Original Research Article

Novel Combination contains probiotic bacterial and Yeast strains to reduce of Aflatoxin M1 in Milk

ABSTRACT

1 2

3

11

Milk and milk products are important contribution to the human diet especially children. However, the presence of aflatoxins as AFM1 in milk and milk products are considered undesirables due to their health risks in consumer's body. For that reason this study aimed to assess the ability of some microbial strains on aflatoxin removal especially the AFM1 in the milk. AFM1 residue was determined by HPLC after different incubation period (12h, 24h, 48h and 72h) of the probiotic bacteria and/or yeasts in PBS as model for AFM1 determination. This study was performed during the period between 2015 and 2017. The combination of nonviable probiotic bacterial and yeast strains (*Lactobacillus plantarum, Lactobacillus acidophilus, Bifidobacterium bifidum, Kluyveromyces lactis* and *Saccharomyces cerevisiae*) succeeded to reduce AFM1 from 50 (ng/ml) during the incubation periods; 12h, 24h, 48h and 72h, into 9.72±1.31, 6.68±0.55, 5.70±0.33 and 4.56±0.15 ng ml-1, respectively. The highest AFM1 removal % was recorded as; 80.56%, 86.64%, 88.60% and 90.88% in the treated milk samples in respective manner. Sensor evaluation was carried out using Yoghurt as model in sample size 50 gm for each sample.

Conclusion: This study concluded that the combination consisting of probiotic bacteria and yeasts could be used in reducing the concentration of the AFM1 in aflatoxin contaminated milk.

Keywords: Probiotic bacteria, Yeasts, Aflatoxin M1, Contamination and Milk.

Comment [SC3]: delete

Comment [SC1]: species

Comment [SC2]: species

15 **1. INTRODUCTION**

16

12 13

14

17 Aflatoxins are a group of mycotoxins which are considered as the most potent carcinogens. 18 Aflatoxins can not only be found as contaminants in the stable diet (cereal grains) but also 19 are found in milk when the dairy animals ingest contaminated feed with aflatoxin B1 and B2 20 [1-3]. Aflatoxin B1 (AFB1) is converted by the normal metabolism process to aflatoxin M1 but 21 aflatoxin B2 (AFB2) is converted to aflatoxin M2 and then aflatoxin M1 and M2 are excreted 22 and occurred in milk so AFM1 and AFM2 are considered as hepatic hydroxlated metabolites 23 of aflatoxin B1 and B2 [4-6]. Milk and dairy products are considered as very important part of 24 human diet food habit in every home with high rate of consumption for all age because milk 25 is high in nutritional value which maintains the human health. However, it may be act as a

26 27 28 29 30 31 32 33 34 35 36	vehicle of contaminants such as aflatoxins which cause various physiological risks effects in human consumers especially the children who are considered more group susceptible than adults to aflatoxins effects as growth retardation, stunning and liver cancer [7-8]. Aflatoxin M1 (a member of aflatoxins) may be found in breast milk, animal milk and different dairy products. AFM1 has linear relationship with the aflatoxin B1 in animal feed that is ingested by dairy animals. AFM1 is stable in raw milk and different processed products from milk which does not destroyed by pasteurization or heat treatments. Cream separation from milk has a small effect on AFM1 amount in skim milk because AFM1 prefers the binding with the casein (milk protein). The maximum concentration acceptable limit of aflatoxin M1 permitted in milk consumption by humans is 0.5 ppb (parts per billion) that is established by Egyptian standard specification (E.S.S) and European standard regulation [9-11].	
37	AFM1 is more specifically a problem of food safety than a problem of Hygiene which cause	Comment [SC4]: hygiene
39	innovative solutions for reducing and inhibiting health risks of aflatoxin and overlook the	
40 41	issue of atlatoxin exposure by using certain problotic strains which can bind with atlatoxin to form the complex problotic-atlatoxin and then improve elimination of this complex from the	
42	gut through feces. So that Therefore, this biological strategy prevent the absorption of these	Comment [SC5] : Therefore, this biological
43 44	aflatoxin in human and animal bodies through gastrointestinal tract, improve aflatoxin decontaminating from body and minimize potential risks of aflatoxin [12-14].	stategy prevents
45		
45 46	in the diet, lower the risks and enhance the health. These biological methods use to	Comment [SC6]: is considered
47	sequestrate the aflatoxin M1 without affecting the nutritional value, taste of the milk products.	
48	Some probiotic strains like Bifidobacterium Bifidum, Lactobacillus plantarum, Lactobacillus	
49 50	acidophilus have the ability to minimize risks of atlatoxin M1 and also some types of yeast as	Commont [SC7]: Klumaramaa
51	M1 from milk and milk products [15-18]. For that reasons this study aimed to find a microbial	Comment [SC7]: Kuyveromyces
52	combination which able to reduced and control the toxicity resulted from afaltoxins in	Comment [SC8]: aflatoxins
53 54	contaminated milk especially the AFM1.	
55	2 MATERIAL AND METHODS	
56		
57	MICROBIAL STRAINS COLLECTION	
58	All the bacterial strains and yeasts were kindely obtained from	Comment [SC9]: kindly
59	microbiological resources centers (Cairo MIRCEN, Egypt)	Comment [SC10]: Change to arial font 10
60 61	2.1 Standard aflatoxin M1 (AFM1) solutions	
62 63	Standard solution of AFM1 (10 µg/ml) was obtained from Sigma-Aldrich (St. Louis, MO, USA) stock standard solution of AFM1 was prepared by dissolving standard in benzene: acetopi	. A trile
64	(98.2, v/v) until used in the test quantitative measurement of aflatoxin M1 in milk and d	airy
65	products as described by AOAC (2000) [19-20]. Another stock standard solution of AFM1 v	was
66 67	prepared by dissolving standard in PBS at concentration 50 ng ml ⁻¹ till used in test of	the
68	solution was packed in amber vials to protect the work concentration from the light and t	hen
69	stored at 4 °C in refrigerator.	
70 71	2.2 Evoluation the ability of some probletic visible station (Lestabasillus statement	
70	Lactobacillus acidonbilus and Bifidobacterium bifidum) on aflatovin M1 reduction	Comment [SC11]: species
73		

Lactobacillus plantarum, Lactobacillus acidophilus and Bifidobacterium bifidum are some of 74 75 probiotic viable strains which were selected based on their use as probiotic cultures in dairy 76 industry on available information concerning their effects on reduction of aflatoxins in aqueous 77 solution. Several types of lactic acid bacteria (LAB) have binding ability with AFM1 in liquid media 78 and milk solution [21]. 79

80 2.2.1 Preparation of probiotic bacterial strains

81

99

82 Each probiotic bacterial strain (Lactobacillus plantarum, Lactobacillus acidophilus and Bifidobacterium bifidum) was cultivated individually in De-Man-Rogosa-Sharpe broth (MRS) 83 84 supplemented with 0.05% L-cysteine at pH 6.5 and incubated at anaerobic conditions at 37 °C for 85 24 h in anaerobic shaker incubator at 200 rpm with 5% CO₂ [22]. Each probiotic bacterial strain 86 was placed in centrifuge at (4000 rpm, 4 °C and 15 min) to harvest its cells in pellet then washed 87 by phosphate buffer saline (PBS) twice. The pellet of each strain was suspended in PBS at pH 6.8 to determine optical density (OD) by using spectrophotometer at wavelength of 600 nm. Then the 88 suspension were adjusted into different starting concentration treatment at $OD_{600} 0.72 \pm 0.03$ equal 1×10^9 CFU ml⁻¹, $OD_{600} 2.16 \pm 0.03$ equal 3×10^9 CFU ml⁻¹ and $OD_{600} 3.6 \pm 0.035$ equal 5×10^9 CFU 89 90 91 ml⁻¹. The suspension was diluted with PBS until reaching the required concentration treatment. 92 Also, the treatment dose of combination probiotic bacterial strains was prepared by taken equal 93 amount from each bacterial strain at 5×10⁹ CFU ml⁻¹ to give 1ml PBS had three probiotic bacterial strains (Bifidobacterium bifidum DSM 20082, Lactobacillus plantarum DSM 20174 and 94 95 Lactobacillus acidophilus DSM 20079) [23-25]. 96

97 2.3 Binding ability of the viable strains of (Lactobacillus plantarum, Lactobacillus Comment [SC12]: species 98 acidophilus and Bifidobacterium bifidum) with aflatoxin M1

100 The adjusted inoculum concentration of collected cells were suspended as viable in eppendroff tube containing 1 ml of phosphate buffered saline (PBS) contaminated with aflatoxin M1 at 101 concentration of 0.05 ug ml⁻¹ (50 ng ml⁻¹). The three different concentration of each inoculum 102 strain (1×10⁹ CFU ml⁻¹, 3×10⁹ CFU ml⁻¹ and 5×10⁹ CFU ml⁻¹) in table (1) were mixed with 1 ml 103 104 PBS supplemented with 50 ng ml⁻¹ of aflatoxin M1 followed by incubation at 37°C for different 105 times (12h, 24h, 48h and 72h).

