Novel Feed Including Olive Oil Mill Wastewater Bioactive Compounds Enhanced the Redox Status of Lambs

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Abstract. Background/Aim: The aim of the present study was to investigate the antioxidant effects of a feed supplemented with polyphenolic additives from olive mill wastewater (OMW) on lambs. Materials and Methods: Lambs received breast milk until the postnatal period, and then they were divided into two groups and received control and OMW feed for 55 days. Redox biomarkers were measured in blood and tissues at days 15, 42 and 70 after feeding. Results: Feed supplemented with OMW reduced thiobarbituric acid reactive species and protein carbonyls and increased total antioxidant capacity, glutathione and catalase activity in both blood and tissues. Conclusion: The administration of OMW-containing feed reinforced the antioxidant defense of lambs, which may improve their wellbeing and productivity. Additionally, this exploitation of OMW may solve problems of environmental pollution in areas with olive oil industries.

Mediterranean countries are responsible for 95% of worldwide olive oil production (1). In Greece, the extraction and manufacture of olive oil is carried out in about 2,400 small scale agro-industrial units scattered throughout the country. These processes produce two waste streams: olive mill residual solids and olive mill wastewaters (OMW). OMW are formed from the water content of the fruit itself and the water used to wash and process them, and amounts to 0.5-3.25 m³ per 1,000 kg of olives (2). Typically, the composition of OMW is water, organics and mineral salts. OMW comprises large amounts of organic (e.g. fats, lipids, polyphenols), inorganic constituents and water; a part of the organic fraction is composed of phenols (1). These compounds are responsible for its black color, toxicity, phytotoxicity and antibacterial properties (3).

It is estimated that around 30 million cubic meters of OMW are generated annually in the Mediterranean area (4). Disposal of OMW causes serious environmental problems, such as soil contamination, water body pollution, underground seepage and odor (5). The environmental problems and potential hazards caused by OMW have prompted many countries to limit its discharge. Varied new technologies for reducing its pollutant power (e.g. physicochemical and biological methods), have been investigated in recent years (1). Technologies such as advanced oxidative processes have been proposed, but these are characterized by high operational costs and frequently require complex maintenance and availability of experienced personnel (6). Thus, an environmentally safe and cost-effective treatment of OMW has not yet been found (7).

Oleuropein, tyrosol and hydroxytyrosol are the main phenolic compounds found in OMW. Other phenolic compounds that are found in olive oil are caffeic acid, vanillic acid, coumaric acid, ferulic acid, gallic acid, hydroxybenzoic acid, kaempherol, apigenin and quercetin (8, 9). Despite toxic effects exhibited at high concentrations, these polyphenols also exhibit antioxidant activity. Therefore, the polyphenolic content of OMW could represent a source of antioxidants in different fields, such as nutrition,
pharmaceutical and cosmetic industries, while reducing the environmental pollution caused by OMW (10).

In recent years, there is evidence that bioactive compounds present in foods possess properties which give them the ability to protect against chronic diseases (11-13). Studies have shown that in farm animals, oxidative stress is involved in various pathological situations, directly related to animal welfare (14). For example, common diseases such as pneumonia (15), enteritis (16) and septicemia in pigs (17) are caused by disorders in antioxidant homeostasis. The administration of antioxidants can protect animals from these diseases or reduce their clinical symptoms (18). In addition, studies have shown that animals at an early age have a low antioxidant system compared to adults (19, 20), and so it is more important to administer antioxidants to young animals for disease protection.

Polyphenols contribute significantly to antioxidant defense of animals by increasing the antioxidant levels of endogenous molecules and enzymes, while enhancing the immune system (21). In particular, polyphenols from OMW have antioxidant properties which are related to prevention of diseases caused by oxidative stress (22-25). According to Paiva-Martins et al. (26), polyphenols of olive oil protected human erythrocytes from oxidative damage. In particular, 3-hydroxytyrosol, an olive oil polyphenol, plays a remarkable role against reactive oxygen species (ROS) that cause cellular damage. Studies in animals and humans have shown that polyphenol components of OMW exhibit important biological activities that can prevent oxidative stress-associated diseases (13, 27). Recent studies of our research group showed broiler chickens and piglets given feed supplemented with OMW exhibited an increased antioxidant capacity in blood and tissues (10, 28).

