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The Scourge of Aflatoxins in Kenya: A 60-Year Review (1960 to 2020)

Timothy Omara,^{1,2,3} Ambrose K. Kiprop,^{1,2} Phanice Wangila,⁴ Alex Paul Wacoo,⁵ Sarah Kagoya⁶, Papias Nteziyaremye,^{1,2} Mark Peter Odero,^{1,2} Caroline Kiwanuka Nakiguli,^{1,2,7} and Samuel Baker Obakiro^{1,2,8}

¹ Department of Chemistry and Biochemistry, School of Sciences and Aerospace Studies, Moi University, Uasin Gishu County, P.O. Box 3900, Eldoret, Kenya.

² Africa Center of Excellence II in Phytochemicals, Textiles and Renewable Energy (ACE II PTRE), Moi University, Uasin Gishu County, P.O. Box 3900, Eldoret, Kenya.

³ Department of Quality Control and Quality Assurance, AgroWays Uganda Limited, Plot 34-60, Kyabazinga Way, P.O. Box 1924, Jinja, Uganda.

⁴ Department of Physical Sciences, University of Kabianga, P.O. Box 2030, Kericho, Kenya.

⁵ Department of Medical Biochemistry, School of Biomedical Sciences, College of Health Sciences, Makerere University, P. O. Box 7026, Kampala, Uganda.

⁶ Department of Quality Control and Quality Assurance, Sweets and Confectionaries section, Kakira Sugar Limited, P.O. Box 121, Jinja, Uganda.

⁷ Chemistry Department, Faculty of Science, Mbarara University of Science and Technology, P.O. Box 1410, Mbarara, Uganda.

⁸ Department of Pharmacology and Therapeutics, Faculty of Health Sciences, Busitema University, P.O. Box 1460, Mbale, Uganda.

Correspondence should be addressed to Timothy Omara; prof.timo2018@gmail.com, prof.timo2018@mu.ac.ke

Abstract

Aflatoxins is endemic in Kenya. The 2004 outbreak of acute aflatoxicosis in the country was one of the unprecedented epidemics of human aflatoxin poisoning recorded in mycotoxin history. In this study, a comprehensive review was done to synthesize the country's major findings in relation to AFs, their etiology, epidemiology, detection, quantification, exposure assessment and control in various matrices. Data retrieved indicate that aflatoxins in Kenya are mainly produced by *Aspergillus flavus* and *A. parasiticus*, with the Eastern part of the country reportedly more aflatoxin prone. The toxins have been reported in maize and maize products (*busaa*, *chan'gaa*, *githeri*, *irio*, *muthokoi*, *uji*, *ugali*), peanuts, rice, cassava, sorghum, millet, yams, beers, dried fish, animal feeds, dairy and herbal products, and sometimes in tandem with other mycotoxins. The highest total aflatoxin concentration of 58,000 µg/kg has been reported in maize. At least 500 acute human illnesses and 200 deaths due to aflatoxins have been reported. The causes and prevalence of aflatoxins have been grossly ascribed to poor agronomic practices, inadequate government legislation, lack of awareness, and low levels of education. Low diet diversity has aggravated the risk of exposure to aflatoxins, because maize as a dietetic staple is aflatoxin prone. Detection and surveillance are only barely adequate, though some exposure assessments have been conducted. There is need to widen diet diversity as a measure of reducing exposure due to consumption of aflatoxin-contaminated foods.

1. Introduction

Mycotoxins constitute a family of secondary metabolites biosynthesized by fungi from genera *Penicillium*, *Aspergillus* and *Fusarium* [1]. They contaminate various agricultural commodities prior to or after harvest [2]. Aflatoxins (AFs), ochratoxins, deoxynivalenol (DON), zearalenone (ZEA), fumonisin (FUM) and T-2 toxins are some of the mycotoxins of toxicological priority in foods [3, 4]. In developing countries, AFs and FUMs poses the greatest threat [4, 5]. At least 4.5 billion people in developing countries are chronically exposed to AFs [6], and the recommended sanitary and phytosanitary standards set for AFs in foods affect the economy of most developing nations [7-9].

Aflatoxins are a group of mycotoxins produced by at least 20 fungal strains of *Aspergillus* section Flavi, *Nidulantes* and *Ochraceorosei* [10, 11]. Their discovery and recognition is traced back to 1960 in which Turkey “X” disease was recorded in England with several poults lost to the toxins after feeding on a contaminated peanut ration [12, 13]. AFs were eventually recovered in East Africa (Kenya and Uganda) in peanut rations that caused substantial losses in ducklings [14, 15]. AFs are chemically polysubstituted coumarins with very similar chemical structures [16]. About 20 different types have been reported and aflatoxin B₁ (AFB₁), aflatoxin B₂ (AFB₂), aflatoxin G₁ (AFG₁), aflatoxin G₂ (AFG₂) [17], aflatoxin M₁ (AFM₁) and aflatoxin M₂ (AFM₂) are of demonstrated toxicological importance. The B-aflatoxins, typically pentanone derivatives, exhibit strong blue fluorescence under ultraviolet light while the G-group (six-membered lactones) fluoresce yellow-green under UV light, hence the B and G nomenclature [2, 18]. AFB₂ and AFG₂ are dihydroxy derivatives of AFB₁ and AFG₁ respectively, and therefore usually only reported in the presence of the latter [19]. AFM₁ and AFM₂ are metabolic derivatives of AFB₁ and AFB₂ that exhibit blue-violet fluorescence. They can be present in urine and milk of animals fed on AFB₁-contaminated rations [20, 21].

Aflatoxins are multiplicatively carcinogenic, genotoxic, haemorrhagic, dermatitic, mutagenic, teratogenic and immunosuppressive [17] in the order AFB₁ > AFM₁ > AFG₁ > AFB₂ > AFM₂ > AFG₂ [22-24] (**Figure 1**). This order reflects the role of epoxidation of the 8,9-double bond, and the unique potency of the cyclopentenone ring in the B-series [25]. The mutagenic and carcinogenic effects of AFB₁ and other AFs possessing double bonds between C₆ and C₉ in the furan ring have been ascribed to their hepatic bioactivation to the intermediate metabolite (AFB₁-8,9-epoxide) [26-30]. The reaction is catalyzed by polymorphic cytochrome P450 enzymes [28, 31]. AFB₁-8,9-epoxide is a known mutagen [32], is highly unstable and covalently interacts with nucleophilic sites of cellular macromolecules such as nucleic acids (principally DNA and RNA), inducing irreversible metabolic, signaling, genetic and cell structure dysregulations [26, 33-36]. Details of how AFs induce mutagenicity and carcinogenicity has been discussed in sufficient details in our previous studies [28, 37].

Mutegi et al. [38] published a review on the prevalence and strategies for mitigation of AFs in Kenya from 1960 to 2018. Since then, more than 15 studies on AFs have been undertaken in Kenya. The current review digests the scourge of AFs in Kenya from 1960 to present, highlighting the progresses in the occurrence, detection, quantification, and exposure assessment. Prevention and control measures as well as evidence-based management strategies are discussed.

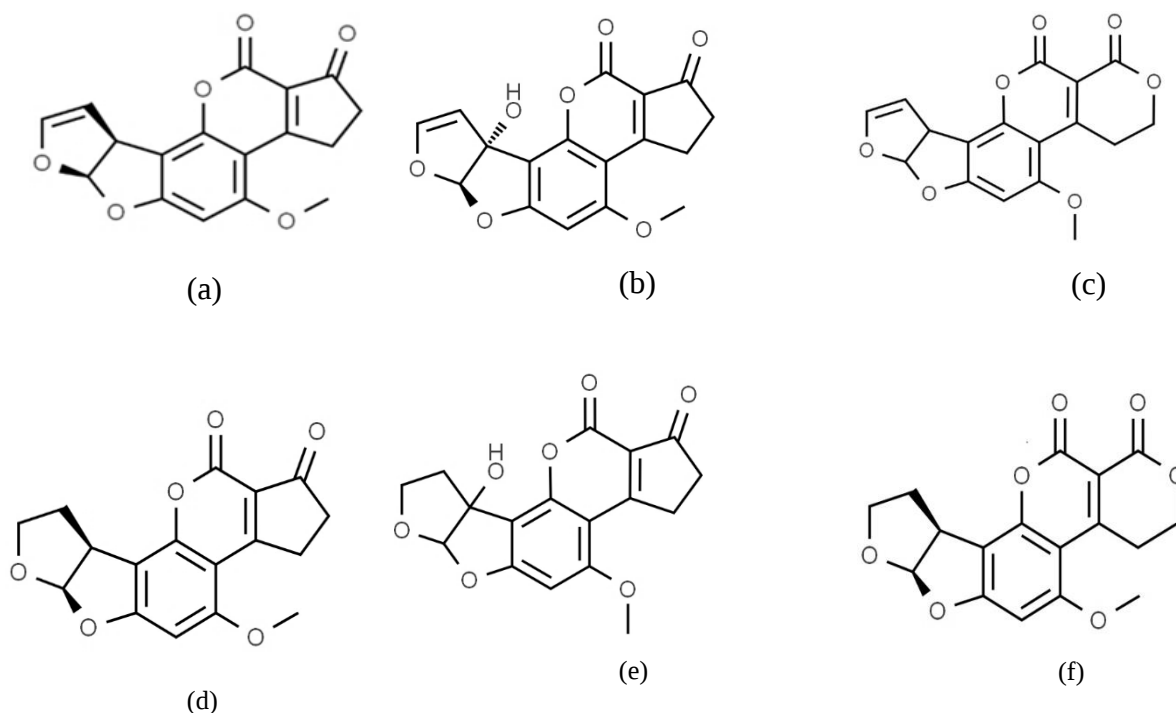


Figure 1: Structure of major AFs of toxicological concern; (a) AFB₁; (b) AFM₁; (c) AFG₁; (d) AFB₂; (e) AFM₂; (f) AFG₂.

2. Occurrence of Aflatoxins in Kenya

2.1 Causative fungi and prevalence of aflatoxins

Aflatoxins in Kenya are majorly produced by *Aspergillus flavus* and *A. parasiticus* [39-49]. *A. flavus* is ubiquitous and produces AFB₁ and AFB₂ plus cyclopiazonic, kojic and aspergillic acids [50]. *A. parasiticus* produces both B and G AFs plus Kojic and aspergillic acids [50-52]. *A. niger*, *A. terreus* and *A. versicolor* were reported in soils and mill dust in Eastern Kenya [40]. Further, the occurrence of *A. caelatus*, *A. alliaceus* and *A. tamarii* in Kenya has been documented [44, 53, 54]. A genetic profiling study reported that *A. minisclerotigenes* in Eastern Kenya exhibited a higher AF biosynthesis potential than *A. flavus* [42]. Though both the L- and S-strain morphologies of *Aspergillus* section Flavi have been reported, probing aetiological studies revealed that aflatoxicoses associated with maize consumption in Kenya has been due to a novel S-morphology fungi previously implicated for the 2004-2006 aflatoxicosis outbreaks [39, 55, 56]. Overall, *A. flavus* is considered the main producer of AFs in most commodities and the optimal growth temperature is 25 °C with a minimum of 0.75 water activity. AF biosynthesis however starts at 10-12 °C [57].

Kenya possesses an erratic tropical climate characterized by periodic droughts, high humidity and high temperatures preceding harvests [58]. The climate varies from tropical along the Coast to temperate Inland, to arid in the North and North Eastern. The country basically have two rainy seasons: the “long rains” from March/April to May/June and the “short rains” from October to November/December. There are four main climatic zones, which can be further subdivided into agroecological zones based on temperature and water requirements of leading

crops. The Central Highlands and the Rift Valley have fertile soils, rainfall of up to 3000 mm per annum and temperatures of 21–26 °C. On the other hand, Western Kenya is hot and remains wet year long. Rainfall is over 1000 mm per annum with temperatures of 27–29 °C. Northern and Eastern Kenya are hot and arid, with annual rainfall of less than 510 mm and temperatures above 30 °C which occasionally reaches 39 °C in some areas [58].

Poor grain conditioning before storage, use of propylene storage bags, drying of grain on bare grounds, insect infestation, poor storage structures (stores with leaking roofs), poor transportation and handling of produce as well as chronic poverty have been criminated for the aflatoxicogenic contamination of Kenyan foods [3, 59-68]. Contamination has also been due to cultivation of maize in ecologically predisposed regions of the country [69-73]. Biophysical factors including soil, host plant susceptibility and genotype, fungal populations (strain specificity and variation, instability of toxigenic properties), low levels of education and awareness, and gender have also favoured the proliferation of AFs in Kenya [60, 67, 74-76]. On toxicological studies, AFB₁ is by far the most studied AF in Kenya, followed by AFM₁ [38]. Thus, most studies reported on the levels of AFB₁, AFM₁ or total AFs. It is worth noting that because of the aflatoxicoses that dawned several times on the country, a number of investigations have been undertaken, with often alarming AF levels reported [38, 63, 70, 77-79].

2.2 Commodities Contaminated

Aflatoxins in Kenya have been reported to contaminate staple foods such as maize (*Zea mays* L.) and its products (*busaa*, *chan'gaa*, *githeri*, *irio*, *muthokoi*, *uji*, *ugali*) [8, 40, 41, 45, 54, 63, 66, 77, 80], sorghum (*Sorghum bicolor* L.) [66, 76, 80], millet (*Eleusine coracana*) [76, 81], pigeon peas, and their local products [80, 82, 83], peanuts (*Arachis hypogaea* L.) and peanut products [53, 61, 62, 80, 84], cassava (*Manihot esculenta* Crantz), rice (*Oryza sativa* L.), dried silver fish (*Rastrienobola argentea*, locally called *omena*) [80, 85], animal feeds [73, 86], dairy products (milk, yoghurt, *Lala*) [57, 66, 87-90] and herbal products [91]. Research on AFs in Kenya has concentrated mostly on maize, peanuts, animal feeds and dairy products, particularly milk [92]. Despite their ubiquitous presence in foods, food processing techniques are not sufficient to completely eliminate AFs from contaminated foods and feeds due to their heat-resistant nature [93].

