FORTIFICATION OF BISCUITS WITH IRON FROM NATURAL SOURCES

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ABSTRACT

Iron deficiency anemia is considered one of prevalent patients in developing countries, whereas it is well known that wheat flour is deficient in iron, hence, in this study wheat flour (72%) extraction fortified with celery seeds and cinnamon bark meal individually as a natural sources of iron at levels 5, 7.5 and 10 g/100 g wheat flour and preparation of biscuit samples. Iron content in wheat flour, celery seeds and cinnamon were determined. Biscuit samples were sensory evaluated and baking quality tested. Total iron and available iron were determined in biscuit samples. Biological evaluation for experimental rats designed and histopathological examination was tested for heart organ of rats. The results showed that wheat flour, celery seeds and cinnamon contained from iron 1.98, 57 and 50 mg/100 g respectively. Total iron and available iron increased in biscuits samples by increasing celery seeds and cinnamon additives compared with unfortified biscuits (control). Sensory evaluation of biscuit samples showed slight decrease in color, crunchiness and appearance while odor and taste significantly improved by increasing celery seeds and cinnamon additives compared with control. Baking quality of biscuit samples indicated increasing in weight, while volume, diameter and thickness slightly decreased by increasing celery seeds and cinnamon additives compared with control. Biological evaluation revealed that mean values of hemoglobin, hematocrit, red cell count, mean corpuscular volume, mean corpuscular hemoglobin, mean corpuscular hemoglobin concentration, serum iron and serum ferritin significantly improved after 8 weeks in groups rats fed on biscuits fortified with celery seeds and cinnamon compared with control. Histopathological overhaul declared amelioration in organ heart for groups rats fed on biscuits fortified with celery seeds and cinnamon compared with anemic control.

INTRODUCTION

Iron is an essential element has several vital function in the body. It serves as a carrier of oxygen to the tissues, the lungs by red blood cell hemoglobin as a transport medium for electrons within cell and as an integrated part of important enzymes in various tissue. Disturbance of iron metabolism are part of the metabolic syndrome which clusters insulin resistance, hyperinsulinemia, hyperglycemia, hypertension and obesity (Sandrine et al 2008).

Iron is an important mineral for our body, primarily used in the formation of hemoglobin which in turn takes oxygen to different parts of body with the help of feritin and transferrin proteins. Thus, iron carries oxygen from the lungs through red blood cells and distributes it to the rest of the body’s muscles and tissue’s (Ekhard et al 2009).

Iron deficiency is common in developing countries, where the diet is largely cereal-based and contains little animal protein and it will take a long time for dietary changes to result in improvements in iron status. One way to ensure the adequate iron nutrition of a population is to fortify commonly consumed foods with iron and provide the foods in relatively predictable amounts. Food fortification and supplementation are generally considered the best approaches for combating iron deficiency in a population (Laura and Beard 2009).

Iron deficiency can be defined as occurring when the body’s iron stores become depleted and a restricted supply of iron to various tissues becomes apparent. The clear consequences of iron depletion are a reduction in oxygen transport ca-
capacity and a reduction in oxidative capacity at the cellular level of functioning. The process by which iron stores are depleted may occur rapidly or very slowly and depends on the balance between iron intake and iron requirements (Barbara et al. 2009).

Iron deficiency is the most common single nutrient deficiency disease in the world and is a major concern for ~15% of the world’s population. The commonly used definition for anemia, regardless of its cause, is a low hemoglobin concentration. If iron deficiency is an underlying etiology, then by definition an individual must have depleted iron stores, low ferritin in plasma or decreased stainable iron in bone marrow and inadequate delivery of iron to tissues as characterized by a low transferrin saturation, a high erythrocyte protoporphyrin concentration and an elevated transferring receptor concentration (Zhenyu et al. 2008).

