Histological Study of the Possible Protective Effect of Lycopene on Glucocorticoid-Induced Osteoporosis in Adult Male Albino Rat

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Abstract

Background: Osteoporosis is one of the most commonly occurring bone diseases characterized by decrease in bone mass and increase bone resorption leading to fractures, severe pain and deformities. Glucocorticoid-induced osteoporosis results from long-term and high dose use. Lycopene is an unsaturated carotenoid that gives tomatoes and other fruits their red color. It was reported that it is a natural antioxidant and was associated with the prevention of many chronic diseases and cancers.

Aim of Study: This work was performed to study the possible protective effect of lycopene against glucocorticoid-induced osteoporosis in adult male albino rat.

Material and Methods: 40 adult male albino rats were used that divided equally into four main groups: Group I (control group) was subdivided equally into two subgroups (I-A & I-B), Group II received lycopene orally 30mg/kg once daily for 8 weeks, Group III received prednisolone orally 20mg/kg once daily for 8 weeks, Group IV concomitantly received lycopene and prednisolone orally once daily at the same previous doses for 8 weeks. At the end of experiment, femur heads of all animals were obtained and processed for histological study and stained with Hematoxylin & eosin, Mallory Trichrome and for scanning electron microscope study.

Results: Light microscopic examination revealed structural changes in femur head of Group III in the form of discontinuous thin trabeculae of cancellous bone with apparent faintly stained matrix with multiple pores inside trabeculae associated with minor fractures, wide bone marrow spaces were seen between trabeculae containing abundant clear areas, the endosteal surface showed irregular eroded surface. Regarding Mallory stained sections; they showed decrease collagen fibers contents with their irregular distribution in bone matrix. Scanning electron microscopic specimens showed discontinuous thin and blind ended cancellous bone trabeculae containing some areas of decreased electron density with multiple pores and minor fractures inside them. Wide bone marrow spaces were seen between trabeculae lined with irregular eroded endosteal surface; which revealed the evident of prednisolone induced osteoporosis. Group IV specimens revealed partial preservation of the normal structure of cancellous bone of femur head to some extent.

Conclusion: Osteoporosis resulted from glucocorticoids in adult male albino rats can be partially minimized by lycopene co-administration.

Key Words: Osteoporosis – Prednisolone – Lycopene – Rat.

Introduction

OSTEOPOROSIS is a skeletal disorder characterized by compromised bone strength. It causes bone resorption and fractures, leading to severe pain, deformities, and in certain cases, secondary complications that may be fatal. Bone strength primarily reflects the integration of bone density and bone quality [1].

Glucocorticoids (GCs) are used extensively as anti-inflammatory and immunosuppressive drugs to treat a variety of diseases such as inflammation, cancer, and autoimmune disorders. However, long-term and large doses of oral treatment with glucocorticoids are associated with an increased risk of osteoporosis or fracture. So, Glucocorticoid-Induced Osteoporosis (GIOP) is the main cause of secondary osteoporosis, as GCs have been reported to exhibit detrimental effects on the proliferation and function of osteoblasts [2-4].

Lycopene is an open-chain, unsaturated carotenoid that imparts red color to tomatoes and other fruits. It was found to be a natural antioxidant and was associated with the prevention of some chronic diseases including cardiovascular diseases, prostate and gastrointestinal tract cancers in human. It was stated that the daily consumption of lycopene decreases oxidative injury, bone resorption and increases intake of calcium, niacin, and vitamins A, D, and K in postmenopausal women. In addition, it was reported that lycopene treatment can inhibit
bone loss and increases bone strength in ovariectomized rats [5-7].

**Material and Methods**

This work was carried out at Histology Department, Faculty of Medicine, Tanta University and at The Electron Microscopic Unit, Faculty of Medicine, Tanta University from 2016-2018.

I- *Experimental animals:*

This study was carried out on 40 adult male albino rats. Their weights ranged between 150-200g. The animals were housed under similar conditions of good illumination and ventilation conditions in the animal house. They were housed in suitable clean properly plastic cages with mesh wire. They were fed on a similar commercial laboratory diet and water. They were carefully selected free of gait disturbance, swollen joints or limping [8]. The animals were acclimatized to their environment at least one week before starting the experiment.

