Histological and Immunohistochemical Study of the Effect of Alendronate on the Submandibular Salivary Gland of Adult Male Albino Rat and the Possible Protective Effect of Propolis

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Abstract

Background: Alendronate is an anti-bone-resorptive drug used in the treatment of post-menopausal osteoporosis. Its long administration induced organ damage especially the submandibular salivary glands. Propolis is a natural resinous substance with an antioxidant and anti-inflammatory actions.

Aim of Study: This work was performed to study the effect of alendronate on the histological structure of the submandibular salivary gland of adult male albino rat and the possible protective effect of propolis.

Material and Methods: 40 adult male albino rats were divided into 4 groups (10 rats each); control group, Group A: Rats received 100mg/kg body weight propolis, Group B: Rats received 2.5mg/kg body weight alendronate and Group C: Rats received propolis 30 minutes before alendronate. Doses were given orally for consecutive 30 days then gland specimens were taken for histological and immunohistochemical studies using light and electron microscopes.

Results: H & E and toluidine blue results of Group B revealed disturbed architecture, separated lobules and acini, cytoplasmic vacuolations, dark nuclei, irregular ducts and congested blood vessels. PAS-AB showed significant increase in its mean color intensity as compared with control group while α-SMA glandular myoepithelial cells immunostaining expressed significant decrease in its mean color intensity in comparison to control group. Electron microscopic examination of acinar and duct cells revealed cytoplasmic vacuolations, rarefaction, abnormal shaped mitochondria with lost cristae, electron dense bodies, dilated RER and Golgi apparatus, apparently increased secretory granules with coalescence, compressed shrunken hyperchromatic nuclei, nuclear membrane irregularities, and dilated perinuclear and intercellular spaces. Group C showed moderate improvement of the previous pathological findings.

Conclusion: Alendronate induced damaging effects on the histological structure of submandibular salivary gland in rats and the administration of propolis could moderately improve such damage through its anti-oxidant and anti-inflammatory effects.


Introduction

PROPOLIS is a natural resinous substance collected by honeybees from buds and exudates of different plants [1]. Its chemical composition is variable depending on the plant sources [2]. Propolis composed of polyphenols as flavonoids that, accompanied by phenolic acid, aldehydes, phenolic aldehydes, and ketones. Besides, volatile oils, aromatic acids as well as waxes. It also contains a number of essential elements like magnesium, calcium, iron, nickel, zinc and vitamins [3].

Propolis has many beneficial effects as anti-inflammatory, immunomodulatory, antioxidant, antimicrobial, as well as antitumor effects. These effects make it widely used in the treatment of various conditions such as microbial infections, wounds, burns, sore throat, ulcers as well as diabetes [4,5]. It is also considered as a rich source of natural antioxidants and can be used as a protective agent against various free radical associated degenerative diseases [6,7].

Alendronate is one of the nitrogen containing bisphosphonates (BPs) which are synthetic analogues of pyrophosphate and are used in the treatment of disorders that involve excessive bone resorption such as osteoporosis, Paget's disease, and hypercalcemia of malignancy [8]. Alendronate mediates its antiresorptive action mainly through inhibiting the action of osteoclasts [9].
Histological & Immunohistochemical Study of the Effect of Alendronate on the Submandibular Salivary Gland

Its long term administration could produce a wide variety of adverse effects as nausea, vomiting, epigastric pain and dyspepsia due to irritation of the mucosa of the gastrointestinal tract. Also, atrial fibrillation and deterioration in renal function may occur, especially in patients with renal insufficiency [10,11]. One of the important side effects is decreasing salivary secretion, so the patient suffers from dry and burning mouth [12].

Salivary secretion is essential for lubrication and coating of the oral cavity. It helps for speech, mastication and swallowing in addition to its antibacterial and antiviral effects. Furthermore, it has an important role in dissolving food particles by stimulating taste receptors. Moreover, saliva initiates digestion of complex carbohydrate by salivary amylase enzyme and triglycerides by lingual lipase [13,14]. Moreover, saliva has an important effect on tooth as it prevents their demineralization by buffering bacterial acids produced during sugar metabolism [15].

From these previous data, we can predict that dry mouth is a big health problem. So this work aims to evaluate the possible protective effect of propolis on alendronate-induced changes on the submandibular salivary glands in adult male albino rats by means of histological as well as electron microscopic studies.

