Effects of ovariectomy, estrogen and soy isoflavones in rats submandibular glands

Luana Marotta Reis VASCONCELLOS¹, Vanessa Avila Sarmento SILVEIRA¹, Raphaella Silveira MEDEIROS¹, Márian Yaktin AMORIM¹, Yasmin Rodarte CARVALHO¹, Renata Falchete PRADO¹

¹ – São Paulo State University (Unesp) – Institute of Science and Technology – São José dos Campos – Department of Biosciences and Oral Diagnosis – SP – Brazil.

ABSTRACT

Objective: A decrease in granular convoluted tubule (GCT) cells and acini occurs in the submandibular glands of castrated female rats, while in rats submitted to hormone replacement and phytotherapy with soy isoflavones, this effect is reversed. This study aimed to elucidate the mechanisms through which these changes occur. Material and Methods: Rats (n=84) were ovariectomized and 21 were sham-operated. Ovariectomized rats were randomly subdivided and orally administered the following: 17β-estradiol (OVX-E; n=21), 15 mg/kg/day of soy isoflavone extract (OVX-I; n=21), 17β-estradiol + soy isoflavone extract (OVX-A; n=21); and water as placebo (OVX; n=21). The rats were euthanized three, five and eight weeks after ovariectomy. The submandibular salivary glands were submitted to histological processing with HE stain and immunohistochemistry was performed using the streptavidin-biotin-peroxidase complex. The cell area and the expression of proliferating cell nuclear antigen and estrogen receptor β were evaluated. Results: The results were statistically analyzed by ANOVA and Tukey test. A decrease in the area of GCT cells in the OVX, was observed, in contrast with an increase in the OVX-E. PCNA in the acinar cells and estrogen receptors were elevated in the OVX-I group. Conclusion: Castration exerts an immediate reductive effect on the volume of GCT cells. Estrogens, soy isoflavones and their combination have different mechanisms of action on the homeostasis of the gland. Estrogens cause an increase in GCT cells area, while isoflavones enhance cell proliferation and the expression of estrogen receptor-β. Their association showed no additional increase in the effect studied.

KEYWORDS

Estrogen; Morphometry; Salivary glands; Soy isoflavones.

RESUMO

Objetivo: Diminuição das células granulares do túbulo contorcido granular e dos ácinos ocorre nas glândulas submandibulares de ratas fêmeas ovariectomizadas, enquanto que nas ratas submetidas a reposição hormonal e fitoterapia com isoflavona da soja, este efeito é revertido. Este estudo tem como objetivo elucidar os mecanismos pelo qual estas alterações ocorrem. Material e Métodos: 84 ratas foram ovariectomizadas e 21 foram sham-operadas. Ovariectomizadas foram aleatoriamente subdivididas e receberam administração oral dos seguintes medicamentos: 17β-estradiol (OVX-E; n=21), 15 mg/kg/day do extrato de isoflavona da soja (OVX-I; n=21), 17β-estradiol + extrato de isoflavona da soja (OVX-A; n=21); e água como placebo (OVX; n=21). As ratas foram eutanasiadas 3, 5 e 8 semanas após a ovariectomia. As glândulas submandibulares foram submetidas a análise histológica por meio do processamento histológico de coloração com HE e imunohistoquímica utilizando o complexo peroxidase-biotina-streptavidina. A área celular e a expressão do antígeno de proliferação celular nuclear e receptor β estrógenico foram avaliados. Resultados: Os resultados foram estatisticamente analisados por meio do teste ANOVA e Tukey. Foi observada diminuição na área de GCT cells no grupo OVX, em contraste com o aumento no grupo OVX-E. PCNA nas células acinares e receptores de estrógeno estavam aumentados no grupo OVX-I. Conclusão: A castração exerce um efeito indutivo imediato no volume das células GCT. Estrógeno, isoflavonas e sua combinação têm diferentes mecanismos de ação sobre a homeostase da glândula. Estrógeno causa um aumento da área de células GCT, enquanto a isoflavona aumenta a proliferação celular e a expressão do receptor-β de estrógeno. A associação destes não mostrou aumento adicional nos efeitos estudados.

