Protective effects of *Cornus mas* fruit extract on methotrexate-induced alterations in mice testicular tissue: Evidences for histochemical and histomorphometrical changes in an animal model study

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**Abstract**

Methotrexate (MTX) as a chemotherapeutic agent, has adverse effects on reproductive organs by enhancing oxidative stress. In this study, the protective effects of *Cornus mas* fruit extract (CMFE) against MTX side effects were evaluated. Forty-eight mature male NMRI mice were divided into six groups: group 1 (control) received 0.10 mL per day of normal saline intraperitoneally (IP), group 2 received MTX (20.00 mg kg\(^{-1}\) per week, IP), group 3 received MTX along with CMFE 250 mg kg\(^{-1}\) per day by oral gavage, group 4 received MTX along with CMFE 500 mg kg\(^{-1}\) per day by oral gavage, group 5 received MTX plus 1000 mg kg\(^{-1}\) per day of CMFE by oral gavage, and group 6 received 1000 mg kg\(^{-1}\) per day of CMFE extract, orally. All animals were treated for 35 consecutive days. Thickness of testicular capsule and germinal epithelium and diameter of seminiferous tubules were measured. Intra-cytoplasmic levels of carbohydrate, unsaturated fatty acid (UFA) and alkaline phosphatase were assessed. Serum level of testosterone and testicular total antioxidant capacity (TAC) were also evaluated. The results demonstrated that MTX administration caused morphometrical parameters except the thickness of testicular capsule were significantly different in comparison to control group and decreased cytoplasmic concentration of carbohydrate in the first three layers of germinal epithelium and increased the UFA levels. Contrariwise, CMFE ameliorates the condition. Moreover, CMFE increased testosterone level and increased the MTX-reduced TAC level. In conclusion, it was revealed that CMFE decreased the cellular atrophy by controlling the energy substrate utilization based on lipids and carbohydrates via provoking the testicular antioxidant status.

**Introduction**

Methotrexate (MTX), a folate antagonist compound, was introduced in 1950 as a chemotherapeutic drug.\(^1\) It is used to control the tumor malignancies including acute and chronic leukemia, lymphoma, bladder cancer, breast cancer, and testicular tumors.\(^2,3\) Moreover, MTX is administrated as an immunosuppressive drug in patients with autoimmune disorders like arthritis rheumatoid and psoriasis.\(^4,5\) A study has shown that MTX could induce degenerative changes in the reproductive system of male Swiss mice.\(^6\) It was also reported that the administration of MTX resulted in a significant reduction in sperm count, an increase in sperm morphological disorders, and sperm DNA disintegration, all associated with the testicular damages caused by MTX.\(^7\)

It has been indicated that the administration of 25.00 mg kg\(^{-1}\) and 50.00 mg kg\(^{-1}\) of MTX in adult rats resulted in elevated damages to seminiferous tubules and increased edema of the interstitial tissue of the testes.\(^8\) It was shown that intramuscular administration of 0.15 and 0.30 mg kg\(^{-1}\) of MTX in both male and female adult mice remarkably reduced spermatogenesis index and follicular growth of ovarian tissue, respectively.\(^9\)

The oxidative stress induced by MTX was considered as the main mechanism for the degenerative impacts of this chemical agent in testicular and brain tissues.\(^10-12\) This oxidative stress in testicular tissue correlates indirectly...
with the lowered endocrine status. Accordingly, diminished Leydig cells distribution, as well as the lowered testosterone levels in serum, are associated with down-regulated antioxidant status in MTX-administrated rats. This clarifies the correlation between defected endocrine activities and enhanced oxidative stress following chronic exposure to MTX.

*Cornus mas* fruit (CMF), commonly known as cornelian cherry, is a deciduous fruit of a shrub or small tree, native to central and southern Europe and western Asia. In Iranian traditional herbal therapy, CMF is used to control inflammation, stomach aches and cramps, diarrhea, different skin infections, intestinal parasites, and hemorrhoids. Biochemical analysis of CMF has shown that it is a rich resource of anti-oxidant and phenolic compounds. In addition, it contains some vitamins including vitamin C, B1, B2, E, as well as anthocyanins, flavonoids, along with very high levels of oxalic acid. Considering the antioxidant and anti-inflammatory properties of CMF, the present study was designed to evaluate the protective effect of CMF on MTX-induced damages to testicular tissue and sperm levels. For this purpose, the histochemical analyses were carried out for detecting any changes in alkaline phosphatase (ALP) enzyme synthesis rate, and intra-cyttoplasmic carbohydrate and unsaturated fatty acid ratios. Biochemical analyses for measuring the testicular antioxidant capacity (TAC) and serum level of testosterone were also carried out.

