Assessment of aflatoxin M$_1$ and B$_1$ in some dairy products with referring to the analytical performances of enzyme-linked immunosororbent assay in comparison to high-performance liquid chromatography

Raghda Mohamed Esam$^1$, Ragaa Shehata Hafez$^2$, Nagwa Ibrahim Mohamed Khafaga$^3$, Karima Mogahed Fahim$^4$ and Lamiaa Ibrahim Ahmed$^1$

1. Department of Food Hygiene and Control, Faculty of Veterinary Medicine, Cairo University, Giza, 12211, Egypt; 2. Department of Food Hygiene, Animal Health Research Institute, Giza, Egypt.

Corresponding author: Lamiaa Ibrahim Ahmed, e-mail: lamiaa_13@cu.edu.eg

Co-authors: RME: raghdaesam8888@yahoo.com, RSH: ragaaghoneim@yahoo.com, NIMK: nagwawarda86@hotmail.com, KMF: dr.karima_fh@cu.edu.eg

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Abstract

Background and Aim: Aflatoxin M$_1$ (AFM$_1$) is a major fungal metabolite found in milk coming from aflatoxin B$_1$ (AFB$_1$) contaminated rations and is subsequently present in milk-based products demonstrating a serious public health hazard. This study aimed to investigate the levels of AFM$_1$ and AFB$_1$, in milk and some dairy products consumed widely by infants and children.

Materials and Methods: This study investigated the incidence of AFM$_1$ in 105 samples of processed cheese, Ras cheese, and raw milk (35 of each) retailed in the Egyptian markets. The degree of sensitivity and accuracy was evaluated using the enzyme-linked immunosororbent assay (ELISA) method followed by the estimation of the positive samples using the high-performance liquid chromatography (HPLC) with fluorescence detection. Mold count was determined in the examined samples by investigating AFB$_1$, content using HPLC.

Results: AFM$_1$ was found in all investigated Ras cheese, raw milk, and 82.86% of the processed cheese samples with mean values of 51.05±6.19, 40.27±3.996, and 10.77±1.39 ng/kg, respectively. Moreover, there was statistically no significant difference between AFM$_1$ levels in the core and crust parts of the tested Ras cheese. AFM$_1$ contaminated Ras cheese and raw milk samples were 48.57% and 25.71%, which exceeded the European and Egyptian tolerance levels. Results showed a comparative correlation between ELISA and HPLC methods with no significant difference (p>0.05). Alternatively, none of the examined samples proved to be contaminated with AFB$_1$, despite the presence of mold with mean counts of 3.79±3.29, 4.39±4.34, and 4.84±4.29 log CFU/g in the examined processed cheese, Ras cheese, and raw milk samples, respectively.

Conclusion: Therefore, it is urgent to regularly inspect the contamination of animal feeds with AFB$_1$, and apply special measures and novel techniques to protect the feed and food from public health hazards.

Keywords: aflatoxin B$_1$, aflatoxin M$_1$, enzyme-linked immunosororbent assay, high-performance liquid chromatography, mold, sensitivity.

Introduction

Recently, consumer concerns about food safety have grown; thus, safety evaluation of milk is vital [1]. Recent studies on chemical contaminants have increased the awareness of the public health hazard of chemical residues, which may be present in dairy food. Among these chemicals, mycotoxins proved to be present in more than 50% of foods [2,3].

Mycotoxins are secondary metabolites produced by mold, with aflatoxins (AFs) representing the most toxic and carcinogenic, which can be found as pollutants in various foods, including dairy cattle feed [4,5]. Aspergillus species (Aspergillus flavus, Aspergillus parasiticus, and infrequently Aspergillus nomius) produce a majority of AFs in warm and humid conditions of tropical and subtropical climatic zones at a temperature range of 20-40°C with optimum growth temperature of 25-30°C and a minimum water activity of ≤0.85 [6].

So far, 18 AFs have been discovered, including aflatoxin B$_1$ (AFB$_1$), B$_2$, G$_1$, and G$_2$, which are the major toxins; AFB$_1$ is notoriously the most potent, as its incidence varies depending on the weather, season, geographical location, and storage conditions [7,8]. From its major metabolites, aflatoxin M$_1$ (AFM$_1$) is excreted in the milk after ingestion of feed contaminated with AFB$_1$ by 12-24 h reaching, a high level after limited days [9,10]. AFM$_1$ is comparatively stable, as it is neither affected by processing (pasteurization, sterilization, and mildly acidic conditions) nor storage conditions; hence, it can be detected in cheese...
and other dairy products made from AFM₃ contaminated milk [11,12].