PALAVRAS-CHAVE

Estrógeno; Morfometria; Glândula salivar; Isoflavona da soja.
INTRODUCTION

The perimenopause or menopause corresponds to the stage of life in which the ovaries cease their function, usually a slow and gradual process, and is responsible for a series of changes in the female organism [1]. Due to estrogen deficiency, the systemic aspects of menopause manifest at this stage. The main issues are hot flashes, excessive sweating and vaginal atrophy [2]. Oral manifestations have been reported, including changes in salivation and taste, gingivitis and recurrent bleeding [3].

Salivary glands are not classic target organs of female sex hormones, but the influence of these hormones on these glands has been studied [4-7]; however, knowledge concerning this interaction remains scarce.

Estrogen receptors (ER) have been identified in human submandibular salivary glands of specimens evaluated by RT-PCR by Leimola-Virtanen et al. [8]. The oral mucosa displays specific receptors that recognize these ovarian hormones connecting them to the cytoplasm or nucleus. A deficiency of sex hormones, particularly estrogen, results in changes in the oral mucosa at the tissue level, since these hormones seem to control the proliferation, differentiation and keratinization of the gingival epithelium and stimulate the proliferation of fibroblasts [9].

Sex hormones seem to regulate salivary gland function. Streckfus et al. [10] analyzed and compared the rates of salivary flow in stimulated and non-stimulated glands, showing a relationship between flow decrease in advancing age and decreasing levels of estrogen production.

The harmful effects of the use of estrogen hormone replacement therapy, such as the increased risk of developing adenocarcinoma, is encouraging researchers to find new treatments for the effects of menopause [11]. Giuca et al. [3] evaluated the efficacy of hormone replacement therapy (HRT) and herbal therapy in postmenopausal women with oral discomfort. According to the authors, oral symptoms abated or decreased in women with both therapies. Both estrogens and phytoestrogens are promising, including the reduction of tooth loss during menopause due to osteopenia and osteoporosis.

In salivary glands, Katz et al. [12] demonstrated that isoflavones (genistein and soy phytochemical concentrate) cause the inhibition of cell proliferation in glandular lineage cultures, derived from a patient submitted to anticancer radiation therapy. Soy isoflavone was extracted by using ethanol as a solvent, with a resulting 49% weight concentration obtained. In a soy isoflavone concentration of 75 μg/ml, which inhibits DNA synthesis by 50%, 6.0 μg/ml (22 μM) of genistein and 4.3 μg/ml (17 μM) of daidzein were found.

In castrated rats, Chaves Carvalho et al. reported a decrease in the volume density of granular convoluted tubule cells and acini in the submandibular glands. They also reported that in rats administered hormone replacement, herbal medicines, or a combination of both, the effect of ovariectomy did not affect the morphometric values of these structures in the glands. The decrease in the ovariectomized group may occur due to a reduction in the volume of the cells or due to a decrease in the total number of cells by apoptosis, for example. Therapies appear to be effective in returning normal cell volume and inducing cell proliferation when administered to rats [4].

To explain the mechanisms by which these phenomena occur, we evaluated the cellular area using histomorphometry and the immunohistochemical expression of proliferating cell nuclear antigen (PCNA). Since the estrogen receptor-β is the predominant estrogen receptor subtype in the human oral epithelium and salivary glands [28], the receptor-β was accessed using immunohistochemistry.
MATERIAL AND METHODS

Animals

One hundred and five 90 days old female adult rats (Rattus norvegicus, Albinus variation, Wistar) weighing approximately 300g were selected. They were maintained in cages at room temperature provided by the vivarium of the Institute of Science and Technology, São José dos Campos Dental School of São Paulo State University (ICT-UNESP) and fed with commercial diets and water ad libitum. The amount of feed ingested per day was predetermined to avoid uncontrolled increase in the weight of the rats. This study was conducted according to the Ethical Principles in Animal Experimentation (COBEA) and was approved by the Research Ethics Committee of the ICT-UNESP, under protocol no. 029/2004/PA-CEP.