Thus, the aim of present study was to exploit OMW byproducts to produce a biofunctional feed of high added value for lambs in order to enhance the animal’s antioxidant defense, welfare and productivity. Moreover, exploitation of OMW to develop biofunctional feeds may reduce associated environmental problems.

**Materials and Methods**

*Olive mill wastewater.* Olive mill wastewater (OMW) was obtained directly from a local olive oil mill in Larissa prefecture (Greece), without any further processing. OMW was characterized by high organic load (chemical oxygen demand: 45-100 g/l, biological oxygen demand: 25-50 g/l), pH: 4.6-5.2, total solids (39.4±1.8 g/l), water content: 960.6±19 g/l, significant concentrations of magnesium, potassium, phosphate salts and organic compounds (29, 30).

*Silage preparation.* OMW was added to lamb feed as silage. The silage contained corn, OMW, water and lactic acid bacteria. Based on previous studies, the proportion of the ingredients was such that the silage contained 60% solids and 40% liquids (10, 28). Standard commercial formulation (11CFT; Pioneer, Buxtehude, Germany) of lactic bacteria was used for the lactic fermentation of corn and the preparation of corn silage. The lactic bacteria had been dissolved in water (10% w/v) by stirring and warmed at 40°C in order to be activated prior to mixing with corn. After activation, lactic bacteria were mixed with corn (1 g of bacteria with 100 kg of corn). For producing the silage, the mixture of lactic bacteria and corn was placed in special airtight-seal plastic bags and was fermented for 3-4 weeks. To prevent the bags from rupturing due to the inflation caused by the carbon dioxide production during fermentation, the material was repackaged in new plastic bags every 2 to 3 days. Finally, the resulting silage, containing 52.5% solids, 7.5% OMW and 40% water, was mixed with other ingredients to make the final lamb feed. The final feed composition before and after weaning is presented in Table I. Milk that was contained in diets before weaning was replaced by wheat and sunflower meal in the diets after weaning (Table I).

**Animals and diets.** The breeding of lambs held at the Research Institute of Animal Science (Paralimni Giannitson, Greece/Hellenic Agricultural Organization – Demeter) by applying diet in normal living and development conditions. Both living conditions and the way that the lambs were sacrificed for blood and tissue collections were performed according to EU Directive 2010/63/EU for animal experiments.

The experiment was reviewed and approved by the Institutional Review Board of the University of Thessaly (no. 89/10.12.2014). Twenty-eight young male lambs of Chios breed were selected from the flock of the Animal Research Institute. When the feeding started at 15 days after birth, the lambs weighed on average 7.99±1.80 kg and were divided into two homogeneous groups (12 lambs per group) as follows: (a) Lambs fed with standard ration (control group) and (b) lambs fed with ration containing silage with OMW (OMW group) for 55 days (i.e. from 15 to 70 days) (Table I). During the age of weaning (i.e. from 15 to 42 days), lambs remained along with their ewes in two separate stalls (one for each group) for breastfeeding and also had access to feed, either to the standard or to experimental feed, alfalfa hay and water for *ad libitum* consumption. The ewes were fed with standard feed without access to the ration supplemented with OMW and they were separated from the lambs at day 42 (Table I). During the experimental trial, lambs were weighed individually weekly and average daily gain (ADG, g/day) was calculated.

**Determination of total polyphenolic content (TPC) of feed rations.** The TPC of control ration and ration supplemented with OMW was determined using Folin-Ciocalteu reagent, as described previously (31). The two rations were originally in a solid state and were liquefied as described previously (32). Briefly, 20 μl of sample was added to a tube containing 1 ml of deionized water (dH2O). The total of 100 μl Folin-Ciocalteu reagent was added to the reaction mixture, followed by incubation for 3 min at room temperature. Subsequently, 280 μl of 25% w/v sodium carbonate solution and 600 μl of dH2O were added to the mixture. Following 1 h of incubation at room temperature in the dark, the absorbance was measured at 765 nm against a blank containing Folin-Ciocalteu reagent and dH2O without the sample. The measurement of absorbance was conducted on a Hitachi U-1900 radio beam spectrophotometer (Hitachi, Tokyo, Japan). The optical density of the sample (20 μl) in 25% w/v solution of sodium carbonate (280 μl) and dH2O (1.7 ml) at 765 nm was also
measured. The results are expressed as gallic acid equivalents using a standard curve prepared from authentic gallic acid (Sigma-Aldrich, Munich, Germany).

