animals (group IV)
showed an increase of the MTPT and a reduction of the MTPD, endorsed histologi­
cally with the decreased inter trabecular connectivity and the increased trabecular
free ends. This findings support the view of Parfitt et al. (1983) who stated that the
reduction of TBV as result of estrogen withdrawal occurs predominantly by a pro­
cess that removes entire structural elements of bone resulting in reduction of the
number of bone trabeculae (MTPD) with slight reduction of their thickness (MTPT)
and converting the continuous trabecular network into a discontinuous one. They
added that an increase in TBV as a result of the treatment of osteoporosis can occur
by a compensatory increase in the thickness of the existing trabeculae without re­
storing the lost ones, which formation stops after fusion of the epiphyseal plate i.e.
once trabecular plates lost, they can never be replaced. Furthermore, Lane et al.
(1999), and Jerome et al. (2001) pointed out that the compression strength of tra­
becular bone depends more on the preservation of connections between the structu­nal elements than on the amount of bone present. They added that the thickened tra­
beculae produced by treatment remain disconnected, so that increasing TBV even to
normal may not restore a biomechanically normal skeleton. Consequently, Lane et
al. (1999) deduced that prevention of bone loss is much more important than at­
tempting to repair the damage, once it has occurred, suggesting a prompt interven­
tion with anti-resorptive agents for preservation of the biomechanical features of the
bone.

From the present work, it is concluded that bilateral ovariectomy results in in­
crease of osteoclastic resorption activity and inhibition of osteoblastic bone forma­
tion with destruction of bone architecture and that carnosine has both a stimulatory
effect on bone formation as well as an inhibitory effect on bone resorption suggest­
ing its role as a pharmacological tool in treatment of osteoporosis. The early carno­
sine intervention is advisable for preservation of the normal bone structural ele­
ments. The complete osteoporotic preventive effect of higher doses of carnosine
needs further study.

SUMMARY

Four groups of adult female hamsters were used in this study, group I (control
group) exposed to sham operation, group II subjected to bilateral ovariectomy,
group III exposed to bilateral ovariectomy followed - next day - by a daily dose of
intramuscular injection of carnosine and group IV subjected to bilateral ovariectomy
followed - 4 weeks later - by a daily carnosine treatment.
The animals of groups I, II and III were sacrificed at intervals of 4, 6 and 8 weeks while those of group IV were sacrificed at intervals of 6 and 8 weeks from the date of the operation. The lumbar vertebrae were removed and prepared for histomorphometric and histological study. The histomorphometric quantification of lumbar vertebrae showed variable and statistically significant reduction of TBV %, MTPD and MTPT in animals of groups II, III, and IV. However, the highest reduction was in the animals exposed to bilateral ovariectomy only (group II). In early carnosine-treated group (III), the values of TBV %, MTPD were higher and those of the MTPT were lower than the corresponding values in late carnosine-treated one (IV).

The histological study revealed that ovariectomy resulted in reduction in the thickness of the compact as well as the trabecular bones. Both showed many osteoporotic cavities and osteoclasts eroding their surfaces. Large cartilaginous areas occupying the bone trabeculae, loss of trabecular connection resulting in increased trabecular free ends with disturbed trabecular architecture and widening of the bone marrow cavities showing fat cells were also demonstrated. Treatment with carnosine resulted in disappearance of the osteoclastic resorption activity and increase the number of osteocytes in the compact and trabecular bones as well as proliferation of the cartilaginous plate with the appearance of osteocytes within the cartilaginous areas occupying the bone trabeculae denoting active process of bone formation. However, early treatment with carnosine (group III) was more effective than the late treatment (Group IV) in preservation of compact bone shell thickness, intertrabecular connectivity and trabecular architecture with reduction of the trabecular free ends.

From the present work, it is concluded that bilateral ovariectomy results in marked increase of bone loss with destruction of bone architecture and that carnosine has both a stimulatory effect on bone formation as well as an inhibitory effect on bone resorption. Early carnosine intervention is recommended for better preservation of the normal bone structural elements.

REFERENCES


دور الكارنوزين في الوقاية والعلاج من المشاكل في الفترات القطنية
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في هذا البحث تم استخدام أربع مجموعات من أنثى الهمستر البالغة: المجموعة الأولى (المجموعة الضبابة) وقد أجريت لها عملية شام والمجموعة الثانية وقـد أجريت لها عملية استئصال للمبيضين والمجموعة الثالثة وقـد أجريت لها عملية استئصال للمبيضين تلاهما في اليوم التالي حقن يومي بالكارنوزين في العضل والمجموعة الرابعة وقد أجريت لها عملية استئصال للمبيضين تلاهما بعد أربعة أسابيع حقن يومي بالكارنوزين.

وقد تم التضحمية بحيوانات المجموعات الأولى والثانية والثالثة بعد أربعة وستة وثمانية أسابيع، أما حيوانات المجموعة الرابعة فقد تم التضحمية بها بعد ستة وثمانية أسابيع من تأريخ العملية حيث أخذت الفترات القطنية وتم تحضيرها للدراسة الهستومورفومترية والهستولوجيـة.

وقد أظهرت القياسات المورفومتريفية انخفاضًا ملحوظًا عن دلالة إحصائية في قيم القياسات لكل من النسبة المئوية لحجم القواطع العظمية (TBV%) ومتـوسط كثافة القواطع العظمية (MPD) ومتـوسط سمك القواطع العظمية (MTPD) في المجموعات الثانية والثالثة ورابعة ولكن الانخفاض الأعلى على الإطلاق ظهر في حيوانات المجموعات الثانية التي تعرضت لاستئصال المبيضين فقط. أما في مجموعة العلاج المبكر بالكارنوزين (المجموعة الثالثة) فقد كانت قيم القياسات TBV% و
وقد أظهرت الدراسة الهستولوجية أن استئصال البيضين أدى إلى انخفاض في سمك العظم الأصلي وأيضا في القواعد العظمية، وقد ظهر في كليهما الكثير من التجاويف الدراسية والخلايا الآكلة للعظام تتلا صدحهما. وظهرت أيضا مساحات خصوية كبيرة في داخل القواعد العظمية. وانفحل في الخطوط الأسمنتية، وفقدان الاتصال بين القواعد العظمية مما أدى إلى زيادة النهايات الحرة للقواعد العظمية واضطراب بنيتها مع اتساع تجاويف النخاع وظهور الخلايا الدهنية. وقد أدى العلاج بالكارنوزين إلى اختفاء نشاط الخلايا الآكلة للعظام وزيادة عدد الخلايا المكونة للعظام في كل من العظم الأصلي والعظم الأسمنتي وأيضا إلى تكاثر الصفائح الخصوية مع ظهور الخلايا المكونة للعظام في داخل المساحات الخصوية التي تحتل القواعد العظمية مما يدل على نشاط عملية بناء العظم. وقد كان العلاج المبكر بالكارنوزين (المجموعة الثالثة) أكثر فاعلية من العلاج المتأخر (المجموعة الرابعة) في الحفاظ على سمك قشرة العظم الأصلي وعلى الاتصال بين القواعد العظمية وعلى بنيتها وفي تقليل النهايات الحرة للقواعد العظمية.

وستخلص من هذه الدراسة أن استئصال البيضين يؤدي إلى زيادة كبيرة في فقدان العظم مع اختلال بنيته وأن الكارنوزين له تأثير على تنشيط بناء العظم وأيضا على تشطيب تأكيل العظم. وينصح بالعلاج المبكر بالكارنوزين للحفاظ على العناصر الطبيعية الإنشائية للعظام.

مجلة التشريح المصرية (26)، يناير 2003 (1)