The rats were randomly divided into two groups: Ovariectomized (OVX) and SHAM. The OVX animals, consisting of 84 rats were submitted to ovariectomy, and subsequently randomly subdivided into four groups, each of 21 females, according to the treatment administered:

a) The estrogen group (OVX-E), administered 1mg/kg/day valerate 17β-estradiol orally;

b) The isoflavone group (OVX-I), administered 15 mg/kg/day of soy isoflavones 40% orally;

c) The associated group (OVX-A), administered 1mg/kg/day valerate 17β-estradiol associated with 15 mg/kg/day of soy isoflavones 40% orally;

d) The placebo ovariectomized group (OVX), administered placebo (water) orally.

The rats of the SHAM group consisted of 21 rats that underwent a faked surgical intervention. They were also administered placebo (water) orally each day. Rats were dosed, by gavage, 7 days per week for 3, 5 and 8 consecutive weeks, starting from the day after ovariectomy.

All rats were weighed at the onset of the experiment and were reweighed on the day of euthanasia, but they were detailed elsewhere [11].

General anesthesia was administered intramuscularly using a solution of 13mg/kg of methyl 2-(2,6-xylidine)-5,6-dihydro-4H-1,3-thiazine (Rompun-Bayer Brazil, São Paulo-SP), sedative, analgesic and muscle relaxant and 33mg/kg ketamine base (Dopalen - Agribands of Brazil Ltda, Paulínia-SP).

Following anesthesia, the rats in ovariectomized group were shaved in the lateral region of the body at the level of the kidneys and below the lowest rib and antiseptis was performed with iodine alcohol. A longitudinal incision was performed measuring 1 cm on average. Following the muscle incision, a dressing was made to contain the bleeding and the ovariaries were exposed and removed bilaterally.

False ovariectomy or sham surgery was performed on the rats from the Sham group, in which the same procedures were followed, including exposure of the ovaries, without the ligature and excision of the organ.

At the end of the procedure, the muscle layer was sutured with resorbable catgut thread no.4 (Cirumédica, Belo Horizonte, Minas Gerais, Brazil), followed by skin suture with 4.0 silk thread (Ethicon, Johnson & Johnson São José dos Campos, SP, Brasil), and repetition of antiseptis with iodine alcohol in the operated region.

Five rats from each group/subgroup of 15 were euthanized at three different time points: three, five and eight weeks post-ovariectomy, using a lethal injection of anesthetic. Their submandibular and sublingual glands were dissected and fixed in 10% formalin for approximately 48 h.

Treatments administered

Pills of valerate17β-estradiol, 1mg (Primogyna - Shering SA São Paulo, Brasil), were used to administer the dose of 1mg/kg/day to rats, dose previously used by Mvondo et al.
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Table 1 - Quantification of the isoflavones according to Herbarium Botanical Laboratory Ltda.

<table>
<thead>
<tr>
<th>Type of isoflavones of the extract 40%</th>
<th>Molecular mass</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycitin</td>
<td>446.41 g/mol</td>
</tr>
<tr>
<td>Glycitein</td>
<td>284.30 g/mol</td>
</tr>
<tr>
<td>Genistin</td>
<td>432.38 g/mol</td>
</tr>
<tr>
<td>Genistein</td>
<td>270.23 g/mol</td>
</tr>
<tr>
<td>Daidzin</td>
<td>416.38 g/mol</td>
</tr>
<tr>
<td>Daidzein</td>
<td>254.24 g/mol</td>
</tr>
</tbody>
</table>

In 2003, Genovese et al. [14] determined that Isoflavine® contained 1.5% α-glycoside, 53% aglycones and 45% β-glycoside. Between isoflavones, 29.7 ± 0.5% was total daidzein, 64.6 ± 0.4% was total genistein and 5.7 ± 0.1% was total glycitein.

