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SANITARY STATUS OF MARKETED FROZEN CHICKEN PRODUCTS EXHIBITED IN PRESENTATION FREEZERS

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SUMMARY

A total of eighty samples of chicken products; 20 each of boneless chicken meat, boneless chicken breast, chicken half legs and chicken drum stick were collected form different markets and examined for the bacteriological status of these products. The obtained results revealed that aerobic plate count, coliforms, enterobacteriaceae and staphylococci mean counts were significantly higher in boneless chicken meat followed by boneless chicken breast then chicken half legs and finally chicken drum stick samples. Salmonella infantis could be isolated only from boneless chicken meat with incidence (5%), while Campylobacter jejuni was detected in both boneless chicken meat and breast (5% of each). The source of contamination to chicken products and the public health importance of the isolated organisms as well as the suggestive measures to minimize microbial contamination of chicken products and also to safeguard consumers were discussed.

INTRODUCTION

The quality of poultry meat is considered optimum immediately after processing and maintainance of acceptable quality depends on the initial microbial levels and measures taken to minimize growth of pathogenic organisms which may under faulty handling lead to a health hazard (Cunningham, 1982).

The spoilage of poultry carcasses stored under cold temperatures was caused mainly by the growth of Psychrophiles (Barnes and Impey, 1968).

S. aureus derives its importance in meat hygiene from its potential production of staphylococcal enterotoxins which leads to food poisoning in human being (Niskanen, 1977; Tolba, 1991 and 1994). Moreover, poultry meat was shown to be the vehicle of disease producing organisms (Khalafalla and Waffiah, 1995 and Tolba et al., 1998).

Spread of Salmonellae in processing plants occurs during processing operations, market poultry becomes contaminated to varying degrees as a result of unsanitary handling and retailing conditions (Bryan, 1968). In addition, contamination of market poultry carcasses depends mainly on the sanitary practices applied during handling, processing, storage, distribution and retailing conditions (Dougherly, 1976 and Khalafalla and Waffiah, 1995).

Clostridium perfringens found in the intestinal tract of chickens, poultry processing operations can spread the organism (Barnes, 1972) which has been isolated from carcasses at various stages of processing.

In recent years, reports have demonstrated the importance of Campylobacter jejuni and Escherichia coli as a source of human enteritis (Garcia et al, 1985). Moreover, investigation of chicken processing plants in different countries have shown that large contamination with Campylobacter jejuni can exist in birds, equipments, hands of processing line workers (Oosterom et al. 1983; Wempe et al., 1983 and Stern et al., 1985).

There are many factors affecting the growth of bacteria, in general food poisoning organisms stop growing at temperatures which still permit the growth of spoilage bacteria, many of which

can multiply slowly at -2°C. However, all bacterial growth ceases when the poultry products becomes frozen, while above freezing point the spoilage organisms grow faster with increasing temperature.

This work was conducted to determine the effect of flactuation of temperatures during storage of chicken products in the presentation freezers.

MATERIAL AND METHODS

A total of eighty random samples of boneless chicken breast, boneless chicken meat, chicken half legs and drum stick (20 samples each) were collected from the finished products of different markets. Collected samples were transferred directly to the laboratory with a minimum of delay. Collected samples were carried out according to the techniques recommended by ICMSF (1980). The following bacteriological examinations were done.

1- Determination of Aerobic Plate Count (APC/g):-

The drop plate technique recommended by ICMSF (1978) was employed. Inoculated plates with control were incubated in a thermostatically controlled incubator at 37±1°C for 48 hours for enumeration of aerobic mesophilic bacteria.

2 - Determination of Enterobacteriaceae count/g:-

The technique applied was that recommended by ISO (1987) using Violet Red Bile Glucose (VRBG) agar. Inoculated plates with control were incubacted at 37°C for 24 hours.

Representative colonies were identified biochemically according to Vernam and Evans (1991).

3 - Determination of Coliforms Count (MPN/g): MPN/g was determined according to ICMSF (1978) and ISO (1975). Inoculated tubes were incubated at 37°C for 24 hours for enumeration of most probable number of coliforms (MPN/g).

