Introduction

In recent years, human nutritionists have recommended an increase in the intake of polyunsaturated fatty acids (PUFA) because of their beneficial effects on health, mainly in the prevention of cardiovascular disease (KRAUSS et al., 2001). Some authors have increased the PUFA content of chicken meat by enriching animal diets in polyunsaturated fats (LIN et al., 1989; AJUYAH et al., 1993; LÓPEZ-FERRER et al., 1999, 2001; GONZALEZ-ESQUERRA and LEESON, 2000). However, a higher PUFA content in poultry meat increases its susceptibility to oxidation (MARASCHIELLO et al., 1999; RUIZ et al., 1999). Part of this variability may be attributed to the analytical methods used to determine α−TA and fatty acid analysis.

In order to improve the oxidative stability and, thus, to increase the shelf life of meat, antioxidants have been successfully added to animal feeds. Among them, α-tocopheryl acetate (α−TA) has demonstrated the highest efficiency in preventing lipid oxidation (LIN et al., 1989; AHN et al., 1995; De WINNE and DIRINCK, 1996; GRAU et al., 2001a,b) and, as a consequence, enhances the development of organoleptic problems compromising meat quality (MALCZYK et al., 1999; GONZALEZ-ESQUERRA and LEESON, 2000; BOU et al., 2001).

The aim of the present study was to determine the effects of increasing amounts of dietary PUFA and different levels of supplementation with α−TA on α−Toc content of broiler breast and thigh meat.

Materials and methods

Animals and diets

The experiment received prior approval from the Animal Protocol Review Committee of the Universitat Autònoma of Barcelona, housing and husbandry of all animals conformed to European Union guidelines.

One hundred and ninety-two Ross strain 308 female broilers at one day of age were randomly distributed into 16 dietary treatments. Three cages by four birds were used for each of 16 dietary treatments. The animals were housed under standard conditions of temperature, humidity and ventilation. The diet was formulated to meet or exceed recommendations of NRC (1994) on the basis of 39% wheat, 34% soybean meal and 13% barley (Table 1). The experimental treatments resulted from the combination of 4 levels of supplementation with α−TA (Rovimix® E-50 Adsorbate. F. Hoffmann-LaRoche Ltd., Basel, Switzerland): 0 (E0), 100 (E1), 200 (E2), and 400 (E4) mg/kg of feed, achieved by blending different proportions of tallow (100, 60, 40 and 0%) and a mixture of linseed and fish oils (0, 40, 60 and 100%) , keeping total added fat constant (9%).

The fatty acid content of the experimental diets is shown in Table 2. More details about fatty acid content of these diets are found in CORTINAS et al. (2004). The increase in the polyunsaturation level of diets, while maintaining the total fatty acid content constant, was achieved at the expense of the saturated and monounsaturated fatty acids. Therefore the PUFA to SFA ratio of diets was 0.4, 1.0, 1.7 and 3.9 for PU15, PU34, PU45 and PU61, respectively.

Feed (all-mash) and water were provided ad libitum during the experimental period. Feed samples were taken during the experimental period for α−Toc and fatty acid analysis.

Sample collection

At the end of the experimental period (44 days of age), two animals per cage were randomly selected and processed in a commercial slaughterhouse. The edible portion of thighs

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and breasts were removed and weighed individually. Thighs were deboned and ground with skin whereas breasts were ground without skin. Thighs were collected with skin because it is usual to commercialize them this way. Tissue samples were freeze-dried, ground and stored at –20°C until further analyses. 

**α−Tocopherol analysis**

The α−Toc from feeds, thighs and breasts was extracted as described by JENSEN et al. (1999) starting from 2 g of feed sample and 100 mg of freeze-dried thigh and breast. α−Toc content was determined by normal-phase HPLC. The α−Toc recoveries were examined in order to observe if the lipid polyunsaturation level of chicken meat affects the α−Toc determination. Two levels of α−Toc standard (10 and 20 µg) (Sigma, St. Louuis, MO 63103, USA) were add−ed to 100-mg aliquots of tissue samples. α−Toc recovery for each α−Toc addition level in thigh and breast meat samples was determined using 4 aliquots.