106 Table 1. Viable and Nonviable probiotic strains and inoculum dose of treatment

Viable and Nonviable probiotic strains Inoculum dose of treatment 1×10⁹ CFU ml⁻¹ 3×10⁹ CFU ml⁻¹ Lactobacillus acidophilus DSM 20079 (A) 5×10⁹ CFU ml⁻¹ 1×10⁹ CFU ml⁻¹ 3×10⁹ CFU ml⁻¹ Lactobacillus plantarum DSM 20174 (B) 5×10⁹ CFU ml⁻¹ 1×10⁹ CFU ml⁻¹ Bifidobacterium bifidum DSM 20082 (C) 3×10⁹ CFU ml⁻¹

Comment [SC13]: species

	5×10 ⁹ CFU ml ⁻¹
Combination of probiotic strains (A+ B+C)	5×10 ⁹ CFU ml ⁻¹
+ ve control	PBS + AFM1
	PBS+ Strain (A) without AFM1
ve control	PBS+ Strain (B) without AFM1
- ve control	PBS+ Strain (C) without AFM1
	PBS+ Combination of probiotic strains without AFM

107 Combination of probiotic strains (A+ B+C) = 333.33 µl of each strain at 5×10⁹ CFU ml⁻¹.

108 2.3.1 Measurement of aflatoxin M1

109 Each sample was centrifuged to separate the cells of probiotic strains from the supernatant fluid for analysis by HPLC. The ability of each strain and the combination of strains (Lactobacillus 110 111 plantarum, Lactobacillus acidophilus and Bifidobacterium bifidum) to be adsorbed or metabolized aflatoxin M1,then the remaining aflatoxin M1 was be determined by HPLC which was unbounded 112 113 to the probiotic bacterial strains after the different incubation times (12h, 24h, 48h and 72h). Then 114 the result of remaining aflatoxin M1 amount compared to the positive control and the negative control to evaluate the ability of each strain individually on aflatoxin M1 reduction and to 115 116 investigate the potential of the interaction or combination of the three strains on aflatoxin M1 117 reduction [26]. 118

119 2.3.2 Derivtization of sample

120

A 100 μl triflour acetic acid with 200 μl N-hexane were added to each sample residues,
 followed by shaking with vortex for 30 second and samples were left for 15 min at room
 temperature. Then 900 μl (Water: Acetonitrile, 9:1) were added and mixed well using vortex.

124 The haxane layer was removed and samples were subjected for HPLC analysis.

125 126

127

6 2.3.3 HPLC-FLD Fluorescence detector analysis and Chromatographic conditions

Determination of aflatoxins (AF) were carried out according to [27] Scaglioni and Badial-Furlong (2014), using HPLC system (Model 6000) a solvent delivery system (Model 720) system controller equipped with Fluorescence detector (Model 274) at 360 Ex, and 450 EM.
The separation was achieved with a symmetry column, (150x 4.6 mm i.d), 5µm at a flow rate of I ml min⁻¹ with an isocratic system composed of 1 % acetic acid: Methanol: Acetonitriel (55: 35:10).

135 2.4 Assessment of the potential of nonviable probiotic bacterial and yeast strains on 136 sequestration of AFM1

137 2.4.1 Evaluation of the efficiency of nonviable probiotic bacterial strains (Lactobacillus plantarum, Lactobacillus acidophilus and Bifidobacterium bifidum) on reduction of aflatoxin M1 140

141 The probiotic bacterial strain (*Lactobacillus plantarum*, *Lactobacillus acidophilus* and 142 *Bifidobacterium bifidum*) were centrifuged at 6,000 rpm for 15 min and the pellets were re-143 suspended in 10 ml PBS buffer followed by heat treatment through autoclaving (121 °C and Comment [SC14]: delete

144 1.5 psi for 20 min) to become nonviable by heat treatment (the viability was tested by the 145 culturing the heated microbes and the samples showed no growth were selected). Pellets 146 were further centrifuged at 6,000 rpm for 15 min, washed twice with distilled water, re-147 suspended in PBS (pH 6.8) and the optical densities were measured at 600 nm to adjust the 148 three different concentrations $(1 \times 10^9 \text{ CFU ml}^{-1}, 3 \times 10^9 \text{ CFU ml}^{-1} \text{ and } 5 \times 10^9 \text{ CFU ml}^{-1}$). The 149 experiment was carried out as described in table 2.

 151
 Table 2. Probiotic bacterial strains (Lactobacillus plantarum, Lactobacillus 152 acidophilus and Bifidobacterium bifidum) and inoculum dose of treatment

 152
 acidophilus and Bifidobacterium bifidum) and inoculum dose of treatment

 Nonviable probiotic strains
 Inoculum dose of treatment

	1×10 ⁹ CFU ml⁻¹			
Lactobacillus acidophilus DSM 20079 (A)	3×10 ⁹ CFU ml ⁻¹			
	5×10 ⁹ CFU ml ⁻¹			
	1×10 ⁹ CFU ml⁻¹			
Lactobacillus plantarum DSM 20174 (B)	3×10 ⁹ CFU ml ⁻¹			
	5×10 ⁹ CFU ml ⁻¹			
	1×10 ⁹ CFU ml⁻¹			
Bifidobacterium bifidum DSM 20082 (C)	3×10 ⁹ CFU ml ⁻¹			
	5×10 ⁹ CFU ml ⁻¹			
Combination of 3 probiotic strains (A, B &C)*	5×10 ⁹ CFU ml ⁻¹			
+ ve control	PBS + AFM1			
	PBS+ Strain (A) without AFM1			
	PBS+ Strain (B) without AFM1			
-ve control	PBS+ Strain (C) without AFM1			
	PBS+ 3 probiotic strains without AFM1			

153 *Combination of 3 probiotic strains (A, B &C) = 333.33 ul of each strain at 5×10 9 CFU ml⁻¹.

 154 2.5 Evaluation the efficiency of nonviable yeast strains (*Kluyveromyces lactis* and Saccharomyces cerevisiae)

156

The yeast strains (*Kluyveromyces lactis* and *Saccharomyces cerevisiae*) were used as nonviable strains by heating 10 min in autoclave in three different concentrations (1×10⁹ CFU ml⁻¹, 3×10⁹ CFU ml⁻¹ and 5×10⁹ CFU ml⁻¹) to assess the potential of these nonviable strains on sequestration of aflatoxin M1. The inoculum strains were mixed with 1 ml PBS supplemented with 50 ng ml⁻¹ I of aflatoxin M1 followed by incubation at 37°C for different

162	times (12h, 24h, 48h and 72h).	The experiment was carried out as described in table 3 [28-			
163	32].				
164	-				
165					
166					
167	Table 3. Nonviable yeast	strains (Kluyveromyces lactis and Saccharomyces			
168	cerevisiae) and inoculum dose	of treatment			
	Nonviable veast strains	Inoculum dose of treatment			

	1×10 ⁹ CFU ml ⁻¹
Kluyveromyces lactis (CBS2359) (D)	3×10 ⁹ CFU ml ⁻¹
	5×10 ⁹ CFU ml ⁻¹
	1×10 ⁹ CFU ml⁻ ¹
Saccharomyces cerevisiae (ATCC 64712) (E	3×10 ⁹ CFU ml⁻¹
	5×10 ⁹ CFU ml⁻¹
Combination of yeast strains (D &E)*	5×10 ⁹ CFU ml ⁻¹
+ ve control	PBS + AFM1
	PBS+ Strain (D) without AFM1
- ve control	PBS+ Strain (E) without AFM1
	PBS+ Combination of yrast strains without Al

* Combination of yeast strains (D &E) = 500 µl of each strain at 5×10⁹ CFU ml⁻¹. 169

170 2.6 Evaluation of the potential of the combination of nonviable probiotic and yeast strains on aflatoxin M1 reduction in PBS 171

172

The combination of nonviable probiotic bacterial and yeast strains (5×109 CFU ml-1) were used in concentration of 5×10⁹ CFU ml⁻¹ at equal volume to evaluate the efficiency of this 173 174 combination on binding of aflatoxin M1. The experiment was carried out as described in 175 table 4. The inoculum strains were mixed with 1 ml PBS supplemented with 50 ng ml⁻¹ of aflatoxin M1 followed by incubation at 37°C for different times (12h, 24h, 48h and 72h) 176 177 178 [29,30].

