

Egyptian Journal of Animal Health

P-ISSN: 2735-4938 On Line-ISSN: 2735-4946 Journal homepage: https://ejah.journals.ekb.eg/

Effect of pesticide contamination on mastitic syndrome: a realistic field study

¹Sultan F. Nagati., ² El touchy ,E.I, ³Fazea A. Sdeek, ⁴Essam Kamel, ⁵Momtaz A. Shahein, ⁶Muhsin Konuk ⁷Ehdaa,O.Hamed and ⁸Hammad O. Hammad

- ^{1, 7}Department of Bacteriology, Animal Health Research Institute (AHRI), Agriculture Research Center (ARC), Egypt.
- ²Department of biotechnology, Animal Health Research Institute (AHRI), Agriculture Research Center (ARC), Egypt.
- ³Central Agricultural Pesticide Laboratory, Agriculture Research Canter (ARC), Egypt.
- ⁴Department of chemistry, Animal Health Research Institute (AHRI), Agriculture Research Center (ARC), Egypt.
- ⁵Department of virology, Animal Health Research Institute (AHRI), Agriculture Research Center (ARC), Egypt.
- ⁶Department of Molecular Biology and Genetics, Uskudar University, Istambul, Turkey ⁷Biochemistry, Toxicology and feed Deficiency. Toxicology Unite, Animal Health Research Institue (AHRI) Agricultural Research Center (ARC), Egypt.

Received in 6/11/2022 Received in revised from 23/11/2022 Accepted in 20/12/2022

.....

Keywords:

mastitis, pesticide, antibiotic resistance, cow, buffalo, milk, fodder

ABSTRACT

his work was carried out to answer a question of multidrug resistance and fodder pesticide residue relations to increasing mastitis syndrome in cattle and buffalos in Egypt. For this, Milk of cattle and buffalos and fodder samples were collected from 3 different localities in El-Fayoum province.

Pesticide residues analysis was carried out using the modified QuEChERS method followed by liquid chromatography coupled to triple quadrupole tandem mass spectrometry (LC-MS/MS). The residue analysis revealed that about 85% of alfalfa fodder samples and 58% of milk samples were contaminated with pesticides in the valley applied pesticide in alfalfa fodder. The pesticides found in this study were chlorpyrifos, cyhalothrin, lufenuron, and malathion. Identification of bacterial isolates was carried out using conventional PCR. Their prevalence rates were as follows: *Escherichia coli* 4, 11, 15 (5, 10, 13.64%) in milk from the farm, non-pesticide applied, and pesticide applied villages, respectively. For the other microorganisms; *Klebsiella* 0, 2, 5 (0, 1.8, 4.5%), *Staphylococcus aureus* 6, 8, 18 (7.5, 7.3, 16.4%), and *Streptococcus agalactiae* 3, 7, 17 (3.75, 6.4, 15.5%). The isolation rates of these identified bacteria from each sampling point were found to be statisti-

Corresponding author: Sultan F. Nagati, Department of Bacteriology, Animal Health Research Institute

(AHRI), Agriculture Research Center (ARC), Egypt

Email: Sultan_farag99@yahoo.com DOI: 10.21608/EJAH.2023.287818 cally significant using the Fischer Exact Probability test (P < 0.05). The results also showed that there was a multidrug resistance to nearly all tested antibiotics in bacteria isolated from milk collected from Ezbet Furqan. The antibiotic sensitivity pattern of bacterial isolates in milk samples collected from a dairy cow's farm in Fayoum province was higher when compared to that of Ezbet Barghout cows. In conclusion, there was a significant correlation between pesticide residues and multidrug resistance.

INTRODUCTION

This research was conducted after observations of veterinarians about increased mastitis cases in Ezbet Furqan, Tamiya city, Fayoum Province Pesticides are widely used in the agriculture sector to avoid or diminish losses from pests. Consequently, this can improve agriculture yield and the quality of the crops, which is important to consumers (Cooper and Dobson, 2007). These pesticides are applied preharvest, which can transfer to animals and accumulate in milk (Shazia and Karam, 2017). The organophosphate insecticide has been reported to bind with human or bovine serum albumin (Ying *et al.*, 2014).

Generally, the source of pesticide residues in milk could be the ambient such as water, soil, and air or fodder or treatment of the animals against disease vectors like ticks, mites, and insects or direct uncontrolled contamination sources such as the dairy utensils that are used during milking or storing it (Özkara et al., 2016). Nevertheless, indirect contamination is a more important way comprising the medication and/or administration of pesticides orally, cutaneous, or via inhalation to the milkproducing animals in closed barns. Whatever the reason does not matter, the active component of the pesticides can be taken into the body, then be metabolized, and finally eliminated into the animal's milk (Fischer et al., 2015).

The mammary gland inflammation is diagnosed as Mastitis and its characteristic is an increase of somatic cell amount in the milk due to the pathology formed in the mammary tissue. Bacteria, mycoplasmas, and fungi are well-known mastitis-causing microorganisms. These can be classified as specific udder pathogens, contagious pathogens, and environmental pathogens.

A well-known problem is that bacteria are gaining antibiotic resistance, and there are several reports related to pesticide residues in the fodders and the antibiotic resistance to cure infectious diseases in domestic animals (Getahun et al., 2007; Fisher et al., 2015 and Wrzecińska et al 2021).

In the light of the limited literature shown above; the presented study aimed to:

determine the microorganisms that caused mastitis in cows and buffalos,

determine the pesticide residues in both milk and fodders

find out any relationship between pesticide residue and antibiotic resistance of mastitiscausing microorganisms.

Material and Methods:

1. Study area:

Fayoum province is located southwest of Cairo with an area of 1,827 km² (CAPMAS 2018). Fayoum is an agricultural province with numerous people in rural dwellings who keep livestock at home. Such a deteriorating situation correlates to some sociodemographic aspects such as high illiteracy rates, poverty, and strong traditional beliefs related to the rural community (HDR 2008; Figure 1).

2. Experimental design:

This study was carried out between October 2021 to April 2022. A cross-sectional study was carried out in two villages (Ezbet Barghout and Ezbet Furqan) and a dairy farm in Taymiyyah city, Fayoum province, Egypt. The samples from dairy farms that grow Egyptian buffalos and local crossbred cows were also collected.