**Fatty acids analysis**

Fatty acid content of the experimental diets was analyzed by gas-chromatography following the direct transesterification method described by SUKHJIA and PALMQUIST (1988). Nonadecanoic acid (Sigma, St. Louis MO, USA) was used as internal standard. Medium and long chain fatty acids

### Table 1. Composition and content of nutrients of basic diet (expressed as % of fresh matter)

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Composition</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wheat</td>
<td>Dry matter</td>
<td>39.3</td>
</tr>
<tr>
<td>Soybean meal 48% CP</td>
<td>Crude protein</td>
<td>34.01</td>
</tr>
<tr>
<td>Barley</td>
<td>Crude fat</td>
<td>13.4</td>
</tr>
<tr>
<td>Added fat</td>
<td>Crude fibre</td>
<td>9.0</td>
</tr>
<tr>
<td>Bicalcium phosphate</td>
<td>Ash content</td>
<td>2.2</td>
</tr>
<tr>
<td>Calcium carbonate</td>
<td>Crude Energy (Kcal/kg)</td>
<td>1.0</td>
</tr>
<tr>
<td>Salt</td>
<td>Metabolizable Energy (Kcal/kg)</td>
<td>0.4</td>
</tr>
<tr>
<td>Vitamin-mineral mix</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>DL-Methionine</td>
<td>0.28</td>
<td></td>
</tr>
<tr>
<td>L-Lysine</td>
<td>0.04</td>
<td></td>
</tr>
</tbody>
</table>

1 Values given in this table are means of 16 dietary treatments, result of a 4 × 4 factorial design with 4 different proportions of tallow, linseed and fish oil, and 4 different levels of dietary supplementation with α−tocopheryl acetate (0, 100, 200 and 400 mg/kg).
2 Vitamin and mineral mix per kg of feed: Vitamin A: 12000 IU; Vitamin D3: 2400 IU; Vitamin K3: 3 mg; Vitamin B1: 2.2 mg; Vitamin B2: 8 mg; Vitamin B6: 11 µg; Biotin: 150 µg; Calcium pantotenate: 25 mg; Nicotinic acid: 65 mg; Mn: 60 mg; Zn: 40 mg; I: 0.33 mg; Fe: 80 mg; Cu: 8 mg; Se: 0.15 mg.
3 Calculated value (FEDNA; http://www.etsia.upm.es/fedna/tables.htm)

### Table 2. Fatty acid composition of the experimental diets, expressed as g per kg

<table>
<thead>
<tr>
<th>Fatty Acid</th>
<th>PU15</th>
<th>PU34</th>
<th>PU45</th>
<th>PU61</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total FA</td>
<td>100.45</td>
<td>98.81</td>
<td>99.57</td>
<td>96.89</td>
</tr>
<tr>
<td>Total SFA</td>
<td>43.75</td>
<td>32.38</td>
<td>26.22</td>
<td>15.74</td>
</tr>
<tr>
<td>Total MUFA</td>
<td>41.30</td>
<td>32.55</td>
<td>28.32</td>
<td>20.31</td>
</tr>
<tr>
<td>Total PUFA</td>
<td>15.40</td>
<td>33.77</td>
<td>45.03</td>
<td>60.84</td>
</tr>
<tr>
<td>C 18:2 ω6</td>
<td>13.16</td>
<td>16.23</td>
<td>17.98</td>
<td>20.17</td>
</tr>
<tr>
<td>C 18:3 ω3</td>
<td>1.55</td>
<td>16.45</td>
<td>24.62</td>
<td>36.27</td>
</tr>
<tr>
<td>C 20:5 ω3</td>
<td>ND</td>
<td>0.81</td>
<td>1.77</td>
<td>3.35</td>
</tr>
<tr>
<td>C 22:6 ω3</td>
<td>ND</td>
<td>0.07</td>
<td>0.18</td>
<td>0.33</td>
</tr>
<tr>
<td>PUFA:SFA</td>
<td>0.35</td>
<td>1.04</td>
<td>1.72</td>
<td>3.87</td>
</tr>
</tbody>
</table>

1 Values given in this table are means of 4 dietary treatments with different level of supplementation with α−tocopheryl acetate: 0, 100, 200 and 400 mg/kg.
2 PU15: 15 g polyunsaturated fatty acids /kg of feed; PU34: 34 g polyunsaturated fatty acids /kg of feed; PU45: 45 g polyunsaturated fatty acids /kg of feed; PU61: 61 g polyunsaturated fatty acids /kg of feed.
3 FA: fatty acids; SFA: saturated fatty acids; MUFA: monounsaturated fatty acids; PUFA: polyunsaturated fatty acids.
ND: Not detected.
were quantified, and the ones containing 2 or more double bonds were included in the group of PUFA.