The exposure of infants, teenagers, and prenatal women to the negative effects of AFM₃ is a serious determinant for their health as it has mutagenic, carcinogenic, and teratogenic properties. International Agency for Research on Cancer (IARC) reclassified AFM₃ from Group 2B (probably carcinogenic to people) to Group 1 (proven to be carcinogenic to people) [13,14].

Different methodologies with the consequence of varying sensitivity and accuracy have been indicated for the determination of AFM₃, including thin-layer chromatography (TLC), high-performance liquid chromatography (HPLC), and mass spectrometry (MS) [15], which have excellent sensitivity and accuracy but necessitate extensive sample preparation, costly equipment, and well-trained personnel. Recently, sample screening is conducted using the enzyme-linked immunosorbent assay (ELISA) technique, which gives qualitative or semi-quantitative results, as it is simple, having hand holding validation, and equipment movability. It is mainly used in routine analysis and is reliable for large-scale sample analysis [2,16].

Several nations have established legal limits for this metabolite in milk and milk products that differ from one country to the next and are influenced by economic concerns. The European Commission regulation (165/2010) and the Egyptian standard permitted a level of 50 ng/kg for AFM₃ in milk or processed dairy products [17,18]. This limit is one order of magnitude lower than the 500 ng/kg limit set by the United States and the Codex Alimentarius [19,20].

This study aimed to investigate the prevalence rate of AFM₃ in processed cheese, Ras cheese, and raw milk samples randomly collected from Egyptian markets, referring to the analytical performance and accuracy of the ELISA method compared to HPLC.

**Materials and Methods**

**Ethical approval**

The study did not involve the use of human subjects.

**Study period and location**

The study was conducted from September 2020 to March 2021. The samples were purchased from several dairy shops and supermarkets at Cairo, Giza, and El-Minia Governorates.

**Collection and preparation of samples**

The study sample consists of 105 samples of processed cheeses, Ras cheeses, and raw milk (35 each). Samples were purchased from several dairy shops and supermarkets at Cairo, Giza, and El-Minia Governorates, then transported to a laboratory in a sterilized insulated icebox at around 4°C and analyzed on arrival.

Ras cheese samples were divided into two parts (core and crust) using a sterilized knife at a distance of one inch from the edge of the surface to assess a comparison between the core and crust, and then, each sample was thoroughly mixed.

The samples were analyzed using ELISA and HPLC methods to determine the correlation between the results for evaluating the accuracy and sensitivity degree of the ELISA technique. Total mold count was also estimated, and contaminated samples with mold were examined for AFB, presence.

**Quantitative determination of AFM₃ by the commercial ELISA**

**Preparation of samples**

Twenty milliliters of raw milk were centrifuged at 3500× g for 10 min at 10°C. The fatty layer was aspirated, and 100 μL of the defatted supernatant was used directly within the ELISA kit to determine AFM₃.

Five grams of finely grated cheese were mixed with 20 mL 70% absolute methanol in a tube with a cap. The mixture was extracted by shaking in a shaker for 30 min at 50°C, then, clarified by centrifugation at 3500 g for 10 min. A glass tube was filled with 2 mL aqueous phase, and a couple of milliliters of hexane were added and shaken for 10 s, centrifuged at 3500 g for 10 min. The top layer of the hexane was scraped off, and 100 μL aliquot was applied within the kit.

**ELISA procedure [21,22]**

To the bottom of each well of a microtiter plate, 100 μL antibody was added, gently mixed, and incubated at 25°C for 15 min; wells were emptied and washed with 250 μL washing buffer 3 consecutive times. AFM₃ standard solutions (5, 10, 20, 40, and 80 ng/kg) or test samples (100 μL/well) were added in duplicate, gently mixed, and incubated for 30 min in the dark at a temperature range of 20-25°C. Wells were emptied and washed 3 times, and 100 μL enzyme conjugate was added, gently mixed, and incubated in the dark at 25°C for 15 min. Washing 3 consecutive times was done, then 100 μL substrate/chromogen was added to each well and mixed thoroughly before incubating for 15 min in the dark. Finally, 100 μL stop reagent was added to each well, and the absorbance was determined at 450 nm in the ELISA reader, using special software, RIDA*SOFT Win.net (Art. No. Z9996FF) (r-biopharm, Darmstadt, Germany). In line with the RIDASCREEN kit (Art. No. R1121 german. zip) (r-biopharm) guidelines, the lower detection limit was 5 ng/kg for milk.

