Cherry laurel fruit extract counters dimethoate-induced reproductive impairment and testicular apoptosis

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Dimethoate is an organophosphorus pesticide used against agricultural insects, which causes oxidative stress and damage in many organs, including the reproductive ones. Cherry laurel (Laurocerasus officinalis Roem.) fruit is rich in vitamins and phenolic compounds with antioxidant effect. The aim of this study was to investigate how effective its extract would be against dimethoate-induced testis and sperm damage in rats. Sixty animals were divided in six groups of 10. Group 1 (control) received only 1 mL of saline (0.9 % NaCl). Group 2 received 7 mg/kg of dimethoate in 1 mL of saline. Group 3 received 4 mg/kg of extract in 1 mL of saline. Group 4 received the extract 30 min before dimethoate administration. Group 5 received vitamin C (positive control, 100 mg/kg in 1 mL of saline) 30 min before dimethoate administration. Group 6 received only dimethoate for the first four weeks and then a combination of dimethoate and extract for another four weeks. All doses were administered daily by oral gavage. After eight weeks of treatment, the rats were euthanised and their reproductive organs removed. We took their body and reproductive organ weights and evaluated testicular oxidative stress, semen characteristics, sperm DNA integrity, apoptosis and histopathological changes. Cherry laurel extract significantly countered many dimethoate-induced adverse effects, both as pre- and post-treatment, including reproductive organ weight, semen parameters, oxidant-antioxidant balance, sperm DNA integrity, testicular apoptosis, and histological structure. Our findings clearly suggest that the beneficial effects of the extract are associated with countering oxidative stress, lipid peroxidation in particular.

KEY WORDS: Laurocerasus officinalis Roem.; organophosphorus pesticide; oxidative stress; sperm DNA damage; testis...
MATERIALS AND METHODS

Chemical agents

Dimethoate (Korumagor 40 EC, 40 % emulsion) was purchased from Koruma Agriculture Co. (Kocaeli, Turkey). Other reagents and materials were purchased from Sigma-Aldrich Chemical Co. (St. Louis, MO, USA), unless otherwise specified.

Plant material and preparation of fruit extract

Cherry laurel fruits were collected from Akçaabat in the Turkish province of Trabzon. A voucher specimen (AEF 26257) has been deposited in the Herbarium of the Ankara University Faculty of Pharmacy, Turkey. We first washed the fruits with distilled water and separated the seed and the pulp. Only pulp was used to prepare CLE by drying it at 40 °C for five days. Twenty grams of dried fruits were macerated with 200 mL of methanol (MeOH) at room temperature with constant magnetic stirring for 8 h, and filtered on Whatman No. 1 filter paper. This procedure was repeated twice with 200 mL of MeOH. The collected filtrates were dried under vacuum using a rotary evaporator at 40 °C. They were then frozen and lyophilised to obtain the extract. Before experiments, the lyophilisate was dissolved in saline (0.9 % NaCl).

Animals and experimental design

For the experiment we used 60 male Wistar rats weighing approximately 200–250 g. The animals were kept in the Experimental and Clinical Research Centre of Erciyes University (Kayseri, Turkey) under standard laboratory conditions (12:12 h light and dark cycle, 22–24 °C temperature, and 55–60 % humidity), and had free access to commercial pellet diet (Optima Food Company, Bolu, Turkey) and water. The experimental procedures were approved by the Ethics Committee for Animal Research at Erciyes University and were carried out in compliance with international standards and national animal welfare legislation.

The animals were randomly divided into six groups of 10 to receive 4 mg/kg bw of CLE and/or 7 mg/kg bw of dimethoate or 100 mg/kg bw of vitamin C by oral gavage every day for eight weeks. The tested dimethoate dose was calculated from the percentage of the active ingredient in the commercial preparation, and the doses of all three substances were based on earlier studies (1, 15). All were freshly dissolved in 1 mL of saline before use in the experiments. We opted for the eight-week treatment to cover the complete spermatogenic cycle in rats, which spans between 49 and 52 days (16).