**Materials and Methods**

**Animals.** Forty-eight mature healthy male NMRI mice were included in the present study. All animals were kept for one week in standard conditions including temperature 22.00 ± 2.00 °C, relative humidity (30.00 to 60.00%) and light/dark cycle (10hr/14hr) in the animal house of Faculty of Veterinary Medicine, Urmia University and fed with standard pellet diet and water.

The study protocol was reviewed and approved by Urmia University Ethical Committee (AECVU-154-2018). All animals received humane care following the “Guide for the Care and Use of Laboratory Animals” prepared by the “National Academy of Sciences” and published by the “National Institutes of Health” (NIH).

**Preparation of hydro-alcoholic extract of *Cornus mas* fruit extract (CMFE).** Fresh CMFs were collected from the rural regions of Qazvin province, Iran and identified by a botanical expert, at the Department of Botanic Sciences, Hamadan Research Center for Agricultural and Natural Resources, Hamadan, Iran. The fruits were dried naturally in room temperature (23.00 - 24.00 °C) for six days and pulverized to powder using an electric blender. Then, 100 g of the plant powder was suspended in 600 mL of 50/50% (v/v) hydro-ethanolic solution for 96 hr at room temperature. The mixture was then filtered using a fine muslin cloth followed by a filter paper (Whatman No 1). The filtrate was placed at 40.00 °C in an oven to dry. The obtained clear extract was used in the study. The extract was kept at −15.00 °C until it was used in the experiment.

**Study design.** Frothy-eight mature male NMRI mice were randomly divided into six groups of eight mice in each group as follows: 1) Control group which received 0.10 mL per day of normal saline intraperitoneally (IP), 2) MTX group (Koçak Farma, Tekirdağ, Turkey) which received 20.00 mg kg⁻¹ per week of MTX, IP, 3) MTX/CMFE/250 group which received MTX in the same way of group 2 plus 250 mg kg⁻¹ per day of CMFE orally via oral gavage, 4) MTX/CMFE/500 group which received MTX plus 500 mg kg⁻¹ per day of CMFE via oral gavage, 5) MTX/CMFE/1000 group which received MTX plus 1000 mg kg⁻¹ per day of CMFE via oral gavage, and 6) CMFE group which received 1000 mg kg⁻¹ per day of CMFE extract via oral gavage. All animals were treated daily for 35 consecutive days.

**Gas chromatography-mass spectrometry (GC-MS) analysis of CMFE.** The GC-MS was equipped with an HP-5MS (30 m × 0.25 mm, 0.25 μm film thickness; Hewlett-Packard, Palo Alto, USA) non-polar capillary column. The Column temperature was set initially at 50.00 °C for 1 min, then gradually increased to 220 °C with a temperature ramp of 4.00 °C per min and finally held at 220°C for 5 min. The carrier gas was helium at a flow rate of 2.94 mL min⁻¹. The injector temperature was set at 240 °C employing the split injection mode and the split ratio of 1:50. The MS scan conditions were: ion source temperature: 200 °C; interface temperature: 250 °C; ionization energy: 70.00 eV; mass scan range: 40.00 - 350 m/z.

**Histological analyses.** After five weeks, all animals were anesthetized by IP administration of ketamine (90.00 mg kg⁻¹; Alfasan, Woerden, Holland) and xylazine (5.00 mg kg⁻¹; Alfasan), then one half of the right and left testes were fixed in 10.00% neutral buffered formalin and after tissue processing, embedded in paraffin. Semi-sections (5.00 μm) were stained by hematoxylin and eosin dyes for histomorphometric analyses. The other half of the right and left testes were fixed for histochemical studies. The histomorphometric parameters (capsular thickness, seminiferous tubules diameter and height of germinal epithelium) were measured using a light microscope (Olympus, Tokyo, Japan) with calibrated graded optical lens. Each grade that observed at 400×, estimated 2.55 μm or 100× magnification estimated 10.89 μm.