**Quantitative determination of AFM₃ by HPLC**

**Sample preparation and extraction**

Milk samples were subjected to chromatographic analysis using the method described by Manetta [15]. The sample was homogenized and centrifuged for 10 min at 3000× g, then, 10 mL aqueous phase was diluted with 10 mL deionized water and purified using a solid-phase extraction-C18 caribagph–4 cartridge, which was conditioned with acetonitrile (5 mL) and deionized water (10 mL). Following the application of the diluted samples and washing with 10 mL water,
20 mL acetonitrile/water (20:80, v/v), and 10 mL n-hexane, AFM was distilled with 6 mL dichloromethane/acetone (95:5, v/v), the elute was evaporated under a gentle stream of nitrogen and the residue dissolved in acetonitrile (200 μL); HPLC analyzed an aliquot (10 μL) of the AFM extract.

Cheese samples were prepared using the method described by Sakuma et al. [23]. Briefly, 10 g of cheese was blended with 40 mL acetonitrile: methanol:water (6:1:3, v/v/v) for 10 min, before being homogenized at 4000 rpm for 5 min, then centrifuged at 3000 rpm for 5 min. Ten milliliters of the supernatant were blended with 30 mL phosphate-buffered saline (PBS; pH 7.4) before filtered using a glass filter (934AH, Whatman plc, Maidstone, Kent, UK). The Immunoffinity column was conditioned with 10 mL PBS, then 20 mL filtrate was loaded onto the column. Finally, the column was washed with 10 mL PBS and 10 mL water. The column was eluted with 1 mL acetonitrile 3 times, and the distill was evaporated under nitrogen gas. Two hundred microliters of trifluoroacetic acid and 200 μL hexane were added to the dried distill, and the mixture was kept for 10 min at 40°C. The mixture was then allowed to dry and eventually dissolved in 1 mL acetonitrile: water (2:8, v/v); then, the solution was filtered through a 0.45 μm filter. The residue was subjected to HPLC.

HPLC procedure

HPLC analysis was applied using an Agilent 1260 series; Agilent Technologies, Waldbronn, Germany. The separation was performed using Eclips C18 column (4.6 mm x 250 mm i.d., 5 μm) (Waters, Milford, MA). Acetonitrile-water (25:75, vol/vol) was delivered to the column at 1 mL/min rate within the mobile phase [24]. The mobile phase utilized isocratic programming. A disposable filter unit (0.45 m) was used to filter the mobile phase. The HPLC system detected AFM$_1$ using a fluorescence detector (RF 20A) at 365 nm (excitation wavelength) and 435 nm (emission wavelength). The injection volume was 10 μL, and the column temperature was 40°C. The detection limit for AFM$_1$ in dairy products was 0.002 μg/L [25,26].

Total mold count was applied according to ISO, 2012 (27)

Total mold count was estimated for all examined samples using Sabouraud’s Dextrose Agar (CM0041B Oxoid™, Belgium). The isolated mold strains on the Sabouraud dextrose slope were subcultured on the Sabouraud dextrose plates using a three-point inoculation technique and incubated at 25°C for 5 days, then, identified macroscopically [28].

Quantifying AFM$_1$ using liquid chromatography-tandem MS

Preparation of cheese samples

Ten grams of cheese were mixed with 60 mL acetonitrile and 50 mL hexane. The mixture was homogenized for 5 min using an Ultra Turrax homogenizer (Sigma-Aldrich, Germany then centrifuged at 4000 rpm for 10 min). The sample was filtered, and the final extracts were dried using nitrogen current. The residue was dissolved in 0.2 mL methanol and filtered into an autosampler vial using a 0.2 mm syringe filter (Pall Gelman Sciences, Ann Arbor, MI, USA) [29].

