EFFECT OF TESTOSTERONE ENANTHATE HORMONES ON SOME PRODUCTION, PHYSIOLOGICAL TRAITS AND THEIR RESIDUE IN THE MEAT OF LOCAL RABBITS IN SULAYMANIYAH, IRAQ


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Abstract. A total of 48 weaned male rabbits were randomly arranged to evaluate the effects of the Testosterone Enanthate (TE) for four treatments: negative control-without injection, positive control-Phosphate-buffered saline PBS, (TE) 4 IU and 8 IU injected intramuscularly (B1 and B2, respectively). Body weight traits measured by weekly weighing for 72 days (slaughter weight) their body weight and average daily gains (ADG) were calculated, and were then slaughtered to assess the TE residues of meat. Blood samples were collected for measuring phosphate-buffered saline (CBC) count, ALT, AST, ALP, Growth Hormone (GH) and Testosterone concentration. The present study observed a significant increase in total BW gain for group B2 (8 IU), compared to negative control, positive control, and B1 group. The blood Testosterone concentration in the B2 group, significantly increased were higher than other treatment groups, at 2.73±0.26 ng/dL. The TE residue in meat there was no significant difference after 30 days end of treatment. In conclusion, TE injection improves BW traits in male growing rabbits when injected with a double dose of TE, with the exception of some CBC counts and has a significant mitigating effect on stress parameters, and increasing effect on serum GH and Testosterone concentration, with no significant changes in the TE residue in the meat.

Keywords: buck rabbits, TE injection, body weight gain, CBC, testosterone concentration, TE residues

Introduction

Domestic rabbits, in recent years, have been identified as economy livestock of meat shortage in high human population developing countries. Animals Meat, including rabbit’s meat, supply a valuable and palatable source of protein. Rabbit meat has a very good nutritive value, being comparatively high in protein, low in fat, calories and sodium, and so could bridge the wide gap in dietary protein intake (Adeyinka et al., 2007). Anabolic androgenic steroids (AAS) are synthetic derivatives of testosterone hormone in the male that have been adjusting to improve their anabolic rather than androgenic activity (Shahidi, 2001). The AAS effects promote protein synthesis, muscle growth and erythropoiesis (Mottram and George, 2000). Anabolic steroids are a class of steroid hormones based on the androgen testosterone and are recognized for their effects on building up muscle and are used as an enhancing drug (Thienpont et al., 1998). Anabolic steroids increase muscle size by the promotion of positive nitrogen balance by stimulating protein production and decreasing destruction (Guan et al., 2010). However, there is
insufficient available information for their use in these animals. Recently, boldenone undecylenate (androgenic steroid) has been used in the growth improvement and conversion of food in food-producing animals. It is also well known for increasing vascularity in preparation for bodybuilding contests. It might also play an important role not only in controlling normal testicular development, but also in maintaining spermatogenesis and normal testicular function (Tousson et al., 2012). The use of the anabolic steroid (BOL) resulted in obvious improvement in the growth rate (Tousson et al., 2012). This effect could be attributed to the promotion of the body tissue building process by protein synthesis indirectly via stimulation of growth hormone, insulin-like growth factor secretion, and animal appetite (Ferreira et al., 1998) or reduction of glucocorticoid receptor levels and sensitivity to endogenous glucocorticoids; therefore, the strong growth promoting potency is based not only on its anabolic activity as an antiglucocorticoid (Melloni et al., 1997; Thienpont et al., 1998). However, there were some reports reviewed that the rabbits’ growth performance was not affected by testosterone injection (Tawfeek et al., 1994). Administration of BOL to male rabbits at a dose of (5 mg/kg body weight) had a positive and significant benefit on growth performance (feed efficiency, total and daily weight gain) and a significant increase in dressing percent (Mohammed et al., 2016). Nahed et al. (2010) reported that using of BOL in rabbit was not sufficient to confirm its use in rabbits although a considerable improvement of total weight gain, feed efficiency and feed conversion ratio were attained after application of BOL in the male rabbit. However, there is no reported in detail study on the effect of BOL on all carcass traits and blood parameters (Abdel-Hamid and Farahat, 2015). The double and the normal recommended dose positively increase serum total protein and globulin, while normal recommended dose apparently increases serum cholesterol and decrease plasma corticosterone level. The European Economic Community (EEC) banned the use of anabolic compounds as growth accelerators in food animals. While the United States Food and Drug Administration (USFDA) permitted the limited use of some hormones with natural origins (such as estradiol and testosterone) and some synthetic hormones (such as Zeranol and trenbolone) in animal husbandry (MacVinish and Galbraith, 1988; Sadek et al., 1998). Little work has been conducted in Iraq regarding the use of testosterone enanthate TE injection as a growth promoter and their residue in edible tissue. A report of Omar (2012), there were no significant differences among the three experimental groups in the TE residue of the cooked meat at 8 months of age, but their values were significantly lower than the fresh meat of Karadi lambs at the same period. A number of authors and official international committees proved that residues of natural steroidal hormones are not dangerous on the health of the consumer (JECFA, 1988; Lone and Van Ginkel, 1997; Galbraith, 2002). Testosterone is usually given in an injection vehicle with oestradiol, and as same with other steroid hormones, the testosterone injection do not differ than the natural hormone inside the body of the organism, and this hormone focused in the liver tissue and kidney, but in fat are the highest concentration of this hormone. When comparing the residues in treated animals and the natural ratios in animals without treatment, they have to be equal or close to it (Omar, 2012). As well as the content of the natural hormone and the testosterone in male sheep and goats is higher than females, although the consumer is preferred the meat of males on other meats. Hence, this study was performed to determine the effects of Two-doses administration of TE on body weight (BW), some blood biochemical parameters and testosterone residual effect of mature local male rabbits.
Materials and Methods