Celery seeds or one of its extractives used in the flavoring of beverages, confections, chewing gums, ice creams, pizza loaf and baked good. Ground cinnamon oil and/or the oleoresin of cinnamon is used commercially in the manufacture of confection, ice cream, beverages, chewing gum, cakes, cookies, pies and other baked good. Moreover, celery seeds and cinnamon rich content in iron (Farrell, 1999).

The intention of this study was carried out to produce biscuits fortified with celery seeds and cinnamon as a natural sources of iron and studying chemical composition and iron content in raw materials, moreover, investigation sensory evaluation and baking quality of biscuits samples, furthermore, evaluation iron blood picture for experimental rats fed on biscuits.

MATERIALS AND METHODS

Materials

Celery seeds (Apium graveolens L.) and cinnamon (Cinnamomum zeylanicum Nees) were obtained in 2009 from Medicinal and Aromatic Plants Research Department, Agriculture Research Center, Giza, Egypt.

Wheat flour (72% extraction) was obtained in 2009 from North Cairo Mills Company, El-Hoda Mill, Shobra El-Kheima, Egypt.

Vitamins mixture were obtained from Sigma Chemical Company, USA.

Minerals mixture were obtained from ADWIC Company Egypt.

Kits were purchased from Randox laboratories LTD. USA. (70 Dorman St., Suite 3 A San Francisco, CA 94124).

Sixty-four male albino rats of wistar strain average weight 85±5 g were obtained from the Laboratory Animal Colony, Vaccine and Immunity Organization, Cairo, Egypt.

Methods

Preparation of celery seeds powder

Celery seeds were milled via laboratory mill (Retsch GmbH5657HAAN) to get the fine celery seeds powder.

Biscuits preparation

Biscuits samples were prepared according to the method of Wade (1988). Biscuits formula was consists of 100 gm wheat flour (72% extraction rate) control or fortification with 5, 7.5 and 10 gm celery seeds and cinnamon, 40 gm sugar, 12 gm butter milk, 18 gm water, 20 gm milk, 3 gm baking powder, 25 gm egg and 1 gm vanilla. Butter, sugar and vanillia were mixed in a dough mixer using the flat beater for 1 minute, then scraped down and continued to mix for 3 minutes at high speed. When flour and baking powder were added to the mixture and mixed at low speed then it was sheeted to 15 mm thickness. Circle pieces cut of dough were formed by using of templates with an outer diameter of 45 mm. The biscuits were baked at 180°C for 12 minutes. Biscuit samples were cooled and packed in bags made from low density polyethylene (Density 0.915g/cm³).

Sensory evaluation

The color, taste, odor, appearance and crunchiness of the biscuit samples were evaluated according to method of Wade (1988).

Baking quality

The baking quality of biscuits test included weight, volume, specific volume, diameter, thickness and expansion factor were determined according to methods of AACC (2002).

Chemical analysis

Moisture, protein, fat, fiber, ash, total iron, available (soluble) iron, phytate phosphorus and tannins were determined according to the methods of AOAC (2005).
Fortification of Biscuits

Biological evaluation

Sixty-four male rats were housed individually in stainless steel cages with wire mesh floor. Distilled water and diets were offered ad libitum. Baseline mean of hemoglobin was found to be about 13.14 g/dl. After 3 days adaptation period rats were divided randomly into eight groups (n = 8 rats) as follow: Group 1 fed on basal diet consist of 10% casein, 10% corn oil, 4% salts mixture, 1% vitamins mixture, 5% cellulose and 70% starch according to AIN (1977) and considered as normal control. Fifty six rats were fed on basal diet free from iron for four weeks until the hemoglobin level decreased to about 9 – 10 g/dl (anemia state). Group 2 fed on basal diet free from iron until end of experimental and considered anemic control. Group No. 3, 4 and 5 fed on biscuits fortified with 5, 7.5 and 10 g/100 gm celery seeds respectively. Group No. 6, 7 and 8 fed on biscuits fortified with 5, 7.5 and 10 g/100 gm cinnamon respectively. After 8 weeks blood was withdrawn from the orbital puncture by a heparinized capillary tube containing heparin as a blood anticoagulant then, blood was centrifuged at 3000 r.p.m (10 min) to obtain the serum which was stored at (4°C) for biochemical analysis. On the final day of the experiment the rat were fasted overnight and anesthetized by using overexposure to diethyl ether then rats were sacrificed by decapitation.