**Histochemical examination of testicular alkaline phosphatase (ALP) activity.** Histochemical measurement of ALP was performed using a modified method described previously. Briefly, tissue sections were deparaffinized in xylene and hydrated through graded descending alcohol series, and incubated for overnight in a solution of 1.00%
magnesium chloride in 100 mm Tris maleate buffer (pH 9.20; Sigma, St. Louis, USA). Tissue sections were then incubated for 2 hr at room temperature in a solution containing ALP substrate, freshly prepared 100 mM Tris-maleate buffer (pH 9.20), 0.20 mg mL⁻¹ naphthol AS-MX phosphate (Sigma) and 0.40 mg mL⁻¹ Fast Red TR. After washing with distilled water, the sections were counterstained with hematoxylin and mounted with Kaiser’s glycerol jelly (Merck, Darmstadt, Germany).

**The assessment of intra-cytoplasmic carbohydrate amount.** To evaluate the carbohydrate amount, a peroxidase Schiff (PAS) kit (Pajohesh Asia, Tehran, Iran) was used. In brief, the paraffin sectioned specimens were deparaffinized and rehydrated. Hydrated slides were oxidized in 5.00% of the periodic acid solution for 5 min. Having rinsed in distilled water, the slides were placed in Schiff reagent for 15 min and washed with lukewarm water. After 5 min of incubation in room temperature, the slides were counterstained with Meyer’s hematoxylin.

**Unsaturated fatty acid (UFA) analyses.** The Oil Red O staining special commercial kit was used for UFA measurement based on manufacturer instructions (Ayandeh Science Co. Urmia, Iran). Briefly, samples that were fixed in 10.00% neutral buffered formalin were sectioned in frozen by cryostat microtome (Bright Instrument Co. Ltd., Huntingdon, UK) to 10.0 µm slices at air-dried conditions. Then, the slides were fixed again in formalin and thoroughly washed with distilled water before being rinsed in 60.00% isopropanol. Then, the slides were stained with freshly prepared Oil Red O solution for 15 min. The slides were rinsed again in 60% isopropanol and lightly stained with hematoxylin. Finally, the slides were rinsed several times with distilled water and mounted with glycerol jelly.

**Blood sampling and hormonal assay.** Blood samples were collected from the mice and serum were separated by centrifugation at 3000 g for 5 min. Testosterone was measured by an immunoradiometric method using WHO/Sigma Asso-RTGC-768/98 kit (Abbott Laboratories, Abbott Park, USA). The intra-assay coefficient of variance and inter-assay coefficient of variance of 10 tests for testosterone were 3.56 and 8.98, respectively.

**Assessment of testicular TAC level.** An amount of 0.30 - 0.40 g of the testicular tissue was homogenized in ice-cold KCl (150 mM) and centrifuged at 3000 g for 10 min. The supernatants were used for evaluating TAC level. The assessment of TAC was carried out based on the ferric reduction antioxidant power (FRAP) assay as described previously.²⁰

**Statistical analyses.** All the histomorphometrical, hormonal and TAC data were expressed as mean ± SD. Data were statistically analyzed using SPSS software (version 20.0; IBM Corp. Armonk, USA) using one-way ANOVA followed by Bonferroni post-hoc test. A p-value less than 0.05 was considered significant.

**Results**

**GC-MS findings.** GC-MS analysis of the CMFE by HP-5MS column revealed the presence of 108 components. The CMFE consisted of 34.26% alcohols (17.59% aliphatic, 8.33% monoterpenic, 5.56% sesquiterpenic, and 2.78% aromatic alcohols), 22.23% hydrocarbons (12.04% aliphatic, 1.85% aromatic, 7.41% sesquiterpenic, and 0.93% diterpenic hydrocarbons), 11.11% esters (2.78% aliphatic, 2.78% fatty acid methyl, 4.62% fatty acid ethyl, and 0.93% terpenoid esters), 10.19% ketones (7.40% aliphatic ketones, 1.85% lactones, 0.93% benzoquinone derivatives), 8.33% aldehydes (2.78% aliphatic, 3.70% monoterpenic, and 1.85% aromatic aldehydes), 7.41% acids (0.93% aliphatic, and 6.48% fatty acids), and 6.48% heterocyclic compounds (2.78% furan derivatives and 3.70% terpenoids). The major components of the extract of the mature fruits were: 16-kaurene, α-terpineol, 1-tridecene, α-cadinene, 1-octen-3-ol, furfural, and 1-pentadecene.