Group 1 (control) received only 1 mL of saline. Group 2 received dimethoate alone. Group 3 received CLE alone. Group 4 received CLE 30 min before dimethoate administration. Group 5 received vitamin C 30 min before dimethoate administration (positive control), and group 6 received dimethoate alone for the first four weeks, and then a combination of dimethoate and CLE for the other four weeks.

Specimen collection

At the end of treatment, the rats were euthanised with an anaesthetic cocktail containing 8 mg/kg bw xylazine (Rompun® 2 %, Bayer, Istanbul, Turkey) and 75 mg/kg bw ketamine (Ketasol® 10 %, Interhas, Ankara, Turkey). Their whole body and absolute reproductive organ weights (testes, epididymides, seminal vesicle, and ventral prostate) were recorded in grams immediately and used to calculate relative organ weights (organ weight x 100 / whole body weight). One testis of each animal was fixed in Bouin’s solution (a mixture of 15 mL picric acid + 5 mL formaldehyde + 1 mL glacial acetic acid) for 16 h for histopathological analysis. The other testis was stored at -20 °C for oxidative stress analysis.

To determine sperm concentration, the right cauda epididymis was cut at the junction with the corpus epididymis, thoroughly dissected into small pieces with a scalpel and scissors into a Petri dish containing 1 mL saline, and the mixture was poured into an Eppendorf tube and left to incubate at room temperature for 4 h in order for all spermatozoa to pass into fluid. The left cauda epididymis was used to determine sperm motility and abnormality in a 5–10 µL suspension of spermatozoa (17).

Analysis of oxidative stress parameters

Frozen testicular tissues were put into 9-fold phosphate buffered saline (PBS), dissected into small pieces and homogenised in cold saline. We measured the levels of malondialdehyde (MDA) as an indicator of LPO and the activities of superoxide dismutase (SOD), glutathione-peroxidase (GPx), and catalase (CAT) as described earlier by Eken et al. (15) using an ultraviolet-visible spectrophotometer (Shimadzu, Tokyo, Japan) at the following absorbances: 532 nm for MDA, 505 nm for SOD, 340 nm for GPx, and 240 nm for CAT. MDA levels were expressed in nmol/mg protein, and antioxidant enzyme activities in U/mg protein.

Analysis of semen parameters

Semen analyses followed the methods described by Türk et al. (17). Sperm concentration in the right cauda epididymis was determined by a haemocytometer. Motility percentage in the left epididymal spermatozoa diluted with Tris buffer solution was visually determined at 400x magnification with the help of light microscope (Nikon Instruments Inc., Melville, NY, USA) with heated stage at 37 °C. Abnormal spermatozoa were identified under the same microscope with 400x magnification on slides containing a mixture of eosin-nigrosin stain and sperm samples. Abnormalities included head (banana-shaped,
broken-, flat-, double-, and curved-headed sperm) and tail defects (curved-, broken-, short-, and double-tailed sperm). A total of 200 spermatozoa were examined per slide, and the findings expressed as the percentage of abnormal spermatozoa. Total abnormality was also calculated as head + tail.