Biochemical analysis

Hemoglobin (Hb): To 20 µl of the whole blood sample, 5 ml of cyaname thenoglobin reagent (Egle diagnostics) was added. After standing 3 min at room temperature, the formed colour was measured at 540 nm against reagent blank (cyaname thenoglobin) using Spiclo Colorimeter. Hb concentration (grams per deciliter) was calculated according to Wintrobe (1965) using the following equation:

\[ Hb \text{ (g/dl)} = \frac{\text{optical density of sample}}{\text{optical density of standard}} \times 36.77 \]

Hematocrit (Ht): Hematocrit determination was done according to Dacie and Lewis (1984) using the micro-method. Heparinized microhaematocrit tubes were used, filled with capillary blood, sealed and centrifuged at 3000 rpm for 5 min.

Red blood cell count (RBCs): Red blood cell were counted on hemocytometer using a light microscope at 40X magnification after diluting the blood samples 200 times with a physiological saline solution (0.9% sodium chloride solution) before counting (Heplar, 1966).

Red corpuscular values: Absolute red cell values were calculated according to Dacie and Lewis (1984) as follows:

- Mean corpuscular volume (MCV) = [hematocrit (%)] \times 10^2 + RBCs (millions/mm^3) as femtoliter (fl).
- Mean corpuscular hemoglobin (MCH) = [Hb (g/dl)] \times 10 + RBCs (millions/mm^3) as pico gram (pg).
- Mean corpuscular hemoglobin concentration (MCHC) % = [Hb (g/dl)] \times 100.

Serum iron (SI) was measured according to method of Dacie and Lewis (1984).

Serum ferritin (SF) was measured by enzyme linked immunosorbent assay according to method of Flowers et al (1986).

Histopathological examination

The heart was kept in 10% formalin vatile emmbeded in paraffin wax, then heart sectioned at the thickness of 6µ and stained in haematoxyli and eosin mixture according to Yoon et al (2001). Tissue section heart examined using ordinary microscope at 200 magnification.

Statistical analysis

The obtained results were analyzed using Statistical Analysis System SAS (2001).

RESULTS AND DISCUSSION

Chemical composition of raw materials

An adequate knowledge of the chemical composition of food is vital to the health, well-being and safety of the consumer.

Raw materials (wheat flour, celery seeds cinnamon) were chemically analyzed and the results were illustrated in Table (1). The results revealed that the moisture, protein, fat, fiber, ash and carbohydrates contents in wheat flour (72%) extraction were 10.95, 10.38, 1.10, 0.65, 0.50 and 87.37% respectively. From the same table it is obvious that the moisture, protein, fat, fiber, ash and carbohydrates contents in celery seeds were 8.45, 17.98, 2.91, 3.22, 3.15 and 72.74% respectively. It can be seen from the same table that the moisture, protein, fat, fiber, ash and carbohydrates contents in cinnamon were 9.83, 4.12, 2.84, 4.81, 2.90 and 85.33% respectively.

Arab Univ. J. Agric. Sci., 18(2), 2010
Table 1. Chemical composition (%) of raw materials (on dry weight basis)

| Samples          | Moisture   | Protein     | Fat       | Fiber    | Ash       | Carbohydrates* |
|------------------|------------|-------------|-----------|----------|-----------|----------------|
| Wheat flour (72%)| 10.95±0.81a| 10.38±0.73b | 1.10±0.28b| 0.65±0.19c| 0.50±0.11b| 87.37±1.23a    |
| Celery seeds     | 8.45±0.96c | 17.98±0.65a | 2.91±0.42a| 3.22±0.23b| 3.15±0.22a| 72.74±1.58c    |
| Cinnamon         | 9.83±0.91b | 4.12±0.31c  | 2.84±0.25a| 4.81±0.37a| 2.90±0.17a| 85.33±1.17b    |
| L.S.D (0.05)     | 0.21       | 0.10        | 0.71      | 0.83     | 0.51      | 0.95           |