**CMFE inhibited the MTX-induced histological damages.** The results demonstrated that MTX caused severe histological degenerations in testes. The animals received MTX, intensive seminiferous tubules atrophy, germinal cells dissociation, and severe edema in the testicular interstitial tissue were observed. However, the administration of CMFE, in a dose-dependent manner, inhibited the MTX-induced damages. Accordingly, the germinal epithelial dissociation in seminiferous tubules, as well as the edema, was reduced in a dose-dependent manner. Comparison of the testis capsule thickness (to check fibrosis) among study groups showed no significant differences. But, the capsular thickness was more in the MTX group in comparison to other groups. Meanwhile, CMFE reduced the capsular fibrosis in a dose-related manner. There were no histological changes in testes of the animals received CMFE (Table 1).

**CMFE balanced the intra-cytoplasmic carbohydrate and fatty acid concentrations.** Histochemical analyses showed that MTX in a single administration reversed the cytoplasmic carbohydrate accumulation. Because of the low levels of carbohydrates, the spermatogonia cells and the spermatocytes (types A and B) exhibited faint reaction for PAS staining. Meanwhile, the same cells in the control group animals showed intensive reaction for PAS staining, representing high carbohydrate levels.

The CMFE and MTX treated animals showed approximately the same staining characteristics to those of the control group. Accordingly, the cytoplasmic carbohydrate level was increased in the first three germinal lineages in CMFE treated animals in a dose-dependent manner (Fig. 1).

Moreover, carbohydrate alterations were calculated in Sertoli and Leydig cells, the results showed that the carbohydrate accumulation was increased in CMFE treated animals versus to MTX group.
Table 1. Testicular histomorphometric parameters in different experimental groups. The values are expressed as mean ± SD.

| Groups           | Germinal epithelium height (µm) | Tubular diameter (µm) | Capsule thickness (µm) |
|------------------|----------------------------------|-----------------------|------------------------|
| Control          | 50.47 ± 2.50^a                   | 217.91 ± 2.58^a       | 12.35 ± 0.68^a         |
| MTX              | 30.47 ± 2.58^b                   | 117.00 ± 2.80^b       | 14.71 ± 0.60^a         |
| MTX/CMFE/250     | 41.75 ± 2.66^a                   | 175.83 ± 2.19^bc      | 12.78 ± 0.57^a         |
| MTX/CMFE/500     | 42.51 ± 2.23^a                   | 196.82 ± 3.69^ac      | 12.93 ± 0.43^a         |
| MTX/CMFE/1000    | 41.15 ± 1.00^a                   | 184.34 ± 8.39^ac      | 14.42 ± 0.47^a         |
| CMFE             | 46.35 ± 2.17^a                   | 201.46 ± 1.94^ac      | 14.63 ± 0.63^a         |

MTX: Methotrexate, CMFE: Cornus mas fruit extract.

abc Different superscripts in each column are representing significant differences between groups (p < 0.05).

Fig. 1. Intra-cytoplasmic carbohydrate amount in testis cross-sections. A) Control group, germinal epithelium is presented normally; B) MTX group, significant decrease in intra-cytoplasmic carbohydrate amount especially in the first three layers of the germinal epithelium; C) MTX/CMFE/250 group; D) MTX/CMFE/500 group; E) MTX/CMFE/500, co-administration of CMFE enhanced carbohydrate storage in all cellular layers (red reactions) and F) MTX/CMFE/1000 group (PAS staining, Bar = 60 µm).

Fig. 2. Accumulation of UFA in testis: A) Control group: The germinal epithelium of the control testis is presented normally; B) MTX group: Decrease in intra-cytoplasmic Oil-Red-O reaction for fatty acids; C) MTX/CMFE/500 group, and D) MTX/CMFE/1000 group. See ameliorated fatty acids accumulation in CMFE group (Oil-Red-O staining, Bar = 60 µm).