Location and duration of the study: The present study was carried out at the Experimental farm of the University of Sulaimani Polytechnic, Halabja Technical Agriculture College, Halabja-Iraq, during spring (April to June) of 2017. A total of (48 bucks) weaned male rabbits (14 weeks old) of Local breed were randomly assigned to a completely randomized design arrangement of treatments (Four treat 12 bucks /treat: negative control-without injection, positive control injected intramuscularly by Phosphate-buffered saline PBS, Testosterone Enanthate (4 IU) injected intramuscularly (B1) and Testosterone Enanthate (8 IU) injected intramuscularly (B2). All rabbits were injected after 14 days of adaptation and repeated once after 30 days of the first injection. Bucks were housed in a semi-closed rabbitry housing system with does and kept in batteries of individual cages (60 x 50 x 35 cm) supplied with feeding hoppers made of galvanized steel sheet and nipples for an automatic drinker. A commercial concentrate pellets ration is introduced to bucks throughout the experiment (Table 1), containing (16.91%) crude protein and (2703 Kcal/Kg DM) ration metabolizable energy (ME) (NRC, 1977). Fresh and clean drinking water was supplied ad-libitum. The feed and the water were offered ad-libitum. All the bucks were grown under identical environmental and feeding conditions as well as the same stocking density. A photoperiod (12L:12D) from 0900 to 2100 h was used throughout the experimental period. The temperature was maintained at a range of 28-32°C inside the house.

Table 1. Ingredient Composition Of The Diet During The Experiment Period

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maize</td>
<td>35</td>
</tr>
<tr>
<td>Wheat Grain</td>
<td>25</td>
</tr>
<tr>
<td>Soybean Meal</td>
<td>10</td>
</tr>
<tr>
<td>Sesame Oil</td>
<td>15</td>
</tr>
<tr>
<td>Wheat Bran</td>
<td>14</td>
</tr>
<tr>
<td>Common Salt</td>
<td>0.6</td>
</tr>
<tr>
<td>Premix</td>
<td>0.25</td>
</tr>
<tr>
<td>DL. Methionine</td>
<td>0.15</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
</tr>
<tr>
<td>Crud Protein (%)</td>
<td>16.91</td>
</tr>
<tr>
<td>Metabolizable Energy Kcal/Kg Dm</td>
<td>2703</td>
</tr>
</tbody>
</table>

Traits Measurements: Traits measured were; body weights: fryer rabbits were weighed weekly until 72 (slaughter weight) days and weights were recorded in grams. Body weight gain: calculated as differences between two successive different weights. Average daily gains ADG: calculated as the differences between two successive different weights at two different periods divided by the number of days between the two weights.

