

## Studies on fat in some chicken products with trial to prevent its oxidation

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**SUMMARY:** A total of 100 samples of frozen half cooked chicken nuggets and shawerma were collected from grocery market in Giza and Cairo governorates (50 of each). The samples were subjected to chemical analysis (Fat %, Cholesterol, g / 100 gm, free fatty acids Acid value %, peroxide value m Eq/ kg and TBA value mg MD/ Kg). The mean values of the aforementioned parameters for nuggets and shawerma were  $17.78 \pm 0.36$ ,  $64.32 \pm 0.94$ ,  $1.88 \pm 0.14$ ,  $0.95 \pm 0.07$ ,  $10.13 \pm 0.52$ ,  $0.71 \pm 0.052$  and  $15.3 \pm 0.28$ ,  $57.98 \pm 0.69$ ,  $1.85 \pm 0.13$ ,  $1.099 \pm 0.17$ ,  $10.912 \pm 0.064$ ,  $0.74 \pm 0.035$  respectively. There were significant decrease of fat % and cholesterol (mg/ 100 gm) at  $P < 0.01$  in chicken shawerma lower than chicken nuggets. Due to **ES (2005) and Codex Standard (1991)**, the rejected samples percent of both nuggets and shawerma were 40 and 43 which exceeded the recommends fat percent. On the other hand, the percent of rejected nuggets and shawerma due to the fat oxidation criteria exceeding limits (Acid value, free fatty acids, peroxide value and TBA) were 6 and 8 respectively. There is significant increase in shelf life at  $P < 0.01$  in the treated chicken nuggets and shawerma with ascorbic more than untreated sample when stored at (-18°C). The results were statistically analysed and it was recommend to use natural antioxidant for preventing the fat oxidation and extending the shelf life during freezing storage.

[El- Shater M.A; Hanan, G. seadawy; and Maha, M. Mohamed. **Studies on fat in some chicken products with trial to prevent its oxidation.** *N Y Sci J* 2012;5(10):56-63]. (ISSN: 1554-0200). <http://www.sciencepub.net/newyork>. 10

**Key wards:** Fat oxidation -Chicken products -Antioxidants

### INTRODUCTION

Egypt's modern poultry industry began in 1964 with establishment of the national General poultry company (G.P.C) which aimed to provide Egypt's fast growing human population with high quality a fordable animal protein. Another important goal for G.P.C was the transformation of poultry production into an industry rather than an agricultural activity, through the introduction of modern technology of poultry processing and skilled mangent to get different chicken product as chicken shawerma, chicken nuggets etc. (FAO, 2006).

Chicken nuggets are one of chicken products consists of chicken meat with seasoning with different shape (each piece  $16 \pm 29$ ), preferred (half cooked) with light golden brown and breaded. It must be free from banned antibiotics, pesticides, residues and sanitizer residues, the product is stored at -18°C (Venky's, 2010).

Shawerma is a well known precooked product in Middle East countries that from beef or chicken meat with animal fat the product kept in freezing state at (-18°C) fat (Khalid, 2002).

For poultry, pre-cooking also represents a "Pasteurization" process to improve the microbiological status-relative particularly to salmonella SPP. Many products are cooked to a center temperature above 70°C. The meat - products

uses an enormous range of cooking methods but for poultry the use of fryer and ovens predominant (Singhal *et al.*, 2001).

Freezing is as excellent process for preserving the quality of poultry and poultry products as well as other meat for long periods. However, deterioration of quality caused by chemical pr physical factors can occur. Many studies have shown that lipid oxidation is one of the primary causes of quality losses of frozen stored poultry and meat products (Gökalp *et al.* 1983).

The most important process in the oxidation of lipids in meat and poultry products is the peroxidation of polyunsaturated fatty acids from cell membrane. The major catalysts are trabsitional metalions, such as  $Fe^{2+}$  and  $Cu^{+}$ . Heam compounds of meat can also contribute to this process due to the participation of heam-iron in accelerating lipid peroxidation (Decker and Welch 1990; Pikul 1992).