2.2.1 Cereals and Cereal-Based Products

Maize, millet and sorghum are Kenyan staple foods depending on the region. Maize is the main dietary staple, contributing 65% of food calories and 36% of the total caloric intake [94, 95]. Small-scale farmers store maize under various sub-optimal conditions for up to 4 months before home use or sale [96]. Maize is often for home consumption as flour or used for making *irio* and *githeri* (a traditional dish of maize mixed with legumes or pulses such as beans, pigeon peas and cowpeas, usually cooked whole), though some may be sold [8]. An estimated 60% of maize is processed by consumers using hammer mills [40, 97]. It was previously echoed that maize consumption is the primary route through which Africans have been chronically exposed to AFs [98-100]. On the other hand, millet and sorghum are grown primarily in the semi-arid regions of the country and are consumed mainly as flours used for preparation of thick porridge

(*ugali*) and thin porridge (*uji*). *Uji* is an ingredient of infant weaner foods and diet for children [9].

In Kenya, maize-meal consumption is estimated at 400 g/person/day with an average total AFs content of 0.132 µg/kg and has been criminated for all aflatoxicoses recorded [71, 101]. In one of the pioneering studies, Kenji et al. [81] reported very high total AFs of 1,120 µg/kg in malted maize with an 86% incidence of AFB₁. AFB₁ ranged from 0-260 µg/kg in malted millet from Thika market (Kenya) though no AFB₂ and AFG₁ were detected. On the other hand, maize flour had AFB₁ ranging from 0-160 µg/kg (from Nairobi) and traces (from Thika) with undetectable AFB₂. In another study, 68% of a maize-based traditional brew (*Busaa*) in the slums of Nairobi was declared to contain AFs in concentrations above 5 µg/kg, 17% of which were above 50 µg/kg [102]. Likewise, the magnitude of AF contamination of 480 maize grains, maize flour and dehulled dry maize-*muthokoi* (362 random environmental samples, 26 cases and 92 controls) samples from Makueni, Kitui, Machakos and Thika districts was assessed [103]. It was reported that 46.4% of the environmental samples, 15% of cases and 29.3% of controls were within the then threshold of 20 µg/kg, implying that 54.6% of the samples could not be used for human consumption. Further, 6.9% of the environmental samples, 57.7% of cases and 21.7% of controls had AF concentrations above 1000 µg/kg. The overall AF contamination of the samples ranged from 0-58,000 µg/kg [103]. Further, Sirma et al. [76] recorded total AF levels of 0.17–5.3 µg/kg from 67% of maize collected from different parts of the Rift Valley region which is the major producer of maize in Kenya. About 92% of millet and 50% of sorghum samples collected in the study were positive for AFs in the ranges of 0.14–6.4 µg/kg and 0.21–210.1 µg/kg respectively.

Later, Muthomi et al. [40] reported that samples of whole maize, mill dust and semi-processed maize in Machakos, Eastern Kenya had more than 20 µg/kg AFB₁ threshold allowed by then in Kenya. The highest AFB₁ level (160 µg/kg) was recorded in whole grains. Mill dust had the highest AF contamination, probably due to dehulling operations and the continuous availability of maize products which are potential substrates for *A. flavus* proliferation. As expected, semi processed grains had the lowest AF contamination and this was speculated to be so due to dehulling of the grains as reported elsewhere [104].

Similarly, a cross-sectional survey to assess the extent of market maize contamination and evaluate the relationship between market maize AFs and aflatoxicosis outbreak was conducted [71]. A total of 65 markets were surveyed, 243 maize vendors interviewed, and 350 samples of maize and maize products were taken from the most affected districts as per previous history of aflatoxicoses. About 55% of the samples had AFs in levels above the then advisory threshold of 20 µg/kg, 35% had levels > 100 µg/kg, and 7% had levels > 1,000 µg/kg (**Table 1**). Makueni district which had the highest number of aflatoxicosis case-patients had evidently higher market maize AF concentrations than Thika district (which had the fewest case-patients) with geometric mean of 52.91 µg/kg versus 7.52 µg/kg. In addition, maize sampled from local farms in the affected areas were more likely to have AFs in concentrations > 20 µg/kg when compared with maize purchased from other regions of Kenya or other countries (*odds ratio* = 2.71; 95% *confidence interval*). Because it was understood that contaminated home-grown maize from local farms in the affected areas infiltrated the distribution system, wild AF contamination of market maize was inevitable, and the contaminated market maize bought by farmers after their home-grown supplies were exhausted was cited as a source of continued exposure to AFs. The

authors stressed that efforts to meaningfully interrupt exposure to AFs during an aflatoxicosis outbreak must always consider the potential role of the market system in sustaining exposure [71].

Table1: Distribution of aflatoxins in maize products collected from agricultural markets in some Kenyan districts following the 2004 aflatoxicosis outbreak

District	Number of samples ^a	Total aflatoxin concentration ^b			
		≤ 20 µg/kg (%)	21-99 µg/kg (%)	100-1,000 µg/kg (%)	> 1,000 µg/kg (%)
Makueni	91	32 (35)	12 (13)	36 (40)	11 (12)
Kitui	73	28 (38)	15 (21)	23 (32)	7 (10)
Machakos	102	50 (49)	26 (25)	23 (23)	3 (3)
Thika	76	50 (66)	13 (17)	10 (13)	3 (4)
Total	342	160 (47)	66 (19)	92 (27)	24 (7)

Excerpted from Lewis et al. [71]. Values shown are the number of samples with AFs and the percentage of total samples within the district. ^a Number of samples analyzed for AFs, did not include samples that were collected but not analyzed. ^b Acceptable upper limit for AFs in grains by then was 20 µg/kg.

Probst et al. [55] reported that in Eastern province (Kitui and Mukueni), Coast (Makueni, Kwale, Kilifi, Tana River and Taita Taveta) and Rift Valley (Marakwet, Keiiyo II, Kajaido, Baringo, Nakuru and Laikipia), total AFs in maize ranged from 219.6 to 426.3 µg/kg, 0.1-120.4 µg/kg and below detection limit (BDL) to 13.4 µg/kg respectively. Indeed, The Aflacontrol Project [105] also reconfirmed this observation. Maize grain sampled between January 2010 and May 2010 from fields (pre-harvest), stores (post-harvest), and from wholesalers, retailers, and open-air vendors were declared to be contaminated with AFs. The highest level of AFs in preharvest maize (n = 281) was 1,455 µg/kg from Mbooni East (Eastern Kenya). No appreciable differences were noted between samples from Western and Eastern Kenya. For example, samples from Homa Bay and Rongo had 37 µg/kg and 54 µg/kg of total AFs vis-à-vis 21 µg/kg for Makueni, 25 µg/kg from Mbeere North and 44 µg/kg reported in Mbooni East. Matter-of-factly, more samples from the Western sites were unfit for human consumption (had total AFs > 10 µg/kg) than those from the Eastern sites. For 241 post-harvest samples, 38% from the Eastern region had AF levels above 10 µg/kg. The plague was most acute in Makueni where 87% of samples were unfit for human consumption and the maximum AF level was 1,777 µg/kg. In Mbooni East and Mbeere North, the proportion of maize with levels above 10 µg/kg were 29% and 7% respectively. In entirety, the proportion of maize unfit for human consumption was higher in the Eastern sites than the Western sites but there was considerable variation across the different areas sampled.

Another study reaffirmed the foregoing. For 306 maize samples collected from markets in Upper Eastern Kenya (n = 101), Lower Eastern Kenya (n = 87), Homabay/Rongo (n = 102) and Kisii Central (n = 21), majority (206) had AF levels below 10 µg/kg. However, the Eastern side had more samples with AFs > 10 µg/kg; with a maximum of 1,633 µg/kg recorded. In another concerted study, Collins et al. [106] reported that maize from Homa Bay and Rongo had mean AF levels of 37.0 µg/kg and 54.0 µg/kg compared to 21.0 µg/kg, 25.0 µg/kg and 44.0 µg/kg in Makueni, Mbeere North and Mbooni East respectively.

In consonance with the aforementioned, [107] evaluated the distribution and contamination levels of *Aspergillus* species (spp) and AFB₁ in soil, maize and maize-based products. Maize

grain (n = 256), semi-processed grain (n = 56), flour (n = 52), hammer mill dust (n = 11), and soils (n = 117) had *A. flavus* in all the samples, though the fungi was prevalent in the grain. AFB₁ was undetected in samples from the humid regions but was present in concentrations in excess of 10 µg/kg in 20% of the samples, with maxima of 136 µg/kg for semi-processed maize, 77 µg/kg for whole grain and 41 µg/kg for flour in open bags. Incidental high temperature and periodic droughts prevalent in the semi-arid regions were criminated for the higher levels of *A. flavus* and AFB₁ recorded. Further, a regional report (cited in [16, 108]) indicated that maize in Kenya is the most contaminated in the East African community with a mean total AF content of 131.7 µg/kg (**Table 2**).

Table 2: Per capita food and aflatoxin contamination patterns in Eastern Africa

Food	Country	Per capita food consumption (g/person/day)	Mean AF content (µg/kg)
Maize	Kenya	405	131.7
	Tanzania	69	49.7
	Uganda	400	9.7
Groundnuts (peanuts)	Uganda		25.1
	Tanzania		15.0
	Burundi	65	12.5
Cassava chips	Uganda		0.5
	Tanzania	214	0.9
Sorghum	Tanzania	40	3.0
	Kenya	750 ml	0.8
Milk	Tanzania	750 ml	0.9
	Rwanda	750 ml	Not detected

From the report by the East African Community's AF working group in April 2013 (Dar es Salaam-Tanzania, EAC/TF/405/2013) cited in [16, 108]. **Boldened** means show exceedance of East African threshold limits.

A total of 54 processed, unprocessed (brands A and B) cattle feed from Agricultural and Veterinary stores and 96 human foods (unprocessed and processed maize, polished and unpolished rice, peanut seeds and flour) samples collected from open market traders in Nairobi County were analyzed [86]. The awareness of the traders on AFs and the associated health effects were assessed using questionnaires. Total AF concentrations recorded were 120.9 ± 27.2 µg/kg (processed feed), 77.6 ± 16.0 µg/kg (brand A), 48.6 ± 12.0 µg/kg (brand B), 49.7 ± 14.7 µg/kg (unprocessed maize), 101.20 ± 21.30 µg/kg (maize flour), 38.2 ± 10.5 µg/kg (unpolished rice), 63.9 ± 14.5 µg/kg (polished rice), 54.6 ± 14.8 µg/kg (peanut seeds) and 120.9 ± 27.2 µg/kg in peanut flour. Higher AF levels were reported in processed foods (mean: 95.0 ± 12.7 µg/kg) than in non-processed foods (mean: 47.5 ± 7.6 µg/kg) and this implied that some food processing techniques used predisposed the foods to aflatoxigenic contamination. Roughly 56.6% of the traders were aware of AF contamination; cattle feed traders were more conversant with AFs (40%) than human food traders (17%). A very small portion of food traders (3.7%) and feed traders (8%) were aware of the health effects of AFs in human and animals respectively. Because the mean AF levels in both feeds and foods were above statutory limits, the author recommended the need for creating traders' awareness on AFs, their effects and practices that favour AF proliferation.

Exposure to AFs through consumption of maize and maize products was evaluated through analysis of 20 samples each of maize kernels, *muthokoi* (dehulled maize grains) and *githeri* (maize meal) randomly sampled from households in Kibwezi district, Makueni County of Eastern Kenya [109]. Uncertainty and variability in dietary exposure was modelled quantitatively. AFs were recorded in 45% of maize kernels (range: 18-480 µg/kg), 20% of

muthokoi (range: 12-123 µg/kg) and 35% of *githeri* (range: 6-30 µg/kg). The mean dietary exposure to AFs in maize kernels, *muthokoi* and *githeri* respectively were 292 ± 1567 , 27 ± 154 and 59 ± 62 ng/kg bw/day. The amount and frequency of consumption of the three corn foods were cited as the relevant contributing factors to the risk of dietary exposure to AFs. Moreover, some maize (n = 268), sorghum (n = 62) and millet grains (n = 39) from households and markets in villages of Nandi county were subjected to AF analysis [76]. Computed 67.7% (72/106), 73.3% (44/60) and 65.7% (67/102) of maize samples collected from Laboret, Kilibwoni and Chepkongony sub-locations were contaminated with AFs (range: 0.17-5.3 µg/kg); 92.9% (13/14), 100% (9/9) and 87.5% (14/16) of millet from Laboret, Kilibwoni and Chepkongony had AFs in the range of 0.14-6.4 µg/kg. However, only 50% (9/18), 36.4% (8/22) and 27.3% (6/22) of sorghum drawn from Laboret, Kilibwoni and Chepkongony respectively had AFs above 10 µg/kg (range: 0.15-210.1 µg/kg).

To check for chronic inadvertent exposure to AFs, maize (n = 75) and maize flour (n = 27) from different parts of Kenya were collected and analyzed [95]. Striking differences in the AF levels of maize grain between the regions and stores from which samples were drawn were reported. Samples from Eastern Kenya had the highest contamination with a mean of 22.54 ± 4.94 µg/kg, while those from Nairobi had the lowest contamination (7.92 ± 1.57 µg/kg). No appreciable differences were observed for total AFs in maize flours from Nairobi, Western and Eastern regions. AFs in maize flours were marginally above European Union (EU) limit of 5 µg/kg, and most of the samples had AFs lower than the statutory limit of 10 µg/kg. The authors attributed this to adherence to good manufacturing practices by the millers. The highest AF level in maize flours from Eastern Kenya was 6.98 ± 0.53 µg/kg [95]. Recently, Obonyo and Salano [63] echoed that maize grain in the greater Eastern Kenya harvested after the long rains (May) had significantly ($p = 0.019$) lower AF levels with variation (5.68 ± 6.31 µg/kg, 100% AFB₁) than that of short rains (10.77 ± 10.14 µg/kg, 72% AFB₁). From the long and short rain seasons, the authors hinted that 16% and 44% of the samples respectively had total AFs above the statutory limit of 10 µg/kg. Another group [110] undertook a cross-sectional survey within three agroecological zones: Kitui (Semi-humid to Semi-arid), Nakuru (Semi-humid) and Kitale (Sub-humid to Semi-humid) to determine the occurrence and distribution of total AFs in 130 stored maize samples and the aflatoxigenicity potential of *A. flavus* in the stored maize. The authors put forward that aflatoxigenic contamination between the sampled sites were markedly different ($p \leq 0.001$), with the highest mean AF of 9.68 µg/kg reported in Kitale district. *A. flavus* was isolated in 70% (n=91) of the samples and the isolates with the highest aflatoxigenicity potential were from Nakuru County with a recorded mean total AF of 239.7 µg/kg.