179

180 Table 4. Nonviable probiotic bacterial and yeast strains in PBS

_	Nonviable microbial strains	Inoculum dose of treatment				
-	Combination of probiotic strains (A, B & C)	5×10 ⁹ CFU ml ⁻¹				
	+ yeast strains (D &E) *					
	+ ve control	PBS + AFM1				

6

Comment [SC15]: species

-ve control

PBS + probiotic strains (A, B &C)

+ yeast strains (D &E) without AFM1

* Combination of probiotic strains (A, B &C) + yeast strains (D &E): The cells were mixed in equal
 volumes in 1ml of PBS media.

183 2.7 Evaluation of the potential of the combination of nonviable probiotic bacterial and 184 yeast strains on aflatoxin M1 reduction in skim milk sample 185

The combination of nonviable probiotic bacterial and yeast strains (5×10⁹ CFU/ ml) were used and incubated in skim milk contaminated with aflatoxin M1 at 50 ng ml⁻¹ to evaluate their sequestration effect after different time (12, 24, 48 and 72 hour) of incubation as described in table 5. The skim milk was evaluated previously to detect its freedom from AFM1 before being used in the test. After the binding times occurred, the tubes of the milk test were centrifuged to separate the milk layer in supernatant than the pellets of microbial strains were taken for analysis of AFM1 residues and to determine the removal of aflatoxin M1 in milk by the nonviable combination of probiotic [31-33].

M1 in milk by the nonviable combination of probiotic [31-33].

195Table 5. Dose culture of nonviable combination of probiotic bacterial and yeast196strains on aflatoxin M1 in milk

Nonviable microbial strains	inoculum dose of treatment				
 Combination of probiotic strains (A, B &C) + yeas strains (D &E) *	5×10 ⁹ CFU ml ⁻¹				
+ ve control	Milk + AFM1				
-ve control	Milk + probiotic strains (A, B &C)				
	+ yeast strains (D &E) without AFM1				

* Combination of probiotic strains (A, B &C) + yeast strains (D &E): The cells were mixed in equal
 volumes in 1ml of milk.

199 2.8 Scanning Electron Microscope analysis (SEM)

200 201 Scanning Electron Microscope analysis was used to detect the characterization of the cell 202 walls of the nonviable probiotic bacterial strains (Lactobacillus plantarum, Lactobacillus acidophilus and Bifidobacterium bifidum), the yeast strains (kluyveromyces lactis and 203 204 Saccharomyces cerevisiae) and the combination of bacterial and yeast strains using Energy-205 Dispersive Analysis X-ray (Joel Jsm 6360LA, Japan). The combination strains in each group (mixed probiotics strains, mixed yeast strains and the combination of bacterial & yeast 206 strains) were mixed in equal volume in 1ml PBS media contaminated with AFM1 (50 ng/ml) 207 208 and incubated for 72 h at room temperature as treated sample and without AFM1 as 209 untreated sample. Each combination from mixed probiotics, mixed yeast and the 210 combination of bacterial & yeast strains were separately spread over a clean glass slide, coated with gold particles and photographed using scanning electron microscope (SEM) [30]. 211 212

Comment [SC16]: 5 x 10⁹ CFU ml⁻¹

213 2.9 Sensory evaluation of treatment yoghurt sample by the best efficient 214 combination of probiotic bacterial and yeast strains on aflatoxin M1 215 sequestration

216

Yoghurt was mixed with the combination of nonviable probiotic bacterial and yeast strains 217 218 (treatment).Whenever, the control sample was only Yoghurt without any microbes. The size of each sample was about 50 gm. Yoghurt was prepared from total milk fat obtained from 219 reputable large milk and dairy products supermarket then was boiled for 20 min (to avoid the 220 221 presence of another microbes in the raw milk). Further, that the milk kept to cool to 43°C then added yoghurt starter (S. thermophiles and L. bulgaricus) obtained from the same 222 223 source of milk with shaking to distribute the starter culture in the milk (control sample). Also, 224 milk was inoculated with the combination of nonviable probiotic bacterial and yeast strains (Lactobacillus plantarum, Lactobacillus acidophilus, Bifidobacterium bifidum, kluyveromyces 225 226 lactis and Saccharomyces cerevisiae) at an inoculum dose of treatment at 5×10⁹ CFU m 227 so we used as (treatment sample). Then the inoculated milk for yoghurt preparation was 228 incubated at 43°C for 5h after that the samples were cooled in refrigerator 4°C until the 229 sensory evaluation. The panel persons of sensory evaluation included 32 members from 230 Food Technology Department, Animal and Fish Production Department and other 231 departments, Arid Lands Cultivation Research Institute (ALCRI), City of Scientific Research and Technological Applications (SRTA-City). The yoghurt samples (control and treatment 232 233 samples) were evaluated for appearance, texture, tenderness, flavor and taste and overall 234 acceptance according to scores from 1-7 whereas 1= Very poor, 2= Poor, 3= Fair, 235 4=Medium, 5=Good, 6= Very good and 7= Excellent was the best score [34,35].

236

237 2.10 Statistical analysis

238

The results were performed by SPSS (Statistical package for social science) softwareprogram version 16 for Statistical analysis.

241 **3. Results and discussion**

242 **3.1** Evaluation the ability of viable probiotic strains on aflatoxin M1 243 reduction

244 Results presented in table 6 shows the effect of different concentration of viable probiotic strains in removal of AFM1 (50 ng ml⁻¹) residues along 72h. It can be seen from the table that Lactobacillus plantarum at 1×10^9 CFU ml⁻¹ had removal effect on AFM1 (50 ng/ ml⁻¹) to 245 246 40.14±1.23, 38.24±1.44, 36.73±11.56 and 33.64±1.25 ng ml⁻¹ during different time 12h, 24h, 48h 247 and 72h, respectively. When the inoculum concentration was increased to 3×10⁹ CFU ml⁻¹, the 248 removal effect of AFM1 was increased from 32.72% to 34.10% with AFM1 residual at 39 ± 11.07 , 37.22 ± 1.64 , 35.74 ± 1.32 and 32.95 ± 1.62 ng ml⁻¹, respectively during the different times. The highest concentration of this strain (5×10⁹ CFU ml⁻¹) with the highest incubation time (72h) 249 250 251 produced the highest removal effect on AFM1 (50 ng ml⁻¹) to 36.90% with AFM1 residual at 252 253 50.23±1.36, 38.95±1.24, 35.78±1.24, 33.69±1.41 and 31.55±1.22 ng ml⁻¹.

Lactobacillus acidophilus at 3×10^9 CFU ml⁻¹ had removal effect of AFM1 (50 ng ml⁻¹) to 34.26±1.53, 30.78±1.62, 29.02±1.35 and 26.53±1.27 ng ml⁻¹ during different time 12h, 24h, 48h and 72h, respectively. However, the concentration of 5×10^9 CFU ml⁻¹ produced more AFM1 reduction from 50 to 25.65±1.76 ng ml⁻¹. Also, it clear from the table that *Bifidobacterium bifidum* was reduced AFM1 concentration to 24.71±1.31 and 21.16±0.87 ng ml⁻¹ at 1×10^9 CFU ml⁻¹, 3×10^9 CFU ml⁻¹ and 5×10^9 CFU ml⁻¹ ,respectively so when the probiotic concentration and incubation time were increased, the effect

Comment [SC17]: species

-{	Comment [SC18]:
-{	Comment [SC19R18]:
1	Comment [SC20R18]:
1	Comment [SC21]: delete, instead paraphrase as
	Comment [SC22]: delete, instead paraphrase as before addition of yorghurt starter cultures
`٦	Commont [SC22]: Kluuwaramwaaa

261 of removal AFM1 was increased from 45.06% at 1×10^{9} CFU ml⁻¹ to 57.68% at 5×10^{9} CFU ml⁻¹ 262 after 72h. The removal effect of *Bifidobacterium bifidum* (57.68%) was more than *Lactobacillus* 263 *plantarum* (36.90%) and *Lactobacillus acidophilus* (48.70%) which was considered the highest 264 viable probiotic strain between other strains.

Moreover, the combination between the three viable probiotic strains (*Bifidobacterium bifidum*, *Lactobacillus plantarum* and *Lactobacillus acidophilus*) at concentration 5×10⁹ CFU ml⁻¹ produced higher removal AFM1 percent (64.62%) than each individual strain. The combination of different probiotic strains had sequestrate effect with AFM1 (50 ng ml⁻¹) in BPS media to became 17.69±1.24 ng ml⁻¹. Some research reported results in agree with results obtained in this study concerning to the binding effect of some bacterial and yeast strains in PBS media, milk and in yoghurt sample.