During food processing and storage, polyunsaturated fatty acids tend to be oxidized. Cholesterol can be oxidized by the same mechanism as fatty acids. Therefore, lipids radicals formed during the processing and storage foods can accelerate the oxidation of cholesterol and produce cholesterol oxidation products (COPs) (Chan *et al.*, 1993).

Aerobic packaging significantly increased cholesterol oxidation products (COPs) and thiobarbituric acid relative substances (TBARS) in cooked poultry and meat products after 7 days storage which were closely related to the fatty acid composition of food (Du *et al.*, 2001).

Acid value test measures free fatty acid as an indication of hydrolytic rancidity. Peroxide value is one of the most widely used tests for oxidative rancidity, peroxide value is a measure of the concentration of peroxides and hydroperoxides formed in the initial stages of lipid oxidation. The 2 - thiobarbituric acid (TBA) method is the most widely used test for measuring the extent of lipid oxidation in muscle foods. The test is believed to measure the breakdown products of unsaturated fatty acids oxidation. Saturated aldehydes, 2 - enals, and 2 - dienals, produced in the termination phase of lipid oxidation, can be detected by reaction with 2 - thiobarbituric acid (Haimd, 2011).

Biologically active compounds can be destroyed and, in some cases, toxic and carcinogenic substances represents by hydroperoxides, acids, etc. accumulate (Balev *et al.*, 2005).

The meats are important sources of fat in the typical diet in the world, many consumers believe that red meat is unhealthful, because is high in saturated fatty acids (SFA) and cholesterol. Recently, it has been recently demonstrated that replacement of red meat with chicken is associated with significant decrease in a polypoprotein B and total cholesterol levels in microalbumin uric type 2 diabetic patients this effect is probably related to the higher PUFA (polyunsaturated fatty acids). Content of chicken meat in comparison to beef (Gross *et al.*, 2002).

This study was planned to study:-

Part I: Chemical criteria of market half cooked frozen chicken nuggets and shawerma including.

- 1- Fat %.
- 2- Cholesterol (mg / 100 gm).
- 3- Fat Lipolysis.
  - 3-1- Free fatty acid % as oleic.
  - 3-2- Acid value % as oleic.
- 4- Fat oxidation criteria.
  - 4-1- Peroxide value m Eq/ kg.
  - 4-2- Thiobarbituric acid value (TBA mg MD/ kg).

Part II

Trial to prevent the lipolysis and fat oxidation through addition of Ascorbic acid as antioxidant (natural) to experimentally produced chicken nuggets and shawerma, then stored at -18 with control samples (without Ascorbic acid).

The experimentally produced chicken nuggets and shawerma (treated with ascorbic acid) and control chicken nuggets and shawerma (untreated with ascorbic) were examined every 2 weeks till spoilage for free fatty acids, Acid value, peroxide value and TBA value.

## MATERIALS AND METHODS

### Par I:-

A total of 100 samples of frozen chicken nuggets and shawerma (50 of each) were collected from market of Giza and Cairo governorates. The samples were transported to the laboratory in ice box without delay for the following examination.

#### 1. Determination of fat %.

The test was carried out according to (AOAC, 2000), in which the traditional technique is the quantitative determination of fat through a solvent extraction with petroleum ether. The soluble material is extracted from dried test samples of sample by a two - step treatment with petroleum ether solvent (Soxhlet extraction procedure).

#### 2. Determination of cholesterol content:

Cholesterol was determined according to Ojiako and Akubugwo (1997) using spectrophotometer. Only 0.1 ml of extracted fat and standards cholesterol dissolved in chloroform in ratio 1: 10 was evaporated to dryness in water bath at 50 °C. Glacial acetic acid (3 ml) and 3.0 ml color reagent was added to each sample and standards, then shaking vigorously to dissolve the sample. Blank contained 2.0 ml chloroform, 3 ml Glacial acetic acid and 3 ml of color reagent was prepared.

After cooling for 30 min. at room temperature, absorbance of blank sample and standards were measured at 560 nm. The cholesterol content in sample were calculated from the standard curve.