**Determination of antioxidant activity of feed rations: 2,2-Diphenyl-1-picrylhydrazyl (DPPH) radical-scavenging assay:** The free-radical scavenging activity (RSC) of both control ration and ration supplemented with OMW was evaluated by DPPH radical assay (33). Briefly, 1.0 ml of freshly prepared methanolic solution of DPPH radical (100 μM) was mixed with the tested samples at different concentrations. The contents were vigorously mixed, incubated at room temperature in the dark for 20 min and the absorbance was recorded at 517 nm. The measurement was conducted on a Hitachi U-1900 ratio beam spectrophotometer (Hitachi). In each analysis, the sample extract alone containing 1 mM of ABTS and 30 μM of H$_2$O$_2$ in 50 mM (PBS, pH 7.5) was used as a blank, while the ABTS** radical solution alone with 10 μl water was used as a control. The percentage inhibition and the IC$_{50}$ values were determined as described above for the DPPH method. All analyses were carried out in triplicate and on at least two separate occasions.

The percentage RSC of the tested extracts was calculated using the following equation: \[ \text{RSC} (%) = \left( \frac{A_{\text{control}} - A_{\text{sample}}}{A_{\text{control}}} \right) \times 100 \]

where \( A_{\text{control}} \) and \( A_{\text{sample}} \) were the absorbance values of the control and the tested samples, respectively. Moreover, in order to compare the radical-scavenging efficiency of the feeds, the IC$_{50}$ value, i.e., the concentration leading to 50% scavenging of the DPPH radical was calculated from the graph plotted of percentage RSC against the extract concentration. All experiments were carried out in triplicate and at least in two separate occasions.

**2,2'-Azinobis-(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS)**

**radical-scavenging assay:** ABTS** radical-scavenging activity was measured as described previously with minor modifications (34). In brief, ABTS** radicals were produced by mixing 2 mM of ABTS with 30 μM of H$_2$O$_2$ and 6 μM of horseradish peroxidase (HRP) (Sigma-Aldrich) in 50 mM phosphate-buffered saline (PBS, pH 7.5). Immediately following the addition of HRP, the contents were vigorously mixed, incubated at room temperature in the dark and the reaction was monitored at 730 nm until stable absorbance was obtained.

**Blood and tissue collection.** Blood and tissues from 28 lambs were collected at three different time-points, at 15, 42 and 70 days after birth. At day 15, blood samples were collected from four lambs, in order to determine the redox status of lambs at a very young age before the administration of ration, at day 42 from 12 lambs (i.e. six lambs from each group) and at day 70 from 12 lambs (i.e. six lambs from each group). For blood collection, 4 ml of blood were collected from the jugular vein and placed in vacutainer tubes with ethylenediamine tetraacetic acid (EDTA). The isolation of plasma and red blood cell lysate (RBCL) and the determination of hemoglobin concentration were performed as described previously (21).

At day 15, tissues were collected from four lambs in total (lambs were divided into two homogeneous groups at 15 days after birth when the feeding started), in order to determine their redox status at an early age before the feed administration. At day 42, tissues were collected from 12 lambs (i.e. six lambs from each group) and finally, at day 70 from 12 lambs (i.e. six lambs from control and six lambs from the OMW group). For tissue collection, lambs were transported to a slaughterhouse and were immediately stunned prior to slaughtering in order to minimize suffering. All relevant procedures were executed by specialized staff according to industry-accepted procedures. Five tissues from vital organs, namely the heart, liver, brain, spleen and quadriceps muscle, were removed as quickly as possible, placed into tubes and snap-frozen in liquid nitrogen. In preparation for biochemical analysis, tissues samples were initially ground using a mortar and pestle under liquid nitrogen. The homogenization process was carried out as described previously (21). Plasma, RBCL and tissues were stored at −80°C until biochemical analysis.