4 - Determination of Staphylococci count /g:-

Surface spread plate method according to ICMSF (1978) was applied. Inoculated Baird Parker plates with control were incubated at 37°C for 24-48 hours for enumeration of S. aureus.

5 - Isolation and Identification of Salmonellae:-

The technique adopted was that recommended by Harvey and Price (1981), suspected colonies were identified biochemically according to Kauffman white scheme (Kauffmann, 1974).

6 - Isolation and Identification of E. coli :-

The applied technique was recommended by ISO (1987). Identification of typical E. coli according to Vernam and Evans (1991).

7- Isolation and Identification of Clostridium perfringens:-

Suspected colonies were identified according to Raper and Fennell, 1965; Zycha et al., 1969; Barnnett and Hunter, 1972 and Samson, 1979.

8- Isolation and Identification of Staph. aureus:-

Suspected colonies were identified morphologically (Cruickshank et al., 1975) and biochemically (ICMSF, 1978).

The obtained results were statistically analyzed using Hypothesis test of mean and correlation coefficient according to Senedecor (1969).

RESULTS AND DISCUSSION

From the results achieved in Table (1) and Fig. (1). It is evident that boneless chicken meat samples as compared with those samples of boneless chicken breast have a significant higher mean values \pm S.E. of Aerobic mesophilic count, Colifoms, Enterobacteriaceae and Staphylococci counts were 3.2 x $10^5 \pm 2.1$ x 10^5 , 6.1 x $10^2 \pm 2.9$ x 10^2 , 1.5 x $10^3 \pm 2.3$ x 10^2 and 2 x $10^3 \pm 2.3$ x 10^3

Table (1): Statistical analytical results of examined chicken products kept in presentation freezers based on determination of their hygienic status.

Type of examined samples	No. of samples	APC	Coliform count	Enterobacteriaceae count	Stophylococci count	
Bondess chicken meat Min Max Meat ± S.E. R Log mean	20	7.2x 10 ³ 4x 106 3.2x10 ⁵ ±2.1x10 ⁵ +0.949 5.51	10 ² 6 x 10 ³ 6.1x10 ² ±2.9x102** + 0.707 2.79	8 x10 ² 4.6 x 10 ³ 1.5x10 ³ ±2.3x102* - 3.18	$ \begin{array}{c} 9 \times 10^{3} \\ 10^{4} \\ 2 \times 10^{3} \pm 6.2 \times 102^{*} \\ +0.944 \\ 3.30 \end{array} $	
Boncless chicken breast Min Max Mean ± S.E. R Log mean	20	6 x 10 ³ 2.6x10 ⁵ 2.1x105±1.4x105 - 5.32	$ \begin{array}{c} 15 \\ 3 \times 10^{3} \\ 2.1 \times 10^{2} \pm 1.5 \times 10^{2} \\ + 0.732 \\ 2.32 \end{array} $	$ \begin{array}{c} 6.4 \times 10^{2} \\ 4 \times 10^{3} \\ 1.4 \times 10^{2} \pm 2.1 \times 10^{2} \\ + 0.380 \\ 3.15 \end{array} $	6 x 10 ² 8 x 10 ³ 1.1 x 10 ³ ± 3.7 x 10 ² 3.04	
Bondess half legs Min Max Meat ± S.E. R Log mean	20	7×10^{2} 2.2×10^{6} $1.1 \times 10^{5} \pm 10^{5}$ 5.04	<3 2 x 10 2 48.4 ± 12* - 1.69	1.1 x 102 3.5 x 10 ³ 9.3x10 ² ±2.1 10 ² * + 0.193 2.97	10 ³ 7 x 10 ³ 7x10 ² ±3.5x 10 ² 2.85	
Chicken drum stick Min Max Mean ± S.E. R Log mean	20	3.1 x 10 ² 3 x 10 ⁵ 3.6x10 ⁴ ±2.1x10 ⁴ 4.56	< 3 90 39± 9.5 + 0.152 1.59	1.3 x 10 ² 2.7 x 10 ³ 6.7x10 ² ±1.6x 10 ² - 2.83	102 5 x 103 5.4 x 10 ² ±2.5 x 10 ² - 2.73	

Correlation coeffecient between APC and S. aureus count in boneless chicken meat.

chicken meat

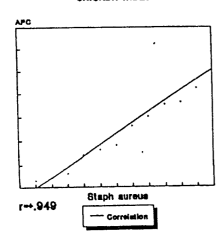
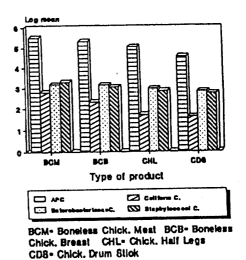


Fig. (1): Log mean of microbiological examination of chicken products.