#### 3. Determination of Lipolysis criteria as follow:

##### 3.1. Acid Value:

It was carried out according IUPAC (1979) using titration technique against alkali (1 - 10 gm dissolved fat). The acid value was calculated according to the equation.

$$\text{Acid value (mg KOH/g)} = \frac{\text{Titration (ml)} \times 5.61}{\text{Wt of sample used}}$$

##### 3.2. Free fatty acids:

The official titration with alkali method recommended by ISO (1980) (Reference method) was used the F.F.A figure in usually calculated as oleic (percent).

#### 4. Determination of Fat oxidation criteria:

##### 4.1 Peroxide Value:

The sample was extracted for fat. To quantify peroxide value, (0.01- 0.05g) was placed in 10 ml screw capped test tube and dissolved in: 1 ml chloroform / acetic acid (2: 3), with addition of 100 ml Fe (II) solution, mix for is sec. on vortex mixer and left it in dark place for 10 min. Deionized water (2 ml) was added and 4 ml of diethyl ether (containing 7 pp BHT). One ml of aqueous phase was mixed with 100 ml of saturated ammonium thiocyanate solution. The sample was measured against a water blank at wave length 470 nm. after 10 min.

The peroxide value was measured using calibration curve. For calibration, a set of Fe (III) concentration in range 0 - 10  $\mu\text{g} / \text{m}$  the method was carried out according to **Shantha and Decker (1994)** the P.V was expressed as mEq / kg.

##### 4.2 Thiobarbituric acid value:

The method due to **Tarlidgis et al (1960)** was used, in which the reaction is applied to a distillate produced under standardized conditions from an acidified macerated food. The results are expressed as malonaldehyde by reference to a standard graph prepared by using 1, 1, 3, 3 - tetra ethoxy propane which yields malonaldehyde by acid hydrolysis

TBA no (as mg malonaldehyde/ Kg) = 7.8 D where S (absorbance against blank at 358 nm).

#### Part II:

##### 1. Production of chicken Nuggets experimentally:

It was carried out according to **Yavas and Bilgin (2010)** using fresh poultry breast meat in addition to seasoning (control), the other group was treated with antioxidant (Ascorbic acid 500 ppm).

The samples (control and treated) were fried in vegetable oil for 2 - 3 min (half cooked). The control and treated chicken nuggets were put in suitable polyethylene bags then put in carton package.

##### 2. Production of chicken shawerma experimentally:

It was carried out according to **Odu and Akamo (2012)**. Fresh bonless chicken breast was sliced, put in a pan, seasoned with salt, curry, thyme, black paper, nutmeg and garlic (control). The treated sample was made as mentioned in addition to (500 ppm) of Ascorbic acid was added. The control and treated prepared shawerma were heated for 10 min (half cooking) after addition of water and vegetable oil. After cooling, control & treated shawerma were wrapped as mentioned in chicken nuggets.

#### 3. Storage of Experimental samples (chicken Nuggets and chicken shawerma):

The Experimental samples were kept at - 18°C and examined every 2 weeks for lipolysis and fat oxidation criteria till spoilage of the products appear as mentioned before.

#### Statistical Analysis:

Data obtained were statistically analysed using **SPSS14 (2006)**.

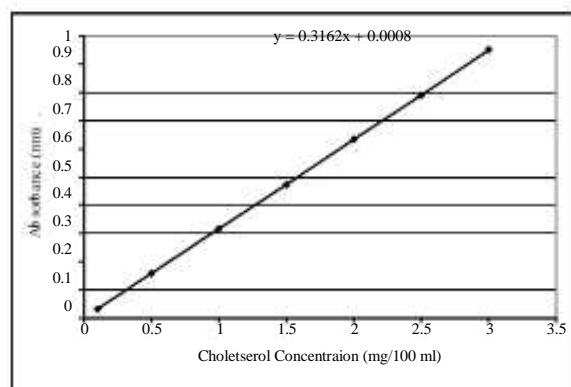


Fig. (1): Cholesterol standard curve.