**Determination of oxidative stress biomarkers.** Reduced glutathione (GSH) was measured according to the method of Reddy et al. (35) In this assay, proteins in the erythrocyte lysate were precipitated twice with 5% trichloroacetic acid (TCA) in order to eliminate protein-linked -SH groups. Briefly, 20 μl of erythrocyte lysate or

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**Table I. Ingredients and nutrient composition (% w/w) of experimental and control diets for lambs before and after weaning.**

| Ingredients                      | Before weaning | After weaning |
|----------------------------------|----------------|---------------|
|                                  | Control group  | OMW group     | Control group  | OMW group     |
| Corn silage*                     | 45.0           | 45.0          | 45.0           | 45.0          |
| Wheat bran                       | 9.0            | 9.0           | 15.0           | 15.0          |
| Wheat meal                       | 0              | 0             | 13.0           | 13.0          |
| Soybean meal 44% CP              | 21.0           | 21.0          | 18.0           | 18.0          |
| Milk replacer                    | 20.0           | 20.0          | 0              | 0             |
| Vitamin and mineral premix (2.5%)| 2.5            | 2.5           | 2.5            | 2.5           |
| Sunflower meal                   | 0              | 0             | 4.0            | 4.0           |
| Salt                             | 0.5            | 0.5           | 0.5            | 0.5           |
| Limestone                        | 1.2            | 1.2           | 1.2            | 1.2           |
| Monocalcium phosphate            | 0.8            | 0.8           | 0.8            | 0.8           |

*Silage contained 60% corn solids and 40% water in control feed, and 52.5% corn solids, 7.5% olive oil mill wastewater (OMW) and 40% water in the OMW group.

**Table II. Total polyphenolic content (TPC) of the two experimental feed types before and after lamb weaning. TPC is expressed as gallic acid equivalents, as measured by Folin-Ciocalteau assay. The results are presented as the means±SEM.**

| TPC                              | Control group | OMW group |
|----------------------------------|---------------|-----------|
| Before weaning                   | 0.170±0.023   | 0.348±0.019*|
| After weaning                    | 0.138±0.028   | 0.272±0.028*|

*Statistically significant different compared to the control (p<0.05).
tissue homogenate (diluted 1:2) treated with 5% TCA was mixed with 660 μl of 67 mmol/l sodium potassium phosphate (pH 8.0) and 330 μl of 1 mmol/l 5,5′-dithiobis-2 nitrobenzoate. Samples were incubated in the dark at room temperature for 10 min and the absorbance was read at 412 nm. The GSH concentration was calculated on the basis of calibration curve made using commercial standards (Sigma-Aldrich, Munich, Germany).

Catalase activity was determined in erythrocyte lysate using the method of Aebi (36). Briefly, 4 μl of erythrocyte lysate (diluted 1:10) or 40 μl of tissue homogenate (diluted 1:2) were added to 2991 or 2955 μl, respectively, of 67 mmol/l sodium potassium phosphate (pH 7.4) and samples were incubated at 37˚C for 10 min. A total of 5 μl of 30% hydrogen peroxide (H2O2) were added to the samples, and the change in absorbance was immediately read at 240 nm for 2 minutes. Calculation of catalase activity was based on the molar extinction coefficient of H2O2 (43.6 M−1 cm−1).

Protein carbonyl determination was based on the method of Patsoukis et al. (37) In this assay, 50 μl of 20% TCA was added to 50 μl of plasma or tissues homogenate (diluted 1:2), and this mixture was incubated in an ice-bath for 15 min and then centrifuged (15,000 × g, 5 min, 4˚C). The supernatant was discarded, and 500 μl of 10 mmol/l 2,4-dinitrophenylhydrazine (in 2.5 N HCl) per sample (500 μl of 2.5 N HCl for the blank) was added to the pellet. The samples were incubated in the dark at room temperature for 1 h with intermittent vortexing every 15 min and were centrifuged (at 15,000 × g for 5 min at 4˚C). Proteins were then precipitated with 10% TCA and washed three times with ethanol-ethyl acetate (1:1 v/v). The supernatant was discarded, and 1 ml of 5 mol/L urea (pH 2.3) was added, vortexed and incubated at 37˚C for 15 min. The samples were centrifuged (at 15,000 × g for 5 min at 4˚C) and the absorbance was read at 375 nm. Calculation of protein carbonyl concentration was based on the molar extinction coefficient of 2,4-dinitrophenylhydrazine (22×103 M−1 cm−1). Total protein was assayed using Bradford reagent (Sigma-Aldrich, Munich, Germany).