**Evaluation of sperm DNA damage**

We relied on the method described by Sarıözkan et al. (18). Briefly, diluted sperm samples were centrifuged (CN 090, Nüve, Ankara, Turkey) at 3000 x g at 4 °C for 10 min and supernatant removed. The remaining spermatozoa were washed with PBS without Ca²⁺ and Mg²⁺ to yield a concentration of 1 x 10⁹ spermatozoa per mL. For DNA damage analysis we used the alkaline comet (single-cell gel electrophoresis) assay. Each microscope slide was pre-coated with a layer of 1 % normal melting point (NMP) agarose in PBS and thoroughly dried at room temperature. Next, we mixed 100 µL of 0.7 % low melting point (LMP) agarose in PBS and thoroughly dried at room temperature. They were then removed from the lysis buffer, drained, and solidified in a moist box at 4 °C for 5 min. Cover slips were placed on top of the first slide layer. The slides were allowed to solidify in a moist box at 4 °C for 5 min. Cover slips were removed and the slides immersed in a freshly prepared cold lysis buffer containing 2.5 mol/L NaCl, 100 mmol/L Na₂-EDTA, 10 mmol/L Tris, 1 % Triton X-100, 10 % dimethyl sulphoxide (DMSO), and 40 mmol/L dithiothereitol (pH 10) at 4 °C for 1 h. The slides were then removed from the lysis buffer, drained, and placed in a horizontal electrophoresis unit filled with fresh alkaline electrophoresis solution containing 300 mmol/L NaOH and 1 mmol/L EDTA (pH 13) for 20 min to allow the DNA to unwind. Electrophoresis (12 V, 250 mA) at 8 °C lasted 20 min. The slides were then washed with a neutralising solution of 0.4 mol/L Tris (pH 7.5) to remove alkaline and detergents, stained with 50 µL of 2 µL/mL ethidium bromide, and covered with a cover slip. All steps were performed under a dim light to prevent further DNA damage. For imaging and image analysis of 50 random nuclei per animal we used the Comet Assay Software Project for Windows (CASP-1.2.2, CaspLab, University of Wrocław, Wrocław, Poland).

**Histopathological examination**

Fixed testicular tissues were cut to 5 µm thick slices and stained with Mayer’s haematoxylin and eosin. Lesions like degeneration, congestion, and haemorrhage were examined semi-quantitatively in 20 seminiferous tubules, which were randomly selected in each section under a high-resolution microscope (Olympus BX51, Tokyo, Japan) at 400x magnification. Two independent histologists analysed preparations in a blind fashion and photographed them with the same microscope.

**Assessment of testicular apoptosis**

Testicular apoptosis was determined using the terminal deoxynucleotide-transferase (TdT)-mediated dUTP nick end labelling (TUNEL) assay with *in situ* Cell Death Detection Kit (Roche Diagnostics GmbH, Mannheim, Germany) following the manufacturer’s recommendations (19). Fixed testicular tissues were embedded in paraffin, sliced at 4 µm thickness, deparaffinised in xylene, dehydrated with graded alcohol, washed in PBS, and treated with 0.05 % proteinase K for 5 min. Followed the treatment with 3 % hydrogen peroxide for 5 min to inhibit endogenous peroxidase. After re-washing in PBS, slices were incubated with the TUNEL reaction mixture containing TdT enzyme and digoxigenin-11-dUTP in a humidified chamber at 37 °C for 1 h, and then with a stop/wash buffer for another 30 min. For visualisation we used a diaminobenzidine substrate. Negative controls were treated with distilled water instead of TdT. For evaluation, 20 seminiferous tubules were randomly selected in each slide and TUNEL-positive apoptotic germ cells were counted on images obtained with the Olympus microscope (at 400x magnification) using the Image J program (Image J, Bethesda, MA, USA).

**Statistical analysis**

For statistical analysis we used the SPSS for Windows v. 22.0 (IBM Corp., Armonk, NY, USA). The inter-group comparisons were made with the non-parametric Kruskal-Wallis and non-parametric Mann-Whitney *U* tests. The results are expressed as means ± standard error of mean (SEM). The level of statistical significance was set at *P*<0.05.

**RESULTS**

**Body and reproductive organ weights**

CLE treatment alone significantly increased only seminal vesicle weight compared to control (*P*<0.001). Dimethoate, in turn, significantly decreased whole body, and organ weights (*P*<0.001). CLE pre- and post-treatment or vitamin C significantly countered these effects of dimethoate in the organs, but not in respect to the whole body weight (Tables 1 and 2).