The animals were divided into four groups as follows:

- **Group I (the control group):** It included 10 rats used for histological study of the bone. They were divided into 2 subgroups (Subgroup I-A, Subgroup I-B):
  - Subgroup I-A: Included 5 rats received no medications.
  - Subgroup I-B: Included 5 rats received distilled water 2.2ml/100gm BW per day (the solvent of both prednisolone and lycopene capsule) orally by gastric tube for the same period as experimental animals.

- **Group II (the lycopene group):** It included 10 rats given lycopene orally at a dose of 30mg/kg BW once daily for 8 weeks [7]. It was purchased as (Lycopene-50 capsules, manufactured and distributed by KRK Supplements in USA, Sold and Fulfilled by Amazon), each capsule contained 50mg lycopene and its content was suspended in 20ml distilled water to be concentrated as 2.5mg/1ml distilled water. So, each rat received 1.2ml of prepared solution/100gm BW per day which was given orally by gastric tube. This dose was found to induce maximal degree of osteoporosis as reported by Ogoshi et al., [9].

- **Group IV (the protective group):** It included 10 rats given both lycopene orally at a dose of 30 mg/kg BW once daily and prednisolone orally at a dose of 20mg/kg BW once daily for 8 weeks.

II- *Specimen achievement:*

At the appropriate time after 8 weeks and 24 hours after the last dose, rats of all studied groups were anesthetized with intraperitoneal injection of sodium thiopental 200mg/kg BW [10,11]. Then, the head of each femur of all animals was carefully dissected out and processed for histological studies. The bodies of sacrificed animals were packed in a special package according to safety precautions and infection control measures and sent with the hospital biohazards.

III- *Histological study of the bone:*

The head of each femur of all animals was excised and cleaned from muscle attached, trimmed or dissected as appropriate then subjected to the following:

- The head of right femur of all animals was taken and immediately fixed in 10% formal saline solution for 5 days. The specimens were washed by tap water for half an hour then they were decalcified in the chelating agent disodium EDTA solution [12]. The head of left femur of all animals was taken and immediately fixed in gluteraldehyde solution then prepared for scanning electron microscope.

**Decalcification method [13]:**

- Decalcification processes was carried by chelating agent, Ethylene Diamine Tetra Acetic acid (EDTA) where it binds with or absorb the calcium ions; it acts slowly but preserves all cellular components.
- EDTA disodium salt (10%) or EDTA tetra sodium salts (14%) are approaching saturation and can be simple aqueous or buffered solutions at a neutral pH of 7-7.4 or added to formalin. The
time required to totally decalcify dense cortical bone may be 6-8 weeks or longer.

- Decalcification lasted for about 4 weeks, during which the solution was renewed every 2 days until the tissues had softened. The time for exposure to the decalcification will vary; closely monitor the progress is needed.

- The decalcified head of femur was cut longitudinally in a coronal plane along the central portion and specimens were processed to form paraffin blocks.

Serial sections of 5 micrometer thickness were obtained and subjected to the following:

A- Histological stained sections:

1- Haematoxylin and Eosin (H & E) stain [14]:
   Was used to demonstrate the general histological structures of rat’s head of femur among various experimental groups. Nuclei were stained dark blue, cytoplasm were stained varying degrees of pink.

2- Mallory’s trichrome stain [15]: For demonstration of collagen fibers. Collagen fibers were stained blue, cytoplasm and nuclei were stained red.

All these sections were examined and photographed using Olympus light microscope (Tokyo, Japan) coupled to an Olympus digital camera (D XC-1850P, Tokyo, Japan) in Histology Department, Faculty of Medicine, Tanta University.

B- Morphometric study and statistical analysis:

   The image analysis system was done by image analysis software program (Image J, version 1.48), it was used to measure the mean area percentage of collagen fiber contents in Mallory Trichrome stained sections in all groups. ANOVA test was done to compare different groups of our study with the control group.