HPLC-MS MS procedure

The mass spectrometric analysis was conducted using the Alliance 2690 Separations Module, (Waters Alliance, USA) with 10 mL of the sample injected into the C18 column (3.5 μm, 2.1 x 100 mm), and a guard column of the same phase. The extracts were distilled at a rate of 0.2 mL/min. The initial conditions were water-acetonitrile (75:25), for 16 min, followed by water-acetonitrile (10:90) for 24 min. The column was pre-conditioned with 25% acetonitrile. The HPLC system was linked to a MicroMass Quattro Micro triple-quadrupole mass spectrometer (Micromass Ltd., Manchester, UK) with a positive-mode electrospray ionization probe. The compounds were identified and quantified using the multiple reaction monitoring mode [15].

Statistical analysis

The data were analyzed using Statistical Package for the Social Sciences (SPSS) V.17 (SPSS, Inc., Chicago, IL, USA) and reported as a percentage, minimum, maximum, and mean±standard error of the mean. The calibration curve and trend line equation were created using Excel. Data obtained from AFM$_1$ estimation using ELISA and HPLC were compared using t-test (t) in SPSS to see a statistically significant difference between the two methods’ mean results. If p-value <0.05, the mean values of the two methods were supposed to be significantly different.

Results

Table-1 and Figure-1 show the analyzed results of 115 processed cheese, Ras cheese, and raw milk samples for AFM$_1$ contamination. Using the ELISA method; AFM$_1$ was found in 82.86% of the tested processed cheese samples with the mean value of 10.77±1.39 ng/kg, whereas all Ras cheese and raw milk samples contaminated with AFM$_1$ had a mean value of 51.05±6.19 and 40.27±3.996 ng/kg, respectively. When nearly half of the positive ELISA samples (18 samples from each product) were estimated by HPLC for comparing the obtained mean results using the two methods, consequently, evaluating the sensitivity of the ELISA method as a screening test; 83.33, 100, and 100% of the examined processed cheese, Ras cheese, and raw milk samples was confirmed for contamination with AFM$_1$, with mean values of 16.92±2.90, 49.58±7.54, and 49.50±7.23, respectively. The results showed no significant difference between the mean AFM$_1$ values estimated using the two methods (t=1.954, p=0.056).

All examined samples were agreed with the prescribed limit of US regulations for AFM$_1$. None of the
**Figure-1:** Aflatoxin M1 (AFM₁) concentrations in the examined samples using high-performance liquid chromatography. (a) AFM₁ standard (50 ppt). (b) Blank sample. (c) Contaminated sample (26 ppt) and (d) contaminated sample (21 ppt).
examined processed cheese samples exceeded the recommended safety limits of the EC and the Egyptian regulation for AFM$_1$. In contrast, 48.57% and 25.71% of Ras cheese and raw milk samples, respectively, were unacceptable (Figure-2). Our study showed that mold was found in 48.57, 45.71, and 65.71% samples with mean counts of 3.79±3.29, 4.39±4.34, and 4.84±4.29 log CFU/g or mL in the examined processed cheese, Ras cheese, and raw milk samples, respectively (Table-2). 51.43% and 54.49% of the examined processed and Ras cheese samples respectively, were compatible with the Egyptian standards regarding their mold count as illustrated in Table-3.

Figure-3 depicts different mold species isolated from cheese and milk samples. All mold contaminated samples were examined for the presence of AFB$_1$ and the toxin was absent (Figure-4). Results of AFM$_1$ and AFB$_1$ verified that there is no direct relationship between the presence of AFM$_1$ and AFB$_1$ in milk and milk products.

Discussion

Prevalence of AFM$_1$ in the examined samples

Mold contamination was considered a quality issue rather than a threat to food safety, later some species ability to produce toxigenic mycotoxins made it a threat to public health, as they can cause human food poisoning outbreaks in addition to their carcinogenic effect. Consequently, the level of mold contamination, identification of the predominant mold species, and the level of mycotoxins are essential for determining the quality and safety of milk and dairy products [30-32].