Material and Methods

This work was carried out from January 2017 till August 2017 at Histology Department, Faculty of Medicine, Tanta University and at the Electron Microscopic Unit, Faculty of Agriculture, Mansoura University.

Experimental animals:

This work carried out using 40 adult male albino rats weighing 120-180 grams. The animals housed in suitable clean properly ventilated cages under similar environmental conditions and fed on a similar commercial laboratory diet and water. The experiments directed following the guidelines on the care and usage of experimental animals of Tanta University, Faculty of Medicine with the authorization of The Animal Experiment Committee of the faculty. Moreover, the bodies of the sacrificed animals were packed in a special package according to the safety precaution and infection control measures then sent with the hospital biohazards.

The experimental groups:

Animals randomly divided into four groups (10 rats each); control group: Subdivided into further two subgroups: Subgroup C1: Rats kept without treatment and subgroup C2: Rats given 1ml distilled water; Group A: Rats given propolis (Sigma, Cairo, Egypt) in a dose of 100mg/kg/day [16,17]; Group B: Rats given alendronate (Merck & Co., Inc., white house station, N.J., USA) in a dose of 2.5 mg/kg/day 30 minutes before breakfast according to [18] and Group C: Rats given 100mg/kg/day propolis before 2.5mg/kg/day alendronate.

Rats received their doses orally through intragastric tube, for 30 consecutive days. On the day after the last dose, animals anesthetized by ether inhalation. Then, the submandibular salivary glands carefully dissected and removed, then, processed for histological, immunohistochemical and electron microscopic examination.

Preparation of specimens for paraffin sections:

Gland specimens fixed in 10% formalin buffered saline for 48 hours. Then dehydrated, cleared, and infiltrated with molten paraffin wax, followed by embedding in hard paraffin. Finally, sections of 3-5 microns obtained and prepared for histological and immunohistochemical examination [19].

1- Haematoxylin and Eosin (H & E) stain: Sections dewaxed, then rehydrated. After that, sections placed in Ehrlich’s hematoxylin for 30 minutes, then at 1% eosin Y for 10 minutes. Lastly, sections were dehydrated, cleared, and mounted.

2- Combined (Periodic Acid Schiff-Alcian Blue) (PAS-AB) stain: Sections dewaxed in xylene, then rehydrated and stained with alcian blue solution for 30 minutes, followed by oxidation with periodic acid. Then, sections covered with Schiff’s reagent for 15 minutes and finally, sections dehydrated, cleared, and mounted with Canada balsam.

3- Immunohistochemical staining for alpha-smooth muscle actin (α-SMA): Sections deparafinated, and hydrated. Then, sections placed in 3% hydrogen peroxide in methanol for 30 minutes at room temperature to inhibit the formation of background staining from the endogenous peroxidase activity. After that, slides rinsed in phosphate buffered saline, then sections incubated with citric acid (pH 6.0) in a microwave oven at 121°C for 5 minutes for antigen retrieval and allowed to cool for 8 minutes at room temperature. Next, the sections incubated with normal goat serum to prevent non-specific protein binding followed by the addition of α-SMA
primary antibody (1:50) (Abcam, USA) overnight at 4°C. Slides then, incubated with biotinylated peroxidase-conjugated secondary antibody for 1 hour and rinsed in PBS buffer. Afterwards, Diaminobenzidine (DAB) solution applied to the sections for 5-10 minutes, followed by washing in PBS, then counterstaining with Mayer's haematoxylin for 1-2 minutes. Finally, sections dehydrated, cleared, mounted in Canada balsam, and examined by a light microscope (Olympus, Japan) [20,21]. For negative controls, the same procedure performed by replacing the step of addition of primary antibody by PBS. While, the positive control is the normal human colon mucosal smooth muscle tissue (abcam124964).

Preparation of sections for electron microscopic study:
The submandibular salivary gland specimens a cut by sharp razor blade into small pieces of 1 mm and prepared for semithin and ultrathin sections according to [22]. Briefly, specimens immersed in 4% glutaraldehyde in phosphate buffer with pH 7.2 at 4°C for 2 hours. Followed by, post-fixation in 1% osmium tetroxide. Then, sections dehydrated, infiltrated in epoxy resins, embedded in resin and polymerized in an oven at 60-70°C for 1-2 days. After that, ultrathin sections 40-150nm obtained using LKB ultramicrotome then, picked up on 200 mesh, copper grids and double stained, with uranyl acetate and lead citrate. Lastly, the specimens examined and photographed using (JEOL-JEM-100 SX electron microscope, Japan) at the Electron Microscopy Unit, Faculty of Medicine, Tanta University and at the Electron Microscopic Unit, Faculty of Agriculture, Mansoura University.