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^{*} Significant at p < 0.005
** Significant at p < 0.001

6.2 x 10^2 for former against 2.1 x $10^5 \pm 1.4$ x 10^5 , 2.1 x $10^2 \pm 1.5$ x 10^2 , 1.4 x $10^3 \pm 2.1$ x 10^2 and 1.1 x $10^3 \pm 3.7$ x 10^2 for later respectively, such counts were slightly decreased to 1.1 x $10^5 \pm 10^5$, 48.4 ± 12 , 9.3 x $10^2 \pm 2.1$ x 10^2 and 7 x $10^2 \pm 3.5$ x 10^2 c.f.u./gm in examined chicken half legs, while they were 3.6 x $10^4 \pm 2.1$ x 10^4 , 39 \pm 9.5, 6.7 x $10^2 \pm 1.6$ x 10^2 and 5.4 x $10^2 \pm 2.5$ x 10^2 in examined drum stick respectively.

The obtained results revealed that the aerobic plate count in most products was over than the permissible limits (105), this may be attributed to the frequent thawing and refreezing of chicken products kept in some presentation freezers as a result of ineffeciency of cooling system which may be due to careless in covering suchfreezers by its cover lids resulted in a partial loss of freezing degrees and subsequently accelerate the growth and multiplication of food spoilage organisms.

In this respect, Sauter et al., (1978) mentioned that the number of salt tolerant bacterial were declined rapidly after freezing storage, while they become relatively constant during thawing of chicken products. This may be attributed to the effect of freezing and subsequent thawing causes tissue disruption releasing nutrients which required to improve the metabolic activity of salt tolerant bacteria. The authors revealed that APC and S. aureus counts of examined frozen and thawed chicken products were 2.1×10^4

and 1.1 x 10³ organism/g respectively. This substantiates the findings reported in the present investigation, while lower findings of APC and enterobacteriaceae counts were reported by Notermans and Kamplemacher, 1975 (3.2 x 10³) and 3.2 x 10²) respectively. Moreover, the obtained results of APC of examined boneless chicken breast and drum stick samples were higher than those obtained by Kotula, 1966 (1.1 x 10³ and 8.7x 10²) respectively. Kraft et al., 1963 found that the examined frozen chicken product samples attaining APC, S. aureus and colifroms counts 3.6 x 10^4 , 6.9 x 10^2 and 5.2 x 10² organism /g. respectively. These results agree with that reported in the present investigation.

Presence of coliforms ogranisms in great numbers indicates contamination from faecal materials and lack of careful handling during processing. This helds the view reported by Pini and Gilbert (1988) and Hudson and Mead (1989). Moreover, Ostovar et al., 1971. revealed that freezing of deboned poultry carcasses resulted in a significant reduction of faecal coliforms.

Weak and direct correlations were noticed between S. aureus and enterbacteriaceae (r=+0.380), enterebacteriaceae and coliforms (r=+0.193) and between staphylococci and coliforms (r=+0.152) in examined boncless chicken breast, chicken half legs and drum stick respectively. However, the correlation coefficient was

stronger between enterobacteriaeae and coliforms in boneless chicken meat and boneless chicken breast than those of other products (r=+0.707 and +0.732), the correlation was higher between staphylococci and each of APC and coliforms in boneless chicken meat (r=+0.949 and +0.944).

On the other hand, significant higher mean value (p<0.005) of enterobacteriaceae and staphytococci and between enterobacteriaceae and coliforms in examined boneless chicken meat and chicken half legs were observed respectively, while strong higher mean value (P<0.001) were noticed between coliforms and staphylococci in booneless chicken meat than the other products.