Thiobarbituric acid-reactive substances (TBARS) assay was used for the determination of lipid peroxidation. TBARS were determined according to a slightly modified assay of Keles et al. (38). In detail, 100 μl of plasma or 50 μl of tissue homogenate (diluted 1:2) was mixed with 500 μl of 35% TCA and 500 μl of Tris–HCl (200 mmol/l; pH 7.4), and incubated for 10 min at room temperature. One milliliter of 2 mol/l Na2SO4 and 55 mmol/l thiobarbituric acid solution was added, and the samples were incubated at 95˚C for 45 min. The samples were cooled on ice for 5 min, and were vortexed after 1 ml of 70% TCA was added. The samples were centrifuged at 15000 × g for 3 min, and the absorbance of the supernatant was read at 530 nm. A baseline shift in absorbance was taken into account by running a blank along with all samples during the measurement. TBARS are expressed in terms of malondialdehyde (MDA) equivalents. The molar coefficient of MDA is 155×103 mol/l.

Determination of total antioxidant capacity (TAC) was based on the method of Janaszewska and Bartosz, (39). Briefly, 20 μl of plasma or 40 μl of tissue homogenate (diluted 1:10) were added to 480 μl or 460 μl, respectively, of 10 mmol/l sodium potassium phosphate (pH 7.4) and 550 μl of 0.1 mmol/l DPPH free radical and samples were incubated in the dark for 60 min at room temperature. Samples were centrifuged for 3 min at 20000 × g, and the absorbance was read at 520 nm. TAC is presented as mmol of DPPH reduced to 2,2′-diphenyl-1-picrylhydrazine (DPPH:H) by antioxidants of plasma and tissue.

Each assay was performed in triplicate and within 3 months of blood and tissue collection. Samples were stored in multiple aliquots at −80˚C, and thawed only once before analysis. All reagents were purchased from Sigma-Aldrich (Munich, Germany). All measurements were conducted on a Hitachi U-1900 ratio beam spectrophotometer (serial no. 2023-029; Hitachi, Tokyo, Japan).

Statistical analysis. Data were analyzed by one-way ANOVA. The level of statistical significance was set at \( p<0.05 \). All results are expressed as the mean±SEM. Data were analyzed using SPSS, version 13.0 (SPSS Inc., Chicago, IL, USA).

Results

Assessment of TPC of experimental feed. The results from the TPC assay indicated TPC was at least 2-fold higher in the feed supplemented with OMW, both before and after weaning, compared to the control group because of the inclusion of OMW in the feed (Table II).

Figure 1. 2,2-Diphenyl-1-picrylhydrazyl (DPPH•) and 2,2′-Azinobis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS•+) radical-scavenging capacity (A) before and (B) after weaning of standard (control) and experimental (OMW) feed. *Significantly different from the control value (\( p<0.05 \)).
Assessment of antioxidant activity of experimental feed. The antioxidant activity of each feed was evaluated according to two radical-scavenging assays, DPPH and ABTS (Figure 1). The results from both assays showed higher antioxidant activity of the feeds supplemented with OMW, both before and after weaning, by at least two-fold, compared to the control feed.

Animal growth performance. Feed supplemented with OMW significantly improved the ADG (Figure 2). Prior to the weaning period (i.e. days 15-42), ADG was increased in the OMW group by 70.2% compared to the control group. In the post-weaning period (i.e. days 42-70), no significant difference was observed in the ADG between the two groups. Overall (i.e. days 15-70), lamb body weight did not differ significantly between OMW and control groups (Figure 2).

Assessment of oxidative stress markers in lamb blood. All oxidative stress biomarkers measured in blood (both plasma and erythrocyte lysate), except catalase activity, indicated that feed enriched with OMW improved the redox status of lambs. Specifically, the GSH level in red blood cell lysate was significantly increased both at days 42 and 70 post-birth by 106.9 and 155.9% in the OMW group compared to the control group (Figure 3A).