In villages, since there was a lack of recorded formal data; the information about the studied animals was collected from the locals. The owners of the studied animals are individual farmers and every farmer had only 2 to 3 cows and/or buffalos. These farmers are mainly keeping their animals in the backyard of their house. In Fayoum, the diet of the animals (home-prepared concentrates) is not formulated according to physiological needs. At home, the concentrates (1-3 kg/animal) are provided once daily, in addition to dry wheat hay. The animals are taken to the field early morning

after milking every day for feeding by green ration and return home just before sunset. In recent years, some farmers spray different pesticides onto their crops which are used to feed their animals. The samples of both milk and grass (Alfalfa fodder) were collected and analyzed between October 2021 and April 2022.



Figure 1: The map of the study area

Animal grouping:

The animals were chosen as fed with pesticide-applied alfalfa fodder and non-pesticide applied alfalfa fodder from dairies located in Ezbet Furqan, and Ezbet Barghout districts. The study was designed as follows:

- a. Both cows and buffalos' milk were tested for subclinical mastitis using the California test. Positive samples were subjected to bacterial isolation and identification and pesticide residue analysis.
- **b.** green grass sample rations were tested for pesticide residue analysis.
- c. All mastitis-positive animals' milk samples were then subjected to further examination to determine the bacterial strains by PCR followed by a susceptibility test for different antimicrobial agents as well as pesticide residue analysis.

Table 1. Type and number of samples collected from a dairy farm and two groups from individual farmer in tow valleys.

Species	No of animal	Milk	Grass (Alfalfa)	Total samples
Cow	230	230	35	265
Buffalo	70	70	10	80
Total	300	300	45	345

Grass (Alfalfa fodder) samples were collected from feeds in front of animals under study at 3 studied districts

Chemicals

Chlorpyrifos, Cyhalothrin, Lufenuron, and Malathion reference standards, were purchased from Sigma-Aldrich. Used chemicals were of HPLC grade, acetonitrile, methanol, n-hexane, formic acid, ammonia solution, and glacial acetic acid were obtained from Sigma-Aldrich (USA). QuEChERS Kits 5982-5650 reagent were obtained from Agilent Technologies (USA). Deionized water was produced by the Millipore system.

Sample Extraction and Cleanup:

All the extractions from both fodder and milk samples were carried out by using a modified method of QuEChERS as explained by Lehotay et al., (2005). In the extraction procedure, 10 g of alfalfa fodder, and 2 ml of milk were employed, and the extracts then were subjected to LC-MS/MS analysis.

Preparation of standard solution.

1 mg/ml of stock solution of the analytic standard was dissolved in acetonitrile as sol-

vent. This was used for fortification of the matrices, and a calibration curve was obtained by serial dilutions. All standard solutions were stored at 4°C until use. The standard calibration curve was created by plotting analytic concentrations versus peak area.

LC-MS/MS system:

HPLC (Agilent) 1200 Series instrument coupled to API 4000 Qtrap MS/MS from AB Sciex with electrospray ionization (ESI) interface in the positive mode, source temperature was 400 $^{\circ}$ C, and ion spray potential was 5500 V. Separation was performed on Agilent C18 column ZORBAX Eclipse XDB 4.6 x 150 mm with 5.0 μ m particle size. The injection volume was 10.0 μ l.

Table 2 shows the used gradient elution program at a 300 μ l/min flow rate. One reservoir contained a mobile phase buffer of 10 mM ammonium formate solution in methanol: water (1:9 v/v) at pH=4 and the other reservoir contained LC-MS grade Methanol. The total run time was 32 minutes.

Table 2. The LC Gradient Elution program.

Time (min)	Mobile Phase Buffer %	Methanol %
0	100	0
13	5	95
21	5	95
28	100	0
32	100	0

Internal Quality Control (IQC):

The IQC is an important item in the technical requirements (ISO/IEC 17025, 2005). To assess the extraction efficiency IQC was carried out using spiked blank samples at $0.05~\mu g/ml$. Tables (3) demonstrate the range of recoveries for tested pesticides varied between 75-116 %, which is acceptable according to SAN-TE and Eurachem guidelines (Magnusson & Ornemark, 2014; SANTE/11945/2015, 2015).

Recovery % = $\begin{array}{c} \mu g \text{ pesticide found in the spiked sample} \\ ------ \times 100 \\ \mu g \text{ pesticide added in the spiked sample} \end{array}$

Methods validation:

The method was validated by the conventional validation parameter, including the level of detection (LOD), level of quantitation (LOQ), and accuracy (recovery%) as recommended by SANCO (2013). LOD was calculated considering 3 times the value of background noise obtained for blank samples.

Bacterial Examination:

California Mastitis Test (CMT):

In this test, the method detailed by Leach et al., (2008) was followed. For this, ~ 2 mL of milk sample was collected from each quart in a plastic oar of four shallow cups marked A, B, C, and D. An equal amount of CMT reagent was added to the milk and mixed then the paddle was rotated after ~ 20s, and the score was read. The test was performed daily to support the data obtained by precise somatic cell counting

Microbiological examination:

MacConkey agar was employed for E. coli detection after the 0.1 ml milk sample smears on it. After 24h of incubation at 37°C, five lactase-positive colonies were marked and selected. The selected colonies were isolated by subculture on blood agar (BA). After 24 h of incubation, the cultures were tested by their oxidase activity (OXI) (PLIVALachema, Brno, Czech Republic). OXI-negative strains and controls were transferred on Simmons citrate agar and Motility Test Medium and incubated for another 24h at 37°C. After their assess-

ment, biochemical identification was carried out.

0.1ml milk sample inoculum on Mannitol Salt Agar was used in detecting S. aureus. After a 36h of incubation at 35°C, typical colonies were subcultured on blood agar media (BA) and incubated for 24h at 37°C, then both catalase and staphytect tests (Oxoid) were determined. Staphytect positive strains were examined by using Voges-Proskauer (VP) test according to the method of Rysanek, et al. (2007). 0.05 ml milk sample as inoculum on BA is used to detect the Streptococcus species. After 24-48h of incubation at 37°C, the β- hemolytic colonies were subcultured on BA and incubated at 37°C for 24 h, and a catalase test was carried out. API 20 Strep (Lancefield grouping) was employed in identification as explained by Rysanek, et al. (2007).

Bacterial strains and growth conditions.