The extract was suspended in distilled water. The laboratory quantified the principal following types of isoflavones (Table 1). Each rat received 15 mg/kg/day of soy isoflavones orally. The 15 mg/kg/d dose of soy isoflavone 40% extract was selected because similar doses are believed to be effective in preventing experimental osteoporosis without causing stimulation in the uterus [15,16].

Tissue processing and immunohistochemistry

Following fixation, the left submandibular glands of each group were dissected, washed, dehydrated, cleared in xylene and embedded in paraffin blocks. Five semi-serial sections were obtained from each rat, with 100 mm intervals. These were stained by conventional hematoxylin and eosin (HE), and subsequently photographed for the histomorphometric analysis.

For immunohistochemical analysis, 3 μm sections were stretched on silanized slides treated with 3-aminopropyl-triethoxysilane (Sigma Chemical Company, Saint Louis USA) for staining to detect estrogen receptor-β and PCNA.

One section of each rat (n=5, antigen retrieval resulted in a reduced number of immunohistochemical stained slides), totalizing 75 slices for each antibody were deparaffinized in xylene, rehydrated in solutions with decreasing concentrations of ethanol and washed in running water. Antigen retrieval was performed in citrate buffer (pH 6.0) in a microwave. The endogenous peroxidase activity was blocked with 6% H2O2 and methanol.

The primary antibodies were: polyclonal anti-estrogen receptor beta antibody (Abcam ab3576 Cambridge, MA, USA) and monoclonal anti-nuclear antigen of cellular proliferation (PC10, DAKO, Carpinteria, USA) incubated 1:100, and 1:75 respectively, overnight at 4°C. A labeled streptavidin-biotin-peroxidase (LSAB) kit (Dako, Carpinteria, USA) was used. The chromogenic substrate used was diaminobenzidine (DAB, Sigma, Saint Louis USA). The sections were counter-stained with Mayer's hematoxylin (Dako, Carpinteria, USA) and mounted with Permount.

Analysis of the results

Histomorphometric and immunohistochemical analysis were performed as blinded randomized controlled trials. In each group of 7 rats, a total of 35 slides were obtained, with a total of 10 granular convoluted tubule (GCT) cells (2 cells per field, one field per semi-serial slice). A total of 10 acini cells were randomly identified in the same fields used to count the GCT cells.

Analysis of the cell area was performed on 5 random microscopic serial fields (HEstained)
obtained using Zeiss Axiophot 2 microscope (Carl Zeiss, Oberkochen, Germany) with a 10X ocular and 40X objective, coupled to an Axiocam MRC5 digital camera (Carl Zeiss, Oberkochen, Germany).

The AxioVisionRel 4.7 (Carl Zeiss, Oberkochen, Germany) software was used to outline randomly selected cell membrane of cells of acini and ducts, and calculate the resulting area(µm2). Ten cells of the granular ducts and other 10 cells of acini were outlined and the mean area obtained.

The results of the immunohistochemical reactions were examined under an optical microscope. Five fields per slide were photographed using the same equipment mentioned above. For estrogen receptor-β, analysis of the percentage of glandular cells was performed, attributing the following scores: 0 = no positive glandular cells; 1 = less than 25% positive glandular cells; 2 = 26 to 50% positive glandular cells; 3 = 51 to 75% positive glandular cells; and 4 = more than 75% positive glandular cells. The scoring processes described above resulted in one value per field and a mean value of ER-β expression per rat.

For the analysis of PCNA, microscopic fields with the highest density [hot-spots] were identified using 100x magnification. These were subsequently photographed at 400x magnification. Three hundred acinar and GCT cells were analyzed with positive brown cells counted and expressed in absolute numbers. The values of all 15 groups were used in the statistical approach (5 values per group).