The results agree with Reference [24], they reported that three strains of lactic acid bacteria; Lactobacillus delbrueckii spp. bulgaricus, Lactobacillus rhamnosus and Bifidobacterium lactis had 272 ed that three strains of lactic acid bacteria; 273 274 removal effects of AFM1 in skim milk. This removal was ranged from 0.5 to 0.442±0.022 and to 0.442± 0.022 ng ml⁻¹ during 30 and 60 min of incubation respectively [23], reported that five 275 276 strains of LAB and bifidobacteria to remove aflatoxin M1(AFM1) from yoghurt. Lactobacillus 277 plantrium was the highest strain capable of removing AFM1. Yoghurt fermented by 50% yoghurt 278 culture (Streptococcus thermophilus and Lactobacillus bulgaricus) and 50% Lactobacillus 279 plantrium recorded the highest reduction in the level of AFM1 at the end of storage period. [36] 280 came to the same conclusion, when they used different strains of LAB, i.e., Lactobacillus casei sp. 281 (ATCC 15088), Lactobacillus acidophilus (ATCC 11975), Lactobacillus sp. GG. ATCC 53103 and Lactobacillus rhamnosus (ATCC 10863). The reduction level by these strains ranged from 26.2% 282 283 to 34.0%, depending upon the bacterial isolates. [37] studied the ability of Lb. bulgaricus to reduce 284 AFM1 from PBS and yogurt. Binding was 40% after 2 h PBS incubation and increased up to 285 87.6% after 14 h. In yogurt the AFM1 binding reached up to 60% after 6 h yogurt incubation. Sarimehmetoğlu and Küplülü (2004) [38] analyzed commonly used yogurt bacteria, Lactobacillus 286 delbrueckii subsp. bulgaricus for its binding ability of AFM1 in PBS and in milk. Binding was better 287 288 in milk (27.6%) than in PBS (18.7%) after 4 h incubation at 37 °C.

Comment [SC24]: These results agree with findings by [24] whereby

Comment [SC25]: Paraphrase sentence as..Similalrly, findings by [23] reported that

Comment [SC26]: Using a different combination of strains of LAB including *Lactobacillus casei sp.* (ATCC 15088), *Lactobacillus acidophilus* (ATCC 11975), similar results were obtained by [36].

Comment [SC27]: Studies by [37] on the ability of Lb bulgariscus to reduce AFMI from PBS and yorghurt established a 40% binding after 2h PBS incubation and a further increase to 87.6% after 14h.

Table 6: Effect of different concentration viable of probiotic strains in removal of AFM1 (50 - Comment [SC28]: insert 'of' 291 ng/ml) by detection AFM1 residual during different time and removal % after 72h.

Type of strain	Inoculum concentration	0 h	12 h	24 h	48 h	72 h	Removal % after 72h
Lactobacillus plantarum	1×10 ⁹ CFU ml⁻ 1	50.17±1.15	40.14±1.2 3	38.24±1.44	36.73±11.56	33.64±1.25	32.72%
	3×10 ⁹ CFU ml ⁻	50.04±1.42	39±11.07	37.22±1.64	35.74±1.32	32.95±1.62	34.10%
	5×10 ⁹ CFU ml ⁻	50.23±1.36	38.95±1.2 4	35.78±1.24	33.69±1.41	31.55±1.22	36.90%
Lactobacillus acidophilus	1×10 ⁹ CFU ml ⁻	50.26±0.56	35±1.10	32.71±1.64	30.95±1.52	29.02±1.29	41.96%
	3×10 ⁹ CFU ml ⁻ 1	50.15±0.66	34.26±1.5 3	30.78±1.62	29.02±1.35	26.53±1.27	46.94%
	5×10 ⁹ CFU ml ⁻ 1	50.16±0.90	33.72±1.2 8	30.29±1.27	27.26±1.43	25.65±1.76	48.70%
Bifidobacterium bifidum	1×10 ⁹ CFU ml ⁻	50.34±0.78	34.61±1.5 1	31.952±1.2 5	31.84±1.24	27.47±1.36	45.06%

	3×10 ⁹ CFU ml ⁻	50.20±0.56	31.84±1.7	27.59±1.62	26.74±1.38	24.71±1.31	50.58%
	5×10 ⁹ CFU ml ⁻	50.22±0.65	26.84±1.5 8	25.29±1.20	23.07±1.43	21.16±0.87	57.68%
CPS-V	5×10 ⁹ CFU ml ⁻	50.22±1.36	22.93±1.1 4	20.06±1.25	18.56±1.23	17.69±1.24	64.62%
+ve control	BPS + AFM1	50	49.99	49.98	49.88	49.85	0.00%
-ve control	BPS +P	0.00	0.00	0.00	0.00	0.00	0.00%

292 CPS-V: Combination probiotic strain viable (*B. bifidum*+ *L. acidophilus* + *L. plantarum*).
 293 Mean and SD of AFM1residual

293 Mean and SD of AFM1residual 294

294

296 **3.2 Evaluation the efficiency of nonviable**

3.2.1. Evaluation the efficiency of nonviable probiotic strains on reduction of
 aflatoxin M1

299 Table 7 shows that nonviable Lactobacillus plantarum reduced AFM1 from 50 ng/ml_to 33.54±1.44, 26.15±1.64 and 24.13±0.95 at 1×10⁹ CFU ml⁻¹, 3×10⁹ CFU ml⁻¹and 5×10⁹ CFU ml⁻¹, 300 301 respectively after 72h. Lactobacillus plantarum had the sequestration effect of AFM1 which produced removal % at 51.74%. On the other hand, nonviable Lactobacillus acidophilus at 3×10⁹ 302 CFU ml⁻¹ reduced concentration of AFM1 from 50 to 17.51±1.28 ng ml⁻¹. However, the 303 304 concentration at 5×10^9 CFU m⁻¹ had reduction effect on AFM1 concentration to 22.65±1.37. 305 20.76±1.11, 17.89±1.33 and 16.04±1.00 ng ml⁻¹ during different times 12h, 24h, 48h and 72h, 306 respectively. The highest concentration of Lactobacillus acidophilus at 5×10⁹ CFU ml⁻¹ gave 307 67.92% removal effect.

Bifidobacterium bifidum was considered higher probiotic effect than other two strains on AFM1 sequestration, which had AFM1 removal % at 70.62% to AFM1 removal %. AFM1 reduced to 21.00 \pm 1.43, 18.37 \pm 1.34, 16.67 \pm 1.64 and 14.69 \pm 1.62 ng ml⁻¹ during 12h, 24h, 48h and 72h, respectively at 1×10⁹ CFU ml⁻¹ of nonviable *Bifidobacterium bifidum*. When the concentration and the incubation time increased the effect of *Bifidobacterium bifidum* was increased to 21.00 \pm 1.31, 18.37 \pm 1.37, 16.67 \pm 1.27 and 14.69 \pm 0.93 ng ml⁻¹ during different incubation period. However, the highest reduction effect of nonviable probiotic appeared by combination, these strains to give removal effect to 79.66% and AFM1 concentration residual became 10.17 \pm 1.03 ng ml⁻¹ after 72h.

316 Assessed that probiotic-yeast coctile; Lactobacillus acidophilus, Bifidobacterium bifidum, Kluyveromyces lactis and Saccharomyce cerevisiae, had the highest effect of aflatoxins (B1, B2, 317 G1 and G2) removal after 72h (95.59%) in PBS media and when applied in contaminated Cerelac 318 319 with aflatoxins, the removal percentage was increased by time 6, 12, 24, 48 and 72h to 8.17, 320 36.12, 44.75, 64.72 and 93.21%, respectively. Also, when these probiotic-yeast coctile were 321 applied in vivo study had a high effective role in the reduction of aflatoxins (B1, B2, G1 and G2) in 322 mother serum rat and also reduction aflatoxins metabolites (M1 and M2) in babies' serum rat 323 serum [39].