Statistical analysis:
A- Estimation of glandular PAS-AB color intensity:
10 different PAS-AB stained images (X400) of each group used by the image J software (National Institute of Health, Bethesda, Maryland, USA), to estimate PAS-AB color intensity of the submandibular salivary gland in the different experimental groups.

B- Evaluation of α-SMA immunostaining color intensity in the myoepithelial cells of the submandibular salivary gland: By image J software (National Institute of Health, Bethesda, Maryland, USA), α-SMA immunostaining color intensity in the myoepithelial cells of the submandibular salivary gland evaluated. Through which, 10 different images (X400) of each group, containing no blood vessels used.

The statistical significant differences between the experimental groups valued using Minitab Statistical Software for Windows (Version 16.1, Minitab Inc., State College, PA, USA). Then, the variances of data analyzed using either two-tailed Student's t-test or the Mann-Whitney's U-test after evaluation by F-test. Through which, \(p\)-value <0.05 considered to be significant.

Results

Histological results:
In the present work, histological and immunohistochemical results of Group A (propolis) are similar to the control group.

A- Light microscopic study:

I- Haematoxylin and eosin stained sections: Through examination of the control group sections, the normal histological structure of the submandibular salivary gland found. By which, it was divided into numerous lobules by connective tissue septa conveying blood vessels and large excretory ducts lined by stratified columnar epithelium. The serous acini appeared spherical in shape, lined by pyramidal cells with a moderate basophilic cytoplasm, and rounded basally situated nuclei. While, the Intercalated Ducts (IDs), seen lined by short cuboidal cells with centrally located rounded nuclei. Also, flattened myoepithelial cells seen partially surrounding the serous acini and IDs. Moreover, the Granular Convoluted Tubules (GCTs) lined by tall columnar cells with eosinophilic cytoplasm and large rounded basally situated nuclei, and the Striated Ducts (SDs) lined by columnar cells with centrally located rounded nuclei and intensely eosinophilic cytoplasm with basal striations Fig. (1A-D).

Conversely, Group B (alendronate group), revealed marked disturbed gland architecture with disorganized widely separated acini, in addition to irregular excretory ducts with narrow lumen and disorganized darkly stained nuclei besides congested blood vessels. Moreover, the acinar cells showed numerous cytoplasmic vacuolations in addition to darkly stained nuclei. While, the GCTs, appeared surrounded with wide space besides, vacuolated, fragmented cytoplasm, disarranged darkly stained nuclei Fig. (2A-D).

As regards Group C (alendronate and propolis group), it showed a moderate improvement when
compared to the control group. Through which, some serous acinar and GCT cells showed cytoplasmic vacuolation besides, few SD cells with flattened nuclei Fig. (2E,F).

**II- Combined Periodic Acid Schiff-Alcian Blue (PAS-AB) results:** Combined PAS-AB stained sections of the control group showed, serous acini with mild PAS-positive reaction and moderately positive alcian blue staining, while, the GCTs showed moderate PAS-positive reaction. Considering IDs and SDs, they showed faint PAS positive staining. Conversely, the alendronate group showed, serous acinar cells with moderate PAS positive reaction and strong alcian blue reaction while, the GCT cells showed an intense PAS positive reaction. The ID and SD cells showed moderate PAS positive reaction as well as accumulation of AB positive material in the duct lumen. As regards the propolis & alendronate group, they displayed few SDs with accumulated AB positive material in their lumen and some GCT cells with an intense PAS positive reaction Fig. (3A-D).

Considering the statistical results of the mean color intensity of PAS-alcian blue, it showed significant increase in the alendronate group as compared with the control group; in contrast to a significant decrease in the alendronate and propolis group in comparison to alendronate group Fig. (4).

**III- Immunohistochemical results for α-SMA:** Immunostained α-SMA sections from the rat submandibular salivary gland of the control group showed, strong positive reaction in the cytoplasm of the myoepithelial cells at the periphery of the serous acini as well as the IDs. While in the GCTs and SDs, there was no observed reaction at their periphery. Furthermore, the smooth muscle cells lining the wall of the blood vessels showed positive immunoreaction. On the contrary, the alendronate group expressed weak positive reaction at the periphery of both serous acini and IDs. Whereas, the GCTs and SDs showed negative reaction. However, propolis & alendronate group exhibited some acini with weak positive immunoreaction Fig. (5A-D).