CMFE reduced ALP enzyme activity. The histochemical analyses for cellular ALP activity showed that the animals in the control group exhibited slight ALP activity only in the endothelial cells. However, animals in the MTX group showed significantly higher ALP positive sites in the germ layers, especially in the spermatocytes, spermatids, and Leydig cells compared to the other experimental groups (p < 0.05). In contrast, CMFE reduced ALP activity in a dose-dependent manner.

Accordingly, MTX/CMCF/1000 group showed the lowest ALP activity compared to MTX/CMCF/250 and MTX/CMCF/500 groups (p < 0.05), (Fig. 3). No considerable changes were observed in ALP activity in the CMFE group.
administration remarkably inhibited the MTX-induced derangements in a dose-dependent manner.

The glucose is considered as the main substrate of energy in the spermatogonia, spermatocytes, and spermatid cell lineages. In vitro experiments demonstrated that the germinal cells use large amounts of lactate and glucose for their mitosis and meiosis. Carbohydrate staining showed that MTX significantly decreased intra-cytoplasmic carbohydrate concentration especially in Sertoli and spermatogonia cells as well as spermatocyte lineages. Indeed, Sertoli cells metabolize glucose and convert it into lactate. Lactate then secrets from Sertoli cells into the tubular medium via stimulation of the follicular stimulating hormone (FSH). Therefore, we could conclude that MTX disrupted the glucose and lactate metabolisms, recognizable by decreased intra-cytoplasmic carbohydrate level, and ultimately affected the spermatogenesis process. Meanwhile, the CMFE up-regulated these metabolic pathways, enhanced the cytoplasmic carbohydrate levels, and so raised the MTX-reduced energy substrate in Sertoli and germlinal cells. Carbohydrate accumulation in the cytoplasm of Leydig cells was decreased significantly in the MTX group, however, it was reversed in MTX/CMFE groups, dose-dependently. Carbohydrate accumulation in the CMFE group was the same as the control group.

More intensively, under conditions that the glucose transport and/or metabolism undergoes a severe derangement, the germlinal cells switch their energy substrates from glucose to UFA. Tissue survival and the activity of spermatocytes and spermatids depend strictly on carbohydrate metabolism both at anaerobic and aerobic pathways for maintaining the ATP synthesis. Therefore, the cells in germinall epithelium use UFAs as a secondary source of energy. Our histochemical analyses showed that all cellular layers exhibited intensive UFA in the animals of the MTX group. However, the MTX and CMFE treated animals, albeit depending on dose, exhibited lower cytoplasmic UFA levels and up-regulated carbohydrate accumulation. It seems that CMFE acted by reducing the UFA levels and up-regulated the carbohydrate levels which was inhibited by MTX treatment. Sertoli cells in MTX animals exhibited the highest UFA positive foci in the cytoplasm. In order to understand this alteration one should note that Sertoli cells play a critical role in the phagocytosis of residuals during spermatogenesis and spermiogenesis.

Our histological analyses showed that the percentage of atrophied seminiferous tubules with germinal epithelium dissociation was increased in the MTX group. In contrast to CMFE animals, the MTX group showed decreased TAC level. The enhancement of abnormal metabolism associated with increased cellular damage and elevated cellular degeneration results in provoking oxidative stress in testicular tissue. Beside this fact, the
down-regulated endocrine status in MTX animals, recognizable by decreased serum level of testosterone, promoted the cellular atrophy and ultimately enhanced the oxidative stress. In fact, Leydig cells control the physiologic interactions of Sertoli cells by secreting testosterone. Therefore, it would be more logical to conclude that MTX down-regulated the endocrine status and negatively affected Sertoli cells’ involvement in intact spermatogenesis and spermiogenesis processes. This could ultimately elevate cellular degeneration. Due to its high antioxidant activity, CMFE improved survival and secretive potential of Leydig cells and indirectly supported Sertoli cells physiologic function. Higher serum level of testosterone in CMFE animals confirmed this hypothesis.