Some mold spores produce AFB$_1$, which directly reaches humans through contaminated food or as a metabolic residue in food of animal origin. The liver metabolizes AFB$_1$ into AFM$_1$ and is secreted in the milk. AFM$_1$ is stable in raw milk and processed milk products, making milk and dairy products the primary vehicle for introducing AFM$_1$ into the human diet. AFM$_1$ is hepatotoxic, mutagenic, and carcinogenic; its carcinogenicity is nearly 2-10% greater than AFB$_1$. This made the IARC to transfer AFM$_1$ from Group 2B

Table-1: Statistical analytical results of aflatoxin M$_1$ concentration (ng/kg) in the examined samples using ELISA and HPLC methods.

| Examined samples | Processed cheese | Ras cheese | Raw milk |
|------------------|------------------|------------|----------|
| ELISA (n=35)     | Positive samples | 29.0 (82.86%) | 35.0 (100%) | 35.0 (100%) |
|                  | Min.             | <5.0       | 100.04   | 108.14   |
|                  | Max.             | 27.75      | 108.14   | 108.14   |
|                  | Mean±SEM         | 10.77±1.39 | 66.97±8.39 | 51.05±6.19 |
| HPLC method (n=18) | Positive samples | 15.0 (83.33%) | 18.0 (100%) | 18.0 (100%) |
|                  | Min.             | 2.0        | 10.0     | 10.0     |
|                  | Max.             | 30.0       | 106.0    | 110.0    |
|                  | Mean±SEM         | 16.92±2.90 | 49.58±7.54 | 49.50±7.23 |

*n=Number of examined samples of each food category; processed cheese, Ras cheese, and raw milk. SEM=Standard error of the mean, ELISA=Enzyme-linked immunosorbent assay, HPLC=High-performance liquid chromatography

Table-2: Statistical analytical results of mold count (log CFU/g or mL) of the examined samples (n=35).

| Examined samples | Positive samples | Min. | Max. | Mean±SEM |
|------------------|------------------|------|------|----------|
| Processed cheese | 17               | 4.4  | 3.79±3.29 |
| Ras cheese       | 16               | 5.54 | 4.39±4.34 |
| Raw milk         | 23               | 5.61 | 4.84±4.29 |

CFU=Colony-forming unit, SEM=Standard error of the mean

Table-3: Compatibility of the examined samples with the Egyptian standards regarding their mold count.

| Examined samples | Egyptian standards | Critical limit CFU/g | Compatible samples |
|------------------|--------------------|----------------------|--------------------|
| Processed cheese | ES: 999-2/2005 [64] | Nil                  | 18.0  51.43 |
| Ras cheese       | ES: 1007-5/2005 [65] | Not>10               | 19.0  54.49 |

Figure-2: Compatibility of the examined samples with different regulation standards regarding their aflatoxin M$_1$ (AFM$_1$) content. ES: The Egyptian standards (ES, 7136/2010) [18], EC: European Commission regulation no. 165/2010 [17]. They indicated that milk and dairy products should not contain AFM$_1$, more than 50 ppt. US: United States regulation which established a maximum limit of 500 ppt for milk and milk products [19].
Aspergillus candidus
Aspergillus fumigatus
Aspergillus niger
Aspergillus flavus
Fusarium
candidum
Geotrichum
candidum
Penicillium
Cladosporium
candidum

Figure-3: Frequency of occurrence of the isolated mold strains in the examined samples.
The mean AFM$_1$ content in processed cheese was similar to that obtained by Ahmed et al. [22] and Tahoun et al. [34] who detected AFM$_1$ with a mean value of 24.53±3.91 ng/kg, while higher values were reported by Amer and Ibrahim [35] and El-Kest et al. [36]. Findings related to Ras cheese were in agreement with Amer and Ibrahim [35] and Aiad and Abo Ei-Makarem [11] who found that the mean AFM$_1$ value in the examined Ras cheese was 56.048±6.629 ng/kg; while higher than Hosny et al. [37], Ahmed et al. [22], and Younis et al. [38] who reported that the examined samples of Ras cheese were contaminated with AFM$_1$ with a mean value of 15 ng/kg. The results were lower than those cited by Nassib et al. [39] and Abdel All et al. [40].

The mean values of AFM$_1$ in the core and crust samples of Ras cheese were 75.10±8.43 and 86.97±8.39, respectively, with no statistically significant difference. These results were parallel to those recorded by Bahout and El-Shawaf [41] who examined 50 Ras cheese and found that the mean AFM$_1$ in cheese surface (at depth <5 mm) (6660 ng/kg) was nearly similar to the cheese interior samples (at depth >5 mm) (6540 ng/kg).