**Statistical analysis**

Statistical analysis was performed on the average values per rat. Data was analyzed descriptively and inferentially using two-way ANOVA. The variables were euthanasia time (3, 5 and 8 weeks) and treatment used (SHAM, OVX, OVX-E, OVX-I, OVX-A). The Tukey test was used for multiple comparisons. A value of 5% was used as the significance criterion.

Five statistical approaches were performed: ANOVA and Tukey test for acinar cells area data, ANOVA and Tukey test for GCT cell area data, ANOVA and Tukey test for PCNA positive acinar cells data, and ANOVA and Tukey test for PCNA positive GCT cells. The latter compared the mean values of scores obtained from each ERβ immunostained slide, according to rat treatment and euthanasia time.

**RESULTS**

The Kappa test was used in the analyses to determine the calibration of the examiner and substantial agreement (0.70) was observed.

**Size of acinar cells**

ANOVA showed that the time of euthanasia, the experimental group and the interaction between them had significant effects (p ≤ 0.05). At the earliest time point, the SHAM group (Figure 1a and Figure 2) presented the largest mean cell area, but this was not significantly different from the OVX (Figure 1b and Figure 2) and OVX-A (figure 1e and figure 2) groups for the same period. At 8 weeks, the OVX-A and OVX-E (Figure 1c and Figure 2) groups presented small averages of acinar cell area. At 8 weeks, both SHAM and OVX-I (Figure 1d and Figure 2) groups presented the highest average acinar cell area.
Figure 1 - Submandibular gland presenting predominantly serous acini. Acinar cells in rat submandibular glands are polarized cells, described as pyramidal, with a wide base facing the basal lamina. HE cuts show oval cells with eccentric nucleus and large basophilic cytoplasm, rich in secretory vacuoles (blue mark). Granular convoluted tubule (GCT) cells containing secretory granules are shown strongly stained by eosin, acquiring an almost red shade. Their lumens frequently display secretory granules (red arrow). Striated duct cells are columnar and contain longitudinal striation. Their lumens are usually well round and demarcated (green arrow). A) SHAM group at 3 weeks. GCT cells are big and completely filled by red granules B) OVX group at 3 weeks. GCT cells present a smaller area and less granules compared to SHAM.
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Figure 2 - Effects of ovariectomy and different treatments on acinar cells area. Bar charts indicate the average and standard deviation for each group. The number of animals evaluated for each cohort was n = 15 (Sham 3, 5 and 8 weeks, n=5), n=15 (OVX 3, 5 and 8 weeks, n=5), n=15 (OVX-E 3, 5 and 8 weeks, n=5), n=15 (OVX-I 3, 5 and 8 weeks, n=5), n=15 (OVX-A 3, 5 and 8 weeks, n=5).

*Values that do not share the same superscript letters are significantly different from each other (p < 0.05).

Cell size of the granular ducts

The morphological appearance of the granular ducts is shown in Figure 1.

ANOVA showed that the time of euthanasia, the experimental group and the interaction between them had significant effects (p ≤ 0.05). The SHAM group showed the largest mean cell area at 3 weeks, similar to the OVX-A group for the same period. At five and eight weeks, mean cell size remained similar between all the groups, except for OVX-A at eight weeks (Figure 3).