C- Scanning electron microscopic specimens:

   The head of femur specimens were subjected to the following steps [16,17]:

1- Fixation and cutting the samples: The specimens were rapidly fixed in 2.5% phosphate buffered glutaraldehyde (pH 7.4) at 4°C for 2 hours followed by three rinses in the phosphate-buffer (each 10min). Then the head of femur specimens were cut longitudinally by a saw cutting device for hard tissue that was done in department of dental material, Faculty of Dentistry, Tanta University. Then, specimens were submerged in fixative (2.5% glutaraldehyde and 1% formaldehyde in phosphate buffer at pH 7.4), initially at 37°C, for 24-48h.

2- Critical Point Drying (CPD): Specimens were placed in a sample holder and subjected to a dehydration process using a graded series of solutions of acetone, or ethyl alcohols in concentrations typically from 15, 30, 50, 70, 90, 95, and 100% to gradually replace the water in the sample. Ethanol was substituted by admitting liquid CO2, at room temperature, to a committed pressure vessel, the CPD chamber. Liquid CO2 was allowed to escape intermittently to flush dissolved ethanol from the chamber. The high-pressure CO2 gas was allowed to escape, leaving the samples dry and ready for mounting and coating.

3- Coating (sputtering of gold): Metal coating was applied to a specimen surface to aid signal generation but also to prevent the buildup of a charge on a non-conducting surface. Charging can cause displacement of the primary beam, resulting in some unsightly imaging artifacts. Samples were coated in a sputter coater to monitor the coating process to determine the thickness of coating being deposited. Gold was used as a coating material. A coating thickness up to a few nm is usually sufficient to form a sufficiently conducting surface, ensuring good distribution of metallic particles across the whole specimen surface.

4- Morphological imaging of the bone: Finally, the specimens were examined and photographed using JEOL-JSM-5200LV scanning electron microscope with microscopy suited camera at Electron Microscopic Unit, Faculty of Medicine, Tanta University.

Histological Results

I- Light microscopic results:

A- Haematoxylin & eosin stained sections:

   Group I: Control group (subgroup I-A & subgroup I-B): Examination of H & E stained sections obtained from the head of femur of both subgroups I-A & I-B were similar and showed the same histological picture which was identical to the well-known histological structure of the epiphysis. There were normal anastomosing thick network of cancellous bone trabeculae having bone lamellae and osteocytes inside their lacunae. Smooth regular continuous endosteal surface of that trabeculae covered by osteoprogenitor and osteoblast cells was observed. Bone marrow spaces containing large number of hemopoietic cells were seen between trabeculae Fig. (1A-D).
**Group II:** Lycopene group: Examination of sections obtained from the head of femur of this group revealed apparently the same histological picture as in the control group.

**Group III:** Osteoporosis induced group by prednisolone: Examination of sections obtained from the head of femur of this group revealed changes of different degrees of osteoporosis which were focal in some sections and diffused in others and were different from rat to other. These changes were in the form of discontinuous thin cancellous bone trabeculae with focal areas of apparent faintly stained matrix and pores. There were blind ended trabeculae associated with minor fissures and fractures. Osteocytes inside their apparently large lacunae and some empty lacunae were detected. Wide bone marrow spaces were seen between trabeculae containing less hemopoietic cells and abundant adipocytes than the control group, their lining endosteum showed irregular eroded surface Fig. (2A-D).

**Group IV:** Protective group (prednisolone + lycopene): Examination of sections obtained from the head of femur of this group revealed relatively less destructive changes than that were observed in Group III (osteoporosis induced group). There was also some degree of preservation of the normal histological structure which was variable from one animal to other. There were relatively thin trabeculae of cancellous bone and wide bone marrow spaces in between. Few small pores appeared inside the trabeculae. Apparent more osteocytes inside their lacunae were occasionally observed more than control group. Regular continuous endosteal surface was noticed Fig. (3A,B).

**B- Mallory trichrome stained sections:**

**Group I:** Mallory trichrome stained sections of both subgroups I-A & I-B showed the same arrangement and distribution of the collagen fibers in the head of femur exhibited by blue coloration of bone matrix which was identical to the well-known histological structure of the epiphysis Fig. (4A,B).