However, similar findings of AFM$_1$ in raw milk samples were reported by Yilmaz and Altinci [25], Tahoun et al. [34] and Younis et al. [38]. In comparison, lower detectable levels of AFM$_1$ were recorded by Lee et al. [42] and Elzupir and Elhussein [43]. A higher incidence of AFM$_1$ was recorded by Kirino et al. [44], Nadira et al. [45], and Kagera et al. [46].

Due to the hazardous nature of AFM$_1$ along with its extreme thermal resistance, most countries established legal regulations for AFM$_1$ in raw milk and dairy products with an admissible limit, which varies from 50 ng/kg recommended by EC regulation and the Egyptian standard [17,18] to 500 ng/kg established by the Codex Alimentarius Commission and National Agency for Food and Drug Administration. Our study showed that all positive processed cheese, Ras cheese, and raw milk examined samples agreed with the mentioned limit of the US regulations. None of the processed cheese examined samples exceeded the prescribed safety limits of the EC and the Egyptian regulation. In contrast, 48.57% and 25.71% of Ras cheese and raw milk examined samples, respectively, were unacceptable.

The acceptability of this toxin in milk and dairy products was studied worldwide, from the results that exceeded the limit set up by many countries of 500 ng/kg [40,43]. Alternatively, other studies exceeded the limits imposed by the EC [20,46,47].

The contamination level of milk and milk products with AFM$_1$ varies widely according to dairy feed quality, environmental factors, variation in the original milk contamination, cheese production technologies, type of cheese, extraction, and analytical methods, including regulatory limits for AFM$_1$ in animal feeds, milk, and dairy products [11,48]. In addition, when cheeses were compared to the milk from which they were made, soft cheeses had a 3-fold greater AFM$_1$ concentration while hard cheeses had a 5-fold greater concentration due to the preferred affinity of AFM$_1$ for casein fraction [49].

The most commonly used analytical methods for the quantification of AFM$_1$ in milk include TLC, HPLC with a fluorescence detector (HPLC-FL), and the ELISA [50]. Notwithstanding its extensive and time-consuming sample preparation that necessitates the use of numerous chemical solvents, HPLC-FL is currently the most accurate method [51-53]; however, ELISA gives quick and sensitive results, cost-effective, and requires small sample volumes and fewer preparation procedures. Therefore, ELISA can be a reliable alternative to HPLC-FL and a preferred method at the routine level and in research studies [53]. HPLC-FL as a reference method is used for confirming the obtained ELISA results. Especially due to cross-reaction interferences, particularly at concentrations <50 ng/L, the ELISA method may not be completely reliable as it is resulting in false-positive or false-negative results [53,54].

There is statistically no significant difference between the mean AFM$_1$ values estimated using ELISA and HPLC methods when nearly half of the positive ELISA samples of processed cheese, Ras cheese, and raw milk were reexamined by HPLC; hence, the mean AFM$_1$ was 36.81±4.25 and 37.98±4.31 ng/kg by ELISA and HPLC, respectively (p>0.05). These results are comparable to those reported by Mwanza et al. [50] and Maggira et al. [53]. The obtained results confirm the high degree of ELISA sensitivity and accuracy.

The examined samples revealed contamination with a carcinogen, which remains relatively stable after pasteurization, storage, and preparation of dairy products and poses a serious threat to children and the elderly who consume it, thus, emphasizing the importance of lowering AFM$_1$ levels in milk to the absolute minimum. Therefore, continuous monitoring surveys should be considered in this regard, including the feedstuff ration being kept away from fungal contamination and checked regularly to be free from AFB$_1$ contamination. Further studies and application of new or modern technologies for AFM$_1$ detoxification is necessary [34,55].

**Total mold count in the tested samples**

Contamination of mold in some of the examined samples could be attributed to the unhygienic milking procedures and equipment used for milking, inadequate refrigeration during storage and distribution, inadequate sanitation during manufacturing and ripening, warm weather, and poor personal hygiene, moreover, air and sackcloth packaging of Ras cheese are considered major sources of fungal contamination. The presence of mold in processed cheese indicates post-processing contamination or the survival of mold spores [30,56,57].
The results of processed cheese and Ras cheese samples were similar to those reported by Mohamed et al. [57]. Higher results were obtained by Abdel-Salam and Soliman [58] who found that the mean mold counts of processed cheese and Ras cheese were 5.83±5.80 and 5.56±5.40 log CFU/g, respectively, and Mohamed et al. [31] who examined Ras cheese and found that the mean mold count was 4.85 log CFU/g. While lower results were obtained by Hameed [59] who examined processed cheese and found that the mean mold count was 1.23±0.4 log CFU/g.