PCNA in acinar cells

The ANOVA test was applied to the value of positive acinar cells of a total of 300 cells and showed as significant effect, the experimental group (p = 0.003). The Tukey test showed no differences between treated groups in 3, 5 and 8 weeks. Group OVX-A presented the highest value at 5 weeks, marginally above most others, with the exception of SHAM at 8 weeks (Figure 4).
Figure 3 - Effects of ovariectomy and different treatments on GCT cells. Bar charts indicate the average and standard deviation for each group. The number of animals evaluated for each cohort was n = 15 (Sham 3, 5 and 8 weeks, n=5), n=15 (OVX 3, 5 and 8 weeks, n=5), n=15 (OVX-E 3, 5 and 8 weeks, n=5), n=15 (OVX-I 3, 5 and 8 weeks, n=5), n=15 (OVX-A 3, 5 and 8 weeks, n=5). *Values that do not share same superscript letters are significantly different from each other (p<0.05).
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Figure 4 - Effects of ovariectomy and different treatments on acinar cell proliferation. Bar charts indicate the average and standard deviation for each group. The number of animals evaluated for each cohort was n = 15 (Sham 3, 5 and 8 weeks, n=5), n=15 (OVX 3, 5 and 8 weeks, n=5), n=15 (OVX-E 3, 5 and 8 weeks, n=5), n=15 (OVX-I 3, 5 and 8 weeks, n=5), n=15 (OVX-A 3, 5 and 8 weeks, n=5).
*Values that do not share same superscript letters are significantly different from each other (p<0.05).

**PCNA in cells of the granular ducts**

In cells of the granular ducts, ANOVA showed that the time of euthanasia and its interaction with the experimental group had significant effects (p = 0.004 and p = 0.015 respectively). At 5 weeks, the OVX-A group presented the highest value, but was statistically similar to OVX-I (Figure 5).

**Estrogen receptor-β**

The expression of estrogen receptor-β, predominantly in granular ducts, is shown in Figure 6.

ANOVA showed a significant effect for experimental group (p = 0.003). At 5 weeks, the OVX-I group showed the highest expression, though this was very similar to the OVX-E, OVX-I, OVX,OVX-A and SHAM for different periods of euthanasia (Figure 7).
Figure 5 - Effects of ovariectomy and treatments on GCT cell proliferation. Bar charts indicate the average and standard deviation for each group. The number of animals evaluated for each cohort was n = 15 (Sham 3, 5 and 8 weeks, n=5), n=15 (OVX 3, 5 and 8 weeks, n=5), n=15 (OVX-E 3, 5 and 8 weeks, n=5), n=15 (OVX-I 3, 5 and 8 weeks, n=5), n=15 (OVX-A 3, 5 and 8 weeks, n=5).

*Values that do not share same superscript letters are significantly different from each other (p<0.05)
Figure 6 - Immunohistochemical location of estrogen receptor-β (ER-β) in light microscopy imaging clearly showing positive immunostaining predominantly in the submandibular gland GCT cells (red arrow). Section was originally photographed with 400 A magnification.
DISCUSSION

Hormonal interference in the phenotypic aspects of rats submandibular glands was the target of this study. The dose of estradiol valerate in this study was based in Mvondo et al. [13] that conducted a study based on various estradiol doses, using a postmenopause-like model of ovariectomized Wistar rats to evaluate the estrogen like effects of an isoflavone and a flavanone derived from E. lysistemon. Also, to determine the dose, we drew from our own experience, gained from previous research [17], as well as from consulting other works [16,18].

Other papers using higher doses were also consulted. Learning and memory in ovariectomized rats appears to be affecting by administration of high exogenous levels of estradiol (4mg/kg BW), due to enhance oxidative stress [19]. Estrogen therapy (weekly injections of estradiol valerate (2 mg/kg; i.m.) in OVX rats improved spatial memory retention in the test to find hidden platform [20]. Doses of 3,9mg/kg diet of 17-β estradiol suppressed the OVX induced body weight increase and maintained bone density at SHAM levels [21].

Phytoestrogens, particularly in isoflavones exhibit both agonist and antagonist actions, depending on serum estrogen concentrations, estrogen receptors, etc[22]. Isoflavones have greater binding affinity for estrogen receptor-β (ERβ) than ERα [18]. The standard isoflavone...
soybean extract at 40% concentration is widely documented in literature [23-25], occasionally being recommended as an alternative to the use of estrogen against menopause signs and symptoms [15,25-27]. We chose isoflavone soy extract at 40% concentration to be evaluated in this research because salivary glands present its most important target, the type-β estrogen receptor [28].