Recently, maize from smallholder farmers' fields in Eastern and South Western Kenya (n = 789) were analyzed for AFB₁ [94]. The authors detected AFB₁ (range: 0.01-9,091.8 µg/kg; mean: 67.8 µg/kg) in 274 of the 416 samples from Eastern Kenya. In South Western, the toxin was detected in 233 of the 373 samples drawn (range: 0.98-722.2 µg/kg; mean: 22.3 µg/kg). Of these, 153 (55.8%) from Eastern and 102 (43.8%) from South Western had AFB₁ surpassing the maximum permissible limit of 5 µg/kg in maize grain. The probable daily intake (PDI) of AFB₁ in Eastern Kenya ranged from 0.07 to 60,612 ng/kg bw/day (mean 451.8 ng/kgbw/day), while for South Western, PDI ranged from 6.53 to 4,814.7 ng/kgbw/day with a mean of 148.4 ng/kgbw/day. The average PDI for both regions exceeded the estimated provisional

maximum tolerable daily intake of AFB₁, which is a health concern for the population in these regions. As such, the study unveiled that preharvest AF contamination of maize were prevalent in both regions and it was advanced that prevention of preharvest infection of maize by toxigenic *A. flavus* strains should be a critical focal point to avert AF contamination and exposure through maize consumption [94].

The prevalence and levels of AFs in freshly harvested maize and freshly milled maize flour (n = 338) from households in Siaya and Makueni Counties were evaluated by Nabwire et al. [77]. All (100%) of the samples had detectable AFs, which ranged from 2.14 to 411 µg/kg. The geometric mean of total AFs in samples from Makueni and Siaya Counties were reported as 62.5 µg/kg and 52.8 µg/kg respectively. This study revalidated the fact that AFs are prevalent in maize and maize products in the studied area. Overall, regional variation in AF contamination of maize in Kenya has been reported with the drought-prone and semi-arid Eastern regions recording higher levels of contamination of up to 58,000 µg/kg [70, 103, 109] compared with the highlands and Western Kenya that have recorded a high of 4,500 µg/kg [72, 78]. Recently, Kenya Bureau of Standards (KEBS) banned a number of maize flour products on the market because of high AF levels [8]. As per the current study, reluctance to dehull maize has been recognized as one of the probable reasons for the high AF concentrations recorded in Kenyan maize flour.

2.2.2 Peanuts (*Arachis hypogaea* L.)

Peanuts (groundnuts) is the only cheap source of dietary proteins in Kenya [65]. It is mainly cultivated in Western Kenya but is sold and consumed countrywide [61, 111]. Peanut productivity has over the years declined due to unpredictable rainfall, lack of disease-resistant peanut varieties, poor agronomic practices as well as poor institutional support accorded to farmers [112-114]. It is primarily for local consumption but is also exported mainly through the World Food Programme [115]. In 2010, FAO statistics indicated production of 99,072 metric tons of peanuts in Kenya harvested from 19,291 hectares [116]. Peanut is rich in proteins (26% to 39%), fats (47% to 59%), carbohydrates (11%), zinc (3.2 mg/100 g), sodium (42.0 mg/100 g), potassium (705.11 mg/100 g), calcium (2.28 mg/100 g), magnesium (3.98 mg/100 g), iron (6.97 mg/100 g), phosphorous (10.55 mg/100 g) and vitamins E and B [117]. In tandem with maize, they are the major portions of the gruel used to make weaning foods in Kenya and these have been shown to be a route of AF exposure [118, 119]. Most of the peanut samples tested in the country had AF levels above recommended regulatory limits set by the KEBS [120]. Fortunately, its consumption is at a lower level, estimated at 1.1 g/person/day [92].

In one of the earlier surveys [61], baseline data on AF levels as well as 384 and 385 peanut samples from Busia and Homabay districts of Western Kenya respectively were collected and analyzed. Total AFs ranged from 0 to 2,688 µg/kg and 0 to 7,525 µg/kg in samples from Busia and Homa Bay respectively. Out of all the samples drawn (n = 769), 87.01% contained < 4 µg/kg of AFs, 5.45% were in the range ≥ 4 and 20 µg/kg while 7.54% surpassed the advisory threshold of 20 µg/kg. There was a highly significant ($\chi^2 = 14.17$; $p = 0.0002$) association between the district of origin of the samples and the analytical total AF concentrations recorded, which was further corroborated by a significant ($\chi^2 = 11.98$; $p = 0.0005$) correlation between total AF levels and agroecological zones. Logistic regression analysis further unveiled

that peanuts from Busia were 2.6 times at risk of contamination vis-à-vis those from Homabay, and that planting improved cultivars could lower the odds of contamination to a half (*odds ratio* = 0.552) those for local landraces. In continuity of the foregoing, the authors [44] reported that the total AF content of 436 peanut samples drawn from Busia and Homa bay districts varied from BDL to 2,687.6 µg/kg and BDL to 1,838.3 µg/kg in about 32% of the samples with detectable AFs. Both the incidence and the number of colonies of *A. flavus* S-strain were significantly and positively correlated with total AF content of the samples. Up to 99.3% of the samples containing < 10 µg/kg of total AFs did not have *A. flavus* S-strain. This corroborated a previous report which confirmed the presence of *Aspergillus*, *Rhizopus*, *Fusarium* and *Penicillium* spp in peanuts with AF contents spanning beyond 100 µg/kg [84].

In another survey by Mutegi et al. [62], peanut and peanut products were drawn from supermarkets and informal markets and analyzed. The authors announced that raw podded peanuts had the lowest AF contamination, with 96% having levels of less than 4 µg/kg and only 4% having more than 10 µg/kg. Irrespective of the provenance, 69% of the samples and 75% of spoilt nuts had total AFs exceeding 10 µg/kg. Though most samples (59%) had AF levels below 4 µg/kg, only 4% of these were acceptable under the KEBS but could be rejected under EU regulations. Of these, 37% of the peanuts were found to be unfit for human consumption as per KEBS and EU regulatory limits. Further, the team [64] evaluated the effect of storage bags, temperature and relative humidity on the quality and AF content of peanut kernels of Homabay Local, Valencia Red, ICGV-SM 12991 and ICGV-SM 99568 varieties stored for 6 months in jute, polypropylene and polyethylene bags. Moisture content, physical damage, rancidity and AF levels were determined before storage and after every 30 days during storage. Moisture content of the peanuts changed remarkably from 3.3 to 6.9% with samples stored in different bag types recording mean values of 5.1% (polypropylene), 5.2% (polyethylene), and 5.3% (jute). Physical damage (range: 0.1 to 9.8%) was influenced by storage temperature and relative humidity, and the type of storage used. Rancidity ranged from 0.8 to 5.3 and increased with storage duration from a mean of 1.5 before storage to a peak of 2.5 after 5 months of storage. There was a reported variation in total AFs (range: 0 to 47.8 µg/kg) of nuts stored in polyethylene bags having 7.3% and 13.4% more contamination than those stashed in polypropylene and jute bags.

Accordingly, it was hypothesized that processing of peanuts in cottage industry could facilitate their contamination by AFs. As such, Ndung'u et al. [53] assessed the AF content of raw and roasted peanuts and peanut butter marketed in Nairobi and Nyanza Provinces of Kenya. Marketers and processors of these were also interviewed on the source of groundnuts and the incidence of *Aspergillus* section Flavi was determined. The authors stressed that the percentage of defective nuts among all unsorted nuts ranged from 0-26.3%. The mean percent defective nuts were higher for Nairobi (imported from Malawi) than Nyanza (home grown) samples. Total AFs in the samples ranged from 0 to 2,377.1 µg/kg with higher mean total levels in raw samples from Nairobi than Nyanza (**Table 3**). The source of groundnuts and defective nuts were positively associated with AF levels. *A. flavus* (L- and S-strains), *A. parasiticus*, *A. niger*, *A. tamari*, *A. alliaceus*, *A. caeleus* and *Penicillium* spp were isolated from the samples.

Table 3: Aflatoxigenic contamination of peanuts and peanut butter from some market outlets in Nairobi and Nyanza Provinces, Kenya.

Nyanza Provinces, Kenya.							
Source	Sample	Sample type	AF level (µg/kg)		Aflatoxin positive samples (%)		
			Range	Mean	≤ 4.0 µg/kg	≤ 10.0 µg/kg	≥ 10.0 µg/kg
Cottage industry	Raw peanuts	Pink regular (n = 3)	BDL-52.4	18.3	60	80	20
		Red regular (n = 1)	NA	5.0			
		Red small (n = 1)	NA	BDL			
Nairobi wholesale outlets	Roasted peanuts	Red regular (n = 8)	2.4-297.7	54.8	25	50	50
	Peanut butter	Paste (n = 11)	BDL-2,377.1	318.3	18	27	73
	Unsorted peanuts	Pink regular (n = 11)	BDL-364.7	111.2	22	26	74
		Red regular (n = 12)	BDL-276.1	89.1			
	Sorted peanuts	Pink regular (n = 4)	BDL-82.4	24.0	36	82	18
		Red regular (n = 5)	2.0-9.2	5.3			
Red small (n = 2)		6.0-7.8	6.9				
Nyanza retail outlets	Unsorted nuts	Pink large (n = 3)	3.7-128.8	71.6	71	75	25
		Pink regular (n = 9)	BDL-229.8	44.9			
		Red regular (n = 9)	BDL-14.0	1.9			
		Red mixed (n = 3)	NA	BDL			

Adapted from [53]. BDL: Below method detection of 0.5 µg/kg limit, NA: Not applicable.

The prevalence and diversity of fungal spp and aflatoxigenic contamination of 228 marketed peanut samples (from 140 formal and 88 from informal markets) in Kericho and Eldoret towns of Kenya were established [115]. *A. flavus* (L- and S-strains), *A. parasiticus*, *A. tamarii*, *A. caelatus*, *A. alliaceus* (members of *Aspergillus* section-Flavi) and *A. niger* as well as *Penicillium*, *Mucor*, *Fusarium* and *Rhizopus* spp were encountered. Total AFs in the nut products ranged from 0 to 2,345 µg/kg in raw peanuts, 0 to 382 µg/kg in roasted coated peanuts, and 0 to 201 µg/kg in roasted decoated peanuts. Altogether, AFs occurred in higher concentrations in samples from informal (mean = 97.1 µg/kg) than formal (mean = 55.5 µg/kg) markets. Meanwhile, a positive and significant correlation ($R^2 = 0.63$; $p \leq 0.05$) was cited between AF levels and the major aflatoxigenic fungi in raw peanuts from formal markets of Eldoret. Further, AFs in raw nuts from informal markets in Kericho positively and strongly correlated ($R^2 = 0.81$; $p \leq 0.05$) with the population of *A. flavus* (both strains). In roasted coated peanuts sampled from Eldoret formal markets, AFs correlated positively and significantly ($R^2 = 0.37$; $p \leq 0.05$) with *A. flavus* S-strain.

Another investigation [121] which compared the oil content and total AF level of peanuts in Busia and Kisii Central districts reported that Valencia red, Uganda local, Homa Bay local and Local red peanut varieties from Busia had lower levels of total AFs except the Local red variety which had the highest total AF of 267 µg/kg with the lowest average oil content of 42.7% (**Table 4**). Peanuts from Kisii Central had higher AF levels and low oil contents. Summed up, there was an increase in total AF levels with decreasing oil contents ($r = -0.496$, $p = 0.031$) except for Uganda local red from Kisii. In continuity of the foregoing study, Menza and Muturi [49] reported the occurrence of five causative *Aspergillus* fungi: *A. flavus* (L- and S-strain), *A. parasiticus*, *A. niger* and *A. tamari*. Overall, the occurrence of *A. flavus* (both strains) was significantly higher than other aflatoxigenic spp identified in the nuts. *A. flavus* L-strain was the most common isolate (58.8%) in samples from Busia while the S-strain dominated (60.2%) in peanuts from Kisii Central. All in all, *A. flavus* S-strain was the most dominant with a mean prevalence of 45.1%.

Table 4: Oil content and total aflatoxins of peanuts from Busia and Kisii Central districts of Kenya

District	Variety	Mean oil content (%)	Mean total AFs (µg/kg)
Busia	Valencia red	47.2	2.3
	Uganda local red	46.7	2.4
	Homa Bay local	43.2	2.8
	Local red	42.7	267.0
Kisii central	Valencia red	46.6	93.0
	Uganda local	45.7	405.0
	Homa Bay local	40.6	101.5

Adapted from [121]. Mean values in **bold** are higher than maximum permissible limits of 10 µg/kg.

From the prevent reports, it can be noted that relatively higher concentrations of AFs have been reported in peanuts in Kenya. A plausible explanation advanced has been that aflatoxigenic fungi contaminate the shells, testa and seeds as the pods grow in the soil. Further, mechanical damage during harvest, drying and storage further increases the chances of fungal contamination and mycotoxin production. This is corroborated by a Tanzanian report which unveiled that grains and oilseeds from maize, sorghum, and sunflower produced in above the ground reproductive structures had relatively lower AF contamination vis-à-vis those produced in geocarpic structures of peanuts and Bambara nuts [122].