Catalase activity in erythrocyte lysate did not differ significantly between the two groups at both these time points (Figure 3B).

Regarding TAC in plasma of lambs, it was significantly higher in the OMW group compared to the control group only at day 42 post-birth (by 10.1%; Figure 4A).

The level of TBARS in plasma was significantly lower in the OMW group compared to the control group by 30.6% at day 42, but there was no significant difference at day 70 between the two groups (Figure 4B).

Finally, there was a significant decrease in protein carbonyl level by 40.3% at day 70 in the OMW group compared to the control group (Figure 4C).

Assessment of oxidative stress markers in lamb tissues. The feed supplemented with OMW enhanced the antioxidant levels and improved the redox status in lambs in almost all the tested tissues. In particular, the administration of feed supplemented with OMW to young lambs significantly increased the GSH level in spleen and brain at day 42 by 110.0 and 36.9%, respectively, compared to the control group (Figure 5A) and by 19.0, 101.5, 40.9, and 12.8% in heart, spleen, quadriceps muscle and brain, respectively, at day 70 (Figure 5A).

The rate of $\text{H}_2\text{O}_2$ decomposition was significantly increased only in heart at day 42 by 17.3%, while at day 70, there was a significant increase in heart, liver and quadriceps muscle by 53.1, 7.9, and 116.7% respectively in the OMW group compared to the control group (Figure 5B).

TAC was significantly increased only in brain at both day 42 and 70, by 56.1 and 32.6% respectively (Figure 6A).

The level of TBARS was significantly lower in all tissues except brain at day 42 in the OMW group by 10.3, 35.2, 55.2 and 11.9% in heart, spleen, liver and quadriceps muscle, respectively, than in the control group (Figure 6B). At day 70 post birth TBARS was decreased only in brain tissue, by
52.0% in the OMW group compared to the control group (Figure 6B).

Finally, protein carbonyl levels were significantly decreased in heart, spleen and liver by 45.2, 34.3 and 50.1% respectively, at day 42, while at day 70, protein oxidation was lower by 20.3% only in quadriceps muscle in the OMW group compared to the controls (Figure 6C).

**Discussion**

OMW is a liquid of violet to dark brown color with a strong smell of olive oil (40). This byproduct is composed of vegetable water from the fruit and the water used in different stages of oil extraction that contains olive pulp, mucilage, pectin, oil, and other suspended components in a relatively stable emulsion (41). The composition and amounts of OMW cause serious environmental problems in areas of olive oil production, and the discharge of large quantities of these pollutants into the sewage system without any treatment is not possible. Different biological and chemical/physical methods have been proposed to reduce the organic matter and tannins present in OMW in order to reduce their effects on the environment. On the other hand,
phenolic extracts from OMW can be used as natural alternatives to commercial synthetic antioxidants with applications in food, as well as in the development of nutraceutical and medical products (8).

The aim of this study was to evaluate the effects of feed supplemented with OMW on the redox status of young lambs and to suggest a new method in order to reduce environmental pollution caused by the high organic matter of byproducts which are produced by olive oil industries. Both in vitro free radical-scavenging assays (i.e. DPPH and ABTS) showed that the feed supplemented with OMW had about 2-fold higher antioxidant activity than that of the control. This strong antioxidant activity of OMW byproducts may mainly be based on their high polyphenolic content (e.g. tyrosol, hydroxytyrosol, oleuropein, caffeic acid, vanillic acid, coumaric acid, ferulic acid, gallic acid, hydroxybenzoic acid, kaempherol, apigenin and quercetin) (8).