**Statistical analysis**

The Mann-Whitney U test was used to determine whether the α−Toc recoveries in thigh (n = 4) and breast (n = 4) meat were affected by the α−Toc addition levels. One-way ANOVA was performed to determine whether the PUFA dietary levels affected α−Toc global recoveries in thigh (n = 8) and breast (n = 8) meat. Effect of a dietary treatment on α−Toc content in chicken thigh (n = 96) and breast (n = 96) meat was tested by multifactorial ANOVA with repeated measurements. Data were treated using the MIXED procedure of SAS (n = 96). The interaction allowed comparing the slopes of the equations. Relationship between α−Toc content in thigh (n = 96) and breast (n = 96) meat, respectively, on the one hand, and dietary α−Toc and PUFA on the other hand, was fitted by using non-linear regressions by means of the NLIN (non-linear regression) procedure of SAS. The equation type was \( y = x_1(a - bx_2) \), where \( y \) is the α−Toc concentration in studied tissues, \( x_1 \) is α−Toc content of the diet and \( x_2 \) is PUFA content of the diet. This data analysis was represented using SigmaPlot 8.02 (2002). In all cases, \( P \leq 0.05 \) was considered significant.

**Results and discussion**

The α−Toc content of the experimental diets is shown in Table 3. It can be observed that the analyzed dietary α−Toc content was approximately 20% higher than the theoretical value.

**Recovery of α−tocopherol**

As hypothesized, the level of dietary supplementation with α−TA did not affect α−Toc recoveries in thigh and breast meat, but the interaction allowed comparing the slopes of the equations. Relationship between α−Toc content in thigh (n = 96) and breast (n = 96) meat, respectively, on the one hand, and dietary α−Toc and PUFA on the other hand, was fitted by using non-linear regressions by means of the NLIN (non-linear regression) procedure of SAS. The equation type was \( y = x_1(a - bx_2) \), where \( y \) is the α−Toc concentration in studied tissues, \( x_1 \) is α−Toc content of the diet and \( x_2 \) is PUFA content of the diet. This data analysis was represented using SigmaPlot 8.02 (2002). In all cases, \( P \leq 0.05 \) was considered significant.

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Table 3. Analysed α−Tocopherol values in experimental diets (mg/kg; mean ±SE).

<table>
<thead>
<tr>
<th>Dietary polyunsaturation</th>
<th>Dietary supplementation with α−tocopheryl acetate</th>
<th>Recovery of α−Toc (α−Toc; %) in meat from broilers fed diets with different degree of polyunsaturation</th>
</tr>
</thead>
<tbody>
<tr>
<td>PU15 (5 ± 0.3)</td>
<td>E0: without supplementation with α−tocopheryl acetate</td>
<td>α−Toc recoveries in thigh (n = 4)</td>
</tr>
<tr>
<td>PU34 (6 ± 0.7)</td>
<td>E1: supplemented with 100 mg/kg α−tocopheryl acetate</td>
<td>α−Toc recoveries in breast (n = 4)</td>
</tr>
<tr>
<td>PU45 (6 ± 0.8)</td>
<td>E2: supplemented with 200 mg/kg α−tocopheryl acetate</td>
<td>α−Toc recoveries in thigh (n = 8)</td>
</tr>
<tr>
<td>PU61 (5 ± 0.6)</td>
<td>E4: supplemented with 400 mg/kg α−tocopheryl acetate</td>
<td>α−Toc recoveries in breast (n = 8)</td>
</tr>
</tbody>
</table>

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Table 4. Recovery of α−tocopherol (α−Toc; %) in meat from broilers fed diets with different degree of polyunsaturation.