2.2.3 Cassava (*Manihot esculenta* Crantz)

Cassava is an important food crop due to its high dietary carbohydrate content. The main food sources are starchy tuberous roots, though the proteinaceous young leaves are also edible [123, 124]. However, cassava contain two cyanogenic glucosides: linamarin and lotaustralin (methyl linamarin) which are normally produced for defence against predators. These cyanogens are distributed widely throughout the plant, with the highest amounts in the leaves and the root cortex and lower amounts in the interior of the root parenchyma [125]. Cyanide inhibits cellular respiration of aerobic organisms by blocking mitochondrial electron transport and preventing oxygen uptake. In addition, cassava is also prone to mycotoxins, particularly AFs.

In a study, dried cassava chips (n = 13) and cassava flour (n = 26) sourced from Nairobi and Mombasa markets were assessed for hydrogen cyanide, AF and moisture contents [126]. Hydrogen cyanide ranged from 27.20 to 42.92 mg/kg and 21.45-37.77 mg/kg in cassava chips, 21.53 to 64.63 mg/kg and 21.70 to 70.03 mg/kg in flour from Nairobi and Mombasa respectively. These were all above 10 mg/kg recommended by the East African standards (EAS 739: 2010 and EAS 740: 2010 respectively). AFs were detected in 2 flour samples from Nairobi (mean levels of 6.60 and 8.89 µg/kg), and a sample from Mombasa (mean: 2.84 µg/kg). Moisture content ranged from 8.62-9.98% and 8.85-11.57% in cassava chips, and 8.50-12.51% and 7.30-11.0% in flour samples from Nairobi and Mombasa respectively. The study revealed that marketed cassava flour though of good aesthetic quality could be mycotoxigenically unsafe for consumption.

There are no reports in open literature on plant products such as sugarcane, spices, beans, wheat and barley in Kenya. *A. flavus* was not isolated from soils under sugarcane cultivation which had *A. niger*, *Fusarium equiseti*, *Trichoderma viride* and *Phanerochaete chrysosporium* [46]. Sugar cane is one of the daily consumables in form of sugar and has been previously reported to house AFs [127]. In addition, no recent studies has reported on the AF content of commercial

beers consumed by Kenyans, yet it is among the most consumed foods that perhaps use all the major cereals: maize, sorghum, and barley as well as cassava. Beers are practically products of mixed-culture fermentations, a process that continues up to consumption time. As such, brewing is an ideal route for exposure to AFs as it offer auspicious conditions for aflatoxigenic fungal growth and creates an avenue for use of contaminated grains as the final consumers will not be able to physically detect [16]. Similarly, no reports exist on AFs in beans. In the neighbouring Uganda where sometimes Kenya import beans, AFs were earlier recorded in excess of 1,000 µg/kg [128].

2.2.4. Animal Products

Aflatoxin contaminated animal products such as blood, eggs, ghee, meat, milk and dairy products present food safety concerns [129]. In Kenya, AFM₁ in bovine milk is the most studied. A list was developed [58] of the regions in Kenya that are at risk of AF outbreaks from milk consumption, and this encompassed all the milk production areas of Kenya. Milk production is mainly from dairy cattle, mostly crosses between dairy and zebu breeds, which produces over 70% of the total national milk output. They are fed on natural forage, cultivated fodder and crop byproducts such as maize stalks and stover. Supplements such as dairy meal, maize germ, maize bran, cottonseed cake, wheat pollard and wheat bran are also sometimes used [74].

A correlative study conducted in four urban centers by Kang'ethe and Lang'a [89] analyzed 613 milk and 830 feed samples for AFM₁ and AFB₁. About 86% (353/412) of the feed samples from farmers were positive for AFB₁ and 67% (235/353) of these exceeded the FAO/WHO limit of 5 µg/kg. About 81% (197/243) of the feeds from feed millers and 87% (153/175) from agrochemical shops were AF positive, with 58% (115/197) and 66% (92/153) of these samples exceeding permissible limits respectively. Approximately 72% (315/439) of the milk from dairy farmers, 84% (71/85) from large and medium scale farmers and 99% (88/89) of the pasteurized marketed milk were positive for AFM₁, and 20%, 35% and 31% of positive milk from dairy farmers, medium and large scale farmers and market outlets respectively exceeded the WHO/FAO limits of 0.05 µg/kg. On the one hand, 67% of the urban smallholder dairy farmers had knowledge that milk could be contaminated with AFM₁ but did not know the possible exposure mitigation strategies. Feed millers, on the other hand, knew about AFB₁ in grains and its excretion as AFM₁ in milk but were not alleviating exposure to animals [89] (Table 5). Similarly, Sirma et al. [130] surveyed 286 households in 37 villages representing four agroecological zones (semi-arid, temperate, sub-humid and humid). They drew 280 samples of bovine milk which were subjected to AFM₁ analysis. AF levels were from 0 to 0.359 µg/kg. Generally, 58.9% of the milk sampled had AFM₁ levels BDL though 9.3% exceeded the WHO/FAO limit of 0.05 µg/kg (Table 6).

497

Table 5. Synopsis of aflatoxins in animal feeds and bovine milk in some Kenyan municipalities (from [89])

	Source/Municipality	% AF positive	> 5.0 µg/kg (%)	Mean	Range
Aflatoxin B ₁ (Feed samples)	Urban small holder dairy farmers				
	Nyeri (n = 118)	68.6	49.2	136.0 ± 10.0	4.0-63.0
	Eldoret (n = 108)	98.1	61.1	23.2 ± 23.2	4.2-178.2
	Machakos (n = 99)	94.9	73.3	27.7 ± 74.9	3.6-595.0
	Nakuru (n = 87)	80.5	58.6	17.4 ± 11.1	1.8-58.0
	Feed manufacturers				
	Nyeri (n = 14)	100.0	42.9	6.4 ± 4.9	1.9-15.8
	Eldoret (n = 18)	88.9	66.7	13.9 ± 12.8	1.9-49.0
	Machakos (n = 1)	100.0	100.0	43.8 ± 0.0	43.8
	Nakuru (n = 171)	77.8	43.3	26.0 ± 44.5	0.9-280.0
	Nairobi (n = 390)	84.6	56.4	13.0 ± 15.9	0.9-280.0
	Agrochemical shops				
	Nyeri (n = 19)	89.5	31.6	8.9 ± 8.5	1.9-28.7
	Eldoret (n = 58)	93.1	72.4	17.0 ± 34.6	1.8-238.0
Aflatoxin M ₁ (Milk samples)	Urban small holder dairy farmers				
	Nyeri (n = 120)	60.8	3.3	33.8 ± 68.7	5.0-46.0
	Eldoret (n = 107)	68.2	10.3	39.9 ± 39.7	5.4-228.0
	Machakos (n = 99)	82.8	24.2	99.7 ± 168.9	5.1-780.0
	Nakuru (n = 110)	77.3	20.9	83.3 ± 129.3	5.2-550.0
	Medium and large scale farmers				
	Nyeri (n = 25)	76.0	0.0	20.2 ± 29.0	5.2-50.0
	Eldoret (n = 16)	68.8	12.5	115.6 ± 202.7	5.5-560.0
	Machakos (n = 7)	100.0	50.0	52.2 ± 34.7	10.9-102.5
	Nakuru (n = 27)	89.9	55.6	65.1 ± 36.7	5.3-165.0
	Nairobi (n = 10)	100.0	50.0	99.8 ± 97.3	10.0-245.0
	Marketed milk				
	Nyeri (n = 10)	100.0	30.0	129.3 ± 198.8	16.5-600.0
	Eldoret (n = 18)	100.0	22.2	36.4 ± 24.5	5.8-74.0
	Machakos (n = 18)	94.4	16.7	33.1 ± 17.0	11.0-67.0
	Nakuru (n = 19)	100.0	36.8	36.1 ± 22.9	8.0-71.0
	Nairobi (n = 24)	100.0	41.7	64.9 ± 76.4	7.9-300.0

498

n = number of samples. Means were presented with errors as standard deviations.

499

Table 6: AFM₁ contamination of bovine milk in some selected agroecological zones of Kenya (from [130]).

County	Agroecological zone	Number of samples	AFM ₁ positive samples (%)		
			< 0.002 µg/kg	≤ 0.002-0.05 µg/kg	≥ 0.05 µg/kg
Tharaka Nithi	Humid	64	34	41	25
Kwale	Sub-humid	29	76	17	7
Bungoma	Temperate	64	53	41	6
Kisii	Temperate	63	65	30	5
Isiolo	Semi-arid	60	77	22	2

500

501 Aflatoxins were detected and quantified in fresh and sun-dried *Rastrineobola argentea* (Dagaa
502 fish) collected from various markets in Luanda, Rongo, Kisumu, Ahero and Maseno of the
503 Winam gulf of Lake Victoria [85]. Fresh samples had no detectable AFs, but the dried samples
504 had mean total AF levels of 0.34 ± 0.09, 0.21 ± 0.00, 0.25 ± 0.06, 0.53 ± 0.11 and 0.11 ± 0.00
505 µg/kg wet weight respectively. It was asserted that the occurrence of AFs in processed *Dagaa*
506 fish (*omena*) could have been due to the fact that the samples were collected from the markets
507 in July 2010 when there were rains and drying was incomplete, thus the sun-dried *Dagaa* fish
508 were packed in sacks when they were incompletely dried which favoured the growth of moulds.

In a bid to assess the AF status of marketed raw milk and associated risk factors in peri-urban Nairobi, raw milk retailers in Dagoretti division were interviewed and milk samples were drawn and tested for AFM₁ [131]. Four types of businesses were found: kiosks (71%), dairy shops (21%), street or mobile vendors (3%) and grocery stands (1%); for 4%, the business type was not identified. Milk was mainly sourced directly from dairy farms (59%) or from intermediate distributors (35%). Although 58% of the retailers had heard about AFs and the majority of them agreed AFs could be present in milk, only 29% believed that “milk safety cannot be solely judged by sight or taste” and only 6% that “milk is not completely safe even after boiling”. Analysis of the milk samples recorded mean AFM₁ of 0.1287 µg/kg (median = 0.0499 µg/kg; maximum of 1.675 µg/kg). In entirety, 55% of the samples exceeded the EU maximum level of 0.05 µg/kg and 6% exceeded the recommended maximum level of the United States Food and Drug Administration of 0.5 µg/kg. Vis-à-vis milk from street vendors, a significantly higher AFM₁ concentration was detected in milk from kiosks and dairy shops, especially when the milk were sourced from farms without an intermediate distributor. Similarly, it was reported that 156 samples out of 185 (150 raw milk and 35 processed milk and milk products) from Bomet County were positive for AFM₁ with an overall prevalence of 84.32% [132]. About 43.8% of these were above 0.05 µg/kg, with raw milk compared to processed milk (52% vs. 8.6%) having more contamination.

In the same manner, AFM₁ was detected in 291 samples of raw, pasteurised and UHT milk, yoghurt and *Lala* [90]. Monthly samples were drawn over a period of 1 year, just as a consumer would purchase them from retailers and traders in a low-income area (Dagoretti), and a major supermarket in a middle/high-income area (Nairobi). More than 50% of the samples had AFs exceeding 0.05 µg/kg, though only 3 exceeded 0.5 µg/kg and the geometric mean AFM₁ level was 0.0619 µg/kg in the 135 samples from Dagoretti while it was 0.0361 µg/kg in 156 samples from Nairobi. The levels varied significantly depending on the time of year, with the lowest levels reported in January. UHT milk had the lowest AF levels, and more expensive milk had lower AFM₁ levels [90].

In a recent study [129], it was pointed that exposure to AFM₁ in milk and the health risks associated with it are not clearly understood and monitored in Kenya. Thus, the team assessed the awareness, knowledge and practices of urban and peri-urban farmers about AFs and evaluated the levels of AFs in on-farm milk in Kasarani sub-county, Nairobi County. In total, 84 milk samples were analyzed, and 90% (83/84) were analytically declared to be contaminated with AFM₁ (mean value of 0.084 µg/kg). About 64% of the samples had AFM₁ levels well above EU limit of 0.05 µg/kg. Though 80% of the farmers had knowledge of AFs, no correlation existed between the farmers’ knowledge and gender with AFM₁ prevalence.

Kang’ethe et al. [87] reported that 45.5% and 98.6% of bovine milk and animal feeds in Kenya were positive for AFs. About 49% and 83% of these had AFM₁ above 0.05 µg/kg and AFB₁ above 10 µg/kg respectively. Similarly, an AF risk mapping study from milk consumption using biophysical and socio-economic data [58] reported a mean AFB₁ content of 9.25 µg/kg in animal feeds and mean AFM₁ content of 0.0265 µg/kg in bovine milk. Higher mean of the logarithmic AFB₁ concentrations were reported in areas with historical aflatoxicosis outbreaks compared to those without outbreak history, a phenomenon that was not true for the mean logarithm of AFM₁ when compared between areas with and those without history of aflatoxicosis outbreaks. Analogously, a cross-sectional study of aflatoxigenic contamination of

bovine milk and dairy concentrates was done in five counties of Kenya representing the agroecological zones: Kwale, Isiolo, Tharaka-Nithi, Kisii and Bungoma [133]. Concentrates and milk were collected twice (during the dry season and rainy season) from 285 farmers in the five counties and analyzed for AFB₁ and AFM₁. Between 0-68% used concentrates, which had AFB₁ ranging from < 1 µg/kg to 9,661 µg/kg with 47.8 to 90.3% positive samples. About 33.3% to 87.5% of the concentrates had more than 5 µg/kg AFB₁ (83.3% to 100% from retailers and 28.6% to 100% from manufacturers). AFM₁ prevalence in milk was lowest in Kwale (13.6%) and highest in Tharaka-Nithi (65.1%). About 3.4% (Kwale) to 26.2% (Tharaka-Nithi) of milk samples had AFM₁ above the WHO/FAO threshold of 0.05 µg/kg, with the highest contamination of 6.999 µg/kg. The study was in consonance with preceding studies which indicated that AFs are prevalent in Kenyan dairy rations and milk.