Notermans and Kampelmacher (1975) revealed that there is no significant difference regarding the regression coefficient between APC and enterobacteriaceae counts. These results are consistent with that reported in the present investigation. In this respect, Cox et al. (1974) found that there is no significant difference in mean value at (P<0.05) of APC and enterobacteriaceae counts in examined boneless chicken breast and chicken drum stick samples. These results agree with that reported in the present investigation.

From the present data it could be concluded that Achromobacter, Citrobacter freundii, Enterobacter agglumerans, Enterobacter aerogenes, Flavobacterium, Proteus vulgaris, Proteus myxo-

Table (2): Frequency distribution of isolated organisms from examined chicken products.

	Type of samples									
Isolated	·Boneless ·chicken			Chicken		Drum		Total No. of		
Organisms	meat		breast		half legs		stick		positive	
	No	%	No	%	No	%	No	%	No	%
Achromobacter	1	5	-	0.0	-	0.0	-	0.0	1	5
Citrobacter C. freundii	2	10	2	10	1	5	1	5	6	30
Enterobacter E.agglumerans E. aerogenes	1 4	5 20	1	5 0.0	2	10 0.0	- 1	0.0	4 5	20 25
Flavabacterium	i	5	-	0.0	-	0.0	-	0.0	1	5
Proteus P. vulgaris P.myxofaciens	1 - 3	5 0.0 15	2 1 2	10 5 10	 - - 1	0.0 0.0 5	- - 1	0.0 0.0 5	3 1 7	15 5 35
E.coli Klebsiella: K.ozonae K.cloacae Clostridium perfringens S. aureus Sal. Infantis	1 1 2 1 1	5 5 10 5 5	1 1 3 -	5 5 15 0.0 0.0	1 1 1 -	5 5 5 0.0 0.0	1 1 1 -	5 5 5 5 0.0	4 4 7 2 1	20 20 35 10 5
Campylobactor jejuni l	1	5	1	5	-	0.0	<u> </u> -	0.0	2	10

faciens, E. coli, Klebsiella ozonae, Klebsiella cloacae, Clostridium pefringens, S. aureus and Campylobacter jejuni could be isolated from the examined samples at different percentages ranged from (5% to 35%), while Salmonella infantis could be isolated from examined boneless chicken meat only (5%). (Table2). The results of salmonella isolation are relatively consistent with that reported by Bryan (1968) who pointed out that the incidence of salmonellea in examined poultry meat and their products from retail markets varied from 1 to 50%. In this respect, Roberts, 1972 and Watson, 1975 revealed that the incidence of salmonellae isolated from poultry meat ranged from 3% to 62%. Moreover, Sadler and Corstvet, 1965 and Glezen et al., 1973 could isolate Sal. infantis from examined chicken meat products. This substantiates the findings reported in the present investigation. On the other hand, Ostvar et al. (1971) could isolate Clostridium perfringens, S. aureus, achromobacter, flavobacterium from examined frozen chicken products with different percentages.

The results of S. aureas isolation recorded here are relatively more or less inaccordance with that previously reported by several investigators (Surkiewicz et al., 1969; Sauter et al., 1978 and Devriese, 1980). Healthy poultry tissues does not support prolonged growth of staphylococci, while bruised tissues will allow persistance of such organism. This agree with hypothesis reported by Genigeorgis and Sadler (1966) and

Cunningham (1982).

Campylobacter jejuni isolation rates were even higher than those of Salmonellae. This agree with that reported by Bruce et al., 1977; Butzler, 1978; severin, 1978 and blaser et al., 1979, while lower incidence of Campylobacter jejuni from examined chicken carcasses was recorded by Elgamal et al.,1992 (4%).

Presence of coliforms in chicken products may be indicative of defective techniques applied during preparation, handling, processing and storage which may lead to economic losses through the development of undesirable changes rendering the product of low quality or even unfit for consumption (Chambers et al., 1976 and ICMSF, 1978). Moreover, some of the isolated organisms have been implicated in food illness e.g. proteus species, citrobacter freundii, enterobacter species and klebsiella species (Krieg and Holt, 1984; Marzouk, 1985; Elmossalami et al., 1988 and Abdel- Aziz, 1993).