The findings showed that the feed supplemented with OMW increased animals’ growth performance in the OMW group compared to controls, especially at a young age. According to Payne and Southern (42), ROS oxidize and damage cellular biological molecules, as a result causing a

Figure 5. Effects on oxidative stress markers glutathione (GSH) (A) and catalase (B) in lamb tissues (heart, spleen, liver, quadriceps and brain) after feeding with standard (control) and experimental (olive oil mill wastewater, OMW) feed at 15 (n=4), 42 (n=6 from each group) and 70 (n=6 from each group) day post birth. *Significantly different from the control group at the same sampling time (p<0.05).
Figure 6. Effects on oxidative stress markers TAC (A), TBARS (B) and protein carbonyl levels (C) in lamb tissues (heart, spleen, liver, quadriceps and brain) after feeding with standard (control) and experimental (olive oil mill wastewater, OMW) feed at 15 (n=4), 42 (n=6 from each group) and 70 (n=6 from each group) day post-birth. *Significantly different from the control group at the same sampling time (p<0.05).
variety of impairments to intestinal membrane integrity. Thus, the improvement in ADG and growth performance of lambs may be due to the experimental feed’s antioxidant properties that resulted in ROS scavenging, reducing intestinal membrane damage, and consequently improving gut functionality.

Regarding the effects of the feed supplemented with OMW on the redox status of the lambs, the results showed that the GSH level was significantly increased in spleen and brain tissues at day 42 post birth, in heart, spleen, quadriceps muscle and brain tissues at day 70 post birth and in erythrocytes at both sampling times in the OMW group compared to control group. Similarly, another study of our research group, showed that the administration of feed supplemented with polyphenols from OMW to piglets and broilers increased the GSH level in erythrocytes and different tissues (10, 28). Another study conducted on humans also showed that consumption of OMW extract increased plasma GSH (43). These results are of particular importance, since GSH, the most abundant non-protein thiol source in cells, is one of the major intracellular antioxidant molecules protecting against oxidative stress. The effect of feed supplemented with OMW on the GSH level may be explained through the modulation of the main enzymes responsible for GSH synthesis, such as GSH synthase and gamma-glutamylcysteine synthase (γ-GCL) (44). The regulation of the expression of the enzymes involved in GSH synthesis or metabolism is mainly mediated through the antioxidant response element, a cis-enhancer sequence regulating the transcription of various antioxidant genes (45). Interestingly, in another study, we showed that administration of feed supplemented with OMW to lambs increased the activity or expression of antioxidant enzymes regulated by the antioxidant response element (46). In addition, other studies have reported that the OMW-induced increase in GSH level might be due to antioxidant response element-mediated increase in γ-GCL and GSH synthetase expression (43, 46).

The conversion of H$_2$O$_2$ to H$_2$O and O$_2$ in erythrocytes is mainly attributed to catalase activity (47, 48). The results showed that catalase activity in erythrocytes was not significantly different between the OMW and control groups. The fact that polyphenols from OMW had no effect on the activity of catalase may be explained by their ability to enhance other antioxidant mechanisms, basically those associated with the GSH system. However, we and other researchers have reported that administration of polyphenolic extract from OMW increased catalase activity in erythrocytes and the rate of H$_2$O$_2$ decomposition in several tissues in rats, mice and piglets (28, 49, 50). Thus, the effect of polyphenols from OMW on catalase may be animal-specific.

Unlike erythrocytes, the decomposition of H$_2$O$_2$ to H$_2$O and O$_2$ in tissues, is attributed not only to catalase but also to other enzymes such as glutathione peroxidase and peroxiredoxins (51). The rate of H$_2$O$_2$ decomposition was significantly higher in liver and brain at day 70 post birth and in heart at both day 42 and 70 post-birth in the OMW group compared to the control group. These results are very important since, if H$_2$O$_2$ is not decomposed to harmless products such as oxygen and water, in the presence of metal ions it may lead through the Haber-Weiss and Fenton reaction to the production of hydroxyl radicals (52). This is crucial for cells and tissues, because hydroxyl radicals can cause severe damage to macromolecules such as DNA, proteins and lipids (53, 54).