**Oxidative stress parameters**

MDA levels and the activities of SOD, GPx, and CAT are presented in Table 3. Dimethoate significantly increased MDA levels (*P*<0.001) and significantly lowered SOD (*P*<0.01) and GPx (*P*<0.05) activities in comparison to control. These effects were significantly countered by pre- and post-treatment with CLE or vitamin C (*P*<0.001 for MDA, *P*<0.01 for SOD, and *P*<0.05 for GPx). CAT activities did not change significantly between the groups.
Semen parameters

CLE alone significantly increased sperm motility and concentration, while dimethoate significantly reduced sperm motility and concentration and significantly increased head, tail, and total abnormal sperm rate compared to control (P<0.001). Again, these effects were significantly countered by pre-and post-treatment with CLE or vitamin C (P<0.001) (Table 4).

Sperm DNA damage

Dimethoate significantly increased sperm DNA damage versus control (P<0.001). This time only CLE pre-treatment significantly countered dimethoate-induced DNA damage (P<0.001, Table 4).

Table 1 Effects of different treatments on whole body and absolute reproductive organ weights in rats

| Groups                  | Body weight (g) | Absolute reproductive organ weights (g) |
|-------------------------|-----------------|---------------------------------------|
|                         | Testis (Right+left/2) | Whole epididymis (Right+left/2) | Right cauda epididymis | Seminal vesicle | Ventral prostate |
| Control                 | 310.30±6.19a     | 1.339±0.003a                          | 0.516±0.035a          | 0.202±0.001a   | 1.137±0.039a    | 0.544±0.019a    |
| Dimethoate              | 233.75±7.64a     | 0.878±0.012a                          | 0.260±0.010d          | 0.111±0.007c   | 0.761±0.007a    | 0.298±0.003d    |
| CLE                     | 308.00±9.22a     | 1.394±0.021a                          | 0.521±0.002a          | 0.210±0.001a   | 1.245±0.002b    | 0.562±0.025c    |
| CLE _prec_+ dimethoate  | 243.67±4.79b     | 1.151±0.027b                          | 0.416±0.008b          | 0.157±0.001b   | 1.090±0.025a    | 0.402±0.002bc   |
| Vitamin C + dimethoate  | 253.70±2.51b     | 1.209±0.003b                          | 0.437±0.008b          | 0.168±0.002b   | 1.142±0.009b    | 0.442±0.007bc   |
| Dimethoate + CLE _post_ | 235.44±2.27b     | 1.012±0.030b                          | 0.344±0.005c          | 0.139±0.001b   | 0.913±0.010b    | 0.374±0.002bc   |
| Median                  | 254.50           | 1.205                                 | 0.430                 | 0.163          | 1.115           | 0.415           |
| First quartile          | 235.22           | 1.015                                 | 0.343                 | 0.139          | 0.917           | 0.375           |
| Third quartile          | 303.00           | 1.340                                 | 0.520                 | 0.203          | 1.240           | 0.540           |
| Interquartile range     | 67.89            | 0.33                                  | 0.18                  | 0.06           | 0.29            | 0.16            |

Data are expressed as mean ± SEM. Different superscript letters (a, b, c, d) in the same column denote significant differences between the groups (P<0.001). CLE – cherry laurel extract; CLE _prec_ – pre-treatment with CLE; CLE _post_ – post-treatment with CLE; SEM – standard error of the mean

Table 2 Effects of different treatments on relative reproductive organ weights in rats