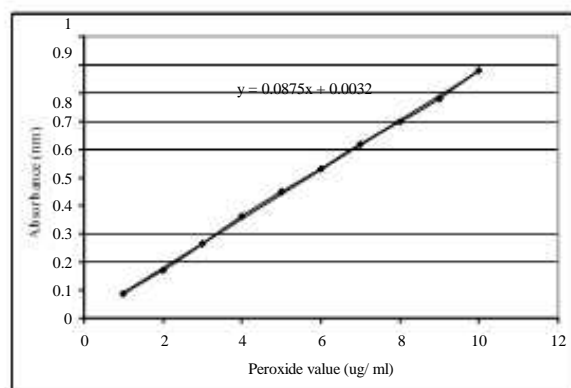


Fig. (2): Peroxide value standard curve.

**RESULTS**

Table (1): Statistical analytical results of chemical analysis (lipolysis and fat oxidation) of market frozen chicken nuggets and shawerma (n = 50 of each).

|                  |          |      | Acid value | Free FA | Peroxide Value | TBA    |
|------------------|----------|------|------------|---------|----------------|--------|
| Chicken Nuggets  | Rejected | No   | 3          | 3       | 3              | 3      |
|                  |          | %    | 6          | 6       | 6              | 6      |
|                  | Accepted | No   | 47         | 47      | 47             | 47     |
|                  |          | %    | 94         | 94      | 94             | 94     |
|                  | Min      |      | 1.02       | 0.56    | 6.2            | 0.52   |
|                  | Max      |      | 5.9        | 2.95    | 23             | 1.44   |
|                  | Mean     |      | 1.8832     | 0.9512  | 10.1284        | 0.7104 |
| SE               |          | 0.14 | 0.07       | 0.52    | 0.025          |        |
| Chicken Shawerma | Rejected | No   | 3          | 5       | 4              | 4      |
|                  |          | %    | 6          | 10      | 8              | 8      |
|                  | Accepted | No   | 47         | 45      | 46             | 46     |
|                  |          | %    | 94         | 90      | 92             | 92     |
|                  | Min      |      | 1.1        | 0.55    | 6              | 0.5    |
|                  | Max      |      | 6          | 9       | 28             | 1.9    |
|                  | Mean     |      | 1.8504     | 1.0988  | 10.912         | 0.7404 |
| SE               |          | 0.13 | 0.17       | 0.064   | 0.035          |        |

Table (2): Fat and cholesterol percent of market frozen chicken nuggets and shawerma.

|          |    | Chicken Nuggets |                         | Chicken Shawerma |                         |
|----------|----|-----------------|-------------------------|------------------|-------------------------|
|          |    | Fat (g/100 gm)  | Cholesterol (mg/100 gm) | Fat (g/100 gm)   | Cholesterol (mg/100 gm) |
| Rejected | No | 42              | 0                       | 43               | 0                       |
|          | %  | 84              | 0                       | 86               | 0                       |
| Accepted | No | 8               | 50                      | 7                | 50                      |
|          | %  | 16              | 100                     | 14               | 100                     |
| Min      |    | 14              | 55                      | 12               | 50                      |
| Max      |    | 24              | 79                      | 19               | 70                      |
| Mean     |    | 17.78           | 64.32                   | 15.3*            | 57.98*                  |
| SE       |    | 0.363           | 0.936                   | 0.282            | 0.693                   |

\*Significant at P < 0.01 using t-student test comparing with Chicken Nuggets

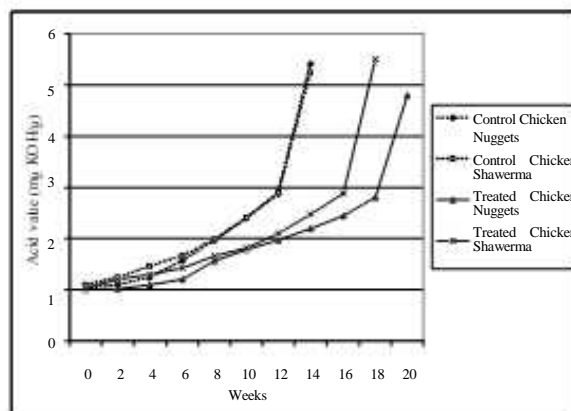


Fig. (3): Changes of acid value of chicken nuggets and shawerma during freezing storage at (-18°C).