**Group II:** Examination of sections obtained from the head of femur of this group revealed apparently the same histological picture as in the control group.
Fig. (2): A photomicrograph of a rat head of femur of Group III (osteoporosis induced group) showing discontinuous thin cancellous bone trabeculae having blind ends and areas of apparent faintly stained matrix (arrows) together with minor fractures (asterisks) and multiple pores (wide short arrows). Wide bone marrow spaces are seen between the trabeculae containing less haemopoietic cells and abundant adipocytes (arrow heads). Notice, endosteum shows irregular eroded surface (wavy arrows) and some osteocytes inside their apparently large lacunae while other lacunae appear without osteocytes (curved arrow) (H & E, 2A, X100, 2B X200, 2C X400 & 2D X1000).

Fig. (3): A photomicrograph of a rat head of femur of Group IV (prednisolone + lycopene group) showing network of relatively thin trabeculae of cancellous bone (arrows) having faintly stained areas or small pores (wide short arrows) and apparent increased osteocytes inside their lacunae (curved arrows). Notice, the regular continuous endosteal surface (bifid arrow) lining the bone marrow cavity (arrow head) (H & E, 3A, X400 & 3B X1000).

Fig. (4): A photomicrograph of a control rat head of femur showing anastomosing thick network of trabeculae of cancellous bone (arrows). Collagen fibers exhibited by blue coloration of bone matrix (C) (Mallory trichrome stain, 4A, X200 & 4B X400).
Group III: Examination of sections obtained from the head of femur of this group revealed irregular distribution of collagen fibers and apparent decrease in their contents as exhibited by faint blue coloration of bone matrix Fig. (5A,B).

Group IV: Examination of sections obtained from the head of femur of this group showed relatively less changes as compared to Group III (osteoporosis induced group). There was irregular distribution of collagen fibers of bone matrix as exhibited by intensity of the blue coloration. Their content was observed to be apparently less than control group but more than that was observed in Group III Fig. (6A,B).

II- Scanning electron microscopic results:

Group I: Examination of scanning electron microscopic specimens obtained from the head of femur of both subgroups I-A & I-B showed anastomosing thick network of cancellous bone trabeculae containing lacunae of osteocytes. Bone marrow spaces were seen between trabeculae lined by smooth regular endosteal surfaces Fig. (7A-D).

Group II: Examination of scanning electron microscopic specimens obtained from the head of femur of this group revealed nearly the same histological picture as in the control group.

Group III: Examination of scanning electron microscopic specimens obtained from the head of femur of this group revealed changes of different degrees of osteoporosis. These changes were in the form of discontinuous, thin and blind ended cancellous bone trabeculae containing areas of decreased electron density with multiple pores and minor fissures and fractures. Wide bone marrow spaces were seen between trabeculae lined with irregular eroded endosteal surface Fig. (8A-D).

Fig. (5): A photomicrograph of a rat head of femur of Group III (osteoporosis induced group) showing discontinuous thin trabeculae of cancellous bone with blind ends (arrows) with irregular distribution of collagen fibers and decreased their contents exhibited by faint blue coloration of bone matrix (C). Notice, minor fractures (asterisks) with multiple pores inside the trabeculae (wide short arrows) and wide bone marrow spaces in-between containing abundant adipocytes (arrow heads). (Mallory trichrome stain, 5A, X200 & 5B X400).

Fig. (6): A photomicrograph of a rat head of femur of Group IV (prednisolone + lycopene group) showing network of relatively thin trabeculae of cancellous bone (arrows) with irregular distribution of collagen fibers and their contents appear less than normal but more than that of Group III as exhibited by blue coloration of bone matrix (C) (Mallory trichrome stain, 6A, X200 & 6B X400).
Fig. (7): A scanning electron micrograph of a control rat head of femur showing anastomosing thick network of cancellous bone trabeculae (arrows) containing lacunae of osteocytes (bifid arrows). Bone marrow spaces (arrow heads) are lined by smooth regular endosteal surface (wavy arrows) (Scanning EM 7A, X35, 7B X100, 7C X200 & 7D X750).