Apart from the enhancement of antioxidant mechanisms, the experimental feed reduced oxidative stress-induced damage such as lipid peroxidation. In biological systems, lipid peroxidation, generates a number of degradation products, such as MDA, and may lead to cell death (55, 56). TBARS in plasma, indicating lipid peroxidation, were significantly decreased at 42 days post-birth in the OMW group compared to the controls. This decrease in TBARS agreed with the corresponding increase in the antioxidant GSH at day 42. These findings are important, since animals at the young age of 42 days have weak antioxidant mechanisms, and so there is a great need for their enhancement (19, 32). Metcalfe and Alonso-Alvarez (57) have suggested that early development is a life stage where oxidative stress levels are high due to the presumed link between the high metabolic activities required for growth and ROS generation. Moreover, lipid oxidative damage was higher in lambs at a young age than in adults, and it was positively related to growth rate over the first 4 months of life (58). Moreover, feed supplemented with OMW reduced TBARS in heart, spleen, liver and quadriceps muscle at 42 days post-birth and in brain at 70 days compared to controls. Unlike in plasma, these results are also in accordance with the corresponding increase in GSH level in these tissues. In previous studies, we have shown that feed supplemented with polyphenols derived from OMW reduced lipid peroxidation in blood and tissues of different farm animals, such as chickens and piglets (10, 28). Major polyphenolics (e.g. tyrosol, hydroxytyrosol) found in OMW have been shown to be responsible for reduction of lipid peroxidation (10, 59). Lipid peroxidation, apart from affecting animal health, may reduce meat quality due to the production of off-flavor products (60, 61). Thus, the decrease in TBARS l in muscle is of particular importance, because it is the main edible part from these animals.

Furthermore, the results showed that feed supplemented with OMW significantly reduced protein oxidation (i.e. protein carbonyl levels) in blood plasma at 70 day post-birth compared to control. In tissues, protein carbonyl levels were also decreased in heart, spleen and liver at day 42 post birth and in quadriceps muscle at day 70 post birth. Similar studies from our research group have shown that feed...
supplemented with OMW reduced these levels both in blood and many tissues of piglets and chickens (10, 28). Zhang et al. have shown that protein oxidation can induce protein polymerization and aggregation, thus affecting their digestibility, which reduces the nutritional value of muscle foods (62). Moreover, they suggested that oxidation of essential amino acids of meat proteins (i.e. cysteine, tyrosine, glycine, histidine, alanine, leucine, lysine) could reduce its nutritional value (62). Thus, the decrease in protein carbonyl levels in quadriceps muscle, an edible tissue, indicates improvement of meat quality.

Finally, the TAC, indicating the total level of antioxidant molecules, was increased in blood significantly at day 42 post-birth in the OMW compared to the control group. This increase was in accordance with the simultaneous increase in GSH and decrease in lipid peroxidation. TAC in tissues of the OMW group did not differ significantly from those of control, apart from brain tissue in which it was increased both at day 42 and 70 post-birth. The increase in TAC in brain may be attributed to the parallel increase in GSH at the same time-points. The absence of increase in TAC in other tissues of then OMW group, which did exhibit an increase in GSH, may be explained by the fact that when some antioxidants mechanisms are increased in living organisms, some others may be reduced as a compensatory mechanism (21, 32). On the other hand, a previous report of our research group showed high TAC in many tissues, and especially in brain, of animals who received feed supplemented with OMW polyphenols (28). Frankel et al. suggested that the increase in TAC levels in pigs after feeding with OMW was due to the direct free radical-scavenging from polyphenols. (8)

In conclusion, as far as we are aware, this is the first study showing that feed supplemented with OMW improved antioxidant mechanisms in lambs and reduced free radicals by reducing lipid and protein oxidation in blood and several vital organs. However, these beneficial effects seemed to be tissue-specific perhaps due to physiological and biochemical differences, as well as to differences in the content of antioxidant enzymes and molecules between different tissues (32). Moreover, an increase in GSH level of lamb blood and tissues seemed to be a main mechanism for the enhancement of antioxidant capacity. Thus, the exploitation of OMW from olive oil industries for making feed with antioxidant properties can be an alternative and cost-effective intervention for the treatment of oxidative stress-induced pathological conditions in lambs. In addition, as mentioned above, OMW causes serious pollutant problems in olive oil production areas and its use for making livestock feed could be a way to protect the environment. Of course, more studies are needed in order to elucidate the molecular mechanisms through which the bioactive compounds of OMW enhance the antioxidant mechanisms and improve the animals’ redox status.

Conflicts of Interest
The Authors declare that there are no conflicts of interest in regard to this study.

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