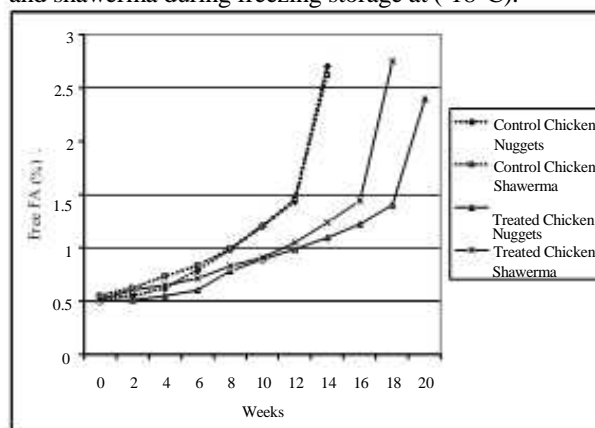


Fig. (4): Changes of free fatty acids of chicken nuggets and shawerma during freezing storage at (-18°C).

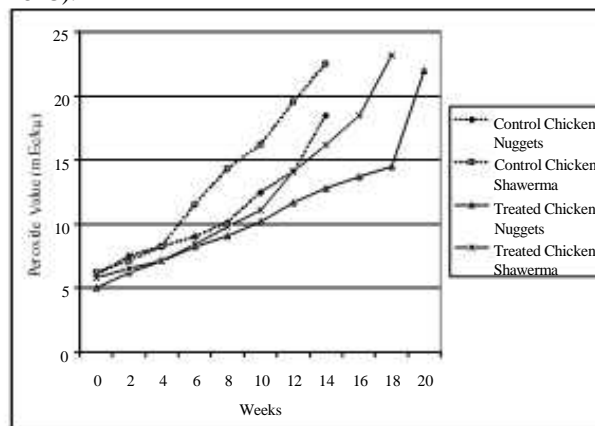


Fig. (5): Changes in peroxide value of chicken frozen nuggets and shawerma during freezing storage at (-18°C).

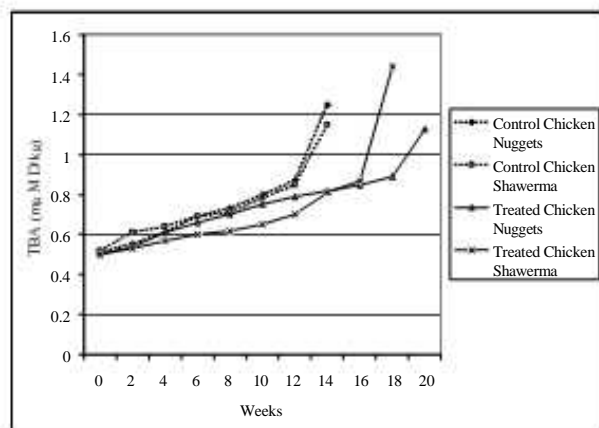


Fig. (6): Changes in TBA value of chicken frozen nuggets and shawerma during freezing storage at (-18°C).

## DISCUSSION:

In present study, the market sample of nuggets had mean values of fat % and cholesterol (mg / 100 gm) were  $17.78 \pm 363$  &  $64.32b \pm 0.936$  while the mean values for chicken shawerma (frozen) were  $15.3 \pm 0.282$  and  $57.98 \pm 0.693$  respectively the fat percent and cholesterol content (mg/ 100 gm) of frozen chicken shawerma were significantly lower than frozen chicken nuggets at  $P < 0.01$ .