As regards, the mean color intensity of α-SMA immunostaining of the myoepithelial cells in the submandibular salivary glandular tissue; it revealed significant decrease in the alendronate group as compared with the control group. Besides, significant increase in the propolis & alendronate group in comparison to alendronate group Fig. (6).

**B- Electron microscopy study:**

1- **Acinar cells:** EM examination of the control group revealed, pyramidal shaped serous acinar cells with basal rounded nuclei (some cells binucleated) and prominent nucleoli in addition to parallel arrays of RER, mitochondria, supranuclear Golgi complex, apical junctional complexes besides, secretory granules with different sizes and microgranular appearance. Moreover, the cells showed normally appeared intercellular space with flattened myoepithelial cells partially surrounding the acini. Alternatively, the alendronate group presented apparently increased granules with basal migration and coalescence of most of them. Along with, dilated Golgi apparatus, and RER, abnormal shaped mitochondria with lost cristae, and electron dense bodies. These all besides, nuclear membrane irregularity, shrunken electron dense nucleus and, dilated intercellular space. Furthermore, propolis & alendronate group revealed, few acinar cells with coalescence of some secretory granules and irregular cellular outlines Fig. (7A-D).

2- **IDs:** The control ID presented with two types of cells; granular and non-granular. Through which, their cytoplasm revealed, rounded centrally located nuclei in addition to junctional complexes and apical microvilli. Regarding the alendronate group, there was wide intercellular space, cytoplasmic vacuolations and shrunken electron dense nuclei. While, propolis & alendronate group revealed, some irregular, destroyed ID cells that separated from its basement membrane Fig. (8A-C).

3- **GCTs:** GCTs of the control group, exposed tall columnar cells with electron dense granules occupying their apical two thirds while, the basal part was occupied by RER and mitochondria surrounding a rounded euchromatic nucleus with prominent nucleolus. On the contrary, GCTs of the alendronate group showed, apparently increased secretory granules, cytoplasmic vacuolation, together with cytoplasmic rarefaction, and nuclear membrane irregularities. Through EM examination of the propolis & alendronate group, the GCTs, showed few cells with cytoplasmic rarefaction Fig. (8A-C).

4- **SDs:** EM examination of control SDs showed tall columnar cells with junctional complexes, and apical microvilli, small clear secretory vesicles located apically and large rounded centrally located euchromatic nuclei. Furthermore, the basal part of these cells showed deep infoldings of the plasma membrane that were parallel to the long axis of
the cell with numerous rod shape mitochondria (palisade arrangement). As regards the SD cells of alendronate group, they displayed cytoplasmic vacuolations and rarefaction, and abnormal distribution of mitochondria around the nucleus, and mitochondria with lost cristae. These were associated with electron dense bodies of variable sizes and irregular nuclear outlines. Even though, the SDs of propolis & alendronate group revealed some changes such as, mitochondrial disarrangement and detachment of some SD cells from the basement membrane Fig. (10A-D).