The ALP is known as the main marker for inflammation in testicular tissue. Accordingly, the germinal cells, especially dissociated and atrophied ones, exhibited higher levels of ALP than normal cells. Pathologically-induced inflammation associated with severe cellular damages positively correlates with oxidative stress. Indeed, increased immune cells and inflammation enhance the ROS generation in testicular tissue. In order to understand the effect of MTX on inflammation and revealing the relation between MTX-induced inflammation and the oxidative stress, we performed ALP staining. Our observations demonstrated that MTX enhanced ALP activity, which is an indicator of intensive inflammation, while CMFE decreased inflammation. However, this study showed that the *Cornus mas* as an antioxidant sometimes maybe has a moderately toxic effect by lowering the level of the testis tissue TAC in healthy animals when administered in a high dose.

The CMFE-treated animals exhibited higher levels of TAC in comparison to the MTX group. It is reasonable to suggest that CMFE could also down-regulate the inflammation due to its antioxidant activity and up-regulate cellular survival. Our findings suggested that CMFE could control energy substrates which were mainly carbohydrates, and inhibit the energy switch into UFA. Moreover, CMFE could improve the testicular endocrine status by enhanced testosterone synthesis. CMFE, as an antioxidant compound, could reduce cellular degeneration, lower inflammation and up-regulate testicular antioxidant status.

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Conflict of interest

The authors have no conflicts of interest to declare regarding the study described in this paper and the preparation of the article.