Blood parameters: Blood samples were collected twice in the experiment, before treatment and after one month after the first injection. The samples were taken in two test tubes, one with an anticoagulant for measuring CBC count levels and the other one without using anticoagulant to separate the serum for measuring ALT, AST, ALP, Growth Hormone (GH) and Testosterone concentration. Plasma and serum samples were stored at -20°C until assayed. ALT, AST, ALP, Growth Hormone (GH) and Testosterone concentration were measured using commercial kits. For CBC levels we used Hematology analyzer (URIT 2900z, Vet Z plus, China), and for enzymes activity, we
used Biochemical Analyser (Floxer pro. S, China), and Hormone concentration by (MINI VIDAS®, France). For each treatment, five samples were assayed.

TE residue in meat: After slaughtering the bucks, 30 days after the last injection of the experiment. 500 gm of muscle were taken and placed in polyethylene bags and stored at -18°C. Semimembranosus (SM) from pelvic limb leg was dissected according to the procedure of Butterfield et al. (1983). The surfaces of the muscles were cleaned from all fat and connective tissue. Quantitative determination of testosterone concentration (μg/kg = ng/mL) was carried out by using Radioimmunoassay (RIA) by using (cobas e 411 analyzer, made in Germany). The analytic sensitivity of the assay was 0.025 ng/mL.

Statistical analysis: Complete Randomized Design (CRD) procedures of XLstat. (7.5.2, 2010) in one-way ANOVA were used to determine the effects of Testosterone Enanthate TE and injection effect on the body weight characteristics, blood parameters, and TE residue. Group differences were determined using Duncan’s multiple range tests at (p≤0.05) (Duncan, 1955).

Results

The effect of Testosterone Enanthate-TE hormone on growth performance parameters of local male rabbits was showed in Table 2. The results revealed that TE injection in male rabbits resulted not significant (p>0.05) different among groups, in initial BW, final BW gain, average daily gain ADG, and the high weight record were seen in B2 group for final BW and ADG (1593.3±41.5 and 6.292±0.384) gm, respectively. But there were a significant (p<0.05) different in total BW gain for group B2 (8 IU), which recorded (377.5±23.0) gm compared to negative control, positive control and B1 group, which were recorded (281.2± 60.8) gm, (271.2± 32.5) gm and (284.1± 37.1) gm, respectively. The best results were for the group (B2) in total BW and ADG increased in the ADG for groups (B1) and (B2) compared with the control.

Table 2. Effect of te hormone on growth performance parameters of male rabbits

<table>
<thead>
<tr>
<th>Parameters</th>
<th>NEGATIVE CONTROL</th>
<th>POSITIVE CONTROL</th>
<th>B1</th>
<th>B2</th>
</tr>
</thead>
<tbody>
<tr>
<td>initial Bw (gm)</td>
<td>1225.0±58.2</td>
<td>1260.8±37.0</td>
<td>1246.6±30.8</td>
<td>1194.1±26.1</td>
</tr>
<tr>
<td>final Bw (gm)</td>
<td>1521.6±44.8</td>
<td>1556.6±25.4</td>
<td>1516.6±36.8</td>
<td>1593.3±41.5</td>
</tr>
<tr>
<td>Total Bw Gain (gm)</td>
<td>281.2±60.8</td>
<td>271.2±32.5</td>
<td>284.1±37.1</td>
<td>377.5±23.0</td>
</tr>
<tr>
<td>ADG (gm)</td>
<td>4.68±1.013</td>
<td>4.521±0.543</td>
<td>4.736±0.620</td>
<td>6.292±0.384</td>
</tr>
</tbody>
</table>

- Negative Control: Without Injection
- Positive Control: Injected Intramuscularly With Bps (0.25 MI)
- B1: Received (4 IU) Intramuscular Injections Of T.E
- B2: Received (8 IU) Intramuscular Injections Of T.E
- Bw: Body Weight; ADG: Average Daily Gain