* Values are mean ± standard error
* Values in the same column with different superscript letters (a, b, ...) are significantly different.
** Calculated by difference

Iron and phytate phosphorus content of raw materials

With respect to iron content of raw materials the results in Table (2) indicated that wheat flour, celery seeds and cinnamon contained from iron 1.98, 57 and 50 mg/100 g respectively. From the same table, the data evinced that wheat flour contained from phytate phosphorus 21 mg/100 g while, celery seeds and cinnamon did not contained phytate phosphorus or tannin. These results are in agreement with those mentioned by Leif and Lena (2000). Meanwhile, Janet et al (2009) reported that several factors affect on iron absorption. For example, tannins and phylate phosphorus in food reduce iron absorption, while ascorbic acid increases it.

Sensory evaluation

Sensory evaluation is considered as an important indicator of potential consumer preferences. Inspite of its short comings it will remain one of the most reliable quality assessment technique for food and food products in general and for bread and bakery products in particular.

With respect to sensory evaluation of biscuit samples the present data in Table (3) appeared that color, crunchiness and appearance slightly decreased in biscuit samples fortified with celery seeds and cinnamon at levels 5, 7.5 and 10 gm compared with biscuits unfortified (control). On the contrary, odor and taste significantly improved in biscuit samples fortified with celery seeds and cinnamon at levels 5, 7.5 and 10 gm compared with biscuits unfortified (control). This significance increasing in odor and taste may be attributed to the volatile oil finding in celery seeds and cinnamon which it is known that enhance the odor and taste. Generally, aromatic plants and spices may improve, enhance or attribute a certain flavor to bakery products. The roasted aroma is one of the attractive flavors. It is the characteristics of all high temperature processing foods including bakery products. Pyrazine compounds are responsible for this roasted aroma (Bassiouny et al 1990).

Baking quality

Biscuits is important bakery items and used all day and considered is one of the major delicate and delicious foods and is not restricted to any particular of the day. It has very long shelf life, as they are not prone to destruction by fingers etc.

Concerning baking quality of biscuit samples, the obtained data in Table (4) elucidated that weight increased in biscuit samples fortified with celery seeds and cinnamon at levels 5, 7.5 and 10 gm compared with biscuits unfortified (control). This increasing in weight might be attributed to the higher fiber content in celery seeds and cinnamon causing higher water holding capacity (Shogren et al 1981). In contrast, volume, specific volume, diameter, thickness and expansion factor slightly decreased by increasing celery seeds and cinnamon levels compared with biscuits unfortified (control). These decreasing may be imputed to increase water holding capacity of the fiber component may have contributed to the higher batter...
Table 2. Iron, phytate phosphorus and tannins content (mg/100 g) of raw materials (on dry weight basis)

| Samples            | Iron  | Phytate phosphorus | Tannins |
|--------------------|-------|--------------------|---------|
| Wheat flour (72%)  | 1.98  | 21                 | N.D     |
| Celery seeds       | 57    | N.D*               | N.D     |
| Cinnamon           | 50    | N.D                | N.D     |