The bacteria detected in the study were *E. coli, Staphylococcus aureus, Klebsiella*, and *S. agalactiae*. All strains were cultured in Tryptic soy broth (TSB; Difco Laboratories, Detroit, Mich.) at 37°C for ~17 h before DNA extraction. Cell numbers were determined by the preparation of serial dilutions of overnight culture in phosphate-buffered saline (PBS) and plating on blood agar (Columbia agar base supplemented with 5% defibrinated sheep blood).

Procedure for isolation and identification of isolates by using conventional PCR DNA extraction:

DNA was extracted from the samples by using the QIAamp DNA Mini kit (Qiagen, Germany, GmbH). For this, a 200 µl of sample suspension was incubated with 10 µl of proteinase K and 200 µl of lysis buffer at 56°C for 10 min. Then, 200 µl of absolute ethanol was added to the lysate. The sample was then washed and centrifuged and then nucleic acid was eluted with 100 µl of elution buffer provided in the kit.

Oligonucleotide Primer:

Primers used were supplied from Metabion (Germany) and are listed in Table 3.

PCR amplification:

Primers were utilized in a 25 μ l reaction containing 12.5 μ l of EmeraldAmp Max PCR Master Mix (Takara, Japan), 1 μ l of each primer of 20 pmol concentration, 5.5 μ l of water, and 5 μ l of DNA template. The reaction was performed in an Applied biosystem 2720 thermal cycler.

Analysis of the PCR Products:

PCR products were run on 1.5% agarose gel

electrophoresis (Applichem, Germany, GmbH) in 1 x TBE buffer at room temperature using gradients of 5V/cm. In the analysis, 15 μ l of the products were loaded in each gel slot. A gene ruler 100-1000 bp ladder (Fermentas, Germany) was also used to determine the fragment sizes. The gel was photographed by a gel documentation system (Alpha Innotech, Biometra) and the data was evaluated by using computer software.

Table 3. Oligonucleotide primers for identification of *E. coli*, *Klebsiella* pneumoniae, *Staphylococcus aureus* and *Streptococcus agalactiae*.

Target agent	Tar- get	Primers sequences	Ampli- fied	Prim. Den.	Amplifi	cation (3 cles)	35 cy-	Final exten-	Refer- ence
	gene		segment (bp)	Den.	Sec. den.	Ann.	Ext.	sion	chec
E. coli	phoA	CGATTCTG- GAAATGGCAAAA G CGTGATCAGCGG TGACTATGAC	720	94°C 5 min.	94°C 30 sec.	55°C 40 sec.	72°C 45 sec.	72°C 10 min.	Hu <i>et al.</i> , 2011
Klebsiella pneumoniae	gyrA	CGC GTA CTA TAC GCC ATG AAC GTA ACC GTT GAT CAC TTC GGT CAG G	441	95°C 5 min.	94°C 30 sec.	55°C 40 sec.	72°C 45 sec.	72°C 10 min.	Brisse and Verhoef, 2001
Staphylococcus aureus	16S rRN A	CCTATAA- GACTGGGATAAC TTCGGG CTTTGAG- TTTCAACCTTGCG GTCG	791	95°C 5 min.	94°C 30 sec.	55°C 40 sec.	72°C 45 sec.	72°C 10 min.	Mason <i>et al.</i> , 2001
Streptococcus agalactiae	cfb	TTTCACCAGCTG- TATTAGAAGTA GTTCCCTGAACAT TATCTTTGAT	153	95°C 5 min.	94°C 30 sec.	50°C 30 sec.	72°C 30 sec.	72°C 7 min.	Konik- kara <i>et</i> <i>al.</i> , 2014

Antimicrobial susceptibility test of different bacterial isolates:

Four or five typical colonies of similar morphological appearance were transferred to a tube containing 5 ml of Muller-Hinton broth and incubated at 37°C for 8 hours until its tur-

bidity exceeds that of the standard McFarland 0.5 barium sulphate tube. A sterile cotton swab was dipped into the standardized bacterial suspension. The dried surface of Muller-Hinton plates were streaked by the swab in 3 different planes. The plate lids were replaced and the inoculated plates were allowed to remain on a

flat and level surface undistributed for 3 to 5 min (not more than 15 min.) Then the disks Clarithromycin (CLR 15μg), Gentamicin (CN 10μg), Amikacin (AK 30μg), Ampicillin + Sulbectam (SAM 20μg), Cefotaxime (CTX 30μg), Amoxicillin (AML 10), Cefepime (CFM 30μg) Amoxicillin+clavulenic acid (AMC 30μg), Spiramycin (SP 100μg) Ampicillin (AM 10 μg) and Sulfa/trimethoprim (SXT 25μg) were applied with a fine pointed forceps on the inoculated plates and incubated in 37°C for 24h. Then measure the sensitivity by measuring the clear zone of inhibition around the

disks and the interpretation was applied according to CLSI (2007).

RESULTS

As seen in Table 4, the recovery percentage was calculated to be between 81-and 98%. The sensitivity was evaluated by determining LOD and LOQ. The calculated LODs were 5, 3, 5, and 1 ppb, while LOQs were 10. 6, 10, and 2 ppb for Chlorpyrifos, Cyhalothrin, Lufenuron, and malathion, respectively (Table 4).

Table 4. Recovery % for spiking detected pesticide residues by LC-MS/MS.

Compounds	Spiking Recovery %	LOD (ppb)	LOQ (ppb)
Chlorpyrifos	87	5	10
Cyhalothrin	92	3	6
Lufenuron	81	5	10
Malathion	98	1	2

In this study, the pesticide detected in Alfalfa fodder were Chlorpyrifos $(4.85\pm1.23 \text{ and } 5.71\pm1.6 \text{ ppm})$ in both non-sprayed and sprayed alfalfa fodder, and Cyhalothrin $(3.22\pm0.79 \text{ ppm})$, Lufenuron (2.76 ± 1.43) , and Malathion (1.78 ± 0.58) in sprayed alfalfa fodder (Table 5).

Pesticide residues in milk collected from dairy animals were shown in Table (5). On a dairy cow's farm, pesticide residues were undetectable.

Dairy cows at the valley were fed with non-pesticide applied alfalfa fodder and the detection and incidences of chlorpyrifos and malathion were 4%, 1.33%, and residue levels of 0.017, 0.017 ppm, respectively. All found pesticides were less than the documented MRL. Otherwise in dairy buffalos, only chlorpyrifos was detected in milk by the incidence of 11.4

% and a mean value of 0.01 ppm.