We tested the hypothesis that ovariectomy decreases the volume of the cells of the acini and granular ducts by histomorphometric analysis of cell area using stereological principles. The results of this research showed that estrogen deficiency caused a reduction in the area of cells of the granular convoluted tubules in accordance with Kouidhi et al. [29] and Sakabe et al. [30] previous statements.

The influence of sex hormones observed in granular convoluted tubule cells did not occur in the acinar cells. This was confirmed by the similarity of cell size in groups SHAM and OVX for all experimental periods, although subtle differences were observed when compared to the other groups. Additionally, the area of granular convoluted tubule cells had similar values at 5 and 8 weeks in both SHAM and OVX. It has been proposed that fatty tissue can provide a source of estrogenic hormones in ovariectomized rats [31], which may justify these findings, as ovariectomy was successfully confirmed by uterine atrophy. When comparing weight, greater gain among rats with hormonal deficiency was observed when compared to the control group.

We observed the effects of ovariectomy on the salivary glands in periods of 3, 5 and 6 weeks. Time was statistically significant for the reduction in the areas of acinar and GCT cells, as well as PCNA rates. Seko et al. [9] performed oral mucosa analysis after periods of 2, 4 and 6 months after ovariectomy. The thinning of oral mucosa epithelium was most obvious in the apex (tip) of the tongue by 6 months after ovariectomy. In contrast, in our study not only OVX groups had a reduction in cell area, SHAM groups also presented significantly smaller values after 5 and 8 weeks compared to 3 weeks.

A passenger effect of estrogen deficiency on cell size of the granular convoluted tubules became apparent, since the statistical difference was observed only in rats euthanized at 3 weeks. SHAM and OVX at 5 and 8 weeks were similar. Administering an association of estrogen and soy isoflavones returned the volume of the cells of the granular convoluted tubules to control values of the sham-operated group. The groups administered isoflavones or estrogen alone showed values superior to the OVX group at 3 weeks. This corroborates the fact that the cells of the granular convoluted tubules are under the multinormonal regulation by androgens, thyroid and adrenocortical hormones [32]. In male mice, such cells are voluminous and contains abundant large secretory granules, while female mice contain fewer secretory granules of smaller volume, thus showing a sexually dimorphic pattern [32].

The alteration in the cell volume of the OVX group reflects the reduction of estrogen-regulated secretory granules [32]. The results showed that estrogen therapy increased the volume of granular convoluted tubules cells compared with the ovariectomized group.

Ovariectomy did not reduce the immunohistochemical expression of estrogen-β receptors in the submandibular salivary glands. Most of the samples showed predominant expression of hormone receptors in the cytoplasm of the granular convoluted tubules cells. The literature affirms that the level of expression of the androgen receptor is variable [33-35]. The nucleus of acinar, granular and intercalated duct cells was the principal site of expression of this receptor in the study by Sawada et al. [34]. Herein we demonstrated that groups receiving soy isoflavones stood out, with high levels of estrogen receptor-β expression at 3 and 8 weeks.

PCNA was influenced by association and estrogen therapies, partially in line with the results found by Gallo et al., which showed
an increase in Ki67 immunoreactivity by 17β-estradiol, although it remained unchanged after soy extract treatment when compared with OVX controls.

Considering the decreased volume of granular cells as a result of ovariectomy (reduced estrogen), we expected to detect reduced expression of estrogen receptor, which did not occur. Apparently, other factors influence cells volume, such as epidermal growth factor (EGF) [36] which appears to regulate the phenotype of granular rodent cells. In males, the expression of EGF is higher than female rats [32]. Pascall et al. [37] showed that the expression of EGF increases in castrated female rats and decreases in castrated male rats. It is possible that the expression of EGF increased in the groups of castrated rats (however, in this research, it was not assessed) and could have affected the volume of the cells of the granular convoluted tubules.