* N.D = not detected

Table 3. Sensory evaluation of biscuit samples

| Samples                      | Color (20) | Crunchiness (20) | Odor (20) | Taste (20) | Appearance (20) |
|------------------------------|------------|------------------|-----------|------------|-----------------|
| Biscuits unfortified (control) | 19.88±0.79a | 19.53±0.56a | 18.35±0.51b | 18.20±0.74b | 19.60±0.45a     |
| Biscuits (5 gm celery seeds)  | 19.75±0.82a | 19.50±0.61a | 18.38±0.42b | 18.27±0.65b | 19.55±0.61a     |
| Biscuits (7.5 gm celery seeds) | 19.63±0.53a | 19.47±0.87a | 18.42±0.65b | 18.45±0.51b | 19.12±0.36a     |
| Biscuits (10 gm celery seeds) | 19.50±0.39a | 19.41±0.88a | 19.50±0.48a | 19.61±0.68a | 18.00±0.47b     |
| Biscuits (5 gm cinnamon)     | 19.71±0.91a | 19.51±0.69a | 18.40±0.39b | 18.33±0.72b | 19.50±0.30a     |
| Biscuits (7.5 gm cinnamon)   | 19.75±0.95a | 19.44±0.97a | 19.67±0.40a | 19.60±0.76a | 19.47±0.45a     |
| Biscuits (10 gm cinnamon)    | 19.86±0.75a | 19.40±0.75a | 19.78±0.51a | 18.11±0.56b | 19.50±0.45a     |

L.S.D. (0.05) | 0.56 | 0.42 | 0.63 | 0.29 | 0.47 |

* Values are mean ± standard error
* Values in the same column with different superscript letters (a, b, ...) are significantly difference.

Table 4. Baking quality of biscuit samples

| Samples                      | Weight (g) | Volume (cc) | Specific volume (v/w) | Diameter (cm) | Thickness (cm) | Expansion factor (Di/Th) |
|------------------------------|------------|-------------|-----------------------|---------------|---------------|--------------------------|
| Biscuits unfortified (control) | 22.51±0.11b | 37.54±0.23a | 1.66±0.16a           | 4.58±0.18a    | 1.51±0.06a    | 3.03±0.09a               |
| Biscuits (5 gm celery seeds)  | 22.94±0.16b | 36.97±0.20a | 1.61±0.25a           | 3.97±0.12a    | 1.37±0.04a    | 2.89±0.09a               |
| Biscuits (7.5 gm celery seeds) | 23.26±0.08a | 36.65±0.29a | 1.57±0.10a           | 3.71±0.17a    | 1.29±0.06a    | 2.87±0.08a               |
| Biscuits (10 gm celery seeds) | 23.88±0.18a | 36.41±0.26b | 1.52±0.18b           | 3.55±0.21b    | 1.25±0.08b    | 2.84±0.11b               |
| Biscuits (5 gm cinnamon)     | 22.89±0.11b | 36.74±0.23a | 1.60±0.34a           | 3.80±0.26a    | 1.32±0.04a    | 2.87±0.06a               |
| Biscuits (7.5 gm cinnamon)   | 22.93±0.08ab| 36.48±0.19a | 1.59±0.27a           | 3.56±0.11a    | 1.26±0.09a    | 2.82±0.11a               |
| Biscuits (10 gm cinnamon)    | 23.18±0.13a | 36.35±0.17b | 1.56±0.13b           | 3.41±0.10b    | 1.22±0.07b    | 2.79±0.08b               |

L.S.D. (0.05) | 0.71 | 0.63 | 0.76 | 0.65 | 0.70 | 0.51 |

* Values are mean ± standard error
* Values in the same column with different superscript letters (a, b, ...) are significantly difference.
viscosity and if these components competed with the starch for water, incomplete gelatinization of the starch could have resulted in reduction of biscuits volume (Shogren et al 1981).

Total, available and availability iron of biscuit samples (In-vitro)

With regard to total, available and availability iron of biscuits samples, the given results in Table (5) demonstrated that total, available and availability iron increased in biscuits samples fortified with celery seeds and cinnamon by increasing celery seeds and cinnamon additives compared with biscuits unfortified (control). Schricker and Miller (1982) mentioned that the baking process had little effect on the relative availability.