**Group IV**: Examination of scanning electron microscopic specimens obtained from the head of femur of this group revealed relatively thin trabeculae of cancellous bone as compared to control group, containing lacunae of osteocytes. Occasionally, fissures and pores were observed in few trabeculae. Bone marrow spaces were seen between trabeculae, their lining endosteal surfaces were smooth with focal areas of irregularity Fig. (9A-D).

Fig. (8): A scanning electron micrograph of a rat head of femur of Group III (osteoporosis induced group) showing discontinuous thin and less electron dense cancellous bone trabeculae with blind ends (arrows) containing minor fissures and fractures (asterisks). Wide interconnected bone marrow spaces are seen between trabeculae (arrow heads) are lined by irregular eroded endosteal surface (wavy arrows). (Scanning EM 8A, X35, 8B X100, 8C X200 & 8D X750).
**Quantitative assessment and statistical analysis:**

Evaluation of collagen fiber contents in Mallory’s trichrome stained sections:

Statistical analysis of the data collected by the image analysis software program (Image J, version 1.48) revealed that there was no significant difference ($p$-value >0.05) in the mean area percentage of collagen fiber contents of sections from the head of femur of Group II and Group IV as compared to control group. But Group III (osteoporosis induced group) showed statistically highly significant decrease ($p$-value <0.001) in the mean area percentage of collagen fiber contents as compared to control group. Also, Group III showed statistically highly significant decrease ($p$-value <0.001) in the mean area percentage of collagen fiber contents as compared to Group IV (protective group). All these data are illustrated in (Tables 1,2) and Histogram (1).

Table (1): The mean area percentage of collagen fiber contents in different studied groups (ANOVA test).

<table>
<thead>
<tr>
<th>Groups</th>
<th>Area percentage of collagen fiber contents</th>
<th>ANOVA</th>
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<tr>
<td></td>
<td>Range</td>
<td>Mean ± SD</td>
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<tr>
<td>I- Control</td>
<td>26.363-37.263</td>
<td>31.207±3.584</td>
</tr>
<tr>
<td>IV- Lycopene + Prednisolone</td>
<td>23.777-37.507</td>
<td>30.003±4.545</td>
</tr>
</tbody>
</table>

Table (2): The mean area percentage of collagen fiber contents in different studied groups (Tukey’s test).

<table>
<thead>
<tr>
<th>TUKEY’S Test</th>
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<tr>
<td>I &amp; II</td>
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<td>&gt;0.05</td>
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SD: Standard Deviation.

$p$-value >0.05: Non-significant.

$p$-value <0.001 (*): Highly significant.
Discussion

Our study was done to evaluate the possible protective effects of lycopene against glucocorticoid-induced osteoporosis in adult male albino rat.

The head of femur was chosen in this work to evaluate the changes in cancellous bone trabeculae because it is the most sensitive part to the effect of glucocorticoid treatment on GIOP [18]. Meanwhile, Kumar et al., [19] stated that trabecular bone is usually affected more severely than cortical bone because of its greater metabolic activity.

The microscopic examination of haematoxylin & eosin stained sections of the femur head, in the osteoporosis induced group showed changes in the form of discontinuous thin trabeculae of cancellous bone with focal areas of apparent faintly stained matrix with multiple pores inside trabeculae. There were blind ends of trabeculae associated with minor fissures and fractures. Wide bone marrow spaces were seen between trabeculae containing less hemopoietic cells and abundant adipocytes than the control group. The endosteum surface appeared irregular and eroded. In addition, osteocytes inside their apparently large lacunae and some empty lacunae were detected.

Examination of Mallory trichrome stained sections obtained from the femur head of this group showed irregular distribution of collagen fibers and decrease their contents exhibited by faint blue coloration of bone matrix. Statistical analysis of the data collected by the image analysis software program revealed that there was statistically highly significant decrease ($p$-value <0.001) in the mean area percentage of collagen fiber contents of sections from the head of femur of this group as compared to control group.