In the valley studied pesticides are used extensively for Agricultural purposes (Table, 6). Since alfalfa was grown intensively and pesticides were used in the region, especially pesticides incidence in milk cows was 57.3, 21.33, 5.33, and 1.33%, as well as detected residues, were 0.037, 0.026, 0.039, 0.019 ppm, for Chlorpyrifos, Cyhalothrin, Lufenuron, and Malathion, respectively.

Buffalos' milk at the same zones were contain higher incidence (60, 51.43, 14.28, and 5.71%) and mean residue levels of 0.041, 0.024, 0.034, 0.017 ppm, for Chlorpyrifos, Cyhalothrin, Lufenuron, and Malathion, respectively.

Table 5. levels of estimated pesticide in Clover bush (Alfalfa fodder) residues comparing with its maximum residue levels.

Group	No of						Pesticid	e (ppm)					
	sam-	(Chlorpyrifos		Cyhalothrin			Lufenuron			Malathion		
	ples	No	%	Mean ± SE	No.	%	Mean ± SE	No.	%	Mean ± SE	No	%	Mean ± SE
Group 1	20	-	UD	-	-	-	UD	-	UD	-	-	UD	-
Group 2	20	3	15	4.85 ± 1.23 (2)	-		UD	-	UD	-	1	5	-
Group 3	20	17	85	5.71 ± 1.6 (13)	7	35	3.22 ± 0.79 (7)	4	20	2.76 ± 1.43	2	10	1.78 ± 0.58
MRL			5 mg/kg*		1 p	pm*, 2 ppr	n**			-			-

Group1: Alfalfa fodder cultivated in farm

Group 2: Alfalfa fodder cultivated in valley did not use specific pesticide for Alfalfa fodder

Group 3: Alfalfa fodder cultivated in valley use pesticide for Alfalfa fodder.

samples exceeded the permissible limits

Table 6. Pesticide residues in milk collected from dairy animals at the three studied zones comparing with MRL.

Group	Spe- cies	No of ani-	Pestio	cide (pp	m)									
		mals	Chlorpyrifos		Cyhalothrin		Lufenuron			Malathion				
		Cow 90	No.	%	Mean ± SE	No.	%	Mean ± SE	No.	%	Mean ± SE	N o.	%	Mean ± SE
Group 1	Cow	80	-		UD	-		UD	-		UD	-		UD
Group 2	Cow	75	3	4	0.017 ± 0.011	-		UD	-		UD	1	1.3	0.017
	Buffa- lo	35	4	11. 4	0.021 ± 0.012	-		UD	-		UD	-		UD
Group 3	Cow	75	43	57. 3	0.037 ± 0.014(15)	16	21.3 3	0.026 ± 0.017	4	5.33	0.039 ± 0.024	1	1.3	0.019
	Buffa- lo	35	21	60. 0	0.041 ± 0.027(21)	18	51.4 3	0.024 ± 0.012	5	14.2 8	0.034 ± 0.022	2	5.7 1	0.017
MRL			0.02 1	mg/kg*, mg/kg**	**			0.2 mg/ kg*			0.15 mg/ kg*			0.02 mg/ kg***

Group 1: Dairy cow's farm Group 2: Animal feeding without spray Clover bush sprayed Clover bush

^{*}Chlorpyrifos, Regulation (EU) 2015/399, Pesticide residue(s) and maximum residue levels (mg/kg),http://www.fao.org/fao-who-codexalimentarius/codex-texts/dbs/pestres/commodities-detail/en/?lang=en&c_id=20

^{**} Australian MRLs of 1 ppm for green animal feeds and 2 ppm for straw fodder (MacLachlan, 2020).

Group 3: Animal feeding on

^{*}Codex alimentations, International Food Standards FAO, WHO, Pesticide Database, http://www.fao.org/fao-who-codexalimentarius/codex-texts/dbs/pestres/pesticides/en/

^{**}Australia Codex (MacLachlan, 2020).

^{***}Malathion, Regulation (EU) 2015/399, Pesticide residue(s) and maximum residue levels (mg/kg), https://ec.europa.eu/food/plant/pesticides/eu-pesticides-database/mrls/?

event=details&pest_res_ids=143&product_ids=&v=1&e=search.pr

^{****}USA Codex (MacLachlan, 2020).

From a microbial presence view, the isolation incidences in milk samples of dairy cows' farms were 5, 7.5, and 3.75% of *E. coli, Staph aureus,* and *Streptococcus agalactiae*, respectively.

In milk samples of both cows and buffalos, the isolated microorganisms were Klebsiella, Staph aureus, Streptococcus agalactiae, and E. col showing the percentage of isolation of 10, 1.8, 7.3, 6.4%,

respectively.

On the other hand, Cows & buffalos milk samples obtained from valley pesticide applied alfalfa showed the incidence of 13.64, 4.5, 16.4, 15.5% of *E. coli, Klebsiella, Staph aureus*, and *Streptococcus agalactiae*, respectively. These results are significantly different with the value of P<0.05 from that obtained in valley non-applied pesticide according to the Fischer Exact Probability Test.

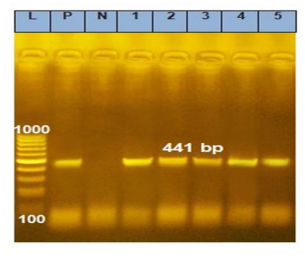
Table 7. Results of isolation of bacteria from milk samples.

Group	Species	No of animal			I	solates fron	n milk samp	ole			
				coli 30)		lla pneu- ae (7)		coccusau- s (30)	Streptococcus aga- lactia (27)		
				No	%	No	%	No	%	No	%
Group1	Cow	80	4	5	0	0	6	7.5	3	3.75	
	Total	80	4	5a	0	0a	4	5a	3	3.75a	
Group2	Cow	75	6	8	1	1.3	6	8	5	6.7	
	Buffalo	35	5	14.3	1	2.9	2	5.7	2	5.7	
	Total	110	11	10b	2	1.8b	8	7.3b	7	6.4b	
Group3	Cow	75	8	10.7	3	4	11	14.7	14	18.7	
	Buffalo	35	7	20	2	5.7	7	20	3	2.7	
	Total	110	15	13.64c	5	4.5c	18	16.4c	17	15.5c	
Total		300	30	10	7	2.33	30	10	27	9	

a, b, c significantly difference at P< 0.05 using Fischer Exact Probability test.