Concerning raw milk samples, mold count agreed with that reported by Gurmesa [60] and Amentie et al. [61], while the mold count was lower than that reported by El-Diasty and El-Kaseh [62] and higher than Talukder et al. [63].

On matching the aforementioned results with the Egyptian specification, it was clear that 48.57% and 45.71% of the examined processed and Ras cheese did not match with the Egyptian standards [64,65] for mold count, respectively. These high counts may result in severe economic losses due to the associated visible signs of spoilage as discoloration and off-flavor, with the possibility of mycotoxin production [30,56].

The isolated mold stains in the study were in agreement with those obtained by other researchers; Elbagory et al. [66] and Seddek et al. [67] who showed that Aspergillus was the most predominant isolated mold from Ras cheese and represented by Aspergillus flavus, Aspergillus niger, Aspergillus ustus, and Aspergillus fumigatus. Another study conducted by Mohamed et al. [57] who showed that the most prevalent mold isolates from processed and Ras cheeses were Penicillium followed by Aspergillus.

These findings highlight the importance of employing more stringent sanitary practices to reduce the risks associated with the fungal contamination of milk and milk products, thereby improving the quality as well as the safety of these products; regulatory intervention, including microbiological standards, enhanced sanitation, and food safety programs, should be developed. Biopreservation and novel packaging are also needed to reduce the incidence of mold spoilage in dairy products [68,69].

Incidence of AFB₁ in the contaminated samples with mold

The presence of AFB₁ in milk and milk products may result from ingestion of feedstuffs containing AFB, that the cow liver has not wholly metabolized to AFM₁, therefore, AFB, is found in milk, as well as the contamination of cheese with mold spores that produce AFB, during processing and storage due to the lack or inadequate hygienic measures applied [70].

The absence of AFB in all examined samples was in agreement with the Egyptian regulations, which stated that AFB₁ should be absent in milk and dairy products [18] and the Directive 2003/100/EC of the European Commission establishing a maximum AFB₁ content of 5000 ng/kg in milk and cheese [71]. Results of AFM₁ and AFB₁ verified that there is no straight relationship between AFM₁ and AFB₁ present in milk and milk products.

Similar results were also reported by Montagna et al. [72] and Embaby et al. [73] who stated that all examined buffalo milk cheese samples were consistently negative for AFB₁. In contrast, positive results were recorded by Abou Ayana et al. [74] and Mao et al. [75].

Conclusion

The detection of AFs in food remains an essential subject in a food safety investigation. The current study revealed that processed cheese, Ras cheese, and raw milk samples were contaminated with AFM₁. In addition, there was statistically no significant difference between AFM₁ levels in the core and crust parts of the tested Ras cheese samples (p>0.05). AFM₁ levels in processed cheese did not exceed the maximum limits set by the Egyptian standards, while 48.57% and 25.71% of Ras cheese and raw milk samples, respectively, were above the imposed limit. The examined samples were contaminated with toxigenic mold strains. However, they did not show AFB₁ contamination. AFM₁ results verified no straight relationship between AFM₁ and AFB₁ presence in milk and dairy products. The results of the comparative evaluation of ELISA and HPLC methods demonstrated a satisfactory correlation between both methods with no significant difference (p>0.05). These results recommend that the rapid ELISA method can be used for routine analysis, while HPLC is still the gold standard for confirmation. Overall, AFM₁ prevalence is considered a significant risk to human health; as a result, all milk products must be kept within the allowed limit. Moreover, integrated surveillance programs should be implemented to continuously monitor AFM₁ levels in milk and dairy products. A novel method should be conducted and applied to ensure the safety of milk and milk products for human consumption by avoiding or reducing the presence of these toxic contaminants.

Authors’ Contributions

RME: Conceptualization, methodology, and drafted the manuscript. RSH: Conceptualization, revised the manuscript, and visualization. NIMK: Visualization, methodology, and supervision. KMF: Investigation, methodology, visualization, and revised the manuscript. LIA: Conceptualization, methodology, original draft preparation, investigation, and revised the manuscript. All authors read and approved the final manuscript.

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Competing Interests

The authors declare that they have no competing interests.

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