In Kisumu, it was cited that 97 randomly selected dairy farmers primarily fed cows on forage and concentrates (62.9%). Levels of AFM₁ in milk collected from these farms ranged from BDL to 0.151 µg/kg (mean of 0.02967 µg/kg) and 26.4% of these exceeded the EU limit. Concentrate feeding was associated with higher AFM₁ levels so that farms feeding concentrates were more likely to record milk AF levels above 0.05 µg/kg [134]. Further, the prevalence of AFM₁ in 96 samples of informally marketed milk from Nairobi, the knowledge of milk traders on AFs as well as the effects of boiling and fermentation on AFM₁ were assessed [135]. By and large, all samples had detectable AFM₁ (limit of detection = 0.005 µg/kg) with a mean of 290.3 ± 0.663 µg/kg. About 64% of the samples had AFM₁ above 0.05 µg/kg while 7.5% exceeded 0.5 µg/kg. Majority of the traders had low (69.8%) or medium (30.2%) knowledge of AFs. The educated and female traders were more knowledgeable, and fermentation of milk to *Lala* (a traditional fermented drink) or yogurt significantly reduced AFM₁ levels by 71.8% (in *Lala* after 15 hour room temperature incubation) and 73.6% in yogurt after incubation at 45 °C for 4 hours. Boiling however, had no appreciable effect on AFM₁ levels [135].

According to Sirma et al. [9] using a quantitative risk model, an equivalent of 5 hepatocellular cancer cases and deaths, and the disability-adjusted life years of 255 for Kenya in 2016 were estimated as due to exposure to AFs in milk. Other than milk, there are no reports in open literature on AF content of other products of animal origin such as blood, eggs, ghee and meat in Kenya.

2.2.5 Animal Feeds

As pointed earlier, farming is one possible exposure route to AFs. For example, maize which is known to be highly susceptible to AF contamination in Kenya is also a major component of livestock and poultry feeds, and therefore, regular indirect human exposure through the consumption of animal products that contain AF residues cannot be underrated. Elevated levels of AFB₁ have been recorded in Kenyan animal feeds [88, 89]. Protein-rich supplements (cottonseed cake, sunflower cake, fish-meal and other oil seed byproducts), cereal grains and their byproducts (maize bran, maize germ, wheat bran) are a rich source of nutrients for moulds [92]. These fungi readily contaminate crop residues and homemade dairy concentrates as a result of poor handling and storage conditions in smallholder farms. The situation is exacerbated by dairy farmers' habit of utilizing spoilt (pest- or mould-damaged, and rotten) grains for formulation of dairy rations [74, 92]. A study carried out on animal feeds in Nairobi

province revealed that AFs ranged from 5.13 µg/kg to 1,123 µg/kg, with the largest proportion lying between 11 µg/kg to 99 µg/kg [78].

Further, 81 fish feeds sourced from 70 farms and 8 feed manufacturing establishments located in Nyeri, Kenya were subjected to AF analysis by Mwihia et al. [73]. Fish were also sampled from 12 farms for gross and microscopic pathological investigation. About 84% of the feeds were AF- positive (range 1.8-39.7 µg/kg, mean of 7.0-8.3 µg/kg, median of 3.6 µg/kg). About 18.5% of the feeds sampled registered total AFs above the statutory limit of 10 µg/kg. Meanwhile, homemade and tilapia feeds had evidently higher AF levels than commercial and trout feeds. Maize bran-based feeds and fish meal recorded higher AF levels than those devoid of these constituents. Microscopy revealed that five trout farms (41.7%) had fish with swollen abdomens, and enlarged livers with white or yellow nodules, large dark basophilic hepatic cells with hyperchromatic nuclei in irregular cords. As such, the authors inferred that aflatoxigenic contamination of fish feeds is a scourge in Nyeri which if left unchecked may cause detrimental health effects in edible fish in the area.

2.3 Co-Occurrence of Aflatoxins with Other Mycotoxins

Several mycotoxins can occur simultaneously in matrices [136]. In 1995, Muriuki and Siboe [45] analyzed 40 samples of flour packed in 90 kg bags, 58 samples of *Ugali* brand and 74 samples of *Jogoo* brand drawn from the Nairobi, Kenya. The samples were analyzed for resident mycoflora and some mycotoxins associated with key fungal spp. *Aspergillus flavus*, *A. sulphureus*, *Fusarium moniliforme*, *Penicillium stoloniferum* and *P. cyclopium* were the reported fungal spp in the samples. Ochratoxin A was the most prevalent mycotoxin, and all the flour brands had AFB₁ and AFB₂ (0.4-20 µg/kg), Ochratoxin A (50-1,500 µg/kg) and ZEA (2,500-5,000 µg/kg). The authors recommended the need for rigorous countrywide monitoring of mycotoxins in maize both at farm and market levels. The foregoing was substantiated by a report by Kedera et al. [137] who reported the presence of *Fusarium* fungi and fumonisin B₁ (FB₁) in maize kernel samples from smallholder farm storages in Bomet, Bungoma, Kakamega, Kericho, Kisii, Nandi, Siaya, Trans Nzoia, and Vihiga districts in the tropical highlands of Western Kenya. Later, Mbugua and Gathumbi [138] affirmed the occurrence of AFB₁, FB₁, ZEA and DON in 36 Pilsner and 39 Tusker beer samples sourced from Nairobi and the surrounding satellite towns. All the samples were negative for AFB₁; the prevalence of DON and ZEA were 100% in both brands while FB₁ incidence was 72%, with incidences in Tusker (76.9%) being markedly higher than in Pilsner (66.7%). The mean values of contamination were 3.29 and 3.57 ng/mL for DON, 0.28 ng/mL and 0.32 ng/mL for FB₁ and 7.84 and 8.50 pg/ml for ZEA in Tusker and Pilsner brands respectively. A positive correlation was reported between DON and FB₁, and DON and ZEA, affirming their co-occurrence to be from *Fusarium* spp. This communication suggested that there were some but safe exposure to *Fusarium* mycotoxins by lager beer consumers of Kenya.

Fumonisin B₁ and AFB₁ in symptomless and rotten maize harvested at different harvest time points after physiological maturity (HTPAPM) from Malava and Tongaren were evaluated [139]. *Fusarium verticillioides* dominated at all HTPAPM though *F. graminearum*, *F. subglutinans*, *A. flavus*, *A. parasiticus* and *Sternocarpella maydis* were also encountered. FB₁ concentrations in symptomless maize ranged between 22 to 1,348 µg/kg with mean levels of 56, 80 and 317 µg/kg respectively at 4, 8, and 12 weeks HTPAPM for Malava in the year

2001. In Tongaren during the same year, mean FB₁ levels of 41, 179 and 590 µg/kg were recorded at 4, 8, 12 weeks HTPAPM respectively. The concentration of FB₁ in rotten maize ranged from 39 to > 5,000 µg/kg and increased with HTPAPM. The highest AFB₁ level was 17.0 µg/kg in rotten maize. The authors hinted that the isolation of *F. subglutinans* and *F. graminearum* was an indication that other mycotoxins (DON, ZEA and moniliformin) associated with infertility and hypoestrogenism could be inevitable in the samples.

In a study scrutinizing commodities, feeds and feed ingredients from Middle East and Africa [140], 48% (12/25) samples from Kenya were positive for B-Trichothecenes (mean: 422 µg/kg, maximum: 3859 µg/kg), none had A-Trichothecenes, 76% (19/25) had FUM (mean: 956 µg/kg, maximum 10,485 µg/kg), 56% (14/25) had ZEA (mean: 67 µg/kg, maximum: 167 µg/kg), 78% (21/27) had AFs (mean: 52 µg/kg, maximum: 556 µg/kg), while 50% (1/2) had Ochratoxin A (mean: 2 µg/kg). A gluten sample from Kenya presented the highest level of FUM found in the whole survey (10,485 µg/kg).

Similarly, maize samples were collected from 30 markets in diverse agroecological zones of Meru, Machakos and Kitui counties during the 2013 harvest [54]. *Fusarium* and *Aspergillus* spp were isolated from the samples. Total AFs in Meru, Kitui and Machakos samples were beyond the threshold of 10 µg/kg. Meru had both the highest and lowest level of AFs detected (115.7 µg/kg and 0.3 µg/kg respectively). FUMs were reported in levels above the acceptable limits in Meru and though detected in Kitui and Machakos, the contamination levels were within acceptable limits. Utilizing a near infra-red single kernel sorting machine, removal of AF and FUM-contaminated kernels was perfected with up to 97.8% efficacy for AFs and 60.8% for FUM. The accepted fractions had statistically lower mycotoxin levels than the rejected maize [54].

The prevalence of AFs and FUM was investigated in maize intended for immediate human consumption in Eastern Kenya. Samples were collected from people who brought their maize for processing at local commercial mills [75]. Interviews and sampling of maize flours was done for 1,500 people who processed maize at 143 mills in 10 administrative districts. Mycotoxin analysis revealed that 39% and 37% of the samples respectively had AFs and FUM in levels above tolerable limits. Samples with AFs above 10 µg/kg were 22-60% across the districts. A higher occurrence of AFs was associated with smaller maize farms, lower grain yield, and monocropping systems. A larger magnitude of the toxin was observed in the sub-humid agroecological zone, in samples with more broken kernels, and less maize ear damage at harvest. Further scrutiny of paired grain samples (visually sorted and unsorted) showed that sorting reduced FUM by 65% to below the advisory threshold of 1,000 µg/kg. Sorting did not, in essence, have any effect on AF concentration [75].

Besides, the presence of AFs, FUM and DON in *Busaa* (a maize-based traditional beer) in Bomet County, Kenya was reported [141]. Of the 61 samples obtained from homesteads involved in brewing in the North Eastern part of Bomet East constituency, 93%, 9.8% and 23% respectively were contaminated with AFs (mean: 5.2 ± 0.2 µg/kg; range: 2.8-11 µg/kg), FUM (mean 1460 ± 188 µg/kg; range: 280 to 4000 µg/kg) and DON (mean: 259 ± 5.2 µg/kg; range: 200-360 µg/kg). About 65.6% of these had AFs above EU limit of 4 µg/kg, but FUM and DON concentrations were all within the tolerable limits of 4,000 µg/kg and 1,750 µg/kg respectively. AFs & FUM, AFs & DON, and AFs, FUM & DON co-occurred in 9.8%, 23% and 3.3% of the samples respectively [141].

Comparably, Mutiga et al. [72] evaluated AFs and FUM in maize from Western Kenya. The study covered 3 agroecological zones, taking samples of milled maize from 985 patrons of 26 hammer mills. AFs were detected in 49% of the samples, with 15% of these being above 10 µg/kg. Estimated 65% of the samples from a drought-prone area were above acceptable limits. In Bungoma County, the authors assessed both AFs and FUM in four maize varieties at harvest and after 2 and 4 months of storage. For this, storage shed grain and milled samples were solicited. Mean AFs were identical for storage sheds and mills at 2.3 µg/kg. About 41% of the samples from mills had detectable AFs, 4% of which were above 10 µg/kg, while 87% had detectable FUM, with 50% above 1,000 µg/kg limit permitted in Kenya. Mean contamination levels did not vary during storage. As such, maize varieties reportedly differed in FUM contamination, with the most popular varieties spotted to be vulnerable to both AFs, FUM and weevils. It was concluded that thorough mycotoxin surveillance is vital for all parts of Kenya, irrespective of past history of mycotoxin poisoning [72].

Samples of 74 animal feeds and 120 milk samples were simultaneously collected from individual cows and actors in the informal sub-value chains of rural and peri-urban dairy systems in Nakuru county, Kenya [142]. AFB₁ was detected in 56 % (41/74) of the feeds in levels above EU limit of 5 µg/kg (range: BDL to 147.86 µg/kg) while DON was identified in 63% (27/43) of the feeds (range: BDL to 179.89 µg/kg). In the peri-urban dairy system, 48.5% (33/68) of the milk samples were contaminated with AFM₁ in levels exceeding EU threshold of 0.05 µg/kg (range: 0.017 to 0.083 µg/kg). Surprisingly, all milk samples from rural dairy system had AFM₁ in levels below EU limit of 0.05 µg/kg (range: BDL to 0.041 µg/kg). Linear regression depicted that there was a correlation between abiotic factors viz; pH, water activity and moisture content of feeds with AFB₁ and DON contamination.

Herbal preparations were sampled from Eldoret (14 Liquid, 2 Oil, and 34 Powder) and Mombasa (12 Liquid, 1 Capsule, 3 Oil, 6 Tablets, and 28 Powder) towns and analyzed for total AFs and FUMs [91]. Reported 32% of herbal products from Eldoret had AF levels less than 0.25 µg/kg, while 34% had AFs between 0.38 to 24 µg/kg. FUM occurred in very low concentrations in more than half of the samples. Samples drawn from Mombasa had AFs in levels lower than those from Eldoret, but the number of AF contaminated samples was higher. About 32% of the samples had < 0.25 µg/kg with 14 µg/kg being the highest. About 80% had < 0.25 µg/kg, and the highest was > 20 µg/kg. Six out of 14 (42.9%) Liquid herbal samples from Eldoret were contaminated with AFs and 3 of the 6 were also contaminated with FUMs. All the 12 (100%) Liquid samples taken from Mombasa were contaminated with both AFs and FUMs. A total of 27 out of 34 (79.4%) Powders from Eldoret were contaminated, 23 with both mycotoxins and 4 with AFs only, while all the tablets (15 samples) and powders (19 samples) from Mombasa were contaminated with both mycotoxins; however, all capsules were free of mycotoxin contamination. All Oily herbal samples (n = 3) from Mombasa were contaminated with both AFs and FUM, while only 1 oil sample from Eldoret was contaminated with FUM [91].