<table>
<thead>
<tr>
<th>Dietary treatments</th>
<th>Thigh with skin</th>
<th>Breast</th>
<th>Recovery of α−Toc (α−Toc; %) in meat from broilers fed diets with different degree of polyunsaturation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>α−Toc standard</td>
<td>Global Recovery</td>
<td>α−Toc standard</td>
</tr>
<tr>
<td></td>
<td>10A (n = 4)</td>
<td></td>
<td>20A (n = 4)</td>
</tr>
<tr>
<td>PU15</td>
<td>86.61</td>
<td>87.49</td>
<td>87.05^a</td>
</tr>
<tr>
<td>PU34</td>
<td>78.41</td>
<td>78.77</td>
<td>78.59^b</td>
</tr>
<tr>
<td>PU45</td>
<td>73.78</td>
<td>76.52</td>
<td>75.15^c</td>
</tr>
<tr>
<td>PU61</td>
<td>59.13</td>
<td>61.17</td>
<td>60.15^d</td>
</tr>
</tbody>
</table>

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1 PU15: 15 g polyunsaturated fatty acids /kg of feed; PU34: 34 g polyunsaturated fatty acids /kg of feed; PU45: 45 g polyunsaturated fatty acids /kg of feed; PU61: 61 g polyunsaturated fatty acids /kg of feed.

2 α−Toc: without supplementation with α−tocopheryl acetate; E1: supplemented with 100 mg/kg α−tocopheryl acetate; E2: supplemented with 200 mg/kg α−tocopheryl acetate; E4: supplemented with 400 mg/kg α−tocopheryl acetate.

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Statistical analysis

The Mann-Whitney U test was used to determine whether the α−Toc recoveries in thigh (n = 4) and breast (n = 4) meat were affected by the α−Toc addition levels. One-way ANOVA was performed to determine whether the PUFA dietary levels affected α−Toc global recoveries in thigh (n = 8) and breast (n = 8) meat. Effect of a dietary treatment on α−Toc content in chicken thigh (n = 96) and breast (n = 96) meat was tested by multifactorial ANOVA with repeated measurements. Data were treated using the MIXED procedure of SAS (n = 96). The interaction allowed comparing the slopes of the equations. Relationship between α−Toc content in thigh (n = 96) and breast (n = 96) meat, respectively, on the one hand, and dietary α−Toc and PUFA on the other hand, was fitted by using non-linear regressions by means of the NLIN (non-linear regression) procedure of SAS. The equation type was \( y = x_1(a - bx_2) \), where \( y \) is the α−Toc concentration in studied tissues, \( x_1 \) is α−Toc content of the diet and \( x_2 \) is PUFA content of the diet. This data analysis was represented using SigmaPlot 8.02 (2002). In all cases, \( P \leq 0.05 \) was considered significant.

Results and discussion

The α−Toc content of the experimental diets is shown in Table 3. It can be observed that the analyzed dietary α−Toc content was approximately 20% higher than the theoretical value.

Recovery of α−tocopherol

As hypothesized, the level of dietary supplementation with α−TA did not affect α−Toc recoveries in thigh and breast meat.
meat samples (data not shown). In addition, the α−Toc recoveries in thigh and breast meat samples were not affected by the level of α−Toc standard addition. Therefore, the global recoveries of α−Toc in meat samples for the different polyunsaturation treatments. On the other hand, global recoveries in breast meat samples were higher than in thigh meat samples, and global recoveries in breast meat samples increased as die-

Table 5. Effect of dietary polyunsaturation and α−tocopheryl acetate supplementation on least-squares means and their pooled SE of α−tocopherol content (mg/kg) in thigh with skin and breast meat

<table>
<thead>
<tr>
<th>Dietary polyunsaturation</th>
<th>Thigh with skin</th>
<th>Breast</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>E0¹</td>
<td>E1</td>
</tr>
<tr>
<td>PU15</td>
<td>2.6²</td>
<td>20.6²</td>
</tr>
<tr>
<td>PU34</td>
<td>0.7²</td>
<td>17.1²y</td>
</tr>
<tr>
<td>PU45</td>
<td>0.4²</td>
<td>12.3²y</td>
</tr>
<tr>
<td>PU61</td>
<td>0.1²</td>
<td>10.8²y</td>
</tr>
<tr>
<td>Grand Mean</td>
<td>0.9D</td>
<td>15.5C</td>
</tr>
<tr>
<td>SE</td>
<td>3.19</td>
<td></td>
</tr>
</tbody>
</table>