Biological evaluation (In-vivo)

In relation to blood picture in groups of rats, the data in Table (6) evinced that mean values of hemoglobin (Hb), hematocrit (Ht), red blood cell (RBC), mean corpuscular volume (MCV), mean corpuscular hemoglobin (MCH) mean corpuscular hemoglobin concentration (MCHC), serum iron (SI) and serum ferritin (SF) significantly improved after 8 weeks in groups rats fed on biscuits fortified with celery seeds and cinnamon at levels 5, 7.5 and 10 gm compared with anemic control group fed on biscuits unfortified. These results are in agreement with those reported by John and Brian (2002) who reported that iron deficiency anemia cause low hemoglobin serum iron (SI), also iron deficiency lead to decrease in mean corpuscular hemoglobin (MCH) and size of red cell. Mark et al (2009) noted that daily iron intake depends on the composition of food consumed and the quantity of iron therein. Several inhibitors and a small number of enhancers of iron absorption are now known to exist. Iron absorption increases in individuals who have depleted iron status and this internal regulator of absorption may be more important than any particular constituents of the food supply. Therewithal Melissa et al (2009) appeared that basal obligatory losses in humans are ≈ 1 mg Fe/d and must be replaced by an equivalent amount of iron derived from the diet. The typical Western diet provides an average of 6 mg of heme and nonheme iron per 4120 kJ of energy intake. The bioavailability of iron is both a function of its chemical form and the presence of food items that promote or inhibit its absorption. Ascorbic acid and meat are known as the most powerful of these enhancers of nonheme iron absorption, whereas the list of inhibitors is much longer. In contrast to heme iron absorption, many factors affect nonheme iron absorption and include bran, hemicellulose, cellulose, pectin, phytic acid which is found in wheat and soy products and polyphenolic compounds. Besides Janet et al (2009) mentioned that one of the causes of nutritional anemia is that the amount of iron absorbed is insufficient to meet the body’s requirements. This insufficiency may be due to both inadequate iron intake from food and to low bioavailability. Most of the iron in the body is present in the red blood cells, mainly as a component of hemoglobin. Much of the rest is present in myoglobin, a compound occurring mainly in muscles and as storage iron or ferritin, mainly in the liver, spleen and bone marrow. Additional tiny quantities are found binding protein in the blood plasma and in respiratory enzymes. In normal circumstances, only about 1 mg of iron is lost from the body daily by excretion into the intestines, in urine, in sweat or through loss of hair or surface epithelial cells.

Histopathological examination

Table (7) and Figures (1), (2), (3) & (4) illustrated histopathological overhaul of organ heart for groups rats fed on biscuits fortified with 10 gm / 100 gm celery seeds (group No. 5 and Fig. 3) and cinnamon (group No. 8 and Fig. 4) compared with normal control (Fig. 1) and anemic control (Fig. 2) groups of organ heart. Briefly, can be said that histopathological overhaul declared amelioration in organ heart for groups rats fed on biscuits fortified with celery seeds and cinnamon compared with anemic control.

In conclusion, from aforementioned results it is evident that fortification of food with iron especially from natural sources is considered the best way for overcome iron deficiency. Natural sources of iron can be used safely because it better than synthetics sources of iron. Therewithal, fortified food can be considered healthy food or functional food which had healthy benefits.
Table 5. Values of total, available and availability iron of biscuit samples

| Samples                  | Total iron (mg / 100 g) | Available iron (mg / 100 g) | Availability iron (%) |
|--------------------------|-------------------------|-----------------------------|-----------------------|
| Biscuits unfortified (control) | 1.92                    | 0.83                        | 43.22                 |
| Biscuits (5 gm celery seeds)      | 3.40                    | 1.97                        | 57.94                 |
| Biscuits (7.5 gm celery seeds)     | 5.85                    | 3.86                        | 65.98                 |
| Biscuits (10 gm celery seeds)     | 8.17                    | 6.42                        | 78.58                 |
| Biscuits (5 gm cinnamon)         | 3.11                    | 1.63                        | 52.41                 |
| Biscuits (7.5 gm cinnamon)       | 4.23                    | 2.60                        | 61.46                 |
| Biscuits (10 gm cinnamon)        | 7.00                    | 5.30                        | 75.71                 |