Into the bargain, a survey covering 116 push-pull and 139 non-push-pull cropping systems was conducted to determine the socio-economic and agronomic factors that influence farmers' knowledge on incidence and contamination of maize by ear rots and associated mycotoxins in Siaya, Kakamega, Kisumu, Migori and Vihiga counties of Western Kenya [143]. Data from smallholder farmers (23-80 years, 50% being female) were collected using questionnaires and

10-20 maize cobs, depending on the size of cob, were collected from the standing crop in the field of each interviewed farmer and analyzed for AFs and FUMs. The authors reported that few farmers had knowledge of AFs and ear rots in maize. Overall, less than 20% of maize samples had AFs co-occurring with FUM, but more samples were contaminated with FUMs (range: 145.3-50,769.2 µg/kg) than AFs (range: BDL-242.3 µg/kg) with maize containing the mycotoxins in levels above permissible limits (10 µg/kg for AFs and 1,000 µg/kg for FUMs) being lower in samples from push-pull cropping system. Age of farmer and county of residence were significantly and positively associated with knowledge of AFs on the one hand. On the other hand, cropping system, county of residence, and level of education were positively associated with knowledge of maize ear rots. In addition, a strong correlation between knowledge of maize ear rots and knowledge of AFs was witnessed. The concentration of the mycotoxins were significantly, and positively associated with the use of diammonium phosphate fertilizer at planting. AF levels were also positively associated with stem borer pest damage, though agronomic practices were not ideally different between push-pull and non-push-pull farmers [143].

2.4 Geographical Distribution of Aflatoxins in Kenya

Kenya was one of the hotspots of AFs recorded [15, 144] with countries such as Uganda, Brazil, Senegal, Mozambique, Swaziland, Nigeria, China, Thailand and Philippines [26, 145]. Kenya is partitioned into about 7 agroecological zones: Humid, Sub-humid, Transitional, Temperate, Semi-arid, Arid and Per arid [146]. AFs tend to be detected in samples from all the different zones [105, 110, 130]. This can be attributed to the similarity in the agronomic, pre-, peri- and post-harvest handling practices and the inter-regional marketing of foods [61, 92, 111, 147]. However, the Eastern part of the country is more aflatoxin-prone, possesses the most toxigenic *Aspergillus* spp, and have been the epicenter of aflatoxicoses recorded in Kenya [46]. Eastern Kenya experiences hotter and drier climatic conditions in comparison to Western Kenya. For this reason, it received characterization as semi-humid to semi-arid while Western Kenya is classified as sub-humid to semi-humid agroecological zone [110]. Environmental conditions have been demonstrated to influence the ability of *Aspergillus* fungi to infect, colonize and survive on crops as well as produce mycotoxins. Further, fluctuations in these conditions also affect the quantities as well as community compositions of aflatoxin-producing fungi [148]. Prevalence of AFs in Eastern Kenya therefore is in congruence with a previous emphasis that mycotoxin infectivity is always multifactorial, but climate is the most important [149].

3. Capacity for Detection and Quantification

Detection and quantification of AFs is key to their mitigation because their distribution in samples is often skewed [150]. The first step for accurate detection and quantification of AFs is sampling i.e. sampling/sub-sampling is the largest source of error in AFs analysis, responsible for over 90% of the error in testing the variance in AF concentrations between the measured sub-sample and the whole, compared to variance from the analysis [151]. For this reason, a representative sample ought to be drawn from the sample lot. For the over 50 KEBS listed laboratories for monitoring mycotoxins in foods, Gafta methods (No. 130, 24:1) and EAS 79 are used as the sampling protocols. However, some clients do the sampling themselves, in

which case the testing laboratories do not question the actual reason for sampling, or where and how the samples were taken. In addition, data from such analyses are always confidential, which does not enhance evidence-based decision making by policy makers [97].

Another emerging challenge in analyses of food toxins in Africa, Asia, America and Europe is “masked mycotoxins” as they are not often identified and detected by the usual analytical techniques [152]. Masked (matrix-associated) mycotoxins are those that are biosynthesized by the toxigenic fungi and later undergoes biomodification by plant enzymes during the infection stages. They may be housed in the vacuoles in the soluble form or bound to macromolecules, and thus remain undetectable [153]. Unfortunately, these modified toxins can hydrolyze and revert back into their toxic forms during processing or digestion [154-156]. A way to circumvent around this analytical problem has been to hydrolyze the modified forms (using enzymes, alkaline, or acidic pre-treatments) [157-159] into their free forms which can then be detected [157, 160]. For this reason, there is paucity of data on masked AFs as usually detection and quantification is done for free AFs in matrices.

The methods for detection of AFs used by studies in Kenya are outlined in **Table 7**. On the whole, AF research in Kenya used laboratory-based enzyme linked immunosorbent assays (ELISA), high performance liquid chromatography (HPLC), thin layer chromatography (TLC), fluorimetry, liquid chromatography tandem mass spectrometry (LC-MS/MS), tandem quadrupole mass spectrometry (TQMS) and Ultra-high pressure liquid chromatography (UHPLC). Lateral flow immunochromatography (LFI) has also been used. There has been a shift in instrumentation for AF analysis, as evidenced by advancement from non-differential TLC in 1982 to relatively fast and differential UHPLC, hyphenated with Triple Quadrupole mass spectrometry (UHPLC-TTQS) in 2017-2020. Overall, the most employed method has been ELISA, which itself has undergone several advancements in the past few years. This could be because it is practically inexpensive, easy to use, is highly sensitive for routine analysis of food products, demands minimum sample clean up and poses no inherent health hazards as it uses enzyme labels. In addition, concurrent analysis of several samples on a 96-well assay platform are possible, thus it has a high sample throughput with low sample volume requirement which offer obvious advantages [28]. In addition, ELISA has lower detection limits than most instrumental techniques which are used for AF determination [28].

However, the drawbacks of the foregoing standard methods are that they are unsuitable for rapid and real-time applications in food and feed sample analysis as they are relatively tedious and require technical know-how to operate. Rapid and robust methods such as polymerase chain reaction (PCR) and non-destructive methods based on fluorescence/near-infrared spectroscopy (FS/NIRS) and hyperspectral imaging (HSI) have emerged for quick and easy detection of AFs [161]. Some studies in Kenya [41-43, 46, 47, 110, 162] have utilized PCR in their analyses. It is of interest to note that at industrial level, agroprocessing companies monitor total AFs in cereals using single-step lateral flow immunoassays utilizing Reveal Q+ test strips that are developed and read on AccuSan Gold readers [163]. The Bright Greenish-Yellow Fluorescence (BGYF) or the Black Light test, which can aptly identify commodities presumed to be contaminated with AFs has been reported in Kenya [54]. This test is relatively cheap and simple especially for detecting AFs in maize where kernels are viewed under an ultraviolet lamp at 365 nm for characteristic bright greenish yellow fluorescence which indicates a

possible presence of aflatoxigenic fungi or the mycotoxin itself [164]. Regulatory bodies in Kenya could develop the capacity to perform this simple detection test for surveillance surveys.

4. Exposure Assessment

4.1. Exposure to Aflatoxins in Kenya

Exposure to AFs occur via periodic ingestion of contaminated plant products or animal products such as meat, milk and blood from animals initially fed on AF-contaminated feeds [20]. Farmers and their workers may also inhale dust generated during processing of contaminated crops and feeds or the toxins may permeate through their skin [165, 166]. Exposure to AFs such as AFB₁ may also be through their endogenous production [32]. In point of fact, AFs are known to cross the placenta, so that exposure to them may start *in utero* and continues in the post-natal period through breastfeeding [92, 167-169].

It is now established that detection and quantification of AFs in foods are not always adequately reflective of the exposure levels as the quantities in foods are not always the same as that ingested. For this reason, epidemiological biomarkers are often used to assess exposure. Biomarkers are more precise for assessing the degree of exposure to AFs, as they are non-subjective and estimate the internal and biologically effective doses. Popular AF biomarkers include the AF-N⁷-guanine adducts excreted in urine (reflect the previous day's exposure), AFB₁ (primarily in breast milk, and reflects exposure over the previous 24 hours) and the aflatoxin-albumin adduct (AF-alb) in plasma or serum with half-life of about 2 months which allows assessment of chronic and routine exposure to AFs [170]. Albumin, the only serum protein that binds AFB₁, forms a high level of adducts [171]. AF-albumin adducts from human blood and urine avail a measure of the biologically effective dose of ingested AFB₁. Both AFB₁ and AFG₁ can be bound by albumin, and are metabolized to AF-8, 9-epoxide [172]. The AF-alb adduct levels are considered as AFB₁ amount ingested as AFG₁ are less prevalent in foods [173]. Thus, the AF-alb biomarker is the most commonly employed as it can be easily detected by ELISA with results in pg AF-alb/mg albumin or pg AF-Lys equivalent/mg alb (**Table 7**) [174]. Quantification of AFB₁-Lys in proteolytic digests of serum with HPLC-FS and LC-MS/MS are also possible [175, 176].

A biopsy material was first utilized in 1967 to illustrate that the Kamba people of Kenya had a frequency of liver cancer that was approximately twice that of the Kikuyu ethnic community [177]. This is partly supported by the fact that subsequent aflatoxicoses were witnessed in Eastern Kenya where the Kamba are the main inhabitants [71, 178]. A dietary AF-liver cancer study in Murang'a district of Kenya reassessed the correlation between AF and the disease incidence rates based on a total of 7 years of cancer registration [82]. The results of the study was however interpreted in combination with a study later done in Swaziland. With consideration of males and females separately, the pooled results of the studies hinted that there

Table 7: Analytical methods used by aflatoxin investigations in Kenya.

Method of analysis	Sample (s)/matrices	Mycotoxin (s) analyzed	Year ^a	Author(s)
ELISA	Maize grain	Total AFs	2020	[179]
Fluorimetry, PCR	Soil	Total AFs	2020	[46]
UHPLC	Maize grain (fresh), maize flour	AFB ₁ , AFG ₁ , AFB ₂ , AFG ₂	2020	[77]
UPLC, PCR	Maize grain	AFB ₁ , AFG ₁ , AFB ₂ , AFG ₂	2019	[42]
Quantitative PCR (qPCR), TLC, HPLC	Maize tissues/grain	<i>A. flavus</i> biomass, AFB ₁ , AFG ₁ , AFB ₂ , AFG ₂	2019	[43]
ELISA	Bovine milk	AFM ₁	2019	[129]
ELISA	Maize	AFB ₁	2019	[94]
ELISA	Maize	Total AFs, FUM	2019	[143]
ELISA	Bovine milk	AFM ₁	2019	[135]
PCR	Soils	<i>A. flavus</i> genotyping	2018	[47]
LC-MS/MS, UHPLC-TTQS, PCR	Maize samples	AFB ₁ , AFG ₁ , AFB ₂ , AFG ₂ , <i>Aspergillus</i> spp genotyping	2018	[48]
PCR, HPLC	<i>Kimere</i> (a fermented milk product)	AFB ₁	2018	[180]
LFI	Maize grain, human sera (children)	Total AFs, AFB ₁ (Lysine adduct)	2018	[181]
ELISA	Raw, pasteurized & UHT milk, yoghurt, <i>Lala</i>	AFM ₁	2018	[90]
ELISA, TLC, HPLC	Maize grain	Total AFs, AFB ₁	2018	[63]
ELISA, PCR	Maize kernels	Total AFs	2018	[110]
ELISA, LC-HRMS/MS	Fish feeds	Total AFs	2018	[73]
ELISA, HPLC	Urine, breast milk, maize flour, sorghum, millet	AFM ₁	2017	[169]
LFI	Maize grain and maize flour	Total AFs	2017	[95]
ELISA	Herbal products	Total AFs, FUMs	2017	[91]
HPLC, UPLC-MS/MS, LC-MS/MS	Human urine, human blood	AFM ₁ , AFB ₁ (Lysine) adducts	2017	[182]
ELISA	Dairy cattle feeds, bovine milk	AFB ₁ , AFM ₁	2016	[58]
ELISA	Bovine milk	AFM ₁	2016	[131]
ELISA	Animal feeds, and bovine milk	AFB ₁ , DON and AFM ₁	2016	[142]
ELISA, HPLC, LC/MS	Maize grain, urine	AFB ₁ , AFM ₁	2016	[183]
ELISA	Dairy cattle concentrates, bovine milk	AFB ₁ , AFM ₁	2016	[133]
ELISA	Bovine milk (raw and processed), dairy products	AFM ₁	2016	[132]
ELISA	Maize, sorghum, & milk	Total AFs & AFM ₁	2016	[66]

Method of analysis	Sample (s)/matrices	Mycotoxin (s) analyzed	Year ^a	Author(s)
HPLC	Peanuts	Total AFs	2016	[121]
ELISA	Maize grain	Total AFs	2016	[184]
ELISA	Cassava (chips and flour)	Total AFs	2015	[126]
ELISA	<i>Omena</i> , maize, sorghum, rice, peanuts, cassava	AFB ₁ , AFM ₁	2015	[80]
ELISA	Maize (grain and flour)	Total AFs, FUM	2015	[72]
ELISA	Maize, sorghum, millet	Total AFs	2015	[76]
HPLC	Human sera (women)	AFB ₁ (Lysine adduct)	2015	[60]
ELISA, qPCR	Human sera (children)	AFB ₁ (Albumin adduct)	2015	[162]
TLC, HPLC	Fresh and sun-dried fish (<i>Rastrineobola argentea</i>)	Total AFs	2015	[85]
ELISA	Cattle feeds, rice, maize, peanuts	Total AFs	2014	[86]
ELISA	Maize grain	Total AFs, FUM	2014	[75]
TLC, HPLC	Maize grains, <i>githeri</i> , <i>muthokoi</i>	Total AFs	2014	[109]
ELISA	Bovine milk	AFM ₁	2014	[130]
ELISA, BGYF	Maize grain	Total AFs, FUM	2014	[54]
LFI	<i>Busaa</i> (a local brew)	Total AFs, FUM, DON	2014	[141]
ELISA	Peanuts (raw and roasted)	Total AFs	2013	[115]
ELISA	Peanuts	Total AFs	2013	[64]
ELISA	Peanuts and peanut products	Total AFs	2013	[62]
ELISA	Peanuts (raw and roasted), peanut butter	Total AFs	2013	[53]
TQMS	Human sera	AFB ₁ (Lysine adduct)	2013	[185]
ELISA	Peanuts	Total AFs	2012	[44]
LC-MS/MS, PCR	Maize kernels	AFB ₁ , AFG ₁ , AFB ₂ , AFG ₂	2012	[41]
TLC	Maize (grains, flour), milled maize-cereal products, dairy cattle feed, oil seed cake	Total AFs	2012	[78]
ELISA	Human plasma (children)	AFB ₁ (Albumin adduct)	2012	[186]
ELISA	Maize (grains, flour, semi-processed), soil, mill dust	Total AFs	2012	[107]
Fluorimetry	Maize grain	Total AFs	2011	[70]
LC-MS, HPLC	Commodities, feeds and feed ingredients	Total AFs, FUM, ZEA, Trichothecenes (A & B), Ochratoxin A	2011	[140]
ELISA	Ground maize, soil	Total AFs	2010	[55]
ELISA	Milk, animal feeds	AFM ₁ , AFB ₁	2009	[89]