Acinar cells showed similar volumes in all groups. Our study confirms the findings reported by Morrell et al. [33], who affirmed that the acinar cells are not affected by sex hormones, either structurally or in their composition.

Soy isoflavones and their association with estrogen caused an increase in the proliferation in acini cells compared with the SHAM group. None of the groups presented statistically significant differences for cell proliferation in the granular convoluted tubules. Soy isoflavones exert different cellular effects that can include inducing the activation of estrogen receptor-α, which promotes cell proliferation, and estrogen receptor-β, which promotes apoptosis. An irregular dose response to isoflavones has been observed, mainly relating to ability of cells to stimulate proliferation. Isoflavones appear to modulate epigenetic mechanisms, including DNA methylation, histone modifying genes and noncoding RNAs [38]. Some of the genes suffering DNA methylation, after exposure to high or low doses of isoflavones [39], regulate the proliferation of mammary epithelial cells. Our study has effectively demonstrated the differential expression of a proliferation marker in castrated rats induced by isoflavones. Epigenetic changes are potentially the mechanism involved in this phenomena. It is important to remember that soybeans, don't present a constant isoflavone content, instead their content is subject to a variety of environmental factors such as temperature, precipitation [40], the location of crops and the season they are planted [41].

The capacity to conjugate isoflavones also fluctuates significantly between mice and rats, as well as between rodents and humans, who have advanced capacity to conjugate isoflavones [42]. Following a dose of 50mg/kg/day of soy isoflavones extract, serum levels in Sprague-Dawley castrated female rats were found to be 0.03µm of free genistein from a total of 0.91µm, and 0.06µm of free daidzein from a total of 1.5µm [16]. This dose was similar to the one we administered in our study.

Therapy with soy isoflavones promoted increased expression of estrogen receptors-β in the submandibular salivary glands. Perhaps this mechanism of increased estrogen receptors, associated with the induction of increased cell proliferation, explain the results obtained by Chaves Carvalho et al. [4] in which isoflavones maintained a volume density in the treated rats similar to that of the SHAM group rats.

The cells of the granular convoluted tubules of rats administered soy isoflavones presented less volume than those of the SHAM group for all the experimental periods. Estrogenic hormones have numerous effects on cells of the salivary glands, differing depending on the cell type.

A clear dose response pattern cannot always be observed [39]. According Blei et al., an isoflavone dose that reflects a lifelong nutritional exposure via a soy rich diet may be sufficient to result in an increase in fat metabolism, protein synthesis and skeletal muscle mass. In cell cultures, the mechanism behind proliferation
and anti-proliferative action of isoflavones is unknown, but non-linear dose response curves have been described, showing proliferation of MCF-7 cells in vitro.

PCNA expression remained similar in the granular convoluted tubules and acini of rats treated with 17β-estradiol. A similar result was reported by Tsinti et al. [5]; however, in their study, the volume of granular cells returned to normal following hormone therapy. This estrogen-dependent mechanism explains the results obtained by Chaves Carvalho et al. [4], who verified similar volume densities in the groups administered estrogen and sham-operated rats.

**CONCLUSION**

Castration exerts almost immediate reductive effect on the cell volume of the granular convoluted tubules of the submandibular gland of rats. Estrogens and soy isoflavones alone and in combination have different mechanisms of action on the homeostasis of the tissue, such that the first causes an increase in the volume of normal cells of the granular convoluted tubules, the second enhances the expression of estrogen receptor-β, while their association showed no combined or additional increase in the effect studied.

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Renata Falchete do Prado (Corresponding address)
Departamento de Biociências e Diagnóstico Bucal
Av. Engenheiro Francisco José Longo, 777,
Jardim São Dimas, São José dos Campos, São Paulo,
CEP 12245000, Brazil.
renatafalchete@hotmail.com

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