Reference [39] tested Lactobacillus gasseri for its ability to remove AFM1 from liquid PBS during 15 to 16 h incubation at 37 °C. Heat killed bacteria had a better AFM1 binding ability than the viable bacteria, 61.5% and 30.8%, respectively and studied the abilities of *Lactobacillus rhamnosus* GG (ATCC 53013), *Lactobacillus rhamnosus* LC-705 and *Lactobacillus rhamnosus* 1/3 to bind AFM1 from PBS. *Lactobacillus rhamnosus* GG bound over 50% of the AFM1 in PBS in all tested forms (precultured, freeze dried, viable and heat killed). Viable *Lactobacillus rhamnosus* LC705 bound around 45–46% and the heat-killed more than 50%. The heat killed *Lactobacillus*

 $\begin{array}{l} \textbf{Comment [SC29]: Non viable L. Plantarum was found to reduce AFMI from50 ng/ml to \\ 33.54\pm1.44, 26.15\pm1.64 and 24.13\pm0.95 at \\ 1\times10^{\circ} \ CFU \ ml^{-1}, 3\times10^{\circ} \ CFU \ ml^{-1} and 5\times10^{\circ} \ CFU \\ ml^{-1}, respectively after 72h (Table 7). \end{array}$

Comment [SC30]: Delete and paraphrase sentence to read..Lactobacillus gasseri was tested by [39] for ita ability to remove AFBI from liquid PBS

Comment [SC31]: Lacto

331 rhamnosus 1/3 strain bound 40% and the viable 18% of the added AFM1. Lactobacillus 332 rhamnosus GG and LC-705 were further tested in skim milk and in full cream milk. Lactobacillus 333 rhamnosus GG bound with limitations: viable cells bound 19% of AFM1 in skim milk and 26% in 334 full cream milk. The heat killed Lactobacillus rhamnosus GG bound 27% of AFM1 in skim milk and 37% in full cream milk. The viable Lactobacillus rhamnosus LC-705 bound over 60% of the AFM1 335 336 in skim and full cream milk when the binding share of heat-treated cells remained at around 30%. While Viable and heat killed Lactobacillus lactis ssp. cremoris (ARH74) strain removed 40.4% and 337 338 38.9% of AFM1, respectively, from PBS [40].

340 Table 7: Effect of different concentration nonviable probiotic strains in removal of AFM1

341 (50 ng ml⁻¹) by detection AFM1 residual during different time and removal % after 72h.

Type of strain	Inoculum concentration	0 h	12 h	24 h	48 h	72 h	Removal % after 72h
Lactobacillus	1×10 ⁹ CFU ml⁻¹	50.00±0.2	39.86±0.3	37.42±1.4	34.52±1.2	33.54±1.4	32.92%
plantarum	3×10 ⁹ CFU ml ⁻¹	1 50.02±0.6 2	1 37.41±0.5 8	0 31.65±1.6 6	3 28.02±1.3 4	4 26.15±1.6 4	47.70%
	5×10 ⁹ CFU ml⁻¹	50.10±1.4 2	34.63±1.6 3	28.41±1.4 1	26.69±1.7 7	24.13±0.9 5	51.74%
Lactobacillus acidophilus	1×10 ⁹ CFU ml⁻¹	50.20±0.2 3	29.81±1.5 2	26.53±1.3 4	23.55±1.3 1	20.17±1.3 2	58.98%
	3×10 ⁹ CFU ml⁻¹	50.11±0.3 4	28.99±1.2 6	21.96±1.4 4	18.99±1.0 6	17.51±1.2 8	64.98%
	5×10 ⁹ CFU ml⁻¹	50.12±0.2 4	22.65±1.3 7	20.76±1.1 1	17.89±1.3 3	16.04±1.0 0	67.92%
Bifidobacterium bifidum	1×10 ⁹ CFU ml⁻¹	50.09±0.5 6	21.00±1.4 3	18.37±1.3 4	16.67±1.6 4	14.69±1.6 2	62.44%
	3×10 ⁹ CFU ml⁻¹	50.15±1.1 0	26.59±1.6 1	21.07±1.2 3	18.19±1.1 8	15.94±1.1 6	68.12%
	5×10 ⁹ CFU ml⁻¹	50.10±0.4 4	21.00±1.3 1	18.37±1.3 7	16.67±1.2 7	14.69±0.9 3	70.62%
CPS-NV	5×10 ⁹ CFU ml⁻¹	50±0.62	19.81±1.5 3	16.53±1.3 4	13.55±1.4 7	10.17±1.0 3	79.66%
+ve control	BPS + AFM1	50.22±0.6 1	50.22±0.6 1	50.22±1.5 3	49.90±1.3 4	49.80±1.4 7	0%
-ve control	BPS +P	0.00	0.00	0.00	0.00	0.00	0%

342 CPS-NV: Combination probiotic strain nonviable (B. bifidum+ L. acidophilus + L. plantarum).

343 3.3 Evaluation the efficiency of some nonviable yeast strains kluyveromyces lactis

344 and Saccharomyces cerevisiae) on reduction of aflatoxin M1

345

346 Table (8) shows the effect of different concentration of It can be seen from table (8) that -Kluyveromyces lactis at 1×10⁹ CFU ml⁻¹ had removal effect on AFM1 (50 ng ml⁻¹) to 25.01±1.06, 347 22.36±1.27, 20.34±1.33 and 19.93±1.25 ng ml⁻¹ during different time 12h, 24h, 48h and 72h, respectively. on the other hand at 3×10^9 CFU ml⁻¹, the AFM1 residues became 24.39±1.52, 348

349

21.08±1.42, 18.97±1.02 and 16.20±1.64 ng ml⁻¹, respectively during the different times (12h, 24h, 48h and 72h, respectively). However, *Kluyveromyces lactis* at 5×10⁹ CFU ml⁻¹ reduced AFM1 to 350 351

was found effective in the removal of AFM1 after 72h (Table 8).

Comment [SC32]: Non-viable yeast strains

³³⁹

22.48±1.39, 18.86±1.64, 16.67±1.92 and 15.43±1.15 ng ml⁻¹, respectively during the different
 times which was more removal effect than low concentration.

354 On the other hand, nonviable *Saccharomyces cerevisiae* reduced AFM1 (50 ng ml⁻¹) to 355 24.30±1.54, 22.61±1.14, 21.73±1.34 and 17.74±1.35 ng ml⁻¹ during 12h, 24h, 48h and 72h, 356 respectively at 1×10^9 CFU ml⁻¹. The effect of *Saccharomyces cerevisiae* was increased to 357 20.76±1.27, 19.63±1.75, 16.96±1.61 and 13.32±1.28ng/ml at 3×10^9 CFU ml⁻¹. This removal effect 358 of *Saccharomyces cerevisiae* was more increased to 16.81±1.61, 13.59±1.56, 12.32±1.27 and 359 10.63±1.01 ng ml⁻¹ at 5×10^9 CFU ml⁻¹ during different incubation time12h, 24h, 48h and 72h, 360 respectively. Also, the results showed that the removal effect of *Saccharomyces cerevisiae* was 361 higher than *Kluyveromyces lactis*.

362 The combination of nonviable yeast strains (Kluyveromyces lactis and Saccharomyces cerevisiae) had a higher removal effect at 5×10^9 CFU ml⁻¹ of concentration with 72h incubation period (85.68%) on AFM1 (50 ng ml⁻¹) than using each yeast strain separately (69.14% for 363 364 Kluyveromyces lactis and 78.74% for Saccharomyces cerevisiae). Reference [28] mentioned that 365 366 Saccharomyces cerevisiae was considered the highest microorganism able to remove AFB1 in vitro study which agrees with our results. However, when used Saccharomyces cerevisiae with 367 368 LAB strains, the AFM1 removal percentage was increased in the milk sample. Also, the researcher detected the increased of incubation time effect positively on the removal percentage 369 370 which near to the results of the present study. [41] assessed that yeast. The highest reduction 371 percentage of AFM1 was observed (65.33%-68.89%)

Comment [SC33]: Findings by [28] on the use of *Saccharomyces cerevisiae* are in agreement with findings from the current study which established it as the most effective species in AFM1 removal.

Comment [SC34]: The highests AFM1 reduction when yeasts were used was in the range 65.33-68.89% [41].

372

373 Table 8. Effect of different concentration nonviable yeast strains in removal of AFM1 (50 ng/ml) by

374	detection A	AFM1	residual	during	different til	me and	removal	% of AFM1	after 72h.	