Scanning electron microscope was chosen in our study to distinguish various bone surface activity status and detect variations in mineralized tissue matrix properties [20,21]. Specimens examined by SEM revealed discontinuous, thin and blind ended cancellous bone trabeculae containing some areas of decreased electron density with multiple pores and minor fissures and fractures inside them. Wide bone marrow spaces were seen between trabeculae lined with irregular eroded endosteal surface.

Therefore, the present study revealed the evidence of prednisolone induced osteoporosis, demonstrated by light and scanning electron microscopes which was identical to what is described by the histopathologists Matthew & Laura [22], who stated that the hallmark of osteoporosis is a reduced bone mass leading to trabecular thinning and perforation resulting in progressive microfractures. Also, it leads to reduction of the trabecular network connectivity and more widely separated than usual.

Yao et al., [23] described that glucocorticoids in animal models impact trabecular bone first, result in significant reductions in its volume and whole bone strength. Similar finding was found by Barbara et al., [24], who reported that with prolonged use of glucocorticoid there is reduction of mean wall thickness of trabecular bone packets, with a consequent decrease in total bone volume. This explain the thin trabeculae with eroded surfaces seen in the present study.

Many previous researches explained the mechanism by which glucocorticoid induced osteoporosis. Goulding and Flower [25] reported glucocorticoids inhibit intestinal absorption of calcium causing secondary hyperparathyroidism, in turn, stimulates osteoclasts to increase bone resorption. In addition, steroid drugs directly inhibit the activity of osteoblasts and stimulate the urinary excretion of both calcium and phosphorus.

The large lacunae seen in the present study were previously explained by Lane et al., [26], who studied glucocorticoid treatment in animal models as a direct effect on osteocytes, resulting in a modification of their microenvironment and the generation of surrounding areas of hypomineralized bone around the lacunae with a reduction of mineral-to-matrix content.

Glucocorticoid induced osteoporosis results from variation in bone turnover. It increases resorption by increasing osteoclast lifespan and
decreases formation by inducing osteoblast and osteocyte apoptosis, that explain the presence of some empty lacunae [27]. In addition, glucocorticoid inhibits the differentiation of mesenchymal precursor cells into osteoblasts by directing the precursors toward adipocyte development, which explains the presence of more adipocytes in bone marrow spaces in osteoporosis group of the present study [28].

During the remodeling process, osteoblasts and osteoclasts interact with multiple molecular agents including hormones, growth factors, and cytokines. Metabolic bone diseases such as osteoporosis result from disturbances in the remodeling process [29]. Moreover, Preedy and Watson [30] reported that osteoporosis is in part due to the actions of oxidative stress on both osteoclasts and osteoblasts; antioxidants counteract these actions. This goes in line with Callaway and Jiang [31], who stated that Reactive Oxygen Species (ROS) may be a key player in the pathogenesis of different pathological conditions, such as osteoporosis, with established positive correlations between the presence of excessive ROS and bone loss. Altindag et al., [32] added that antioxidant supplementation has been proposed as a novel therapeutic strategy in this disease. This was in agreement with Rao et al., [33], who explained that under normal physiological conditions, oxidative stress can be prevented by promoting antioxidant defenses. Exogenous antioxidants from dietary sources present in fruit and vegetables are also utilized to combat oxidative stress.

Although the severity and consequence of glucocorticoid-induced osteoporosis, it is preventable, treatable and potentially reversible. The prevention and treatment of GIOP include some general measures, calcium and vitamin D supplementation and treatment with bone anabolic and antiresorptive agents as bisphosphonates [34]. In addition, there is an increasing demand for complementary and alternative medicine for the prevention and treatment of osteoporosis due to the adverse side effects of hormonal therapy and the bisphosphonates in the management of osteoporosis; these include diet, exercise, and nutritional supplements. Recent trials of antioxidant supplementation are an effective therapeutic tool in the prevention of GC-induced bone damage [35].

In the present study, co-administration of lycopene minimized the structural changes induced by prednisolone together with preservation of the normal histological structure which was variable from animal to animal. However, there were relatively thin trabeculae of cancellous bone and wide bone marrow spaces were seen between trabeculae. Few small pores inside trabeculae and apparent more osteocytes inside their lacunae were observed. Mallory trichrome stained sections, they showed relative preservation of collagen fibers content as exhibited by blue coloration of bone matrix. Statistical analysis of the data collected by the image analysis software program revealed that there was no significant difference (p-value >0.05) in the mean area percentage of collagen fiber contents of sections from the head of femur of this group as compared to control group.