Microorganisms that have been implicated in food poisoning outbreaks attributed to poultry meat are mainly Salmonellae, S. aureus, Campylobacter and E. coli Poor hygienic conditions during processing as well as fluctuation of storage temperature in presentation freezers may allow the organisms to survive and multiply or reinfect the cooked food and leads to food poisoning when the food is consumed.

Improving the sanitary status of chicken products and safeguard the consumers from receiving contaminated chicken products can be achieved by prolonging the durability of the product through application of a strict hygienic measures during preparation, storage and handling which is helpful in reducing its bacterial load which in turn protect the product from being spoiled in the retail markets. Moreover, it protects the consumer from pathogens which may be present in chicken products.

On the manufacturing side to test the raw materials and to control the processing conditions, Good manufacturing practices (GMP) should be followed by the codex Alimentarius Commission (CAC) (1976); Code of hygienic practices for processed products.

REFERENCES

- Abd El. Aziz, A. S. (1993): Sanitary improvement of packed meat in a meat plant. Ph. D. Thesis Fac. Vet. Med., Cairo University.
- Bahy El-Gamal, G.; Refaie, R. S. and Abou El-Ailla, A.
 A. (1992): Occurrence of compylobacter in poultry carcasses. Assiut Vet. Med. J., 26, 52:110-113.
- Barnes, E.M. (1972): Food poisoning and spoilage bacteria in poultry processing. Vet. Rec. 24: 720-722.
- Barnes, E.M. (1976): Microbiological problems of poultry at refrigerator temperature. A review. J. Sci. Fd. Agric., 27: 777.

- Barnes, E.M. and Impry, C.S. (1968): Psychrophilic spoilage bacteria of poultry. J. Appl. Bacterial., 31:97.
- Barnnett, H. L. and Hunter, B.B. (1972): Illustrated genera to imperfect fungi. 2nd. Ed. Burgress pub. Co.
- Blaser, M.J.; Cravens, J.; Power B.W.; Laforce, F. M. and Wang W.L.L. (1979): Campylobacter enteritis associated with unpasteurized milk. Am. J. Med. 67: 715 - 718.
- Bryan, F. L. (1968): What the sanitation should know about staphylococci and Salmonellae in non dairy products. II Salmonella. J. Milk Food Technol. 31: 131 146.
- Bruce, D.; Zochowsky, W. and Ferguson, J. R. (1977): Campylobacter enteritis. Brit. Med. J. pp. 1219.
- Butzler, J. P. (1978): Infection with Campylobacter. pp. 214-239. In J. D. Willams (ed.) Modern topics in infection. W. Heinemann. London.
- CAC (1976): Recommended International Code of hygienic practice for processed meat products. FAO/WHO. P. 264.
- Chambers ,J. V.; Brechbill, D.O. and Hill , D.A. (1976): A microbiological survey of raw ground beef in Ohio. J .Milk Food Technol . 39 , 530
- Cox, N. A.; Mercuri, A. J.; Juven, B. J.; Thomson, J. E. and Chow, V. (1974): Evaluation of succinic acid and heat to improve the microbiological quality of poultry meat. J. Food Sci. 39: 985.
- Cruickshank, R.; Duguid, J.; Mormian, B and Swain, R. (1975): The practice of medical microbiology, 12th. Ed. Churchill Livingstone, Edinburgh.
- Cunningham, F.E. (1982): Microbiological aspects of poultry and poultry products- An update. J. Food Prot. 45: 1149.