Results of PCR bacterial isolates and sensitivity test:

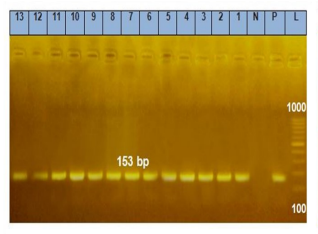
Figures 2-5 show the result of molecular typing of *Klebsiella, E. coli, Streptococcus agalactiae and Staphylococcus aureus* by using PCR following the gel running



1000 720 bp

Fig (2): Results of molecular typing of Klebsiella pneumonia gyrA gene by PCR. L-100bp DNA marker.P-control positive Klebsiella pneumonia strain N-control negative Klebsiellapneumonia strain 1 -5 Positive isolates Klebsiellapneumonia at 441 bp

Fig (3): Results of molecular typing of *E. coliphoA* gene by PCR. L-100bp DNA marker.P-control positive *E. coli* strain N-control negative *E. coli*strain 1 -12-Positive isolates of *E. coli* at 720 bp



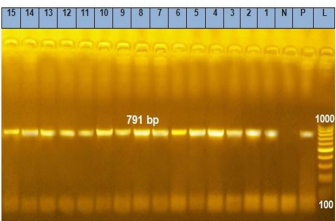


Fig (4): Results of molecular typing of *Strept* agalactiae *Cfb*genebyPCR. L-100bp DNA marker.P-control positive *Strept* agalactiaestrain N-control negative *Strept* agalactiae strain 1 -13-Positive isolates of rRNA at 153bp

Fig (5): Results of molecular typing of *Staphylococcus aureus16S rRNA* gene by PCR. L-100bp DNA marker. P-control positive *Staphylococcus aureus* strain N-control negative *Staphylococcus aureus* strain 1 -15-Positive isolates of 16SrRNA at 791bp

Tables, 8-11 antibiotic sensitivity pattern of bacterial isolates in milk samples collected from dairy cow's farm, cows & buffalos feeding non-applied clover at Ezbet Barghout as well as cows & buffalos feeding on sprayed

Clover bush at Ezbet Furqan, Fayoum show province. These tables show the increased tested bacterial resistance by sequential order

Table 8: Antibiotic sensitivity pattern of bacterial isolates in milk samples collected from Animal feeding without spray Clover bush group 1 (dairy cow's farm) and group 2 (Ezbet Barghout), Fayoum province.

Antimicrobial		E. coli(n = 15)		Klebs	siella pneu (n=2)	ımonia	Staphy	clococcus (n = 14)	aureus	Strepto	ococcus ag (n = 10)	alactia
	R (%)	I (%)	S (%)	R(%)	[(%)	S (%)	R (%)	I (%)	S (%)	R (%)	I (%)	S (%)
Clarythromycin (CLR 15µg)	4 (26.7)	3(20)	8 (53.3)	0(0)	1 (50)	1(50)	2 (14.3)	5 (35.7)	5 (35.7)	0(0)	2(20)	8(80)
Gentamycin (CN 10 μg)	5 (33.3)	3(20)	7 (46.7)	0(0)	1 (50)	1(50)	2 (14.3)	2 (14.3)	10 (71.4)	2(20)	1(10)	7(70)
Amikacin (AK 30μg)	3(20)	5 (33.3)	7 (46.7)	1(50)	0(0)	1(50)	2 (14.3)	4 (28.6)	8 (57.1)	2(20)	3(30)	5(50)
Ampicil- lin+Sulbactam (SAM20µg)	2 (13.3)	5 (33.3)	8 (53.3)	0(0)	1 (50)	1(50)	0(0)	2 (14.3)	12 (87.5)	2(20)	4(40)	4(40)
Cefotaxim (CTX 30µg)	3(20)	5 (33.3)	7 (46.7)	0(0)	0(0)	2 (100)	1(7.1)	0(0)	13 (87.5)	1(10)	4(40)	5(50)
Amoxicillin (AMX 10µg)	7 (46.7)	5(25)	3(50)	1(50)	0(0)	1(50)	2 (14.3)	4 (28.6)	8 (57.1)	3(30)	3(30)	4(40)
Cefepime (CFM 30 μg)	0(00)	1(6.7)	14 (93.3)	0(0)	0(0)	1(50)	0(0)	2 (14.3)	6 (100)	2(20)	6(60)	2(20)
Amoxicillin +Clavulenic acid (AMC 30µg)	2 (13.3)	1(6.7)	12 (80)	0(0)	1 (50)	1(50)	4 (28.6)	4 (28.6)	6(50)	3(30)	4(40)	3(30)
Spiramycin (SP 100 μg)	3(20)	2 (13.3)	10 (63.6)	1(50)	0(0)	1(50)	3 (21.4)	2 (14.3)	9(75)	4(40)	2(20)	4(40)
Ampicillin (AM 10 μg)	6 (45.4)	5 (33.3)	4 (26.7)	1(50)	0(0)	1(50)	2 (14.3)	4 (28.6)	9 (62.5)	5(50)	4(40)	1(10)
Sulphamethoxazole + Trimethoprim (SXT 25µg)	7 (46.7)	3(20)	5 (33.3)	2 (100)	0(0)	0(0)	3 (21.4)	4 (28.6)	7(50)	4(40)	6(60)	0(0)
Average Resistance		25.45 %			27.27 %)		13.63 %			25.45 %	

R=Resistance, I=Intermediate, S=Sensitive, n=number

Table 9: Antibiotic sensitivity pattern of bacterial isolates in milk samples collected from animal feeding on sprayed Clover bush, Ezbet Furqan, Fayoum province.