Examination of scanning electron microscopic specimens of protective group confirmed the results obtained by the light microscope and revealed many lacunae of osteocytes inside the thin trabeculae of cancellous bone. Occasionally, some fissures and pores were observed in few trabeculae. The lining endosteal surfaces of the bone marrow spaces were smooth with focal areas of irregularity.

Plant derivatives such as lycopene from tomato extract is a natural antioxidant helping in contrast- ing the progress of some pathological events whose origin resides in the generation of free radicals [36]. The protective effect of lycopene can be attributed according to epidemiological studies of Sahni et al., [37], who demonstrated the role of lycopene as an antioxidant in the prevention of oxidative stress-related osteoporosis and correlation with bone mineral density in postmenopausal women. This goes in line with Mackinnon et al., [6], who stated the role of lycopene in reduction either oxidative stress or levels of bone turnover markers in postmenopausal women, thus resulting potentially beneficial in reducing the risk of osteoporosis.

It was found that lycopene has stimulating effect on osteoblast cell proliferation as well as a stimulatory effect on alkaline phosphatase activity in human osteoblast [38]. It is also able to prevent and repair the damaging effects of oxidative stress. Also, Yang et al., [39], claimed that lycopene may shift the bone metabolic equilibrium towards an increase in the anabolic state leading to decrease in bone resorption levels in osteoporosis.

The effects of lycopene on osteoclasts have been also reported by Krishnadev et al., [40], who demonstrated that lycopene can inhibit multinucleated osteoclast cell formation as well as the formation of ROS-secreting osteoclasts in bone marrow cells from rat femur. The protective role of lycopene in human disorders such as osteoporosis may be due to its ability to suppress biomarkers of oxidative stress which play important role of the pathogenesis.
of osteoporosis, because oxidative stress controls the functions of both osteoclasts and osteoblasts [41].

The protective role of lycopene according to Batra et al., [42] could be attributed to the involvement of other important processes such as stimulation of gap junction communications, induction of the production of detoxifying enzymes leading to modulation of cell cycle and apoptosis. Since intercellular communications through gap junctions are key events of normal osteoclast and osteoblast metabolism, consequently, on bone remodeling. Lycopene may modulate gap junction communication to suggest that this may also represent a potentially target of action of lycopene on bone cells.

It was observed that lycopene, at levels found in plasma after the ingestion of lycopene-containing products, leading to decrease osteoclast differentiation, but did not affect cell density/survival; calcium-phosphate resorbing ability was also decreased. On the other hand, osteoblast proliferation (via a decrease on apoptosis) and differentiation was increased in the presence of lycopene which may contribute to the promotion of a proper health status of bone tissue. They concluded that, the previous information might be relevant for the prevention and delay in the progression of osteolytic bone conditions [43].

Furthermore, Iimura et al., [44] showed that lycopene intake by ovariectomized rats facilitates the increase of bone mineral density and bone strength. This was in consistent with Ardawi et al., [45], who stated that, in an ovariectomized mice model, lycopene can help against the loss of bone mass, that primarily suppressed bone turnover to restore bone strength and microarchitecture, improving bone biomechanical properties with an increase in both antioxidant enzyme activity and osteoblast activity.

Lycopene can be used either as a dietary alternative or as a complement to the pharmaceuticals used in the form of capsules 30-70mg/day to exert equivalent antioxidant potency in reducing the risk of osteoporosis in postmenopausal women [46]. Similar finding was reported by Jiuhong et al., [47] based on the evidence from observational studies, meta-analysis supported the hypothesis that higher dietary total carotenoids intake might be potentially associated with a low risk of hip fracture.

On the other hand, it was found a non-significant association between serum lycopene and hip fracture risk among elderly women [48]. Furthermore, Wolf et al., [49] reported that there is no significant protective association at any bone mineral density site with serum carotenoids. Sahni et al., [50] also showed a null association of dietary lycopene with hip fracture risk among women.