- Devriese, L. A. (1980): Pathogenic staphylococci in poultry, J. worldís Poul. Sci., 36, 4:227-235.
- Dougherly, T. J. (1976): A study of salmonella contamination in broiler flocks. J. Poultry Sci. 55:1811-1815.
- Edwards, R. P. and Ewing, W. A. (1972): Identification of Enterobacteriaceae. 3rd Ed. 38 Burgess Publ. Comp., Minneapolis.
- Elmossalami. E.; Saad ,S. M.; Niazi , Z. M. and Mohamed , H.M. (1988): Sanitary status of meat in rural areas of kalubia governorate . Vet. Med. J. 36, 3:385.
- FAO/ WHO (1991): Codex committee on Food Hygienic (CAS) General HACCP definitions and procedures for use by codex. FAO/WHO. 25th Session. Washington D.C.
- Garcia M. M.; Lior, H.; Stewart, R. B.; Ruckerbauer, G. M.; Trudel, J.R.R. and Skljarevski, A. (1985): Isolation, characterization and serotyping of Campylobacter jejuni and Camplyobacter coli from slaughter cattle. Appl. Environ Microbiol. 49: 667 672.
- Genigeorgis, C. and Sadler, W.W. (1966): Characterization of strains of S. aureus isolated form livers of commercially slaughtered poultry. Poultry Sci., 45: 973-980.
- Glezen, W.: Martin, P.; Hines, D.V.; Mildred, B.S. (1973): Salmonella in two poultry processing plants, 148,5: 550 552.
- Harvey, R. W. and Price, T. H.(1981): Comparison of Selenite F., Mquller Kauffman tetrathionate and Rappaports medium for Salmonella isolation from chicken giblets and after per-enrichment in buffered peptone water, J. Hyg. Camb. 87,219.
- Hudson, W. R. and Mead, G.C.(1989): Listeria contamination at a poultry processing plant. Letters in Applied

Microbial. 9,211

- International Commission of Microbiological specification for Foods "ICMSF" (1980): Microbiol. Ecology of Food, 1: Factors affecting life and death of microorganisms. Academic Press. London.
- International Commission on Microbiological Specification for Foods "ICMSF" (1978): Micoroorganisms in Food. Their significance and methods of enumeration. 2nd Ed. University of Toronto Press, Toronto, Buffalo, London.
- International Standards Organization ISO (1975): Meat and meat products, detection and enumeration of presumptive coliform bacteria (reference method). International standards ISO / Dis / 2811.
- International Standards Organization ISO (1987): Microbiology General guidance for enumeration of Bacillus cereus- colony count technique at 30°C. ISO 7932. Geneva, Switzerland.
- Kauffmann , F. (1974): Kauffmann white scheme. WHO BO / 72 , L. Rev .I. Acta. Path . Microbiol. Scand . 61 , 385.
- Khalafalla, F.A. and Waffiah A. (1995): Bacteriological study on ducks and squabs. J. Egypt. Vet. Med. Assoc. 55,1,2,: 629 634.
- Kotula, A.W. (1966): Variability in microbiological sampling of chickens by the swab method. Poultry Sci., 45: 233 236.
- Kraft, A. A.; Ayes, J. C.; Weirs, K. F. and Marion, W. (1963): Effect of method of freezing on survival of microorganisms in turkey. Poultry Sci. 42: 128-136.
- Krige, N. R. and Holt, J.G. (1984): Bergey's manual of systematic bacteriology .Vol. 1 ., Williams and Willkins . Baltimore , USA .

- Marzouk, I.(1985): A study on microbial etiology of infantile diarrhoea related serum electrolyte changes and agent related efficiency of antisecretory drugs (chlor-promezine). M.M.D., pediatrics, Fac. Med., Alexandria University.
- Sauter, E. A.; Peterson, C.F. and Parkinson, J. F. (1978): Microfloral comparison of fresh and thawed frozen fryers. J. Poultry Sci., 57:422 - 424. Severin W. H.L. (1078) Campylobacter enteritis. Ned. T. That a cinh ide it (Gonecosk, 122, 15, 1499 - 504.
- food poisoning, production, properties and detestion of oils 113231(1111) 11312 VIRILIAN Food poisoning, production, properties and detestion of oils 113231(1111) 1111123 enterotoxins. Technical Research Center of Finland. Stern N.i., Rothenberg, P.J. and Stone, J.M. (1985): Enu-Materials and Processing Technology, Publication 10 Niskanen, A. (1977): Staphylobocotal dampdoxides and NO Materials and Processing Technology, Publication 19, Espoo pp. 83.
 - try and red meat . J . Food . PROT . 48: 606 610.