Fig. (1): Effect of propolis on H. & E. stained sections of alendronate-induced changes in the submandibular salivary gland of rats' control group. A) Numerous lobules divided by connective tissue septa (*) conveying blood vessels (→) and excretory ducts (4) (H. & E. X400). B) A large excretory duct with wide lumen (*) and lined with stratified columnar epithelium (→). (H & E X1000). C) ID with low cuboidal cells having central rounded nuclei (thin arrow) and myoepithelial cells partially surrounding the ducts (4), serous acinar cells with basophilic cytoplasm, rounded basally situated nuclei (bifid arrow) and myoepithelial cells partially surrounding it (wavy arrow) (H. & E. X1000), D) GCTs lined by tall columnar cells with eosinophilic cytoplasm and basally situated nuclei (curved arrow), SDs lined by tall columnar cells with rounded nuclei located near to cell apex (4). (H. & E. X1000).
Fig. (2): Effect of propolis on H. & E. stained sections of alendronate-induced changes in the submandibular salivary gland of rats' Group B (alendronate) & Group C (Propolis & alendronate). A) Alendronate group, showing marked disturbed gland architecture with disorganized widely separated acini (*) and acinar cell cytoplasmic vacuolations (→) (H. & E. X 400). B) Alendronate group showing, irregular excretory ducts with narrow and obliterated lumen (wavy arrow). (H. & E. X1000) C) Alendronate group, showing large excretory duct with disorganized darkly stained nuclei (→), and congested blood vessels (*). (H. & E. X1000). D) Alendronate group, showing serous acini with cytoplasmic vacuolations (4), GCTs with vacuolated (*), and fragmented cytoplasm (curved arrow), darkly stained nuclei (→), and wide space around the tubules (S). (H. & E. X1000). E) Propolis & alendronate group, showing few serous acini with cytoplasmic vacuoles (→) (H. & E. X400). F) Propolis & alendronate group, showing few GCT cells with cytoplasmic vacuolation (4) and few SD cells with flattened nuclei (double arrow). (H. & E. X1000).
Fig. (3): Effect of propolis on PAS-AB stained sections of alendronate-induced changes in the submandibular salivary gland of the rats' experimental groups. A) control group showing, mild PAS positive, moderate positive alcian blue staining of serous acini (→), moderate PAS positive reaction of the GCT cells (4) and faint PAS positive staining of the ID (curved arrow) and SD cells (wavy arrow). B) Alendronate group showing, moderate PAS positive and strong positive alcian blue staining of serous acini (→), and intense PAS positive reaction of the GCT cells (4). C) Alendronate group showing, ID (→) and SD (4) cells with moderate PAS positive reaction with accumulation of AB positive material in the duct lumen (*). D) Propolis & alendronate group showing, few SDs with accumulated secretory material in their lumen (→) and some GCT cells with intense PAS positive reaction (4). (PAS-AB X400).

Fig. (4): The color intensity of PAS-AB of the submandibular glandular tissue of the different experimental groups expressed as mean ± SEM.

# : p>0.05 non-significant relative to control.
* : p<0.05 significant relative to control.
@ : p<0.05 significant relative to alendronate group.
Fig. (5): Effect of propolis on alendronate-induced α-SMA expression changes of the glandular myoepithelial cells of the rats' experimental groups. A) Negative control of α-SMA immunostained sections from rat submandibular salivary gland showing no reaction. B) Control group showing, strong positive α-SMA immunoreaction at the periphery of the serous acini (curved arrow), ID (thin arrow), and the wall of the blood vessels (wavy arrow), negative immunoreaction at the periphery of GCTs (*) and SDs (thick arrow). C) Alendronate group showing, serous acini (thick arrow), and ID (thin arrow) with α-SMA weak positive immunoreaction at their periphery, GCTs (wavy arrow), and SDs (curved arrow) with negative reaction. (D) Propolis & alendronate group showing, some acini with α-SMA weak positive immunoreaction (→). (α-SMA immunostaining X400).

Fig. (6): The color intensity of α-SMA immunostaining of the myoepithelial cells of the submandibular salivary glandular tissue of the different experimental groups expressed as mean ± SEM.

* : p>0.05 non-significant relative to control.
@ : p<0.05 significant relative to control.
$: p<0.05$ significant relative to alendronate group.
Fig. (7): Effect of propolis on submandibular salivary gland acinar EM changes induced by alendronate. A) EM of the control group showing, acinar cells with basal rounded two nuclei and prominent nucleoli (N), RER (→), mitochondria (thin arrow), secretory granules with microgranular appearance (*), apical junctional complexes (bifid arrow), basement membrane (wavy arrow), intercellular space (thick arrow), and myoepithelial cell partially surrounding the acini (curved arrow) (Mic. Mag. X2000). B) Alendronate group showing, acinar cells with apparently increased secretory granules with basal migration and coalescence of most of them (*), dilated Golgi apparatus (thin arrow), electron dense bodies (curved arrow), nuclear membrane irregularity (wavy arrow), and dilated intercellular space (thick arrow) (Mic. Mag. X2000). C) Alendronate group showing, acinar cells with fused secretory granules (*), dilated RER (→) and shrunken electron dense nucleus (N) (Mic. Mag. X2000). D) Propolis & alendronate showing, acinar cells with fusion of some secretory granules (*), and irregular cellular outlines (→) (Mic. Mag. X2500).