| Groups                  | Testis (Right+left/2) | Whole epididymis (Right+left/2) | Right cauda epididymis | Seminal vesicle | Ventral prostate |
|-------------------------|-----------------------|----------------------------------|-----------------------|-----------------|------------------|
| Control                 | 0.433±0.009a          | 0.167±0.005ab                    | 0.065±0.001ab         | 0.369±0.017a    | 0.176±0.007a     |
| Dimethoate              | 0.379±0.014d          | 0.113±0.007c                     | 0.047±0.003c          | 0.328±0.012c    | 0.129±0.005b     |
| CLE                     | 0.456±0.013abc        | 0.171±0.006a                     | 0.069±0.002a          | 0.408±0.014ab   | 0.184±0.010a     |
| CLE _prec_+ dimethoate  | 0.474±0.041ab         | 0.171±0.005a                     | 0.064±0.001ab         | 0.448±0.014a    | 0.166±0.003a     |
| Vitamin C + dimethoate  | 0.477±0.016e          | 0.172±0.004d                     | 0.066±0.001ab         | 0.451±0.007a    | 0.175±0.003a     |
| Dimethoate + CLE _post_ | 0.430±0.014c          | 0.146±0.002b                     | 0.060±0.001b          | 0.388±0.004b    | 0.166±0.003a     |
| Median                  | 0.442                 | 0.161                             | 0.063                 | 0.390           | 0.165            |
| First quartile          | 0.417                 | 0.146                             | 0.060                 | 0.375           | 0.156            |
| Third quartile          | 0.469                 | 0.174                             | 0.068                 | 0.443           | 0.183            |
| Interquartile range     | 0.060                 | 0.028                             | 0.008                 | 0.063           | 0.025            |

Data are expressed as mean ± SEM. Different superscript letters (a, b, c, d) in the same column denote significant differences between the groups (P<0.001). CLE – cherry laurel extract; CLE _prec_ – pre-treatment with CLE; CLE _post_ – post-treatment with CLE; SEM – standard error of the mean. * calculated by formula: absolute organ weight (g) x 100 / whole body weight (g)
**Histopathology**

Figure 1 shows the microphotography of histopathological findings in the testes of each experimental group. The control group (Figure 1A) and the CLE alone group (Figure 1C) showed normal testis histology with regular seminiferous tubular morphology and germ cells with normal polarity at various levels of differentiation. The Sertoli and Leydig cells also showed normal structure and polarity. Dimethoate (Figure 1B) deteriorated the histological architecture of the testes, which included abnormal seminiferous tubules with damaged walls, capillary congestion, and haemorrhage in the testis. Compared to the dimethoate-treated group pre- and post-treatment with CLE (Figures 1D and F, respectively) and vitamin C (Figure 2E) resulted in fewer atrophic and degenerative changes in the tubular epithelium. In these experimental groups, the epithelium of the germ cell layer was well coordinated, and the interstitial tissue contained blood vessels with normal structure.

**Apoptosis**

Figure 2 and Table 4 show the presence of apoptotic cells in the testes of each group. Apoptotic germ cell count in the dimethoate-treated group was significantly higher than in all other groups (P<0.001). Pre- and post-treatment with CLE or vitamin C significantly reduced this count (P<0.001).
DISCUSSION

Our findings confirmed the adverse effects of subchronic dimethoate exposure on the reproductive system in male rats. In line with other reports (20–23), it significantly decreased body and reproductive organ weights. These losses are likely to be associated with lower food intake (22) and lower testosterone (23) and thyroid (21) secretion. Although we did not measure hormone levels to demonstrate this association, it is well-known that thyroid hormones (T₃ and T₄) are necessary for the growth (21), while testosterone governs the development and function of male reproductive organs (24). Both are affected by oxidative stress evidenced in our study.

Our dimethoate-induced oxidative stress findings are also in line with previous reports, indicating overproduction of free radicals or reactive oxygen species (ROS) that react with membrane lipids, producing cell and tissue damages (20, 25, 26). Dimethoate administration has been reported to stimulate oxidative stress in rats (20), as evidenced by increased LPO and decreased tissue activities of SOD, CAT and GPx (27). In our study, MDA levels significantly increased, while SOD and GPx activities significantly decreased after dimethoate administration. The same is true for semen parameter findings, which support earlier findings for organophosphorus pesticides such as acephate (27), diehylthiophosphate (28), and cypermethrin (29). An earlier study (23) reported some histopathological lesions, such as oedema, haemorrhage, degenerated seminiferous tubules, atrophy, sloughing, degeneration and spermatogenic arrest in testes after dimethoate treatment. As the reason for the oxidant/antioxidant imbalance in testes the authors of that study suggested overproduction of free radicals and ROS in response to dimethoate administration. Besides, deteriorations in semen parameters, sperm DNA and testicular histology are likely to be associated with dimethoate-induced increase in LPO.