W. (1969): Statistical methods . 4th. Ed. The

- Notermans, S. and Kamplemacher, E. H. (1975): Studies bacterial counts from frezen broners J. Appl. Hast. 1382
 - Surkiewicz, B.F.; Johnson, RIW. Moram, A. B. and of different sampling methods for the determination of the learning of the determination of the determination of the learning of the determination of the de chicken eviscerating plants. Food Technol. 23: 1066
- : 25 31 المرب ب بالأسواق وقد أظهرت النظائج أن أعل Oosterom J.; Notetmans, St. Karman, H. and Engels, G. B. (1983): Origin and prevalance of Campylobactor jeju
 ni in poultry processing. J. Food. Prot. 46: 339 - 344.
 - Tolba, K. (1991): Staphylococoal Enteroloxide ein_meat products from urban and rural areas (Ph.D. Thesis Fac.
- Poultry product quality, 5. Microbiological evaluation of mechanically debonded poultry, meat. J. Food Sci.
 - meat . Vet .Med. J. 42: 99 105

Correlation cocff 1998 - 2001 : 36 Pini, P.M. and Gilbert, R.J. (1988): The occurrence in the UK of Listeria species in raw chickens and Inter . J. Food Microbiol. 6:317.

- Tolba, K.; Elmossalami, M. K.; El-Neklawy, E.M. and Zicnab Niazi (1998): Batteriological quality of Turkey.
- Raper, K.B. and Fennell, D. (1965): The genus Aspergillus--Williams and Wilkins Co., Baltimore.
- Duck and Kabbircareasses 4th World Congress. Foodsoft cheeses a lielyborne infections and intoxications 1, S-B 31: 493 - 501.
- Roberts, D. (1972): J. Hyg. Camb. 70:565. Cited after Watson, W. A. (1975).
- Vernam, A.H. and Evans, M.G. (1991): Food borne pathogens. Wolfe, Publishing Ltd., England.
- Sadler, W.W. and Corstvet, R. E. (1965): Second survey of market poultry for salmonella infection. Appl. Microbiol 13:348-351.
- Watson, W. A. (1975): Salmonella infection and meat hygiene: Poultry meat. Vet. Rec., 96:351 - 353.

- Samson, R. A. (1979): A complication of the Aspergillus described since 1965. Studies in mycology No. 18.
- Wempe, J.M.; Genigeorgis, C.A.; Faver, T. B. and Yusufu, H. I. (1983): Prevalece of camplylobacter jejuni in two California chicken processing plants. Appl. Environ. Micorbiol., 45: 355 - 359.
- Zycha, H.; Siepmann, R. and Linneman. G. (1969): Mucorales Eine Beschreibung aller Gattungen und Arten dieser pilzgruppe . Verlage Von. J. Crameer .

الحالة الصحية لمنتجات الدواجن المجمدة المعروضة بفريزارات العرض بالأسواق المعروضة بفريزارات العرض بالأسواق Sanitary status of marketed frozen chicken products exhibited in presentation freezers د. خالد شوقي على طلبه

أجريت هذه الدراسة على ٨٠ عينة من منتجات الدواجين المختلفة (الدجاج المسحب-صدور فيليه- الأفخاذ - السيقان) بواقع ٢٠ عينة من كل منتج تم تجميعها من الأسواق المختلفة لتحديد الحالة البكتيريولوجية لهذه المنتجات أثناء عرضها بفريزارات العرض بالأسواق وقد أظهرت النتائج أن أعلى نسبة للعد البكتيرى الكلى وعد الميكروبات القولونية والمعوية وكذلك الميكروب المكور العنقودي وجدت في عينات الدجاج المسحب يليها الفيلية فالأفخاذ ثم السيقان وتم عزل ميكروب السالمونيلا فقط من الدجاج المسحب بنسبة ٥% أما ميكروب الكامبيليوباكتر فتم عزله من نفس المنتج وكذلك الفيليه بنسبة ٥% لكل منها.

وتم تحليل النتائج إحصائياً باستخدام Test (t) وتحديد معامل الارتباط وتم تحليل النتائج إحصائياً باستخدام Correlation coefficient بين المنتجات وتمت مناقشة الأهمية الصحية للميكروبات المعزولة وكذلك الاشتراطات الصحية الواجب توافرها لتقليل نسبة تلوث بهذه المنتجات بغرض حماية صحة المستهلك.

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