Type of strain	Inoculum	0 h	12 h	24 h	48 h	72 h	Removal %
	concentration						after 72h
	1×10 ⁹ CFU ml⁻¹	50.21	25.01±1.0	22.36±1.27	20.34±1.33	19.93±1.25	60.14%
1.1		±1.0	6				
kiuyveromyces	3×10 ⁹ CFU ml ⁻¹	50.09±0.8	24.39±1.5	21.08±1.42	18.97±1.02	16.20±1.64	67.60%
lactis		8	2				
	5×10 ⁹ CFU ml ⁻¹	50.19±1.3	22.48±1.3	18.86±1.64	16.67±1.92	15.43±1.15	69.14%
		0	9				
	1×10 ⁹ CFU ml ⁻¹	50.23±1.6	24.30±1.5	22.61±1.14	21.73±1.34	17.74±1.35	64.52%
		2	4				
Saccharomyce	3×10 ⁹ CFU ml⁻¹	50.32±1.4	20.76±1.2	19.63±1.75	16.96±1.61	13.32±1.28	73.36%
s cerevisiae		2	7				
	5×10 ⁹ CFU ml⁻¹	50.14±1.2	16.81±1.6	13.59±1.56	12.32±1.27	10.63±1.01	78.74%
		2	1				
CYS-NV	5×10 ⁹ CFU ml ⁻¹	50.19±1.0	14.34±1.4	13.65±1.63	10.46±1.83	7.16±0.90	85.68%
		6	7				
+ve control	PBS + AFM1	50.25±1.2	50.20±1.4	49.98±1.36	49.88±0.98	49.85±1.13	0%
		1	6				
-v econtrol	PBS +P	0.00	0.00	0.00	0.00	0.00	0%

375 CYS-NV: Combination yeast strains non-viable (S. cerevisiae +k. lactis).

376 **3.4** Evaluation the efficiency of some nonviable bacterial and yeast strains

(Lactobacillus plantarum, Lactobacillus acidophilus, Bifidobacterium bifidum,
 Kluyveromyces lactis and Saccharomyces cerevisiae) on reduction of
 aflatoxin M1 in PBS

380 Data presented in table (9) revealed that the combination of probiotic (Lactobacillus plantarum, 381 Lactobacillus acidophilus and Bifidobacterium bifidum) and yeast strains (Kluyveromyces lactis 382 and Saccharomyces cerevisiae) had the highest removal effect of AFM1 (87.92%) after 72h of 383 incubation. Also, the table shows the AFM1 residues to 13.98±1.34, 10.53±1.26, 8.49±0.63 and 384 6.04±0.15 during different incubation period at 12h, 24h, 48h and 72h, respectively. Another 385 research by [42] reported that Lactobacillus Casei TD4 had AFM1 reduction percentage (91.91%). 386 Lactobacillus bulgaricus had 87.6% and Streptococcus thermophilus had 70% removal of AFM1 387 however, the efficiency of removal was increased by using the yeast with the bacterial strain. [43] 388 reported that Bifidobacterium bifidum, Lactobacillus spp. and Lactobacillus spp. had binding ability 389 with AFM1 in solution media. [44] mentioned that probiotic strains in yoghurt had removal effect 390 (49%) of AFM1 at the end of storage period. [45] evaluated that Lactobacillus acidophilus 391 removed 90% of aflatoxin M1 contaminated in yoghurt samples during the first day then the 392 removal increased by the storage time. [38] used a yogurt mixture (Streptococcus thermophilus 393 and Lactobacillus delbrueckii subsp. bulgaricus) to study the AFM1 binding during yogurt 394 fermentation. The mixture bound only 15% of the AFM1 added to the yogurt. [36] studied the ability of yogurt culture mixture Streptococcus thermophilus and Lactobacillus delbrueckii subsp. 395 396 bulgaricus) to remove AFM1 from PBS and yogurt. In both matrices binding increased during 6 h incubation and reached approximately 45% of AFM1 removal level. In PBS the incubation was 397 continued up to 14 h and the binding share of the mixture reached almost 65%. 398 399

Table 9. Effect of nonviable combination of probiotic bacterial and yeast strains in PBS to removal of AFM1 (50 ng ml⁻¹) during different time and removal % of AFM1 after 72h.

Type of strain	Inoculum	0 h	12 h	24 h	48 h	72 h	Removal %
	concentration						after 72h
CPYS-NV	5×10 ⁹ CFU ml⁻¹	50.23±1.4	13.98±1.3	10.53±1.2	8.49±0.63	6.04±0.15	87.92%
		2	4	6			
+ve control	BPS + AFM1	50.00±1.1	50.00±1.3	49.95±1.1	49.77±1.0	49.30±0.8	0%
		6	0	1	8	1	
-v econtrol	BPS +P+Y	0	0	0	0	0	0%

402 CPYS-NV: combination non-viable strains (*B. bifidum+L. acidophilus+L. plantarum +S.* 403 *cerevisiae+ k. lactis*).

404 3.5 Evaluation potential of the combination of nonviable probiotic bacterial and 405 yeast strains on aflatoxin M1 reduction in milk

406 Table (10) shows The effect of the highest effective combination in PBS (combination of probiotic bacterial and yeast strains nonviable) for sequestration of AFM1 (50 ng ml⁻¹) in milk as
 408 experimental media and distribution the removal % of AFM1 during different times (0h, 12h, 24h, 409 24h, 48h and 72h) is demonstrated in Table 10.

410 It shows from the table that the combination of nonviable probiotic bacterial and yeast strains sequestrate of AFM1 (50 ng ml⁻¹) during different times (12h, 24h, 24h, 48h and 72h) with low 411 AFM1 residues as 9.72±1.31, 6.68±0.55, 5.70±0.33 and 4.56±0.15ng ml⁻¹, respectively and with 412 413 high removal % of AFM1 to 80.56%, 86.64%, 88.60% and 90.88%, respectively in milk sample. 414 [24] when used three strains of lactic acid bacteria (Lactobacillus delbrueckii spp. bulgaricus, 415 Lactobacillus rhamnosus and Bifidobacterium lactis) with Saccharomyces cerevisiae (killed by heat), the AFM1 residues decreased to 0.042± 0.003 ng ml⁻¹ during 30 while during 60 min there 416 was no AFM1 residues detected (0 ng ml⁻¹), when these LAB strains used with Saccharomyces 417 cerevisiae (killed by heat) the AFM1 residues decreased to 0.042± 0.003 ng ml⁻¹ during 30 while 418 419 during 60 min there was no AFM1 residues detected (0 ng ml⁻¹).

Comment [SC35]: Delete

Comment [SC36]: Add this at the end of sentence

420							
421	Table (10)	: Effect of the hig	ghest effec	tive combination	of (probic	otic bacterial a	nd yeast
422	strains no	onviable) for seque	estration of	FAFM1 (50 ng ml ⁻	¹) in milk a	s experimental	media and
423	distributio	on the removal % of	of AFM1 du	uring different tim	ies (0h, 12	h, 24h, 24h, 48h	n and 72h).
Туре	e of strain	Inoculum	0 h	12 h	24 h	48 h	72 h

5 1	concentration					
CPYS-NV in	5×10 ⁹ CFU ml⁻¹	50.10±1.10	9.72±1.31	6.68±0.55	5.70±0.33	4.56±0.15
Milk						
+ve control	Milk + AFM1	50.21±0.32	49.90±1.14	49.87±1.05	49.76±1.16	49.33±1.21
-ve control	Milk + CPYS	0	0	0	0	0
Removal %	5×109 CFU/ml	0%	80.56%	86.64%	88.60%	90.88%

424

425 CPYS-NV: Total combination non-viable strains (*B. bifidum+L. acidophilus+L. plantarum* +S. 426 *cerevisiae+ k. lactis*).

427

428 3.6 Scanning Electron Microscope (SEM) of different combination from different probiotic

429 bacterial and yeast strains with AFM1

430 Figure 1 (P) shows Scanning Electron Microscopy (SEM) results of nonviable combination of 431 probiotic bacterial strains control and treatment are illustrated in Fig1(P). It is clear from the figure the difference in the cell wall of probiotic bacterial strains (Lactobacillus plantarum, 432 433 Lactobacillus acidophilus and Bifidobacterium bifidum) in control sample and in treatment one which had spots on their cell wall after adsorption of AFM1 in these spots of cell wall. Figure 1 (Y) 434 435 shows scanning Electron Microscope (SEM) of nonviable combination of yeast strains control 436 (yeast strains without AFM1) and treatment (yeast strains with AFM1) by using magnification at 437 500 x. It is clear from the figure the difference in the cell wall of yeast strains (kluyveromyces lactis -438 and Saccharomyces cerevisiae) of the control sample to the treatment yeast sample which had spots on their cell wall after sequestration with AFM1 in these spots on the cell wall. Figure 1 439 440 (P+Y) shows scanning Electron Microscope (SEM) of nonviable combination of probiotic bacterial and yeast strains (control and treatment) by using magnification at 500 x. It is clear from the figure 441 442 that the cell wall of both probiotic bacterial and yeast strains (Lactobacillus plantarum, 443 Lactobacillus acidophilus, Bifidobacterium bifidum, kluyveromyces lactis and Saccharomyces 444 cerevisiae) in the cell wall in the control sample appeared without this spots on their cell wall while the gical reduction of AFM1. The probiotic-aflatoxin complex and also, yeast-aflatoxin complex 445 imtreatment sample bind or sequestrate with AFM1 in their cell wall spots which act as a good 446 bioloproved the reduction of aflatoxin M1 higher than using probiotic bacterial or yeast strains 447 individually because sequestration sites were became more in the using case of probiotic bacterial 448 449 with yeast strains.