Method of analysis	Sample (s)/matrices	Mycotoxin (s) analyzed	Year ^a	Author(s)
ELISA	Maize, soils, mill dust	AFB ₁	2009	[40]
ELISA	Peanuts	Total AFs	2009	[61]
ELISA	Maize grain	AFB ₁ , FB ₁	2009	[139]
HPLC/ Fluorimetry	Maize grain	AFB ₁	2007	[39]
ELISA	Peanuts	Total AFs	2007	[65]
HPLC	Maize kernels, maize flour, <i>muthokoi</i>	Total AFs	2005	[71]
Fluorimetry	Maize grain	Total AFs	2005	[69]
Fluorimetry	Maize grain and maize products	Total AFs	2005	[103]
ELISA	Pilsner and Tusker beers	AFB ₁ , FB ₁ , DON, ZEA	2004	[138]
TLC	Peanuts	Total AFs	2004	[84]
TLC	Weaning foods	Total AFs	2004	[118]
TLC, HPLC	Malted millet, maize flour	AFB ₁ , AFB ₂	2000	[81]
ELISA, HPLC-FS	Human sera	AFB ₁ (Lysine adduct)	1990	[187]
HPLC	Breast milk, human sera, neonatal cord blood, blood (pregnant women)	AFB ₁ , AFB ₂ , AFG ₁ , AFG ₂ , AFM ₁ , AFM ₂ , aflatoxicol	1989	[188]
HPLC	Human urine	AFB ₁ (Guanine adduct)	1987	[189]
TLC	Local beer, food (maize, millet, sorghum, pigeon peas and yam components)	Total AFs	1973	[82]

848 Years cited represent the years the data were published, with most data collected in over 2 months prior to publication. *Muthokoi* are maize kernels with the outer hull removed. *Lala* also called
 849 *Maziwa Lala* or *Mala* is a locally fermented milk product. *Busaa* is a socio-cultural maize-based traditional brew mostly consumed during events such as male circumcisions, weddings and
 850 funerals, made from raw maize flour and semi-ground finger millet malt [141]. LC-HRMS/MS: Liquid Chromatography High Resolution Mass Spectrometry. LFI: Hoffmann et al. [181] used
 851 Romer AgraStrip rapid test; Kirui et al. [141] used Envirologix Quick Tox kits.

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was a high degree of positive correlation between the calculated ingestion levels of AFs (X) and the adult incidences of hepatocellular carcinoma (Y) for the two studied populations and for both males and females. With the assumption of a wet intake diet of 2 kg/day and a mean body weight of 70 kg, the relationship for adult females was: $Y = 4.14 \log_{10} X - 0.80$. With a further assumption of a daily intake of native beer of 2 liters/day, the regression equation for adult males was $Y = 21.96 \log_{10} X - 11.17$. The regression data were reported to corroborate those reported by previous researchers [82].

To validate the assertion, another team [189] evaluated if there were any synergistic effect of hepatitis B virus and AFB₁ on the incidences of hepatocellular cancer. The study encompassed various parts of Kenya with different liver cancer incidences so as to establish the rate of exposure to AFs and the prevalence of hepatitis infections. It turned out that of all the tested participants, 12.6% were positive for AF exposure as shown by urinary excretion of AF-N⁷-guanine adduct and the highest exposure to the toxins was in the Western Highlands and Central Province. The incidence of hepatitis infection nationwide as measured by the presence of the surface antigens was 10.6% with a marked regional variation. Execution of multiplicative and additive regression analysis suggested that the two were not a synergetic combination in the etiology of liver cancer, though a moderate degree of correlation between AF exposure and liver cancer was observed when the study was limited to certain ethnic groups [189].

Further, Maxwell et al. [188] undertook a study in Kenya, Sudan, Ghana and Nigeria to evaluate the extent of AF exposure by breast-fed infants, and to investigate the possibility that AFs cross human placental membrane. In this study, breast milk, cord blood and maternal blood were analyzed for AFs which were detected in 28% of 191 Kenyan, 37% of 99 Sudanese, and 34% of 510 Ghanaian breast milk samples (**Table 8**). Blood drawn from 101 babies in Kenya, 282 babies in Ghana, and 78 babies in Nigeria had AFs in 37%, 31% and 12% of the samples respectively. In Kenya, the rate of detection was higher in the wet (52%) than dry (23%) season. Maternal blood sampled at delivery in 83 Kenyan cases and 77 Nigerian cases recorded AFs in both maternal and cord blood specimens in 14 Kenyan and 7 Nigerian instances. These confirmed that infantile exposure to AFs occurs, and demonstrated the ability of AFs to cross the human placental membrane [188].

Table 8: AF content of breast milk and cord blood from Kenya, Sudan, Ghana and Nigeria

Sample	Country	Number of samples	Number of AF positive samples	Positive samples (%)
Breast milk	Kenya	191	53	28
	Sudan	99	37	37
	Ghana	510	163	32
	Kenya	101	37	37
Cord blood	Nigeria	78	9	12
	Ghana	282	86	30.5

Adapted from [188]. AFM₁ was detected in 121 milk samples (range: 5-1379 ng/L), AFM₂ in 103 (range: 3-6368 ng/L), AFB₁ in 41 (range: 150-55,792 ng/L), AFB₂ in 10 (range: 49-623 ng/L), AFG₁ in 4 (range: 1890-5180 ng/L), AFG₂ in 3 (range: 10-87 ng/L) and aflatoxicol in 6 (range: 14-270 ng/L). In cord blood, AFM₁ (range: 25-8942 ng/L) and AFM₂ (range: 15-732 ng/L) were detected frequently in 63 and 47 samples; AFB₁ (range: 185-43822 ng/L) and AFB₂ (range: 10-925 ng/L) were detected in 20 and 19 samples. AFG₁ was detected 4 times (range: 611-2086 ng/L), AFG₂ once (37 ng/L) and aflatoxicol thrice (177, 214, & 280 ng/L).

Differently, a survey which recruited adults from Kenya, Thailand, The Gambia and France was used to validate the measurement of AF-albumin adducts by three methods [187]. Levels of 7 to 338 pg AF/mg alb were observed in the first three countries while no adducts were detected in samples from France. Another cross-sectional serosurvey in Kenya confirmed

regional influence on AF exposure patterns [185]. Randomly selected 600 serum specimens stratified by province from a 2007 Kenya AIDS Indicator Survey were analyzed for AFs. About 78% of the sampled group had exposure to AFs and this varied by province. The highest were in Eastern (median = 7.87 pg/mg alb) and Coast (median = 3.70 pg/mg alb) provinces, while Nyanza (median = < limit of detection) and Rift Valley (median = 0.70 pg/mg alb) provinces recorded the lowest exposures. According to the authors, age group, sex, marital status, religion and socioeconomic characteristics did not influence exposure.

In another study, random samples of weaning flours were obtained from 242 households with 3 to 36-month-old children (43.6% males and 53.4% females) in Kisumu district, Kenya and analyzed for AFs [118]. The types of weaning foods, handling and storage of the foods were captured. The nutritional status of the children were also determined along with heights and lengths. About 29% (70/242) of the samples were positive for total AFs (range: 2-82 µg/kg). Malnutrition was 34% for stunting, 30% for underweight and 6% for wasting. About 53.8% of the wasted children were being fed on AF-contaminated weaner flour vis-à-vis 27.7% of the normal children. The contaminated flours (n = 70) were being stored in plastic containers (63%), polyethylene bags (20.4%), metal buckets (3.7%), manila sacks (1.9%), earthen pots (1.9%) and reed baskets (7.4%) for 1 day to 2-3 weeks. These all had effects on aflatoxigenic contamination as those with AFs had higher mean moisture content (13.6%) than those devoid of AFs (12.5%). *Aspergillus* spp (including *A. flavus* and *A. parasiticus*), *Fusarium*, *Mucor*, *Rhizopus nigricans*, *Trichoderma viride*, and *Candida* spp were isolated from the flour samples [118].

Agreeably, Leroy et al. [60] collected socio-economic data to quantify the extent to which socio-economic characteristics could explain the differences in serum AFB₁-Lysine adduct levels in 100 women from Eastern province of Kenya. The correlation between serum AFB₁ level and number of households, farm, and individual characteristics were assessed for 884 mothers (pregnant or with a child under 24 months). AF was detected in all women with a median level of 7.47 pg/mg alb. Higher exposure levels were correlated with poverty: predicted serum AF levels in women living in the worst socio-economic conditions were 4.7-7.1 times higher than those with the best socio-economic status.

Further, samples of *Rastrienobola argentea* (n = 50), polished rice (n = 31), peanuts (n = 22), cassava (n = 37), maize (n = 41), and sorghum (n = 28) were collected from Kibuye wholesale, Kibuye open air, Ahero, Oile and Mamboleo markets in Kisumu County and analyzed for AFs [80]. Processed bovine milk samples (n = 50) were collected from supermarkets along with raw bovine milk samples (n = 30) from 3 market milk bazaars in the same markets. Analytical results indicated that AFs in the solid foods ranged from 0 to 34.5 µg/kg AFB₁, 0.012 to 0.127 µg/kg AFM₁ in processed milk and 0.0002 to 0.013 µg/kg AFM₁ in raw milk. Only cassava among the scrutinized food items had detectable AFs below the regulatory limit of 10 µg/kg AFB₁ by then. Daily AF consumption ranged from 4.43 ng/kg bw/day in a combination of maize flour and milk to 110.4 ng/kg bw/day in a combination of sorghum and raw milk for 6-month old children (average weight: 7.9 kg) with a daily consumption of 60 g of mixed cereal flour and 500 ml of milk per day. These results emphasized that weaning children in Kisumu county are chronically exposed to high AF levels for the fact that the analyzed food items are common ingredients of weaning foods in the area [80]. In addition, the calculated AF consumption of 0.6 ng/kg bw/day for a child at 6 months

weighing 7.9 kg was higher than that indicated by the Codex Alimentarius Committee (0.1 ng/person/day) AFM₁ through milk for the Africa region. Weighted mean concentration of 0.05 µg AFM₁ in milk and a consumption of 0.25 ng/kg bw/day have been associated with a prevalence of between 3.2 to 20 cancer cases/year/10⁶ [190]. The exposure was much higher than estimated because most children in Kenya are breastfed until at least the latter part of the second year and yet they begin to receive cereal-based gruel before the age of 3 months [191]. Further, the results of the study corroborated a previous report which estimated that about 40% of foods from farmers in the Nyanza province had AF levels above the statutory limit of 10 µg/kg [65].

Another cross-sectional study was undertaken involving 204 low-income households randomly selected in two low-income areas (Korogocho and Dagoretti), Nairobi, Kenya [66]. Demographics, a 24-hour dietary recall and anthropometric measurements were conducted in children aged 1–3 years. Height-for-age Z-scores (HAZ), weight-for-age Z-scores (WAZ) and weight-for-height Z-scores (WHZ) were calculated for each child using WHO growth standards reference data. Maize (n = 99 & 87), sorghum (n = 53 & 36) and milk (n = 76 & 52) samples from the households or retailers from Korogocho and Dagoretti respectively were analyzed for total AFs and AFM₁. As a whole, 98% of food samples collected were AF positive (maize: mean-6.7, range: 0.0–88.83; sorghum: mean-8.07, range: 0.1–194.41, and milk: mean-0.132, range: 0.007–2.56 for samples from Korogocho; maize: mean-2.97, range: 0.0–20.0; sorghum: mean-2.59, range: 0.2–14.47, and milk: mean-0.093, range: 0.002–0.64 for samples from Dagoretti). About 41% of the children had stunted growth; boys were more stunted than girls (*p* = 0.057) and Korogocho had more stunted children than Dagoretti (*p* = 0.041). The average AF exposure was 21.3 ng/kg bw/day. Exposure to AFM₁, location and sex were significantly associated with HAZ, with boys and children from Korogocho having lower HAZ, and AFM₁ was negatively associated with HAZ (*p* = 0.047), suggesting that AFM₁ was associated with stunting. No correlation was statistically found between total AFs and HAZ, WAZ and WHZ. The authors reiterated that there was a high prevalence of malnutrition (stunting) in the studied low-income urban sites, and this was most pronounced in the high-density area. It was stressed that the association between AFM₁ and growth impairment warranted further investigations [66].

Kang'ethe et al. [169] reported that with a maize consumption of 0.1 to 0.25 kg/person/day in Nandi and Makueni counties, an AF exposure rate of 0.011 and 0.49 µg/kg bw/day respectively were recorded in children younger than 5 years. Exposure to AFM₁ through milk consumption in this study were 4×10^{-4} and 1×10^{-4} µg/kg bw/day respectively. Breast milk nursed children on the other hand had exposure of 6×10^{-3} and 1×10^{-6} µg/kg bw/day in Makueni and Nandi respectively. Children younger than 30 months in Makueni had 1.4 times higher levels of AFM₁ in urine than those of the same age in Nandi. The stunting and severe stunting rates in Makueni and Nandi were 28.7%, 18.5% and 30.7%, 16.5% respectively.