References

1. Peters GJ, Van der Wilt CL, Van Moorsel CJ, et al. Basis for effective combination cancer chemotherapy with antimitabolites. Pharmacol Ther 2000; 87(2-3): 227-253.
2. Choudhury RC, Ghosh SK, Palo AK. Cytogenetic toxicity of methotrexate in mouse bone marrow. Environ Toxicol Pharmacol 2000; 8(3):191-196.
3. Jensen SB, Mournidten HT, Reibel J, et al. Adjuvant chemotherapy in breast cancer patients induces temporary salivary gland hypofunction. Oral Oncol 2008; 44(2):162-173.
4. Chow M, Koo J, Nq P, et al. Random population-wide genetic damage induced in replicating cells treated with methotrexate. Mutat Res 1998; 413(3):251-264.
5. Gisondi P, Girolomoni G. Biologic therapies in psoriasis: a new therapeutic approach. Autoimmu Rev 2007; 6(8):515-519.
6. Padmanabhan S, Tripathi DN, Vikram A, et al. Cytotoxic and genotoxic effects of methotrexate in germ cells of male Swiss mice. Mutat Res 2008; 655(1):59-67.
7. Chiang HM, Fang SH, Wen KC, et al. Life-threatening interaction between the root extract of Pueraria lobata and methotrexate in rats. Toxicol App Pharmacol 2005; 209(3):263-268.
8. Shrestha S, Dhungel S, Saxena AK, et al. Effect of methotrexate (MTX) administration on spermatogenesis: an experimental animal model. Nep Med Coll J 2007; 9: 230-233.
9. Chelab KG, Majeed SK. Histopathological effects of methotrexate on male and female reproductive organs in white mice. Basra J Vet Res 2009; 8(2):120-124.
10. Nouri HS, Azarni Y, Movahedin M. Effect of growth hormone on testicular dysfunction induced by methotrexate in rats. Andrologia 2009; 41(2):105-110.
11. Vardi N, Paralakpınar H, Ates B. Beneficial effects of chlorogenic acid on methotrexate-induced cerebellar Purkinje cell damage in rats. J Chem Neuroanat 2012; 43(1):43-47.
12. Armagan A, Uzar E, Uz E, et al. Caffeic acid phenethyl ester modulates methotrexate-induced oxidative stress in testes of rat. Hum Exp Toxicol 2008; 27(7):547-552.
13. Demir F, Kalyoncu IH. Some nutritional, pomological and physical properties of cornelian cherry (*Cornus mas* L.). J Food Eng 2003; 60(3):335-341.
14. Eyde RH, Quiyun X. Fossil mastixioid (*Cornaceae*) alive in eastern Asia. Am J Bot 1990; 77(5):689-692.
15. Eshaghi M, Zare S, Banighabib N, et al. Cardioprotective effect of *Cornus mas* fruit extract against carbon tetrachloride induced-cardiotoxicity in albino rats. J Basic Appl Sci Res 2012; 2(11):11106-11114.
16. Hassanpour H, Yousef H, Jafar H, et al. Antioxidant capacity and phytochemical properties of cornelian cherry (*Cornus mas* L.) genotypes in Iran. Sci Hort 2011; 129(3):459-463.
17. US Department of Health Education and Welfare. Guide for the care and use of laboratory animals. Department of Health Education and Welfare; 2011; 41-104.
18. Jayaprakasam B, Olson LK, Schutzki RE, et al. Amelioration of obesity and glucose intolerance in high-fat-fed C57BL/6 mice by anthocyanins and ursolic acid in Cornelian cherry (Cornus mas). J Agric Food Chem 2006; 54(1):243-248.
19. Yoshiki S, Umeda T, Kurahashi Y. An effective reactivation of alkaline phosphatase in hard tissues completely calcified for light and electron microscopy. Histochemie 1972; 29(4):296-304.
20. Benzie IF, Strain JJ. Ferric reducing/antioxidant power assay: Direct measure of total antioxidant activity of biological fluids and modified version for simultaneous measurement of total antioxidant power and ascorbic acid concentration. Methods Enzymol 1999; 299: 15-27.
21. Nakamura M, Okinaga S, Arai K. Metabolism of pachytene primary spermatocytes from rat testes: pyruvate maintenance of adenosine triphosphate level. Biol Reprod 1984; 30(5):1187-1197.
22. Razi M, Sadrkhanloo RA, Malekinejad H, et al. Testicular biohistochemical alterations following experimental varicocele in rats. Iran J Reprod Med 2012; 10(3):209-218.
23. Mita M, Hall PF. Metabolism of round spermatids from rats: lactate as the preferred substrate. Biol Reprod 1982; 26(3):445-448.
24. Robinson R, Fritz IB. Metabolism of glucose by Sertoli cells in culture. Biol Reprod 1981; 24(5):1032-1041.
25. Grootegoed JA, Oonk RB, Jansen R, et al. Metabolism of radiolabelled energy-yielding substrates by rat Sertoli cells. J Reprod Fertil. 1986; 77(1):109-118.
26. Nakamura M, Okinaga S, Arai K. Studies of metabolism of round spermatids: glucose as unfavorable substrate. Biol Reprod 1986; 35(4):927-935.
27. Malekinejad H, Mirzakhani N, Razi M, et al. Protective effects of melatonin and Glycyrrhiza glabra extract on ochratoxin A-induced damages on testes in mature rats. Hum Exp Toxicol 2011; 30(2):110-123.
28. Mita M, Price JM, Hall PF. Stimulation by follicle-stimulating hormone of synthesis of lactate by Sertoli cells from rat testis. Endocrinology 1982; 110(5): 1535-1541.
29. Muñoz EM, Fogal T, Dominguez S, et al. Ultrastructural and morphometric study of the Sertoli cell of the viscacha (Lagostomus maximus maximus) during the annual reproductive cycle. Anat Rec 2001; 262(2):176-185.
30. Razi M, Sadrkhanloo RA, Malekinejad H, et al. Varicocele time-dependently affects DNA integrity of sperm cells: evidence for lower in vitro fertilization rate in varicocele-positive rats. Int J Fertil Steril 2011; 5(3):174-185.
31. Akkoyunlu G, Erdoğru T, Seval Y, et al. Immuno-localization of glial cell-derived neurotrophic factor (GDNF) and its receptor GFR-α1 in varicocele-induced rat testis. Acta Histochem 2007; 109(2):130-137.
32. Shan L, Hardy DO, Catterall JF, et al. Effects of luteinizing hormone (LH) and androgen on steady state levels of messenger ribonucleic acid for LH receptors, androgen receptors, and steroidogenic enzymes in rat Leydig cell progenitors in vivo. Endocrinology 1995; 136(4):1686-1693.
33. Razi M, Malekinejad H, Sadrkhanloo RA, et al. Histological impact of long term varicocele-induction on right and left testes in rat (Evidence for the reduction of sperm quality and mating abilities). Vet Res Forum 2011; 2(3):189-201.
34. Savikin K, Zdunic G, Jankovic T, et al. In vitro cytotoxic and antioxidative activity of Cornus mas and Cotinus coggygria. Nat Prod Res 2009; 23(18): 1731-1739.