Fig. (8): Effect of propolis on submandibular salivary gland ID EM changes induced by alendronate. A) EM of ID control group showing, two types of cells; granular cells with electron dense granules (→) and non-granular cells without granules (curved arrow), rounded centrally located nucleus (N), junctional complexes (wavy arrow) and apical microvilli (bifid arrow) (Mic. Mag. X1500). B) Alendronate group showing, ID cells with cytoplasmic vacuolations (→), wide intercellular space (4) and shrunken electron dense nucleus (N) (Mic. Mag. X2500). C) Propolis & alendronate group showing, some IDs with irregular destroyed cells, separated from its basement membrane (→) (Mic. Mag. X1500).
Fig. (9): Effect of propolis on submandibular salivary gland GCT EM changes-induced by alendronate. A) Control GCT showing, cells with electron dense granules occupying their apical two thirds (*), mitochondria (→), RER (curved arrow) and basal rounded nuclei with prominent nucleoli (N) (Mic. Mag. X2500). B) Alendronate group showing, GCT cells with apparently increased granules (*), cytoplasmic rarefaction (→), and vacuolation (bifid arrow), and nuclear membrane irregularity (wavy arrow) (Mic. Mag. X2000). C) Propolis & alendronate group showing, some GCT cells with an area of cytoplasmic rarefaction (→) (Mic. Mag. X1500).

Fig. (10): Effect of propolis on submandibular salivary gland SD EM changes-induced by alendronate. A) SDs of the control group showing, cells with apical junctional complexes (bifid arrow), and microvilli (4), apical small clear secretory vesicles (→), basal rod shaped mitochondria with palisade arrangement (curved arrow) and large rounded centrally located euchromatic nuclei (N) (Mic. Mag. X2500). B) Alendronate group showing, SD cells with areas of cytoplasmic rarefaction (*), and vacuolations (→), and mitochondrial disarrangement (bifid arrow) (Mic. Mag. X2000). C) Alendronate group showing, SD cells with electron dense bodies of variable sizes (4), mitochondria with lost cristae (curved arrow) and irregular nuclear outline (wavy arrow) (Mic. Mag. X2500). D) Propolis & alendronate group showing, SD cells with mitochondrial disarrangement (→) and detachment of cells from the basement membrane (curved arrow) (Mic. Mag. X2500).
Discussion

Results of this work revealed disorganized glandular structure in the form of widely separated acini, irregular excretory ducts with narrow lumen and numerous cytoplasmic vacuolations, fragmented cytoplasm, either in the acini or GCTs. These changes were clarified by some authors. Through which, alendronate treatment is associated with the release of pro-inflammatory cytokines that led to multiple degenerative changes [23,24]. Another researcher indicated that, accumulation of the secretory material within the cells could contribute to cellular degenerative changes with subsequent cell death, duct shrinkage and irregularity [25].

Additionally, alendronate has an effect on histamine-forming enzymes leads to increased histamine synthesis as well as its release [26,27]. The latter results in increased capillary permeability, and the hydrostatic pressure and consequentially leading to tissue edema [28]. So, this could be the cause of wide space in between and around the acini as well as the ducts.

Regarding the occurrence of cytoplasmic vacuolations of acinar and duct cells; many authors documented that alendronate reduce the mitochondrial membrane potential leading to generation of Reactive Oxygen Species (ROS) and free radicals resulting in lipid peroxidation and damage of cell membranes as well as membranes of the cell organelles ending in failure of the energy-dependent Na+, K+ ion pumps in the cell [29-31].

As well, these changes lead to accumulation of Na+ inside the cells with entry of water expressing cellular swelling and vacuolations. Therefore, this could explain the dilated RER, and Golgi apparatus as well as cytoplasmic rarefaction that detected by EM examination. Additionally, the attributed accumulation of the unprenylated proteins in the organelle cisternae due to interference with their transport and modification in Golgi apparatus could also explain its dilatation [32].

In the present research, the statistical results of the combined PAS-AB revealed, significant increase in the mean color intensity of PAS-AB stains in the alendronate group as compared with the control group. These all might be due to exocytosis interruption by alendronate, consequently, the amount of secretory granules is increased [18].

As regards the immunohistochemical results of the present work. It showed significant decrease in the mean color intensity of α-SMA immunostaining of the myoepithelial cells of alendronate group when compared to the control group. This confirmed the interruptive effect of alendronate on the secretory process of the gland, which was due to disruption of the actin cytoskeleton by alendronate [33]. Alendronate could inhibit Farnesyl Pyrophosphate Synthase (FPPS) enzyme involved in the mevalonate pathway which is essential for the production of Geranylgeranyl Pyrophosphate (GGPP) and Farnesyl Pyrophosphate (FPP) [34]. These metabolites are necessary for the prenylation of small GTPases; Rab and Rho, which act as regulators of intracellular vesicular trafficking and secretory function of the salivary glands [18]. Additionally, mevalonate pathway is associated with reduction of coenzyme Q 10 which causes mitochondrial dysfunction with ATP depletion so; the cells had no energy for the secretory process to take place. These previous mechanisms could explain the most prominent finding detected by EM which was in the form of increased secretory granules with their basal migration, and coalescence of most of them in the serous acini as well as GCT cells [35,36].