Effects of dose of TE and the injection effect on complete blood picture levels parameters were summarized in Table 3. Statistically, no significant differences (p>0.05) were recorded before treatment for complete blood picture levels parameters. While, there were significant differences (p<0.05) were observed for control groups (negative and positive) compared to TE hormone injection groups after treatment for Hb (g/dL), RBC (10⁶xUL) and HCT (%). The hormone injection groups (TE injection) significantly
The results of Table 4 demonstrated that changes in Leukocytes levels changed significantly (p<0.05) after TE hormone treated. Before treatment there were no significant differences (p>0.05) in the leukocytes levels, on the other hand, we recorded significant differences (p<0.05) in WBC count, Lymphocyte %, Monocytes %, Neutrophil%, Mean platelet volume (MPV%) and platelet distribution width (PDW). The TE injection increased the WBC count significantly (p<0.05), which high count recorded in B1 (4 IU, TE), and the highly significant value for lymphocyte and monocyte percent were recorded in B2 (8 IU) (61.0±1.0 % and 0.75±0.02 %, respectively). Conversely, the neutrophils count significantly (p<0.05) higher in BPS injection (Positive Control) (59.6±3.0 %). Also, the significant level was seen in MPV and PDW for B2 injection, which was 6.05±0.02 % and 9.10 ±0.44 %, respectively. As well, B1 injection significantly differentiated too in PDW (8.60±0.01 %), when they compared to negative and positive control (7.50 ±0.01 and 7.63 ±0.02, respectively).
The comparison between the injection groups and negative control in the concentration of ALT, AST, ALP, Growth Hormone GH and Testosterone levels in the local rabbit are summarized in Table 5. Indeed, no significant differences were seen before treatment in all parameters, except in B1 treatment for AST and ALP levels. The AST recorded high level in B1 (39.0±1.6 IU/L). And ALP recorded less level in B1 (79.0±1.2 IU/L) when they compared to other treatments. After treatment, the parameters level change, the significant record seen in ALT, GH and testosterone concentration for B2 (8 IU) TE injection. The significant level for ALT recorded (69.5±9.4 IU/L), as compared to negative control, positive control and B1 injection. The GH concentration recorded significantly (p<0.05) higher level (0.32±0.01 ng/dL) at B2 (8 IU) TE injection when it compared to negative control and positive control (0.24±0.01 and 0.28±0.01 ng/dL, respectively). As well as, the Testosterone concentration in the B2 group, significantly (p<0.05) increased and higher than other treatments groups, which recorded (2.73±0.26 ng/dL). And the other groups recorded significant differences for negative control, positive control and B1(4 IU) TE injection, which were (0.60±0.06, 0.83±0.10 and 1.28±0.15 ng/dL).

Table 4. Changes in the leukocyte levels in different groups under study

<table>
<thead>
<tr>
<th>Items</th>
<th>Negative Control</th>
<th>Positive Control</th>
<th>B1</th>
<th>B2</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC (10^6 xul)</td>
<td>3.33±0.63^a</td>
<td>2.76±0.49^a</td>
<td>4.50±1.07^a</td>
<td>4.75±0.85^a</td>
</tr>
<tr>
<td>Lym. (%)</td>
<td>75.5±1.8^a</td>
<td>56.4±9.2^a</td>
<td>57.6±11.9^a</td>
<td>64.3±4.5^a</td>
</tr>
<tr>
<td>Mon. (%)</td>
<td>7.86±3.02^a</td>
<td>0.87±0.67^a</td>
<td>0.27±0.08^a</td>
<td>0.50±0.14^a</td>
</tr>
<tr>
<td>Neu. (%)</td>
<td>16.0±1.1^a</td>
<td>32.2±9.8^a</td>
<td>47.9±11.3^a</td>
<td>34.0±4.41^a</td>
</tr>
<tr>
<td>MPV (%)</td>
<td>6.26±0.06^a</td>
<td>6.05±0.71^a</td>
<td>6.35±0.05^a</td>
<td>6.47±0.31^a</td>
</tr>
<tr>
<td>PCT</td>
<td>0.133±0.003^a</td>
<td>0.144±0.030^a</td>
<td>0.331±0.142^a</td>
<td>0.111±0.026^a</td>
</tr>
<tr>
<td>PDW</td>
<td>9.50±0.10^a</td>
<td>9.85±1.11^a</td>
<td>10.48±0.15^a</td>
<td>10.62±0.70^a</td>
</tr>
</tbody>
</table>

-ANegative Control: Without injection.
-Positive Control: injected intramuscularly with BPS (0.25 ml).
-B1: received (4 IU) intramuscular injections of T.E.
-B2: received (8 IU) intramuscular injections of T.E.
-WBC: white blood cell; Lym.: lymphocyte; Mon.: monocyte; Neu.: neutrophil; MPV: Mean platelet volume; PCT: procalcitonin test; PDW: platelet distribution width.
-abc Means in the same row with different superscripts are significantly different at (p<0.05).