P values

PUFA *** ***
α−TA level *** ***
PUFA × α−TA level * **

1 Values given in this table were obtained from multifactorial ANOVA (n = 96)
2 PU15: 15 g polyunsaturated fatty acids /kg of feed; PU34: 34 g polyunsaturated fatty acids /kg of feed; PU45: 45 g polyunsaturated fatty acids /kg of feed; PU61: 61 g polyunsaturated fatty acids /kg of feed.
3 ε0: without supplementation with α−tocopheryl acetate; E1: supplemented with 100 mg/kg α−tocopheryl acetate; E2: supplemented with 200 mg/kg α−tocopheryl acetate; E4: supplemented with 400 mg/kg α−tocopheryl acetate.

* = significant effect at P ≤ 0.05; ** = P ≤ 0.01; *** = P ≤ 0.001.

Table 6. Dependence of α−Tocopherol content (y; mg/kg) in breast meat and thigh meat, respectively, on dietary α−tocopherol (x1; mg/kg) and on polyunsaturated fatty acid content in feed (x2; g/kg)

<table>
<thead>
<tr>
<th>Independent variable¹</th>
<th>Dietary treatments²</th>
<th>Thigh with skin</th>
<th>Breast</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Equation</td>
<td>r²</td>
<td>P</td>
</tr>
<tr>
<td>α−Tocopherol content in feed [mg/kg]</td>
<td>PU15, PU34</td>
<td>y = 0.114x1</td>
<td>0.93 ***</td>
</tr>
<tr>
<td></td>
<td>PU45, PU61</td>
<td>y = 0.059x1</td>
<td>0.94 ***</td>
</tr>
<tr>
<td>PUFA content of feed [g/kg]</td>
<td>E0</td>
<td>y = 3.03-0.06x2</td>
<td>0.56 ***</td>
</tr>
<tr>
<td></td>
<td>E1</td>
<td>y = 24.43-0.24x2</td>
<td>0.49 **</td>
</tr>
<tr>
<td></td>
<td>E2</td>
<td>y = 39.68-0.42x2</td>
<td>0.50 ***</td>
</tr>
<tr>
<td></td>
<td>E4</td>
<td>y = 63.55-0.58x2</td>
<td>0.50 ***</td>
</tr>
</tbody>
</table>

1 Values obtained by analysis of feeds.
2 PU15: 15 g polyunsaturated fatty acids /kg of feed; PU34: 34 g polyunsaturated fatty acids /kg of feed; PU45: 45 g polyunsaturated fatty acids /kg of feed; PU61: 61 g polyunsaturated fatty acids /kg of feed; ε0: without supplementation with α−tocopheryl acetate; E1: supplemented with 100 mg/kg α−tocopheryl acetate; E2: supplemented with 200 mg/kg α−tocopheryl acetate; E4: supplemented with 400 mg/kg α−tocopheryl acetate.

** = P ≤ 0.01; *** = P ≤ 0.001.
tary PUFA increased. The differences in the recoveries of α−Toc may be due to the different level of PUFA in the tissues. Increasing the level of dietary polyunsaturation caused an increase in the accumulation of PUFA in thigh (PU15: 17.91, PU34: 38.54, PU45: 47.70, PU61: 55.66 g/kg) and breast (PU15: 3.48, PU34: 5.39, PU45: 5.98, PU61: 8.48 g/kg). More details about fatty acid content of these tissues are found in CORTINAS et al. (2004). In fact, thigh meat had levels of PUFA that were 7 fold higher than those of breast meat. This is due to the fact that skin was included in the thigh meat samples but was excluded from the breast meat ones. The higher PUFA content of thigh meat, together with its higher oxidative potential corresponds with a higher susceptibility to oxidation, that under the prooxidant conditions during saponification may increase lipid oxidation rate, which is in turn, neutralized by the α−Toc present in the sample. The net result is a reduction of total α−Toc in the sample. Therefore, the degree of polyunsaturation, and in turn, the level of oxidation, may negatively affect the recovery of α−Toc from the samples during the analytical determination. A correct use of a combination of antioxidants (e.g., citric acid + pyrogallol + butylated hydroxytoluene) for the hydrophilic and lipophilic phases during the analysis of α−Toc in meat may be able to avoid or minimize differences in α−Toc recoveries (BOU et al., 2004).