Table 6. Blood picture in groups of rats

| Sample                          | Hb (g/dl) | Ht (%) | RBC (millions/mm³) | MCV (fl) | MCH (Pg) | MCHC (%) | SI (µmol/l) | SF (µg/l) |
|---------------------------------|-----------|--------|-------------------|----------|----------|----------|-------------|-----------|
| Normal control (Group. No 1)    | 13.65     | 42.37  | 4.67              | 90.72    | 29.22    | 32.21    | 11.68       | 13.72     |
| Anemic control (Group. No 2)    | ±0.73a    | ±0.60a | ±0.25a            | ±1.15a   | ±0.31a   | ±0.20a   | ±0.17a      | ±0.23a     |
| Group No. 3                    | 10.87     | 38.11  | 4.30              | 88.62    | 21.34    | 24.08    | 9.53        | 9.91       |
| Group No. 4                    | ±0.63c    | ±2.70c | ±0.23a            | ±1.16c   | ±0.30c   | ±0.21c   | ±0.12a      | ±0.16c     |
| Group No. 5                    | ±0.70b    | ±1.18b | ±0.27a            | ±1.22b   | ±0.38b   | ±0.19b   | ±0.15a      | ±0.20b     |
| Group No. 6                    | ±0.41a    | ±1.87a | ±0.20a            | ±1.37a   | ±0.28a   | ±0.23a   | ±0.23a      | ±0.21a     |
| Group No. 7                    | ±0.67c    | ±2.23c | ±0.16a            | ±1.18c   | ±0.36c   | ±0.15c   | ±0.10ab     | ±0.13c     |
| Group No. 8                    | 11.43     | 41.11  | 4.60              | 89.54    | 27.36    | 30.56    | 11.29       | 12.17      |
| Group No. 9                    | ±0.82c    | ±0.75b | ±0.13a            | ±1.23b   | ±0.39c   | ±0.26c   | ±0.19a      | ±0.22b     |
| Group No. 10                   | 12.92     | 43.35  | 4.62              | 89.50    | 27.96    | 31.24    | 11.52       | 12.86      |
| L.S.D (0.05)                   | 0.40      | 0.85   | 0.15              | 0.67     | 0.22     | 0.23     | 0.20        | 0.17       |

* Values are mean ± standard error
* Values in the same column with different superscript letters (a, b, ...) are significantly different.
** Group No. 3, 4 and 5 fed on biscuits fortified with 5, 7.5 and 10 g/100 gm celery seeds respectively.
*** Group No. 6, 7 and 8 fed on biscuits fortified with 5, 7.5 and 10 g/100 gm cinnamon respectively
Hb = hemoglobin, Ht = hematocrit, RBC = red blood cell, MCV = mean corpuscular volume, MCH = mean corpuscular hemoglobin, MCHC = mean corpuscular hemoglobin concentration, SI = serum iron, SF = serum ferritin.

Table 7. Lesion score for heart organ of groups rats

| Lesion       | Normal control (G. No. 1) | Anemic control (G. No. 2) | G. No. 5 | G. No. 8 |
|--------------|---------------------------|---------------------------|----------|----------|
| Congestion   | −                         | ++                        | −        | −        |
| Hemorrhage   | −                         | +++                       | −        | +        |
| Inflammation | −                         | +++                       | +        | +        |

− = Absence of lesion
++ = Presence of lesion by mild degree
+++ = Presence of lesion by moderate degree
++++ = Presence of lesion by considerable degree
++++ = Presence of lesion by severe degree

Arab Univ. J. Agric. Sci., 18(2), 2010
Fig. 1. Photomicrograph of heart for normal control (group No. 1).

Fig. 2. Photomicrograph of heart for anemic control (group No. 2)

Fig. 3. Photomicrograph of heart for (group No. 5)

Fig. 4. Photomicrograph of heart for (group No. 8)
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