Cherry laurel has strong radical (hydroxyl and superoxide) scavenging and antioxidant activities because it contains high levels of pectin, phenolic compounds, vitamins (A, C, D), and minerals (10–14). Since its antioxidant effect is similar to or higher than vitamin C (11), vitamin C was used as reference antioxidant in this study. Eken et al. (15) have reported that CLE significantly counters dimethoate effects in rats by lowering increased liver MDA levels, pathological lesions, and DNA damage and by significantly increasing low SOD, GPx, and CAT activities in the liver. We too observed that CLE showed a significant antioxidative effect in comparison with dimethoate. Our dimethoate-induced oxidative stress findings are in line with in vivo CLE study by Yılmaz (30), which demonstrated improvements in total sperm count, vitality and motility and mitigation of oxidative stress and histopathological lesions against the side effects of a chemotherapy drug methotrexate.

To date, no other study investigated CLE’s potential against dimethoate-induced reproductive damage, and our study provides significant new evidence on this issue. Pre-
Table 4. Effects of different treatments on semen parameters, DNA damage, and testicular apoptosis in rats

| Groups                        | Semen parameters (million/ right cauda epididymis) | Sperm DNA damage (%) | Testicular apoptosis (%) |
|-------------------------------|-----------------------------------------------------|----------------------|-------------------------|
|                               | Motility (%) | Abnormal sperm rate (%) | Head | Tail | Total | Head | Tail | Total |
| Control                        | 74.3±1.09b   | 3.60±0.43d             | 6.10±0.43c           | 20.13±0.72c            | 113±40.18c | 8.47±0.10b |
| Dimethoate                     | 31.25±2.52e  | 46.25±4.22b            | 22.63±0.42c          | 42.76±0.96a            | 10.12±0.14a | 23.3±0.40a |
| CLE + dimethoate               | 83.17±1.28a  | 121.40±1.08c           | 7.67±0.44a           | 16.45±0.40c            | 10.82±0.16b | 6.07±0.42a |
| Vitamin C + dimethoate         | 54.44±1.39c  | 56.67±1.31c            | 9.22±0.22c           | 17.55±0.90c            | 11.33±0.14 | 2.33±0.40c |
| Dimethoate + CLEpost           | 50.19±1.07a  | 71.11±1.14c            | 8.89±0.43c           | 20.22±0.63c            | 12.33±0.10c | 2.77±0.36 |
| Median                         | 55.00       | 76.28                  | 4.00                 | 10.00                 | 10.00       | 0.00 |
| First quartile                 | 75.84       | 112.00                 | 11.00                | 11.00                 | 10.00       | 1.00 |
| Third quartile                 | 26.25       | 35.72                  | 7.00                 | 4.00                  | 10.16       | 6.00 |

Data are expressed as mean ± SEM. Different superscript letters (a, b, c, d) in the same column denote significant differences between the groups (P<0.001). CLE – cherry laurel extract; CLE pre – pre-treatment with CLE; CLEpost – post-treatment with CLE; SEM – standard error of the mean.

The improvement in oxidant/antioxidant balance, semen parameters, sperm DNA integrity, testicular apoptosis, and histology observed in our study are possibly associated with the strong radical scavenging and antioxidant activity of CLE. Sperm plasma membrane is rich in polyunsaturated fatty acids, which are particularly sensitive to free radical attacks (25). The lowering of LPO in sperm plasma membrane by antioxidative action of CLE seems to entail improvements in semen parameters, lowering of DNA damage and apoptosis, and restoration of normal testicular histology. Besides, improved histological structure after CLE treatment seems to explain increase in testicular weight. The increase in epididymis weight may be associated with increased sperm count, and the increase in accessory sex gland weights with increased fluid content.