α−Tocopherol content of thigh and breast meat

The α−Toc content of thigh and breast meat, expressed as mg/kg of tissue is shown in Table 5. At dietary levels of 200 mg α−TA/kg of feed, α−Toc content ranged from 14.9 to 37.3 mg/kg in thigh meat and from 9.0 to 17.3 mg/kg in breast meat. Similarly, at this dietary level of α−TA supplementation, some authors found similar α−Toc content in thigh meat with skin (RUIZ et al., 1999; GRAU et al., 2001a,b), as well as in breast meat (GAVIN et al., 1993; MORRISSLEY et al., 1997; O’NEILL et al., 1998).

Furthermore, α−Toc content in thigh meat was 1.8-2 fold higher (P≤0.001) than in breast meat. Despite the fact that in our study thigh meat contained skin, these results agree with other authors who observed lower α−Toc content of thigh meat without skin were 1.1-2.2 fold higher than those of breast meat (LIN et al., 1989; AHN et al., 1995; CHERIAN et al., 1996; DE WINNE and DIRINCK, 1996; NAM et al., 1997; O’NEILL et al., 1998; MALCZYK et al., 1999). Differences among tissues may result from the different metabolic functions of each tissue. So, thigh meat of broilers, apart from having a more developed vascular system (DE WINNE and DIRINCK, 1996; LIN et al., 1989), also have higher lipid content than breast meat (MALCZYK et al., 1999; CRESPO and ESTEVE-GARCIA, 2001). In the animals from the present study, although PUFA expressed as percentage of total fat was higher in breast meat, the absolute amount was higher in thigh meat, since the total fat content of thigh meat was 6.5-7.8% times that of breast (total FA content in breast was 18.1 g/kg whilst in thigh meat ranged between 141.2 and 116.8 g/kg, as the dietary polyunsaturation level increased see CORTINAS et al., 2004). Therefore, this higher content of PUFA may cause a greater need for α−Toc to prevent lipid oxidation. When the protective potential of α−Toc in muscle was estimated based on the total PUFA content, it revealed that the α−Toc concentration in breast meat (139.5, 305.8 and 436.3 mg/100 g of total PUFA for E1, E2 and E4, respectively) was 3-4 fold higher than in thigh meat (51.5, 83.9 and 135.2 mg/100 g of total PUFA for E1, E2 and E4, respectively). These results agree with those of JENSEN et al. (1997) who observed that α−Toc to PUFA ratio in breast meat was 2 fold higher than in thigh meat.

Dietary α−Toc supplementation and polyunsaturation level influenced α−Toc content of thigh and breast meat. In relation to dietary α−Toc supplementation, several authors have reported that α−Toc content in chicken tissues is well correlated with its dietary supplementation (KLAUSS et al., 1995; JENSEN et al., 1999; FLACHOWSKY et al., 2002). Relationships between α−Toc concentration in feed (supplemented as α−TA) and tissues are shown in Table 6. The α−Toc content of thigh and breast meat significantly increased as the dietary α−Toc (values obtained by analysis) increased (P<0.001). Furthermore, the rate of α−Toc deposition was influenced by dietary polyunsaturation level (P<0.01). However, comparing the slopes of the equations (PU15 and PU34 vs. PU45 and PU61), an interaction between the dietary polyunsaturation level and dietary supplementation with α−TA was observed (P<0.01). Thus, linear regression analysis showed that, in the more saturated treatments (PU15 and PU34), α−Toc content in thigh and breast meat increased at a rate of 0.114 mg/kg and 0.071 mg/kg, respectively, for each mg increase of α−Toc per kg feed, whereas in the more polyunsaturated treatments (PU45 and PU61), this increase was 0.02, 0.11 mg/kg and 0.12 mg/kg, respectively. Therefore, the rate of α−Toc deposition in thigh meat was 1.9 to 2.0 fold higher than that in breast meat (P≤0.001). This agrees with the results of LIN et al. (1989) who observed that α−Toc deposition in thigh meat was approximately 50% higher than in breast meat. The different rate of deposition may result, as explained above, from the higher metabolic rate of thigh muscles.