Comment [SC37]: Insert at the end of sentence Comment [SC38]: Delete the striked through and add the infomation in green

Comment [SC39]: Kluyveromyces



Figure 1. Scanning Electron Microscope (SEM) showing a nonviable combination, control and treatment by using magnification at 500 x.

450

453 3.7 Sensory evaluation the best efficient combination of strains on aflatoxin 454 M1 sequestration applied in yoghurt

- 455 Table 11 shows the The mean and standard deviation of sensory evaluation scores of yoghurt
- 456 was treated with the combination of nonviable probiotic bacterial and yeast strains are illustrated 457 in Table 11. It is clear from the table that control yoghurt sample was taken scores $6.15\pm0.76_{\odot}$

Comment [SC40]: Delete the striked through and add the information in green

458 6.18±0.64, 6.00±0.91, 6.00±0.87 and 5.93±0.87 while inoculated yoghurt sample (inoculated with 459 combination of nonviable probiotic bacterial and yeast strains) (B. bifidum+L. acidophilus+L. 460 plantarum +S. cerevisiae+ k. lactis) was taken scores 5.84±1.11, 5.75±1.16, 5.84±1.11, 5.96±1.33

461 and 5.96±1.23 (good score) regarding to appearance, texture, tenderness, flavor (odour & taste) 462 and overall acceptance, respectively.

463

464 Table 11. Sensory evaluation scores of treatment yoghurt sample.

Sensory evaluation parameter	Control yoghurt sample Treatment yoghurt sample					
Appearance	6.15±0.76	5.84±1.11				
Texture	6.18±0.64	5.75±1.16				
Tenderness	6.00±0.91	5.84±1.11				
Flavour (odour & taste)	6.00±0.87	5.96±1.33				
Overal acceptance	5.93±0.87	5.96±1.23				
	(Good score)	(Good score)				

465

466 32 panel members Maximum score = 7.

467 Figure 2 shows preparation The results on the sensory variables of yoghurt with nonviable 468 combination of probiotic bacterial and yeast strains (Treatment yoghurt sample) or without (Control yoghurt sample) are illustrated in Figure 2. Treatment yoghurt sample prepared with Comment [SC41]: Delete the striked through 469 nonviable combination of probiotic bacterial and yeast strains (Lactobacillus plantarum, Lactobacillus acidophilus, Bifidobacterium bifidum, Kluyveromyces lactis and Saccharomyces 470 471 cerevisiae) to compare to the control yoghurt sample prepared without these strains in 472 473 appearance, texture, tenderness, flavour and overall acceptance.

and add th info in green.



474 475 Yoghurt models prepared by nonviable combination compared with the control Figure 2. 476 yoghurt. 477

478 Table 12 shows The distribution of sensory evaluation scores for yoghurt sample was treated with 479 a nonviable combination of probiotic bacterial and yeast strains are illustrated in Table 12-480 Treatment yoghurt sample was excellent (score 7) in overall acceptance of (46.87%) of the 481 samples, in appearance (34%), texture (29.41%), tenderness (46.87%) and flavor (50%) by the 482 panel members. On the other hand, the control yoghurt sample was excellent in overall 483 acceptance of (28.12%) with (34%), (29.41%), (28.12%) and (29.41%) in appearance, texture, 484 tenderness and flavor, respectively.

Comment [SC42]: Delete the striked through and add the inforation in green

Sensory evaluation	Appea	irance	e Texture		Tenderness		Flavor (odour & taste		Overall acceptance	
parameter	С	Т	С	Т	С	Т	С	Т	С	Т
Excellent (7)	11 34%	11 34%	10 29.41%	10 29.41%	9 28.12%	15 46.87%	10 29.41%	16 50%	9 28.12%	15 46.87%
Very good (6)	16 50%	10 29.41%	18 56.25%	9 28.12%	17 53.12%	6 18.75%	14 43.75%	7 21.87%	14 43.75%	6 18.75%
Good (5)	4 12.5%	7 21.87%	4 12.5%	10 29.41%	4 12.5%	4 12.5%	6 18.75%	3 9.37%	7 21.87%	8 25%
Medium (4)	1 3.12%	3 9.37%	ND	2 6.25%	1 3.12%	6 18.75%	2 6.25%	5 15.62%	2 6.25%	2 6.25%
Fair (3)	ND	ND	ND	ND	1 3.12%	1 3.12%	ND	ND	ND	ND
Poor (2)	ND	1 3.12%	ND	ND	ND	ND	ND	1 3.12%	ND	1 3.12%
Very poor (1)	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND

485 Table 12. Sensory evaluation scores for yoghurt sample treated either by nonviable 486 combination of probiotic bacterial or yeast strains.

487

491

488 C = Control sample of yoghurt

489 T = Treatment inoculated sample of yoghurt with B. bifidum+L. acidophilus+L. plantarum +S.

cerevisiae+ K. lactis (nonviable combination of probiotic bacterial and yeast strains) (CPYS). 490

CONCLUSION 4. 492

493 In conclusion, probiotic bacteria and yeast strains are able to make detoxification for aflatoxin M1 494 in contaminated milk. But a combination from probiotic bacteria and yeast could be good for removal and eliminating of aflatoxins M1 from milk. Moreover, probiotic bacteria and yeast could [Comment [SC43]: elimination 495

496 be used as food additives to reduce the bioavailability of the aflatoxins in dairy products.

497

ACKNOWLEDGEMENTS 498

499 The authors thank the City of Scientific Research and Technologicals for support of this work. The 500 authors also thank the junior staff and technicians of the Arid Lands Cultivation Research Institute 501 at the City for Scientific Research and Technological Application, Alexandria, Egypt for their help 502 in this study.

503

504 **COMPETING INTERESTS**

505 507

506 Authors have declared that no competing interests exist.

508 REFRENCES

- 509
- 510 1. Saddig A. Antifungal and prophylactic activity of pumpkin (Cucurbita moschata) extract 511 against Aspergillus flavus and aflatoxin B1. African Journal of Microbiology Research. 512 2012; 6(41): 6941-7.
- Strosnider H, Azziz-Baumgartner E, Banziger M, Bhat R, Breiman R, et al. Public Health 513 2. 514 Strategies for reducing aflatoxin exposure in developing countries: A workgroup Report:

- 515 Environmental Health Perspectives. Journal of Environmental Health Perspective. 2006; 516 114(12): 1898-903.
- Kangethe E, Langa k. Aflatoxin B1and M1 contamination of animal feeds and milk from urban centers in Kenya. Journal of Africa Health Sciences. 2009; 9(4): 218-26.
- Duarte S, Almeida A, Teixeira A, Pereira A, Falcão A, Pena A, et al. Aflatoxin M₁ in marketed milk in Portugal: Assessment of human and animal exposure. Journal of Food Control. 2013; 30(2): 411-7.
- Londono V, Boasso A, Paula M, Garcia L, Scussel V, Resnik S, et al. Aflatoxin M1survey on randomly collected milk powder commercialized in Argentina and Brazil. Journal of Food Control. 2013; 34: 752-5.
- 525 6. Yu J. Current understanding on aflatoxin biosynthesis perspective in reducing aflatoxin
 526 contamination. Journal of Toxins. 2012; 4: 1024-57.
- 527 7. Scaglioni P, Becker-Algeri T, Drunkler D, Badiale-Furlong E. Aflatoxin B1 and M1 in milk.
 528 Journal of Analytica Chimica Acta. 2014; 829: 68–74.
- Raiola A, Tenore G, Manyes L, Ritieni A. Risk analysis of main mycotoxins occurring in food for children: An overview. Journal of Food and Chemical Toxicology. 2015; 84:169-80.
- Iha M, Barbosa C, Okad I, Trucksess M. Aflatoxin M1 in milk and distribution and stability of aflatoxin M1 during production and storage of yoghurt and cheese. Journal of Food Control. 2013; 29 (1):1-6.
- 535 10. Campone L, Piccinelli A, Celano R, Pagano I, Russo M, Rastrelli L. Rapid and
 536 automated analysis of aflatoxin M1 in milk and dairy products by online solid phase
 537 extraction coupled to ultra-high-pressure-liquid-chromatography tandem mass
 538 spectrometry. Journal of chromatography A, 2016; 1428:212-9.
- 539 11. Egyptian standards regulation (E.S). Maximum levels for certain contaminants in foodstuffs. E.S.7136/2010, Egyptian organization for standardization and quality control, 2010.
- 542 12. Vinderola G, Ritieni A. Role of probiotics against mycotoxins and their deleterious
 543 effects. Journal of Food Research. 2015; 4 (1): 10-21.
- 544 13. Armando M, Dogi C, Pizzolitto P, Escobar F, Peirano M, Salvano M, et al.
 545 Saccharomyces cerevisiae strains from animal environmental with aflatoxin B1
 546 detoxification ability and anti-pathogenic bacteria influence in vitro. Journal of World
 547 Mycotoxin. 2011; 4 (1): 59-68.
- 548 14. Bilandzic N, Varenina I, Solomun B. Aflatoxin M1 in raw milk in Croatia.
 549 Journal of Food Control. 2010. 21, 1279-81.
- 15. El-Kest M, Hariri M, Khafaga N, Refai M. Studies of contamination of dairy products by aflatoxin M1 and its control by probiotics. Journal of Global Biosciences. 2015; 4: 1294-312.
- 16. Pizzolitto R, Bueno D, Armando M, Cavaglieri L, Dalcero A, Salvano M. Binding of aflatoxin B1 to lactic acid bacteria and saccharomyces cerevisiae in vitro: A useful model to determine the most efficient microorganism. Journal of Aflatoxin-Biochemistry and Molecular Biology, 2013; 16: 323-46.
- 17. Armando M, Dogi C, Pizzolitto P, Escobar F, Peirano M, Salvano M, et al.
 Saccharomyces cerevisiae strains from animal environmental with aflatoxin B1
 detoxification ability and anti-pathogenic bacteria influence in vitro. Journal of World
 Mycotoxin. 2011; 4(1): 59-68.
- 18. Vinderola G, Ritieni A. Role of probiotics against mycotoxins and their deleterious
 effects. Journal of Food Research. 2015; 4(1): 10-21.
- 19. Association of Official Analytical Chemists "AOAC": Official Methods of the AOAC
 International Analysis. 13th Ed., Horwitz. W; (Editor), Academic Press, Washington D.C,
 USA. 2000.
- 566 20. Daniel W. Biostatistis. A foundation for analysis in the health science. 6th edition, NY:
 567 John Willey and sons, Inc.; 1995.