CONCLUSION

Our findings in the rat model provide evidence that the protective effects of CLE against dimethoate-induced reproductive damage in rats are similar to those of vitamin C and suggest that consumption of CLE by humans and animals exposed to organophosphorus pesticides like dimethoate may improve protection of male reproduction. Before we jump into any conclusion about human risk assessment, however, we should consider the metabolic differences between rat and human metabolism. In addition, this study tested the effects of a single pesticide and CLE concentration, and further in vivo research should involve a broader dose range.

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Conflict of interest

None to declare.

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Figure 2 Apoptotic germ cells (arrows) in testicular tissues (H&E, 400x). (A) the control group; (B) in the dimethoate alone group, many TUNEL-positive apoptotic cells were found; (C) cherry laurel extract (CLE) alone group; (D) CLE pre-treatment decreased the number of TUNEL-positive apoptotic cells in rats receiving dimethoate; (E) vitamin C-treated group receiving dimethoate; (F) CLE post-treatment group receiving dimethoate.
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Ekstrakt lovorvišnje ublažava štetno djelovanje dimetoata na reproduktivnu funkciju i apoptozu stanica u testisu štakora

Dimetoat je organofosforni insekticid koji uzrokuje oksidacijski stres i oštećuje mnoge organe, uključujući reproduktivne. Plod lovorvišnje (Laurocerasus officinalis Roem.) bogat je vitaminima i fenolnim spojevima s antioksidacijskim djelovanjem. Cilj je ovoga istraživanja bio utvrditi djelotvornost njegova ekstrakta protiv oštećenja testisa i spermija u mužjaka štakora izloženih dimetoatu. Njih 60 podijeljeno je nasumce u šest skupina od deset životinja. Prva je (kontrolna) skupina primala samo 1 mL fiziološke otopine (0,9 % NaCl), druga skupina 7 mg/kg dimetoata u 1 mL fiziološke otopine, treća skupina 4 mg/kg ekstrakta lovorvišnje u 1 mL fiziološke otopine, četvrta skupina ekstrakt 30 minuta prije primjene dimetoata, peta skupina vitamin C (pozitivna kontrola, 100 mg/kg u 1 mL fiziološke otopine) 30 min prije primjene dimetoata, a šesta skupina primala je dimetoat u prva četiri tjedna, potom kombinaciju dimetoata i ekstrakta lovorvišnje u sljedeća četiri tjedna. Sve navedene doze davane su svaki dan gavažom. Nakon osam tjedana primjene, štakori su eutanazirani, a njihovi reproduktivni organi odstranjeni. Izmjerena im je ukupna tjelesna masa i masa reproduktivnih organa te ocijenjena oksidacijski stres u testisima, značajno smanjio masu tijela i organa, gibljivost spermija, aktivnost superoksid dismutaze i glutatijon-peroksidaze, a značajno smanjio lipidnu peroksidaciju. Dimetoat je značajno smanjio masu tijela i organa, gibljivost i koncentraciju spermija te aktivnost superoksid dismutaze i glutacion-peroksidaze, a značajno smanjio lipidnu peroksidaciju. Oštećenja DNA spermija, apoptozu u testisima te uzrokovalo histopatološke promjene tkiva. Ekstrakt lovorvišnje značajno ublažio te štetne učinke, bilo da je davan prije ili nakon dimetoata. Naši rezultati jasno upućuju na blagotvorno djelovanje toga ekstrakta, koje je povezano sa zaštitom od oksidacijskoga stresa, napose od lipidne peroksidacije.

KLJUČNE RIJEČI: Laurocerasus officinalis Roem.; oksidacijski stres; organofosforni pesticidi; oštećenje DNA spermija; testis