The results of TE residues in bucks’ meat after 30 days of treatment are presented in Figure 1. Significant differences were not seen in the testosterone residue of the fresh meat, while there was no significant (P> 0.05) increase in the TE residue in the fresh meat of B1 and B2 group intramuscular injections of TE with 4IU and 8IU (0.093 and 0.134 μg/kg) and as compared to the negative and positive groups which were 0.086 and 0.061 μg/kg, respectively Figure 1.
Table 5. Changes in the concentrations of alt, ast, alp, growth hormone and testosterone levels in different groups under study

<table>
<thead>
<tr>
<th>Items</th>
<th>Negative Control</th>
<th>Positive Control</th>
<th>B1</th>
<th>B2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Before treatment</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT (IU/l)</td>
<td>59.0±1.6*a</td>
<td>55.0±10.2*a</td>
<td>56.0±1.2*a</td>
<td>54.6±1.2*a</td>
</tr>
<tr>
<td>AST (IU/l)</td>
<td>34.0±4.4*ab</td>
<td>29.0±2.4*b</td>
<td>39.0±1.6*a</td>
<td>35.9±1.4*ab</td>
</tr>
<tr>
<td>ALP (IU/l)</td>
<td>158.3±3.8*a</td>
<td>112.3±22.2*a</td>
<td>79.0±1.2*b</td>
<td>174.4±34.3*a</td>
</tr>
<tr>
<td>GH (ng/dl)</td>
<td>0.26±0.02*a</td>
<td>0.32±0.02*a</td>
<td>0.15±0.02*a</td>
<td>0.27±0.01*a</td>
</tr>
<tr>
<td>Testosterone (ng/dl)</td>
<td>0.45±0.02*a</td>
<td>0.86±0.12*a</td>
<td>0.60±0.12*a</td>
<td>0.70±0.14*a</td>
</tr>
<tr>
<td><strong>After treatment</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT (IU/l)</td>
<td>41.0±2.7*b</td>
<td>39.9±1.1*ab</td>
<td>50.8±3.0*b</td>
<td>69.5±9.4*a</td>
</tr>
<tr>
<td>AST (IU/l)</td>
<td>25.6±0.9*a</td>
<td>25.5±0.7*a</td>
<td>25.1±2.1*a</td>
<td>28.7±3.0*a</td>
</tr>
<tr>
<td>ALP (IU/l)</td>
<td>108.6±4.0*a</td>
<td>117.0±16.1*a</td>
<td>107.7±13.9*a</td>
<td>115.1±12.0*a</td>
</tr>
<tr>
<td>GH (ng/dl)</td>
<td>0.24±0.01*c</td>
<td>0.28±0.01*b</td>
<td>0.29±0.01*ab</td>
<td>0.32±0.01*a</td>
</tr>
<tr>
<td>Testosterone (ng/dl)</td>
<td>0.60±0.06*bc</td>
<td>0.83±10*bc</td>
<td>1.28±0.15*b</td>
<td>2.73±0.26*a</td>
</tr>
</tbody>
</table>

-Negative Control: Without injection.
-Positive Control: injected intramuscularly with BPS (0.25 ml).
-B1: received (4 IU) intramuscular injections of T.E.
-B2: received (8 IU) intramuscular injections of T.E.
-ALT: alanine aminotransferase; AST: aspartate aminotransferase; ALP: alkaline phosphatase level; GH: growth hormone.

abc Means in the same row with different superscripts are significantly different at (p<0.05).

Figure 1. Concentration residue of Testosterone Enanthate in meat of local rabbit bucks (Negative Control: Without injection; Positive Control: injected intramuscularly with BPS (0.25 ml); B1: received (4 IU) intramuscular injections of T.E; B2: received (8 IU) intramuscular injections of T.E.) Means in the same row with different superscripts are significantly different at (p<0.05)

Discussions

The growth performance improved in treated groups (B1 and B2) proportional to the control groups (Table 2), which is consistent with antecedent reports (Thabet et al., 2010). The same observation was described by Tousson et al. (2012) and Mohammed et al. (2016) when they use anabolic steroid (BOL, boldenone undecylenate), who stated that the use of the (BOL) give rise to an obvious improvement in the growth rate. This effect could be imputed to the promotion of the body tissue building process by protein synthesis.
indirectly via stimulation of growth hormone, insulin-like growth factor secretion, and animal appetite (Ferreira et al., 1998) or reduction of glucocorticoid receptor levels and sensitivity to endogenous glucocorticoids; therefore, the strong growth promoting potency is based not only on its anabolic activity as an antiglucocorticoid (Melloni et al., 1997; Thienpont et al., 1998). However, Tawfeek et al. (1994) reported that the rabbits’ growth performance was not affected by testosterone injection.