Moreover, the abnormal shaped mitochondria with lost cristae and dilated intercellular space were observed as an EM results in the present work. These findings may be attributed to the release of ROS, causing disturbance of the plasma membrane integrity, and consequently beside edema, could disrupt the intercellular junctions [29,37].

Too, electron dense bodies of variable sizes and heterogeneous consistency in the cytoplasm of acinar and duct cells were recorded. These most probably are secondary lysosome. As the release of ROS besides occurrence of lipid peroxidation requires activation of lysosomal hydrolytic enzymes [37-39].

Furthermore, shrunken nuclei with dense chromatin were noticed as an EM result of the acinar and duct cells. Since Alendronate induced mitochondrial dysfunctions, this could induce causing in the mitochondrial membrane permeability with the release of cytochrome c as well as other pro-apoptotic factors like caspase-9 and -3 causing apoptotic cell deaths. Subsequently, DNA fragmentation and chromatin condensation ensues [35,40,41].

In the present research, propolis administration displayed moderate improvement of the previous pathological findings of the submandibular salivary gland induced by alendronate. It was documented that propolis has protective effect against dry mouth induced by parotid salivary gland dysfunctions due to exposure to radiation [42]. This could attribute
to the protection of cell membranes with subsequent preservation of the intracellular signal transduction involved in salivary secretion.

Moreover, propolis is able to scavenge free-radicals and its antioxidant properties are due to its polyphenol composition \[46,47\]. So, propolis could inhibit the activation of the apoptotic pathways in normal cells exposed to oxidants \[48\].

It was reported that propolis could inhibit the production of pro-inflammatory cytokines such as Interleukin IL-1 \[46,47\], Tumor Necrosis Factor (TNF)-\[a\], and Monocyte Chemotactic Protein (MCP)-\[1\]. It also reduces the pro-inflammatory cyclooxygenase (COX2) and lipoxygenase enzyme, important for leukotriene production. This potent anti-inflammatory action could inhibit the release of histamine so decreasing vascular permeability as well as edema \[46,47\].

Recently, the anti-inflammatory action of propolis is through the reduction of the nuclear factor kappa-light-chain-enhancer of activated B cells (NF-KB) stimulation as well as suppression of the synthesis of ubiquitin units; preventing its action on the cell proteins \[48\].

**Conclusion:**

Based on the current study, it can be concluded that, propolis improve alendronate-induced damaging effects on the rat submandibular salivary gland through its antioxidant and anti-inflammatory actions. In this light, propolis is recommended to for people who take alendronate for a long period to minimize its harmful effects.

**References**


24- MORITA M., IWASAKI R., SATO Y., KOBAYASHI T., ALJAYER A., EL-BAZ D. and BASHIR M.: "Impacts of nandrolone in rat submandibular salivary glands of streptozotocininduced diabetes in rats: Histo-


26- DENG X., FUNAYAMA H., SHOJI N., SASANO T., IWAKURA Y., et al.: "Mutual augmentation of the induction of the histamine-forming enzyme, histidine decarboxylase, between alendronate and immuno-


32- ZAFAR S., COATES D.E., CULLINAN M.P., DRUMMOND B.K., MILNE T. and SEYMOUR G.J.: "Zoledronic acid and geranylgeraniol regulate cellular behaviour and angiogenic gene expression in human gingival fibro-


35- TRICARICO P.M., CROVELLA S. and CELSI F.: "Me-

36- HALAWA A.M., MOHAMED D.G. and OBEID R.F.M.A.: "Capsaicin induced histological and ultrastructural changes in the submandibular salivary gland of albino rats."