According to **ES (2005)** concerning the fat %, it was found that 8 samples of chicken nuggets (16%) and 7 samples of chicken shawerma (14%) were only accepted due to maximum fat recommended by **ES (2005)** table (2).

Microbial activities are considered the major factor of foods alteration during storage and manipulation, when these activities are under efficient control, the foods alteration will be of chemical nature, lipid oxidation is a principal chemical changes of foods, which depends on the level of oxygen, (energy/ hear) and metals. Lipid oxidation products are responsible for the development of rancidity by the production of low molecular weight compounds that cause undesirable flavor (**Frankel, 1985; Frankel et al., 1987**), thus affecting the quality and limiting shelf life of food products. Cholesterol is also oxidized in similar reaction mechanisms to those observed of fatty acids. Many of the cholesterol oxidation products have adverse effect such as cytotoxicity and modifications of enzyme activity (**Erickson et al., 1978; Sevanin and Petreson, 1986; Bosinger et al., 1993**) atherosclerosis (**Kumar and Singal, 1991**) carcinogenicity and mutagenicity (**Ansari and Smith, 1979**).

In this study there was positive correlation between the fat content in both chicken nuggets and

shawerma and their cholesterol content. This result was in accordance with that reported by **Dinh et al (2008)**.

Major sources of cholesterol in the human diet are meat from domestic livestock, although seafood is also rich in cholesterol content in this study in both nuggets and shawerma of chicken ranged from (55-79) and (50-70) mg / 100 g. Nearly similar results were reported by (**Chizolini et al. 1999; Mourot and Hermier 2001; Pironen et al., 2002; Valsta et al., 2005; Bragagnolo, 2009** and **Honikel, 2009**) who recorded that the cholesterol content in poultry products ranges from 40 - 90 mg / 100 gm. Higher concentration of cholesterol content was recorded by (**USDA, 2011**) for cooked chicken dark meat.

Table (1) illustrated the acid value, free fatty acids, peroxide value and TBA for both chicken nuggets and chicken shawerma. The chicken nuggets mean values of the for mentioned parameters were  $1.88 \pm 0.14$ ,  $0.95 \pm 0.07$ ,  $10.13 \pm 0.52$  and 0.71, while for chicken shawerma were  $1.8504 \pm 0.13$ ,  $1.1 \pm 0.17$ ,  $10.91 \pm 0.064$  and  $0.74 \pm 0.35$  respectively.

According to the maximum TBA (mg MD/kg) recommended by **ES (2005)** (0.9 mg MD/ kg). In was found that 3 samples (6%) of chicken nuggets and 4 samples (8%) of chicken shawerma were rejected due to rancidity.

The acid value and the free fatty acid of control and treated chicken nuggets and shawerma were shown in Figs (3 & 4).

The treated chicken nuggets and chicken shawerma exceeded the acid value recommended by codex standard limit at the weeks 20, 18 weeks respectively.

It was noticed that the control chicken nuggets and shawerma were within limit (0.5 - 1.2) percent as oleic recommended by **IUPAC (1979)** till the week (12) of frozen storage while treated nuggets and shawerma till the week 18 and 16 respectively.

Free fatty acids are the products of enzymatic or microbial degradation of lipids and determination of FFA gives information about the stability of fat during storing storage, this agrees with reported by (**Das et al., 2008**). The free fatty acids of antioxidant treated nuggets and shawerma were significantly lower than control (Non-treated) throughout the storage period, this result agreed with that reported by (**Yavas and Bilgin 2010**).

The experimentally produced chicken nuggets and chicken shawerma (with antioxidant) were stored with experimentally control (without antioxidant) at -18°C till spoilage criteria of lipolysis and fat oxidation appear. The acid value of chicken nuggets and chicken shawerma of both experimentally control and treated at zero day were

1.04, 1, 1.1, 1.04 respectively. The acid value of chicken nuggets and chicken shawerma of control samples exceeded the maximum limit recommended by **Codex (1999)** (max 4.0 mg KOH/ g fat) at week (14).