Antimicrobial	E. coli (n¥¥5)				Kellebsiella pneu- moniae (n=5)			elococcu s (n = 18		Streptococcus agalac- tiae (n = 17)		
	R (%)	I (%)	S (%)	R (%)	I (%)	S (%)	R (%)	I (%)	S (%)	R (%)	I (%)	S (%)
Clarythromycin (CLR 15µg)	9(60)	4 (26.7)	2 (13.4)	2 (40)	2 (40)	1 (20)	13 (72.2)	2 (11.1)	3 (16.7)	11 (64.7)	2 (11.8)	4 (23.5)
Gentamycin (CN 10 μg)	8 (53.4)	2 (13.3)	5 (33.3)	2 (40)	1 (20)	2 (40)	8 (44.4)	2 (11.1)	8 (44.4)	7 (41.2)	2 (11.8)	8 (47.1)
Amikacin (AK 30μg)	7 (46.6)	4 (26.7)	4 (26.7)	3 (60)	1 (20)	1 (20)	5 (27.8)	5 (27.8)	8 (44.4)	6 (35.3)	0(0)	11 (64.7)
Ampicillin +Sulbactam (SAM 20µg)	8 (53.4)	4 (26.7)	3 (23.1)	2 (40)	2 (40)	1 (20)	6 (33.3)	7 (38.9)	5 (27.8)	4 (23.5)	6 (35.3)	7 (41.2)
Cefotaxim (CTX 30µg)	6 (40)	2 (13.3)	7 (46.6)	2 (40)	1 (20)	2 (40)	9 (50)	3 (16.7)	6 (33.3)	8 (47.1)	2 (11.8)	7 (41.2)
Amoxicillin (AMX 10µg)	12 (80)	2 (13.4)	1 (6.7)	3 (60)	1 (20)	1 (20)	14 (77.8)	2 (11.1)	2 (11.1)	9 (52.9)	4 (23.5)	4 (23.5)
Cefepime (CFM 30 µg)	4 (26.7)	3 (23.1)	8 (53.3)	2 (40)	2 (40)	1 (20)	8 (44.4)	3 (16.7)	7 (38.9)	10 (58.8)	3 (17.6)	4 (23.5)
Amoxicillin + Clavu- lenic acid (AMC 30μg)	7 (46.6)	5 (33.3)	3 (23.1)	0 (0)	2 (40)	3 (60)	7 (38.9)	3 (16.7)	8 (44.4)	6 (35.3)	4 (23.5)	7 (41.2)
Spiramycin (SP 100 μg)	9 (60)	4 (26.7)	2 (13.3)	0 (0)	3 (60)	2 (40)	3 (16.7)	5 (27.8)	8 (44.4)	4 (23.5)	3 (17.6)	10 (58.8)
Ampicillin (AM 10 μg)	12 (80)	3 (20)	0 (0)	4 (80)	1 (20)	0 (0)	12 (66.6)	4 (22.2)	2 (11.1)	10 (58.8)	4 (23.5)	3 (17.6)
Sulphamethoxazole + Trimethoprim (SXT 25µg)	6 (40)	5 (33.3)	4 (26.7)	2 (40)	2 (40)	1 (20)	9 (50)	5 (27.8)	4 (22.2)	11 (64.7)	4 (23.5)	2 (11.8)
Average Resistance	verage Resistance 53.33 %		,		40.0 %		47.47 %			45.99 %		

R=Resistance, I=Intermediate, S=Sensitive, n=number

		Klebsiella	Staphylococcus	Streptococcus aga-
	E. coli	pneumoniae	aureus	lactiae
Animal feeding without spray Clover	25.45 %	27.27 %	13.63 %	25.45 %
Animal feeding on sprayed Clover	53.33 %	40.0 %	47.47 %	45.99 %

Table 10: Average percentage of studied antimicrobial resistance for isolated microorganisms.

a, b, c significantly differences against higher litter using Fischer Exact Probability test at P < 0.05.

DISCUSSION

Mastitis is caused mostly by bacteria. There are several different antibiotics for mammary infection therapeutic and preventive applications. These antimicrobial compounds are usually injected into the infected part after the infection happened

In this work, it was to comprehend that pesticide usage in agricultural areas should be considered as they extend to the milk obtained from grazing animals including cows and buffalos. The effect of pesticides on antibiotic sensitivity causes a big problem in the treatment of mastitis and antibacterial resistance is increasing.

The LOD and LOQ values in this study were lower than the MRLs established by Codex (Codex Alimentarius Commission, 2019) and Australian MRLs (MacLachlan, 2020) for milk and alfalfa fodder samples (Tables, 3 through 5), and these results give high precision to our conclusion.

In the valley where extensively, pesticides were applied in agriculture, especially on alfalfa, the detected pesticides were chlorpyrifos, cyhalothrin, lufenuron, and malathion (Table 4). While in non-sprayed alfalfa fodder was Chlorpyrifos only. In contrast, alfalfa fodder cultivated on the dairy farm was free of any pesticides. Data analysis proved that percentage of pesticide residues was higher and higher than MRL in group 3 than that recorded in group 2. This could be because of using pesticides in Alfalfa culture as well as other pesticides used in cultivated crops. This was augmented by alfalfa fodder taken from cultivated farms that did not contain any pesticide residues.

Although the application of pesticides did not recommend for alfalfa in Egypt (Agricultural Pesticide Committee Recommendation), its usage is still a fact in some areas. In addition, no residue data were provided for alfalfa fodder and there is insufficient information to suggest MRLs for alfalfa fodder (FAO/WHO, 2020).

In this work, estimation of pesticide residues in milk is made based on the tendency of pesticides to transfer to milk with expected alfalfa dietary exposure. Otherwise, some pesticides estimated in this study registered for use on other crops than alfalfa but find their ways to alfalfa.

Pesticide residues in milk show nearly the same pattern in alfalfa fodder (Table, 5). In milking animals fed with pesticide-applied alfalfa fodder (Ezbet Furqan), 4 pesticides were detected in milk such as chlorpyrifos, cyhalothrin, lufenuron, and malathion, by descending order. Milking buffalos in that zones contain higher incidence and pesticide residue levels than milking cows. This observation could be regarded as higher fat percentages in buffalos' milk than that of cow's milk. Also, buffalo's immunity system could be more sensitive to pesticides than cows, this notice needs further studies to prove this phenomenon.

Chlorpyrifos is an organophosphate insecticide used for the control of several insects in harvests. It is enumerated in Egypt on many pastures to control various pests. Generally, it is not grazed or cut for stock food for 2 days after application (MacLachlan, 2020). The maximum transferred factor for cattle feeding at 30 ppm Chlorpyrifos in the feed was 0.007 ppm for milk (JMPR, 2000). Expected Chlorpyrifos residues from feeding alfalfa fod-

der with residues of 5.17 ppm at 100 % of the diet 5.17 ppm $\times 0.007 = 0.036$ ppm, nearly similar to that detected in this study (0.037 & 0.041 in cattle and buffalos, respectively), where these animals depending mainly on alfalfa fodder in their diet during the studied winter season.