38- NAGANO Y., MATSUI H., SHIMOKAWA O., HIRAYAMA A., NAKAMURA Y., TAMURA M., et al.: "Biphosphonate-induced gastrointestinal mucosal injury is medi-

39- GHADIALLY F.N.: "Ultrastructural Pathology of the Cell and Matrix: A Text and Atlas of Physiological and Path-

40- MÖNKKÖNEN H., AURIOLA S., LEHENKARI P., TRICARICO P.M., EPATE A., CROVELLA S. and CELSI F.: "Me-


42- MOTALLEBNEJAD M., ABEDI S.M., SEYEDMAJIDI M., MOGHADAMNIA A.A., ASHRAPFOUR M., SALEHI M., et al.: "A new endogenous ATP analog (ApppI) inhibits the mitochondrial adenine nucleotide translocase (ANT) and is responsible for the apoptosis induced by nitrogen-containing bisphosphonates." British Journal of Pharma-

43- DALEPRANE J.B. and ABDALLA D. S.: "Emerging roles of propolis: Antioxidant, cardioprotective, and antiang-

دراسة هستولوجية وهستوكليميائية مناعية لتأثير الأليافيدونات على الغدة اللعابية تحت الفكية لدى الجرذان الأبيض بالغ وتأثير الوقائي المحتمل للبروبوليس

المقدمة: يعد الأليافيدونات دواء مضاد للتآكل العظام ويستخدم في علاج هشاشة العظام بعد سن اليأس. وأن إعطاءه في المدى الطويل يؤدي إلى تلف الأعضاء وخاصة الغدة اللعابية تحت الفكية. يعد البروبوليس مادة صناعية طبيعية لديها تأثيرات مضادة للأكسدة ومضادة للالتهابات.

الهدف: أجريت هذه الدراسة لدراسة تأثير الأليافيدونات على التركيب الهيستولوجي للغدة اللعابية تحت الفكية في ذكر الجرذان الأبيض البالغ والتأثير الوقائي المحتمل للبروبوليس.

المواد وطرق البحث: تم تقسيم 40 دود من ذكور الجرذان البيضاء البالغة إلى أربع مجموعات (10 جرذان لكل مجموعة) المجموعة الضابطة مجموعتان تم إعطاءها 100 مجم/كلم من زنجبيل البروبوليس، المجموعة B تم إعطاءها 20 مجم/كلم البنيدرونات، وتم إعطاءها البروبوليس 20 مجم/كلم في الأليافيدونات. ويتم إعطاء الجرذان عن طريق الفم لمدة ثلاثون يومًا متواصلًا ثم تأخذ عينات من الفم لاستخدامها في الدراسات الهستولوجية والهستوكليميائية مناعية باستخدام الميكروسكوب الفضي والكروتيوني.

النتائج: كشفت نتائج الدراسات استجابة الكائنات البكتيريا ورئتين الزرقاء لجبلية ب تركيب غير منتظم، أطعمة مغلفة وعوالات مفصولة عن بعضها، فيروسات سيتويبلازمية، أنواع دايت، أنواع غير متطورة وعمرية محظورة، وأظهر البسات-أثيران الأزق زيادة ذو دالة إجتماية في متوسط الكثافة الوعائية بالموطع الضابطة بينما أظهرت الرسوم البيانية الناتجة من تحليل الحمض المخلص للخلايا الظهارية في المنطقة ذات إصبعية في مستوى الكثافة الوعائية بالموطع الضابطة. وقص درجة الكثافة الوعائية بالموطع الضابطة في الغدة تحت الفكية نصفيًا في منطقة إصبعية في مستوى الكثافة الوعائية بالموطع الضابطة. وقد أظهرت التغييرات الفيسيولوجية لخلايا العنانات والخلايا ودود فوسفات وكثافة وثبات سنتوبرازمية، وتأثيراتها غير منتظمة الشكل مع فقدان الأعراس، إزالة دايت تفاعل الفيسيولوجية الخشنة وجهاز زيادة الحجمات الإفرادية مع إتاحة أدوية مضادة منكمية فائقة الإصبع، عدم إتاحة الفيسيولوجيا النمو، إتاحة الفيزيولوجيا التفاعلات بين الخلايا. وقادها مجموعات جست متوسط التسجيلات البدنية والباسات بين الخلايا. وقد أظهرت هذه المجموعة جست متوسط الناتج المرضي السابق.

الاستنتاج: الأليافيدونات قد أحدث تأثيرات ضارة على التركيب الهيستولوجي للغدة اللعابية تحت الفكية لدى الجرذان، وأن إعطاءه مع البروبوليس قد أدى إلى تحسن معتدل لهذا الضرر من خلال تأثيراته المضادة للأكسدة والمضادة للالتهابات.