The initial peroxide value of the control and treated chicken nuggets and shawerma stored at -18°C at zero week were 6, 5 and 6.2 and 5.8 respectively. The peroxide value gradually increased throughout the storage period. At weeks 2, 4, 6, 8, 10, 12 and 14, the peroxide value of chicks nuggets control and treated were 7.5, 8.2, 9.5, 10.1, 12.5, 14.1, 18.5 and 6.2, 7.1, 8.2, 9.1, 10.2, 11.7, 12.8 respectively. It was noticed that 4th week the peroxide value of the control nuggets exceeded the maximum limit recommended by **Codex Standard (1999)** (max. 12.5 millieq of active O<sub>2</sub>/ kg). The treated nuggets exceeded this limit at 20 week.

On the other hand, the peroxide value of chicken shawerma exceeded the maximum limit for control at the 10<sup>th</sup> week while at 16<sup>th</sup> for treated shawerma (Fig., 5). The peroxide value is the most commonly used parameter to measure of lipid hydroperoxides. It also named primary lipid oxidation products. Chicken products have considerably fat and particularly vulnerable to lipid oxidation which leads to quality and nutritional value deterioration (**Olafsdottir et al., 1997; Yimaz, 1998**). The peroxide value were significantly increased at  $P < 0.05$  during storage period in both nuggets and shawerma, this result in current study is in accordance with that of **Sallam (2007) and Yavas and Bilgin (2010)**. Nutritionist and buyers had arbitrarily established maximum peroxide value levels of between 5 - 20 meq. O<sub>2</sub>/ kg of fat (**Hamilton and Kirestein, 2008**).

The TBA (mg MD/ kg) of control nuggets and shawerma at weeks 0, 2, 4, 6, 8, 10, 12, 14, were 0.5, 0.22, 0.61, 0.69, 0.73, 0.8, 0.87, 0.121 and 0.52, 0.61, 0.64, 0.69, 0.71, 0.78, 0.85, 1.15 respectively. The TBA exceeded the maximum limit (0.9 mg MD/ kg) recommended by **ES (2005)** at 14th week in both control chicken nuggets and shawerma.

The use of antioxidant (Ascorbic acid) extend the shelf life of treated chicken nuggets and shawerma to 18th, 16th weeks at which the TBA were with the permissible limit recommended by **ES (2005)**. At the week 20 and 18 the treated chicken nuggets and shawerma had TBA 1.13 and 1.44 mg MD/ kg which exceeded the maximum limit of TBA respectively (Fig., 6). The treatments of chicken nuggets and shawerma had statistically significant effect ( $P < 0.05$ ) on TBA value during storage at -18°C. The lowest TBA in treated chickens nuggets and shawerma with ascorbic were 0.5 mg/ MD/ kg for each, while the highest TBA values were 1.13 and

1.44 mg MD/ kg. TBA values were significantly lower ( $P < 0.05$ ) in all treated samples of nuggets and shawerma samples as compared to the control. This results agreed with that reported by **Ozgul and Cemalettin (2010)**.

In industrial processing, mainly synthetic antioxidant such as butylated hydroxytolvene, butylated hydroxyinsol and propylated gallate are used prolong the storage stability of food. However the demand of natural antioxidants has recently increased because of natural antioxidants has recently increased because of the toxicity and the carcinogenicity of synthetic antioxidants (**Juntachote et al 2006**). In addition, synthetic antioxidants have limited application because of their low water solubility (**Bekhit et al, 2003**). BHA, however, has a high solubility in animal fats (**Allen & Hamilton 1994; Coppen 1994**). In meat processing, the Ascorbic acid is used as antioxidant (natural antioxidant) for preventing the off - flavor (**Bauernfiend 1985**) as well as maintaining the cured colour.

The anti-oxidation mechanism of ascorbic acid is to activate the primary antioxidants, to be an oxygen scavenger, and to inactivate antioxidants (**Bauernfiend and Pinkert, 1970**). At low concentrations, ascorbic acid promotes lipid oxidation; however, at high concentrations it inhibits lipid oxidation (**Decker and XU, 1998**).

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8/14/2012