Conclusion:

Based on the current study and from all previously mentioned histological and statistical results, it could be concluded that the osteoporosis resulted from administration of glucocorticoids in adult male albino rats can be partially minimized by lycopene co-administration. So, it is recommended that lycopene can be used as a dietary alternative to drug therapy or as a supplement to people at risk for osteoporosis.

References

Histological Study of the Possible Protective Effect of Lycopene on GIOP is in Adult Male Albino Rat


دراسة هستولوجيّة التأثير الوقائي المحتمل لمادة الليوكونين على هشاشة العظام المستحدثة بواسطة الجلوكوكورتيكويد في ذكر الجلد الأبيض البالغ

المقدمة: يعتبر مرض هشاشة العظام واحد من أكثر أمراض العظام شيوعًا، وتتميز بإخفاقات والأشكال المختلفة للعام مع حدوث مضاعفات مثل الكسور والأمراض المصاحبة والتشوهات. يعتبر الجلوكوكورتيكويد من أهم الأسباب لمرض هشاشة العظام نتيجة الاستخدام طويل الأمد والجرعات العالية. مركب الليوكونين هو من مجموعة الكورتيكولونات غير المشبعة، والمستخلص من اللون الأحمر في الطماطم والفواكه الأخرى، وقد أظهر أنه له دور وقائي في بعض الأمراض المزمنة والسرطانات نتيجة خصائصه المضادة للأكسدة.

الهدف من العمل: هو دراسة التأثير الوقائي المحتمل لمركب الليوكونين على هشاشة العظام المستحدثة بواسطة الجلوكوكورتيكويد في ذكر الجلد الأبيض البالغ.

المواضيع والطرق: تم استخدام 40 من ذكر الجلد البضائع البالغة والتي قسمت إلى أربع مجموعات متساوية: المجموعة الأولى (الضبيطة).

وتتم تقسيمها إلى مجموعتيتين فرعية متساويتين (1-6). المجموعة الثانية أُعطيت 20 مجم/كم من الليوكونين عن طريق الفم مرة واحدة يوميا لمدة 8 أسابيع. المجموعة الثالثة أُعطيت 20 مجم/كم من البرينيديزولون عن طريق الفم مرة واحدة يوميا لمدة 8 أسابيع. وفي نهاية التجربة تم الحصول على عينات من رأس عظام الفخذ ومعالجتها لدراسة هستولوجيّة عن طريق المجهر الضوئي بعد صياغتها باستخدام صبغات الهيماتوكسيلين والبيبين، وال蔫اوي ثلاثي الألوان وكذلك فحصها عن طريق المجهر الإلكتروني الماسح.

النتائج: كشف الفحص المجهر الضوئي لعينات رأس عظام الفخذ في المجموعة الثالثة غير محترفة في النسيج العظمي على هيئة ترقق للعظام الإسفنجي بشكل واضح مع وجود فراغات عظمية وكسور رقيقة مع وجود إنسجام كبير لل الأجسام التي تحتوي على الخلايا العظمية كما ظهر تأكل في السطح العظمي. وقد أوضحت صبغة الثلاثيات إناث الألوان التوزيع غير منتظم لنياف الكولاجين مع نقص كبيثا. كما كشف الفحص المجهر الإلكتروني الماسح لعظام الفخذ تراکك وتأكل في العظام الإسفنجية مع عدم إنقاص للعظام العظمية ووجود كسور عظمية رقيقة وكذلك إنسجام كبير لل الأجسام التي تحتوي على الخلايا العظمية مما يثبت وجود هشاشة للعظام ناتجة عن إسهام مقار البرينيديزولون.

كما أظهرت المجموعة الرابعة حفاظ جزئيًا للشكل الهستولوجي للعظام الإسفنجي لرأس عظام الفخذ.

الاستنتاج: هشاشة العظام الناتجة عن إعطاء الجلوكوكورتيكويد في ذكر الجلد الأبيض البالغ يمكن التقليل منها جزئيًا بالتناول المصاحب لمركب الليوكونين.