There is a wide range of variability in the α−Toc content of chicken meat obtained by different authors with similar levels of α−TA supplementation. This variation can be due in part to the tissues studied and to the different analytical methods used by each author to determine α−Toc in tissues. Furthermore, most studies do not determine α−Toc in feed, and the real α−Toc content in diet does not always coincide with the estimated α−Toc from dietary supplementation. In addition, probably factors related to the PUFA composition of the diet cause important differences in α−Toc content of chicken meat.

In relation to dietary polyunsaturation level, α−Toc content of thigh and breast meat was reduced as the inclusion of dietary PUFA increased (Table 6). Thus, the decreased rates of α−Toc in thigh meat when PUFA/kg of feed was increased by 1 g were 0.06 mg/kg, 0.24 mg/kg, 0.42 mg/kg and 0.58 mg/kg for E0, E1, E2 and E4, respectively. Decreases of α−Toc in breast meat when PUFA/kg of feed was increased by 1 g was lower in comparison with thigh meat, and were 0.02, 0.11 mg/kg, 0.17 mg/kg and 0.32 mg/kg for E0, E1, E2 and E4, respectively. It has been shown that fat and oil sources varying in the polyunsaturation degree give variable results with respect to α−Toc accumulation in chicken meat. Dietary polyunsaturation entails higher content of PUFA in chicken meat, which is more susceptible to oxidation (GRAU et al., 2001a,b). Data concerning the oxidation level measured as TBARS values of thigh meat of the animals from this experiment are described in CORTINAS et al. (2005). The relative quantities of α−Toc required to protect a fatty acid are higher as the number of double bonds in the molecule increase (WITTING and HORWITT, 1964). For this reason, supplementing chicken diets with fish oil rich in long chain PUFA with a high number of double bonds, produces a reduction in the α−Toc content of chicken tissues (MILLER and HUANG, 1993; HUSVÉTH et al., 2000; SURAI and SPARKS, 2000; ZANNINI et al., 2003). However, using vegetable oils with high levels of PUFA with a lower number of double bonds, did not affect α−Toc content of chicken tissues (CHERIAN et
al., 1996; LAURIDSEN et al., 1997; MALCZYK et al., 1999; RUIZ et al., 1999; GRAU et al., 2001a,b).

Since in the present experiment α-Toc content in thigh meat and breast meat, respectively (y; mg/kg) increased linearly with dietary α-Toc content (x1; mg/kg) and simultaneously decreased linearly with dietary PUFA (x2; g/kg), the multifactorial regressions were calculated for thigh meat with skin, y = x1(0.1473 - 0.0014x2), and for breast meat, y = x1(0.0746 - 0.0007x2) (P ≤ 0.001). The estimated response surface (Figure 1.A) for α-Toc content in thigh and breast meat is shown in Figure 1.

In conclusion, tissue retention of α-Toc varies considerably among tissues, being higher in thigh than in breast meat. α-Toc content in meat increases linearly as dietary α-Toc supplementation increases. Furthermore, as the dietary polyunsaturation level increases, α-Toc content of chicken meat decreases.