Lambda-cyhalothrin is a synthetic pyrethroid used to control various insects in crops. It is registered on pasture for control of many pests in Egypt. It is residues in cattle fat degenerated with a half-life of 7-9 days. The recommendation for its application for stock food is 7-14 days after application. Presume the residues of cyhalothrin in alfalfa fed to animals 3.22 ppm (obtained result, Table, 4), cyhalothrin, residues in milk fat were 0.5 ppm (Cyhalothrin, 1999) and total fat for milk is 0.02 ppm, thus cyhalothrin, anticipated maximum residues in whole milk would be $3.22 \times 0.5 \times 0.02 = 0.032$ mg/kg (MacLachlan, 2020).

Regarding Malathion (organophosphorus pesticide) and <u>Lufenuron</u> (agricultural pesticide), insufficient data were located to offer a confidential view of livestock residue risks (MacLachlan, 2020).

As stated by Fisher et al., (2015), pesticide residues in milk might have some potential sources. Whatever the source does not matter and pesticides reach into the milk of the lactating animal.

Out of 300 milk samples of cows and buffalo cows, 4 microorganism isolates could be identified (Table, 6) with different incidences. The isolated microorganisms were *E. coli, Klebsiella, and Staph. aureus* and *Streptococcus agalactia*.

To get reliable results on microbial etiology in mastitic animals PCR confirmation occurred (Figures 1-3). Since cases of mastitis in the current study were sampled under pesticide polluted feed criteria and originated from pesticide applied alfalfa, the relation among them was evaluated. The two most common udder pathogens, *E. coli S. aureus*, and *Streptococcus agalactiae* were almost found in other investigations (Ericsson *et al.*, 2009).

The percentage of microorganisms recovered from dairy animals feeding on sprayed clover bush was significantly higher than that from animals feeding without spray clover

bush subsequently higher than milking cows on a closed farm. This could be <u>regarded</u> as restricted biosafety and biosecurity roles applied in farms compared with that in rural animal husbandry. *E. coli* causes mastitis in dairy cows and buffalos around parturition and early during lactation with arresting local and sometimes severe clinical symptoms (Christian *et al.*, 2003).

In this concern, many management factors affect dairy buffalos or cow's conditions during late pregnancy. Pesticides detected in animal feed might be the cause of metabolically immunocompromised buffalos and cows that cause lower resistance to stress-induced by parturition and early lactation, so, animals become highly susceptible to environmental pathogens. There is a tendency to believe that low milk SCC fails to protect the udder from environmental pathogens (Suriyasathaporn *et al.*, 2000). This assumption is based on epidemiological data (Shuster *et al.*, 1996).

Cow and Buffalo mastitic milk (Tables 8-10), showed antimicrobial susceptibility and a resistant pattern of the mastitis-causing organism. Generally, bacterial isolated from pesticide residues milk demonstrated the highest resistance rates to Gentamycin, Amikacin, Ampicillin +Sulbactam, Cefotaxime, Amoxicillin, and Ampicillin. This pattern of resistance could be regarded as excessive use of antibiotic and/or pesticide residues detected in milk. The relevance of pesticides to AMR development is alarming. More evidence of pesticides as agents disturbing bacterial antibiotic susceptibility and generating transient adaptive responses is imperative. The role of pesticides, not only as toxins, but as pathways to AMR must be further evaluated to address the current crisis of antibiotic resistance and to raise awareness of the need for environmental monitoring and regulation (Malagón-Rojas et al

Occurrence of the highest number of *E. coli* and *Staph. aureus* agrees with those reported by Akram et al. who reported a higher isolation rate of E. coli and Staphylococcus in studied milk samples in India (Akram *et al.*, 2007). Higher *E. coli* and *Staphylococcus* incidence in this work could be a reference to the poor hygienic practices in the dairy environment at this valley, while these organisms initiated from the

milking environment and contaminated the udder via the teat canal. In the case of environmental mastitis, the contagion of the ending of the teat is a major predisposing factor (Akram et al., 2007, Sarne et al., 2018). In addition, these observations might be due to harboring the organism in the skin, udder, and milk of the infected gland which acts as a reservoir.

Taking into consideration that all studied animals were taken from the same city but with different districts, the significant differences in antimicrobial resistance patterns could be regarded as excessive pesticide exposure to dairy animals.

In conclusion, this study proved that pesticide pasture and/or feed contamination cause easier access for environmental pathogens to the udder through their open teats during calving and/or drop-in udder immunity. More studies must be needed to prove the observed correlation between pesticide-contaminated milk and antimicrobial resistance. Despite the great efforts made by the Agricultural Pesticides Committee in Egypt, more control overuse the pesticides in Egyptian villages and hamlets must be applied

REFERENCES:

- Akram M, Shahid M, Khan AU. (2007). Etiology and antibiotic resistance patterns of community-acquired urinary tract infections in J N M C Hospital Aligarh, India. Ann Clin Microbiol Antimicrob.; 6:4. doi: 10.1186/1476-0711-6-4.
- Brisse, S. and Verhoef, J. (2001): Phylogenetic diversity of *Klebsiella pneumonia* and *Klebsiellaoxytoca* clinical isolates revealed by randomly amplified polymorphic DNA, *gyrA* and *parC* gene sequencing and automated ribotyping. *Int J SystEvolMicrobiol*. 2001;51:915-924.
- **CAPMAS (2018).** CAPMAS Egypt UN Demographic Yearbook.
- Christian, B., Valérie Van M., Jalil M., Araceli D., Luc D. (2003): Severity of E. coli mastitis is mainly determined by cow factors. Vet Res; 34(5):521-64., doi: 10.1051/vetres:2003023.
- CLSI (2007). Performance Standards for Antimicrobial Susceptibility Testing; Seventeenth Information Supplement. CLSI Document M100-S17 (M2-A7 and M7-A7), Clinical and laboratory Standards institute, Wayne, 27(1).