The obtained findings in Table 3 agree with the results of Battista et al. (2003) and Liewellyn (2006) who founded that, testosterone dosage and its entrance to the body caused an increase in hematocrit %. Moreover, Gagnon et al. (1994) founded that, the raised hematocrit and hemoglobin persist for extended periods after the cessation of androgenic steroids use. Similarly, Ahmed (2014) recorded significant increase RBCs, Hb, and PCV in boldenone administered groups as compared to the control one. These results may be imputed to that; anabolic steroids could stimulate erythropoiesis (Gabr et al., 2009) through the direct positive effect of androgenic steroids on erythropoietin production in renal tissues (Liewellyn, 2006). This reaction is mainly driven by the androgen receptor stimulation in renal tissue, leading to the stimulation of erythropoietin production directly. Androgens may also influence the stem cells directly, perhaps by enhancing the stem cell’s responsiveness to erythropoietin (Snyder, 2008).

Anabolic steroids could spur erythropoiesis, a mechanism that may occur by stimulating erythropoietic-stimulating factor (Gabr et al., 2009). These results (Table 4) are in agreement with Urhausen et al. (2003) and Gabr et al. (2009) who adduced that the liver and kidney functions increased significantly after intramuscular BOL undecylenate injection on weaned male lambs. Similarly, current results are in conformity with Dickerman et al. (1999) and Tousson et al. (2011a, 2011b) who noticed that the anabolic steroid-induced hepatotoxicity. Following the same opinions, Istasse et al. (1988) reported that 17b-estradiol increased nitrogen retention and decreased blood urea nitrogen concentrations. Injection of the anabolic steroid BOL induced changes in oxidative stress biomarker levels and antioxidant defense systems in rabbit liver and kidney (El-Moghazy et al., 2012).

In agreement with our findings (Table 5), in treated groups with anabolic androgenic steroid, the serum testosterone levels were significantly higher than that in the control group (Urhausen et al., 2003; Takahashi et al., 2004; Gabr and Shaker, 2006; Gabr et al., 2009; Ishak and Omer, 2014). Simontacchi et al. (2004) reported that administration of testosterone derivatives alone did not induce plasma testosterone levels. Also, Shimomura et al. (2005) showed that the treatment of rats with ethinyloestradiol alone the testosterone levels decreased significantly in serum and the testis. Our results are in agreement with Omar (2012) who studied the effects of Testosterone Enanthate injection on testis function, live weight gain and carcass traits of Karadi lamb rams. The significant increase in testosterone observed in (B1 and B2) of mature rabbits treated by boldenone may be in accordance with the findings of Urhausen et al. (2003), Takahashi et al. (2004) and Gabr and Shaker (2006) who founded that, serum testosterone levels in treated groups with androgenic steroids were significantly higher than that in the control group. These results shore up former reports (Gabr et al., 2009; Tousson et al., 2012) who mentioned that an increase of testosterone may be attributed to the synthesis of a substrate related to the primary male sex hormone.

In the present study, results of TE residue in the fresh meat after 30 days stopped injection, showed no significant difference between the groups (Figure 1), these results are in contrast with the finding of Paris et al. (2006), when they found a significant
difference between the treated and non-treated lambs of the same age group, for example, the anabolic steroids in the muscle, liver, kidneys and adipose tissue. While there was an increase of TE residue in the fresh meat of the male rabbit treated with 4IU TE and those treated with 8IU TE as compared to control groups. The main possible mechanism for these results could be the testicular hormone especially testosterone which let the male be superior in TE residue than the control groups (Omar, 2012). The exogenous testosterone has also been considered as the other reason for elevating TE residue in fresh meat of the intact animal as compared to the castrated. To our knowledge obtained from available literature, no other previous study dealt with TE injection on rabbit male and their residue in meat. These differences may be related to exogenous testosterone in addition to testicular hormone especially testosterone which produce mainly by leydig cells. As well as increasing TE residue in fresh meat, which could be reduced in the cooking process (Omar, 2012).

Conclusion

From the obtained results it could be concluded that, although intramuscular injection of Testosterone Enanthate espesioly B2 (8IU) to male local rabbits enhances body gain and increased some hematological parameters, it increased oxidative stresses biomarkers levels, pro-inflammatory cytokines and creatinine indicating side effects on the liver and kidneys. And a no significant deferences in TE residue was seen in the fresh meat after 30 days of injection.

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