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Summary
One hundred and ninety-two female broiler chickens were randomly distributed into 16 experimental treatments (three replicates each) as a result of the combination of 4 levels of dietary polyunsaturated fatty acids (PUFA: 15, 34, 45, 61 g/kg) and 4 levels of supplementation with α-tocopheryl acetate (α-TA: 0, 100, 200 and 400 mg/kg), in order to determine the modification of the α-tocopherol (α-Toc) content of chicken thighs and breast meat. Dietary PUFA content influenced the α-Toc recoveries in thigh with skin and breast muscle tissue. Dietary α-Toc and polyunsaturation level influenced (p<0.001) α-Toc content in meat of chickens at the age of 44 days. α-Toc content of thigh increased linearly as the dietary α-Toc supplementation increased. Thus, it increased at a rate of 0.114 mg/kg (P ≤ 0.001) and 0.071 mg/kg (P ≤ 0.001) when α-Toc increased 1 mg/kg of feed in the most and the least saturated treatments, respectively. Furthermore, α-Toc content of thigh decreased linearly to the inclusion of dietary PUFA. When PUFA content in feed was increased by 1 g/kg, α-Toc content in thigh meat decreased in a rate of 0.06 mg/kg, 0.24 mg/kg, 0.42 mg/kg and 0.58 mg/kg for 0, 100, 200 and 400 mg/kg of dietary supplementation with α-TA, respectively. A similar response was observed in breast meat but with rates of α-Toc incorporation 1.9-2.0 fold lower than in thighs.

Key words
Chicken, nutrition, α-tocopherol, dietary polyunsaturation, thigh meat, breast meat

Zusammenfassung
Einfluss des Grades der Ungesättigung der Fettsäuren im Futter auf den Gehalt an α-Tocopherol im Hühnerfleisch

Das Ziel der vorliegenden Untersuchung war, den Einfluss des Grades der Ungesättigkeit der Fettsäuren im Futter auf den α-Tocopherol-Gehalt von Schenkel- und Brustfleisch von Masthühnern zu bestimmen. Hierzu wurden 192 weiße Broiler zufällig auf 16 Behandlungsgruppen mit je 3 Wiederholungen verteilt. Die Behandlungen waren: 4 Grade der Ungesättigkeit (PUFA: 15, 34, 45, 61 g/kg), 4 Zulagestufen für α-Tocopherol (α-TA: 0, 100, 200, 400 mg/kg).

Die PUFA-Gehalte im Futter beeinflussten die Aufnahme von α-Toc in das Schenkel- (mit Haut) und in das Brustfleisch. Der Gehalt an α-Toc im Futter und der Grad der Ungesättigkeit hatten einen hoch signifikanten Einfluss auf den α-Toc-Gehalt im Fleisch der 44 Tage alten Broiler (P<0,001). Der α-Toc-Gehalt im Schenkelfleisch nahm mit der Zulage im Futter kontinuierlich zu. Beim geringsten und höchsten Grad der Ungesättigkeit des Futters erhöhte sich der Gehalt im Schenkelfleisch um 0,114 mg/kg (P≤0,001) bzw. 0,071 mg/kg (P≤0,001) je g α-Toc-Zulage zum Futter. Ferner ging der α-Toc-Gehalt im Schenkelfleisch linear mit zunehmendem PUFA-Gehalt im Futter zurück. Jede Erhöhung des PUFA-Gehaltes im Futter um 1 g/kg führte zu einem Rückgang im α-Toc-Gehalt des Schenkelfleisches in der Höhe von 0,06 mg, 0,24 mg, 0,42 mg und 0,58 mg/kg für die α-Toc-Zulagen von 0, 100, 200 und 400 mg/kg Futter. Für das Brustfleisch wurden ähnliche Beziehungen beobachtet, wobei die Einlagerungsrate von α-Toc 1,9 bis 2-fach geringer war.

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Stichworte
Broiler, Fütterung, α−Tocopherol, PUFA, Schenkelfleisch, Brustfleisch

References


Jensen, S.K., R.M. Engberg and M.S. Hedemann, 1999: All-β−α−tocopherol acetate is a better vitamin E source than all-α−α−tocopherol succinate for broilers. J. Nutr. 129, 1355-1360.

Klauss, A.M., H. Fuhrmann and H.P. Sallmann, 1995: Per− oxidative and antioxidative metabolism of the broiler chicken as influenced by dietary linoleic acid and vitamin E. Arch. Geflügelk. 59, 135-144.


Miller, E.L. and Y.X. Huang, 1993: Improving the nutritional value of broiler meat through increased n-3 fatty acid and vitamin E content. Pages 404-411 in Proceedings of the 11th European Symposium on the Quality of Poultry Meat. Tours, France.


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