- Codex Alimentarius Commission (2019). Procedural Manual Twenty-seventh edition, Secretariat of the Codex Alimentarius Commission Joint FAO/WHO Food Standards Programme Food and Agriculture Organization of the United Nations Viale delle Terme di Caracalla 00153 Rome, Italy.
- Cooper J and Dobson H. (2007): The benefits of pesticides to mankind and the environment. Crop Prot.; 26: 1337–1348.
- Cyhalothrin, 'Environmental health criteria' (1999) 1.Pyrethrins adverse effects 2.Pyrethrins toxicity I. Series ISBN 92 4 154299 3 (NLM Classification: WA 240) ISSN 0250-863X
- Ericsson <u>U.H., Lindberga, A., Persson Wallera, K., Ekman, T., Artursson K., Nilsson-Öst M., Bengtsson</u> B. (2009). Microbial aetiology of acute clinical mastitis and agent-specific risk factors. Vet Microbiol, 137, 1–2, 90-97
- FAO/WHO (2020): Pesticide residues in food 2019 Joint FAO/WHO Meeting on Pesticide Residues, Rome. https://www.who.int/foodsafety/areas_work/chemical-risks/

JMPR_2019_Sep_Report.pdf

- Fisher, W.J.; Schilter, B., Tritscher, A.M. (2015): Contaminants of Milk and Dairy Products: Contamination Resulting from Farm and Dairy Practices. Reference Module in Food Sciences. Elsevier, pp. 1–13. doi: http://dx.doi.org/ 10.1016/B978-0-08-100596-5.00698-3,
- Getahun K., Kelay B., Bekana M. and Lobago, F. (2007). Bovine mastitis and antibiotic resistance patterns in Selalle smallholder dairy farms, central Ethiopia. Trop. Anim. Health Prod., 40 (4). 261-268.
- HDR (2008). Human Development Report, United Nations Development Programme, https://hdr.undp.org.content.human-development-Jan 1, 2008.
- Hu, Q.; Tu, J.; Han, X.; Zhu, Y.; Ding, C. and Yu, S. (2011): Development of multiplex PCR assay for rapid detection of *Riemerella anatipestifer, Escherichia coli*, and *Salmonella enterica* simultaneously from ducks. Journal of Microbiological Methods 87 (2011) 64–69.
- **ISO/IEC 17025, 2005.** General requirements for the competence of testing and calibration laboratories. Geneva: ISO. pp: 05-15.
- JMPR (2000). Joint FAO/WHO Meeting on Pesticide Residues. https://www.who.int/ groups/joint-fao-

- who-meeting-on-pesticide-residues-(jmpr)/publications/reports
- Konikkara, K.P.; Baliga, S.; Shenoy, S. Bharati, B. (2014): Evaluation of Culture, Antigen Detection and Polymerase Chain Reaction for Detection of Vaginal Colonization of *Group B Streptococcus (GBS)* in Pregnant Women. Journal of Clinical and Diagnostic Research; 8(2):47-49.
- Leach, K.A., M.J. Green, J.E. Breen, J.N. Huxley, R. Macaulay, H.T. Newton, and A.J. Bradley. 2008. Use of domestic detergents in the California mastitis test for high somatic cell counts in milk. Vet Rec, 163:566–570.
- Lehotay, S.J., K. Mastovska and A.R. Lightfield, (2005): Use of Buffering and Other Means to Improve Results of Problematic Pesticides in a Fast and Easy Method for Residue Analysis of Fruits and Vegetables. Journal Of AOAC International.
- MacLachlan, D. (2020): Pesticide risk profile for the grazing of pasture and/or cutting of hay and feeding to cattle and sheep. Residues and Food Safety, Last reviewed: June 2020, Australian Quarantine and Inspection Service, Australian Governorate. https://www.safemeat.com.au/globalassets/safemeat/residue-risk-pdfs/pasture_feb2010-5.pdf
- Magnusson, B. and U. Ornemark, 2014. Eurachem Guide: The Fitness for Purpose of Analytical Methods A Laboratory Guide to Method Validation and Related Topics, ISBN 978-91-87461- 59-0. Available from www.Eurachem.org.
- Malagón-Rojas JN, Parra-Barrera EL, Lagos L (2020). From environment to clinic: the role of pesticides in antimicrobial resistance. Rev Panam Salud Publica. 2020;44:e44. https://doi.org/10.26633/RPSP.2020.44
- Mason, W.J.; blevins, J.S.; beenken, K.; wibowo, N.; ojha, N. and Smeltzer, M.S. (2001): Multiplex PCR Protocol for the Diagnosis of Staphylococcal Infection. JOURNAL OF CLINICAL MICROBIOLOGY, Vol. 39, No. 9, p. 3332–3338.
- Özkara, A., Akyıl, D., and Konuk, M. (2016). Pesticides, Environmental Pollution and Health, in Environmental Health Risk Hazardous Factors to Living Spe-

- cies, 1-27 http://dx.doi.org/10.5772/63094 Rysanek, D., Babak, V. and M. Zouharova (2007). Bulk tank milk somatic cell count and sources of
- Bulk tank milk somatic cell count and sources of raw milk contamination with mastitis pathogens. Veterinarni Medicina, 52, 2007 (6): 223–230.
- **SANCO (2013).** European Commission Health & Consumer Protection Directorate-General. Safety of the food chain Chemicals, contaminants, pesticides 19 November 2013 rev. 0 Guidance document on analytical quality control and validation procedures for pesticide residues analysis in food and feed. SANCO/12571/2013 Supersedes SANCO/12495/2011 Implemented by 01/01/2014
- SANTE/11945/2015. (2015). European commission, Guidance document on analytical quality control and method validation procedures for pesticides residues analysis in food and feed.
- Sarne D., Ian O., Sofie P. (2018). Management and prevention of mastitis: A multifactorial approach with a focus on milking, bedding and data-management. Journal of Integrative Agriculture 2018, 17(6): 1214–1233
- Shazia A. and Karam A. (2017): Pesticides Residue in Milk and Milk Products: Mini Review. Pak. J. Anal. Environ. Chem. 18 (1): 37 45
- Shuster D.E., Lee E.K., Kehrli M.E. (1996). Bacterial growth, inflammatory cytokine production, and neutrophil recruitment during coliform mastitis in cows within ten days after calving, compared with cows at midlactation, Am. J. Vet. Res. 14. 1569–1575.
- Suriyasathaporn W., Schukken Y.H., Nielen M., Brand A. (2000): Low somatic cell count: a risk factor for subsequent clinical mastitis in a dairy herd, J. Dairy Sci. 83. 1248–1255.
- Ying, Lv., Xuefen, Li., Zongyi, W., Han, Z., Qi, Z., Ran H., Xiangning, C. and Tao, H. (2014): Short communication: Interaction of bovine milk protein with chlorpyrifos. Journal of Dairy Science, 97 (4): 2056-2060

Egyptian Jour	rnal of Anima	l Health 3, 1	! (2023),	<i>67-82</i>
---------------	---------------	---------------	-----------	--------------