- 568 21. Bovo F, Corassin C, Rosim E, Oliverira F. Efficiency of lactic acid bacteria strains for
 569 decontamination of aflatoxin M1 in phosphate buffer saline solution and in skim milk.
 570 Journal of Food Bioprocess Technology. 2012; 5: 1-5.
- 571 22. Kabak B, Ozbey F. Aflatoxin M1 in UHT milk consumed in Turkey and first assessment
 572 of its bioaccessibility using an *in vitro* digestion model. Journal of food control. 2012;
 573 28:338-44.
- 574 23. Elsanhoty R, Salam S, Ramadan M. Detoxification of Aflatoxin M1 in yoghurt using
 575 probiotics and lactic acid bacteria. Journal of food control. 2014; 43:129-34.
- 576 24. Corassin H, Bovo F, Rosim R. Oliveira C. Efficiency of Saccharomyces cerevisiae and lactic acid bacteria strains to bind aflatoxin M1 in UHT skim milk. Journal of food control. 2013; 31(1):80-3.
- 579 25. Nada S, Amra H, Deabes M, Omara E. Saccharomyces Cerevisiae and probiotic
 580 bacteria potentially inhibit aflatoxins production *in vitro* and *in vivo* studies. The Internet
 581 Journal of Toxicology. 2009; 8(1): 63-7.
- 582 26. Teniola D. Addo A. Brost M. Farber P. Jany D. Alberts F. et al. Degradation of aflatoxin
 583 B1 by cell-free extracts of *Rhodococcus erythropolis* and *Mycobacterium*584 *fluroranthenivorans* sp. nov. DSM44556T. Journal of Food Microbiology. 2005; 105:111585 7.
- 586 27- Scaglioni T, Becker-Algeri T, Drunkler D, Badiale-Furlong E. Aflatoxin B1 and Aflatoxin
 587 M1 in milk. Journal of Analytica Chimica Acta. 2014; 829:68-74.
- 588 27. Salim Abou-Baker, Zohair A, Hegazy A, Said A. Effect of some strains of probiotic bacteria against toxicity induced by aflatoxins *in vivo*. Journal of American Science.
 590 2011; 7(1): 772-83.
- 591 28. Rayes A. Removal of aflatoxin B1from experimentally contaminated whole milk using a
 592 pool of probiotic strains of lactic acid bacteria and baker's yeast Saccharomyces
 593 cerevisiae. New York Science Journal. 2013; 6(8):84:90.
- Hamad G, Zahran E, Hafez E. The efficacy of bacterial and yeasts strains and their
 combination to bind aflatoxin B1 and B2 in artificially contaminated infant's food. Journal
 of Food Safety. 2017; DOI: 10.1111/jfs.12365.
- 597 30. Haskard C, El-Nezami H, Kankaanpää P, Salminen S, Ahokas J. Surface binding of
 598 aflatoxin B1 by lactic acid bacteria. Journal of Applied and Environmental Microbiology.
 599 2001; 67(7): 3086:91.
- 31. Hernandez-Mendoza A, Garcia H, Steele J. Screening of Lactobacillus casei strains for
 their ability to bind aflatoxin B1. Journal of Food and chemical Toxicology. 2009;
 47:1064:68.
- 32. Fernandes M, Correa B, Rosim E, Kobashigawa, E, Oliveira F. Distribution and stability
 of aflatoxin M1 during processing and storage of Minas Frescal cheese. Journal of Food
 Control. 2012; 24:104-8.
- 33. Hamad G, Taha T, Hafez E, El Sohaimy S. Physicochemical, molecular and functional characteristics of hyaluronic acid as a functional food. American Journal of Food Technology. 2017; 12 (2): 72-85.
- 34. Cadena R, Caimi D, Jaunaren I, Lorenzo I, Vidal L, Ares G, Deliza R, GiménezA.
 Comparison of rapid sensory characterization methodologies for the development of functional yogurts. Journal Food Research International.,2014, 64: 446–55.
- 55. Maryamma, K. I., Rajan, A., Gangadharan, B., Ismail, P. K., Valsala, K. V., &
 Manomohan, C. B. Reduction of aflatoxin in milk by fermentation into curd. Journal of
 Veterinary and Animal Sciences, 1990; 21(2), 102-7.
- 616 36. El Khoury, A., Atoui, A., Yaghi, J. Analysis of aflatoxin M1 in milk and yogurt and AFM1
 617 reduction by lactic acid bacteria used in Lebanese industry. Food Control. 2011;
 618 22:1695–9.

- 37. Sarimehmetoğlu, B., Küplülü, Ö. Binding ability of aflatoxin M1 to yoghurt bacteria. Ank.
 Univ. Vet. Fak. Derg. 51, 195–198 (Available from: http://dergiler.ankara.edu.tr/dergiler/11/214/1752.pdf. Accessed 14 August 2014), 2004.
- 38. Hamad G, Taha T, Hafez E, El Sohaimy S, Ali S. supplementation of Cerelac baby food
 with yeast-probiotic cocktail strains induces high potential for aflatoxin detoxification both
 in vitro and in vivo in mother and baby albino rats. Journal of the Science of Food and
 Agriculture. 2018; 98 (2): 707-18.
- 39. Pierides, M., El-Nezami, H., Peltonen, K., Salminen, S., Ahokas, J. Ability of dairy strains
 of lactic acid bacteria to bind aflatoxin M1 in a food model. J. Food Prot. 2000; 63:645–
 50.
- 40. Siavash K, MohammadhoseinM. Reduction of Aflatoxin M1 in Milk Using Kefir Starter.
 Iranian Journal of Toxicology. 2017; 11(6): 27-31.
- 41. Isakhani S, Marhamatizade M, Ebrahimi M. The assessment of reducing aflatoxin M1 in kefir by Saccharomyces kefir and Lactobacillus Casei TD4 by ELISA method. Journal of Trends in life Sciences. 2014; 3(4): 268-74.
- 42. Elgerbi M, Aidoo E, Candlish G, Williams G. Effects of lactic acid bacteria and
 bifidobacteria on levels of aflatoxin M1 in milk and phosphate buffer. Journal of
 Milchwissenschaft. 2006; 61: 197-9.
- 43. Montaseri H, Arjmandtalab S, Dehghanzadeh G, Karami S, Razmjoo M, Sayadi M ,et al.
 Effect of production and storage of probiotic yogurt on aflatoxin M1 residue. Journal of
 Food Quality and Hazards Control. 2014; 1: 7-14.
- 44. Adibpour N. Soleimanian-Zad S, Sarabi-Jamab M and Tajalli F. Effect of storage time
 and concentration of aflatoxin M1 on toxin binding capacity of L. acidophilus in
 fermented milk product. Journal of Agriculture Scientific Technology. 2016; 18: 1209-20.
- 643