In a recent study [181] which enrolled 1230 unborn children, 881 (72%) were included in LAZ and 798 (65%) in the serum AFB₁ analysis. A cluster randomised controlled design was used (28 intervention and 28 control clusters). The intervention arm received a swapping (contaminated maize was replaced with safe maize) and a stockist intervention (households were encouraged to purchase from a stockist supplied with clean maize). Women in the fifth to final month of pregnancy were invited to enrol in the study. Outcomes were child LAZ, the

prevalence of stunting and child serum AFB₁-Lysine adduct level 24 (end-line, primary outcomes) and 11 to 19 months (midline, secondary outcomes) after trial commencement, respectively. The intervention was reported to considerably reduce end-line In serum AFB₁-Lysine adduct levels (intervention effect was 0.273, 95% CI 0.547 to 0.001; one-sided $p = 0.025$) but had no effect on end-line LAZ or stunting (mean LAZ at end-line was -1.64). At midline, the intervention increased LAZ by 0.16 (95% CI -0.009 to 0.33; one-sided $p = 0.032$) and reduced stunting by 7% points (95% CI -0.125 to -0.007; one-sided $p = 0.015$) but had no effect on serum AFB₁ levels [181]. It was inferred that the midline analysis suggested that AFs may affect linear growth at younger ages.

An overall average estimation of exposure rates based on annual consumption, as is appropriate for cancer risk because of the cumulative nature of this response, indicate that AF exposure was 3.5 to 14.8 ng/kg/day in Kenya for about 67% of the population [92, 192]. No study in Kenya has examined the relationship between AFM₁ in breast milk samples and growth impairment in infants.

4.2 Co-Exposure to Aflatoxins with Other Mycotoxins

Aflatoxin poisoning could be compounded by the occurrence of AFs in combination with other mycotoxins such as FUM, trichothecenes, ochratoxins, ZEA and DON [16, 193, 194]. This is supported by the occurrence of mycotoxin producing fungi simultaneously in the same batch of food/matrix and the faculty of some toxigenic fungi to produce more than one mycotoxin in a given matrix. For example, *Fusarium* (*F. verticillioides*, *F. proliferatum* and *F. oxysporum*) [54] and *Penicillium* spp were reported with *Aspergillus* fungi in Kenya [41, 45, 53, 84, 115, 195, 196], sometimes in soils and mill dust around maize stores [40]. *Fusarium* spp are known for the production of FUMs [197].

The current review did not identify any reports evaluating co-exposure to AFs in combination with other mycotoxins and the potential adverse health outcomes. An explanation for this could be due to the underdevelopment of the valid biomarkers [198]. Mycotoxin-specific biomarkers for other mycotoxins (notably FUM in maize and DON in wheat) have been developed and validated only very recently [199, 200] and their utilization in human exposure and health risk assessments can be tagged as nascent. In the neighboring Tanzania, co-exposure to AFs with other mycotoxins utilizing individual biomarkers was recently investigated. Children (6-14 months old) were recruited at a maize harvest season and followed up twice at 6-month intervals. The children were reported to be chronically exposed to AFB₁, FB₁ and DON [201, 202]. Blood AF-alb [201] and urinary DON levels [202] steadily increased over the 12 months, which likely corresponded to increased food intake that is possible as the child grows. A linear trend was not apparent for urinary FB₁ as the mean level at 6 months was significantly lower than mean levels at recruitment and at 12 months [201]. It was deduced that the lower exposure levels observed 6 months post-harvest could be reflective of reduced maize stocks, resulting in lower maize consumption [201]. Though no significant correlation was appreciated between AF exposure and stunted child growth, increased FUM exposure was evidently associated with reduced length-for-age Z-scores [201].

In addition, co-exposure to mycotoxins *in utero* is also wanting, as observations elsewhere reported AF-alb in 36% of the blood samples with urinary AFM₁ and DON present in 47% and 68% of samples from pregnant women in their third trimester co-exposed to AFs and DON

[203]. About 41% of the pregnant women were concurrently exposed to both AFs and DON. Thus, assessment of co-exposure to AFs in Kenya with other mycotoxins is warranted.

5. Infantile Stunting Due to Aflatoxin Exposure in Kenya

The first 1, 000 days of life (from conception to about 36 months) is a critical window for healthy growth and development. Dietary intake of AFs during pregnancy plays a fundamental role in the child's future health status [188, 198]. In Sub-Saharan Africa, and particularly Kenya, malnutrition and child growth impairment are major public health burdens [80, 118, 181]. Intake of low, daily doses of AFs over long periods result in chronic aflatoxicosis expressed as impaired food conversion, stunting in children, immunosuppression, cancer and reduced life expectancy [6, 204-206]. The WHO defined stunting as a height-for-age Z-score (HAZ), of < -2 , being underweight as a weight-for-age Z-score (WAZ), of < -2 , and wasting as a weight-for-height Z-score (WHZ), of < -2 [207]. Stunting of infants in some aflatoxin-prone areas of Kenya are shown in **Table 9**.

Table 9: Aflatoxin levels in foods and stunting in some aflatoxin hotspots of Kenya

County	Stunting (%)	Highest reported AF levels in foods ($\mu\text{g/kg}$)	Author (s)
Urban Nairobi	22.7	4,593.93 (maize and maize products), total AFs	[78]
Nairobi (Korogocho and Dagoretti)	41.0	88.83 (maize), 194.41 (sorghum), total AFs	[66]
Kisumu	33.1	82.0 (cereal-based weaner foods), total AFs	[118]
Homa Bay	37.0	1,000 (peanuts), total AFs	[65]
Makueni	33.5	5,400 (maize), total AFs	[71]
Kitui	47.4	25,000 (maize), total AFs	[71]
Machakos	31.3	3,800 (maize), total AFs	[71]
Embu	23.7	21.0, total AFs	[106]
Kakamega (Malava)	34.2	17.0 (rotten maize), AFB ₁ ; FB ₁ $> 5,000 \mu\text{g/kg}$	[139]
Tongaren (Bungoma)	52.1	17.0 (rotten maize), AFB ₁ ; FB ₁ was $> 5,000 \mu\text{g/kg}$	[139]
Kisii South	35.3	3,442; total AFs	[106]

Adapted from Obade et al. [80].

It was advanced that AF exposure may disrupt insulin-like growth factors (IGF) pathway through liver toxicity. In a study in Kenya [162], AF-alb concentrations were inversely associated with IGF1 levels ($p = 0.039$) and IGF binding protein 3 levels ($p = 0.046$) in a sample of 199 school children from Yumbuni in the West and Matangini (Lower Manganalete) in the East. A path analysis showed that lower IGF1 levels explained about 16% of the effect of AFs on child height ($p = 0.052$). Both IGF1 and IGFBP3 were significantly associated with child height and weight ($p < 0.01$). Children in the highest tertile of AF-alb exposure ($> 198.5 \text{ pg/mg}$) were shorter than those in the lowest tertile ($< 74.5 \text{ pg/mg}$), after adjusting for confounders ($p = 0.043$). To further investigate this putative mechanistic pathway, human hepatocyte line 16 (HHL-16) cells were treated with AFB₁ at 0.5, 5.0 and 20.0 $\mu\text{g/mL}$ for 24-48 hours. IGF1 and IGFBP3 gene expression measured by quantitative PCR and protein in culture media showed a significant down-regulation of IGF genes and reduced IGF protein levels. The study concluded that AF-induced changes in IGF protein levels could contribute to growth impairment where AF exposure is high [162].

Aflatoxin-child growth impairment may also be due to the immunosuppressive effect of AFs that increases neonatal infection susceptibility, consequently impairing nutritional status

through appetite suppression and reduced nutrient absorption [208]. It is also argued that exposure to AFs may promote intestinal damage through protein synthesis inhibition, consequently leading to a reduction in the absorption of essential nutrients and subsequent impaired growth [209]. AFs has also been implicated in the aetiology of other liver diseases including jaundice, cirrhosis and hepatomegaly [186, 210, 211]. A study in Kenya by Gong et al. [186] reported that the prevalence of hepatomegaly, a firm form of liver enlargement, increased in children with higher AF exposure. This is in complete agreement with the knowledge that the liver is the key target organ for aflatoxin toxicity.

6. Aflatoxicoses in Kenya

Since the discovery of AFs, Kenya has been one of the countries with devastatingly severe human exposure to AFs [109, 212, 213]. Exposure to AFs occur primarily through ingestion of contaminated food. Ingestion, however, at very high concentrations (> 6000 mg) results in liver failure and death within 1–2 weeks of exposure (acute aflatoxicosis) [214]. Aflatoxicosis is typified by oedema, convulsions, vomiting, jaundice, abdominal pain, sudden liver failure and lastly death [215]. In practice, acute toxicities associated with exposure to elevated AF levels are not very common globally; cases occur and are concentrated in high-risk regions such as the Makueni County of Kenya [77] (**Table 10**). In humans, acute toxicity due to exposure to high dietary doses of AFs (2,000–6,000 $\mu\text{g/day}$) in contaminated maize was reported in Western India in 1974 with a case fatality rate of 10% [216, 217]. In Taiwan, 26 members of 3 families were victims of consumption of about 200 $\mu\text{g/kg}$ of AFs in mouldy rice. Three (3) of the victims died [218]. In the neighbouring Uganda, a 15-year old boy also succumbed to death following ingestion of cassava containing 1,700 $\mu\text{g/kg}$ of AFs, leaving behind a brother and a sister who survived very narrowly [219]. Recently, consumption of AF-contaminated maize triggered aflatoxicosis in humans with a case fatality rate of 50% in Tanzania [220].

In Kenya, aflatoxicosis was first witnessed in 1960 which recorded death of at least 16,000 ducklings [82]. In 1981, Kenya witnessed its first serious recorded outbreak of human aflatoxicosis [178]. It was found that after 7 days of consumption of maize grain containing 3.2–12 mg/kg of AFB_1 , symptoms of abdominal discomfort, anorexia, general malaise, and low-grade fever were exhibited in 20 cases of patients between 2.5 and 45 years of age. Hepatic failure developed in 12 of the 20 patients, all of whom eventually died 1–12 days following hospital admission. The most unprecedented episode of human aflatoxicosis in history was witnessed in Kenya in 2004 with 317 reported cases of which 125 were fatal [39]. This outbreak, which occurred in the Eastern Province, recorded a case fatality rate of 39% and out of the 308 patients for whom age data were available, 68 (22%) were < 5 years, 90 (29%) were 5–14 years, and 150 (49%) were > 15 years. Children younger than 14 years, representing 51% of the children population, were thus presumed to have had a greater predisposition to aflatoxicosis risk. Case fatality rate was significantly higher in Makueni district than in Kitui district [69, 70, 178, 221, 222].

Since 2004, outbreaks among subsistence farmers have recurred annually in Eastern Province and it is right to assert that the magnitude of exposure to AFs could be higher than reported due to dearth of robust monitoring systems [63, 109]. **Table 10** summarizes some of the fatal aflatoxicoses recorded in the history of Kenya since the discovery of AFs. It is worth noting

that several studies on AF poisoning in humans have shown that low-level chronic intake may be more devastating than one-time high-level intake (that leads to aflatoxicosis) as it is linked to the development of hepatocellular carcinoma [30, 82, 128, 223-232]. During the aflatoxicosis outbreak that occurred in 2010, the levels of AFB₁ sera reported in Kenya were among the highest ever recorded in the world [233]. As can be traced from **Table 10**, most areas that have been hit by aflatoxicosis in Kenya are in the Eastern and some Central part of the country.

7. Prevention and Control

7.1 International, Regional and Statutory Efforts

Appreciable efforts have been advanced towards AF control in Kenya through countrywide awareness creation [97, 234]. The regional mycotoxin facility at the Kenya Agricultural and Livestock Research Institute (KALRO) in Katumani offer various categories of training to extension officers drawn from public and private sectors.

After the fatal aflatoxicosis in which dogs fed on contaminated rations died between 1970-1980s, KEBS came up with a standard for dog feeds in 1985. Standards for maize grain, other grains and their products that have been in existence were also revised. For example, total AFs was initially at 20 µg/kg; this has been revised to 10 µg/kg, with 5 µg/kg as the threshold for AFB₁ in 2007 [235]. At least 25 standards aimed at regulating AFs have been drafted and are in full use, and encompasses key parameters such as moisture, mouldy grains, pest damage, filth, broken kernels/seeds, foreign matter and discoloured grains. Most of these standards have been harmonized with the East African Standards by the Eastern Africa Grain Council (EAGC) in collaboration with KEBS through the Eastern Africa Grain Institute with its Kenyan headquarters at Nairobi, Kenya [236]. Between 2015 and 2018, the duo have trained maize exporters, traders, farmer based organizations and warehouse handlers on understanding the integrated East African maize standard (EAS 2:2013), food standardization, comparison of East African standards with international standards, standard maize sampling methods, maize grading, mycotoxins and the available methods for mycotoxin analysis [236].

Following its launch wayback in 2006, EAGC has been among the lead in the fight against AFs in East Africa as a whole. It has advanced several interventions to reduce the incidence of AFs, including (1) harmonization of AF control measures and improving the regulatory environment, (2) running AF control training programs, (3) furnishing moisture analyzers and tarpaulins for safe drying and storage of grains, (4) sourcing for cheaper field-based AF testing kits and methods for measuring AFs, (5) conducting field surveys, regular analysis and random sampling during harvesting at farm level to assess the prevalence and extent of contamination, (6) working with East African Community to increase AF testing and surveillance in maize, (7) participation in the development of the Partnership for Aflatoxin Control in Africa (PACA) strategy 2013-2022 as well as advising on the East African Community AFs communication strategy [236]. In addition, AF surveillance and capacity has been enhanced through the PACA Curated Africa Aflatoxin Information Management System (Africa-AIMS) in seven member states: Kenya, Malawi, Nigeria, Senegal, Tanzania, The Gambia and Uganda.

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Table 10: Aflatoxicosis outbreaks reported in Kenya since the discovery of aflatoxins in 1960

Affected group	Case patients/Number affected	Area	Toxin source	Recorded effects	Year	Author (s)
Humans, dogs	None confirmed	Eastern Kenya (29 districts)	Suspected contaminated maize	Price spiral down, grain trade breakdown, unconfirmed dog deaths in Nairobi	2010	[237]
Humans	5	Kibwezi, Kajiado, Mutomo	Maize	3 hospitalized, 2 deaths	2008	[40]
Humans	4	Kibwezi, Makueni	Maize	2 deaths in Makindu town of Mukueni county	2007	[3]
Humans	20	Makueni, Kitui, Machakos, Mutomo	Contaminated maize	Acute poisoning, 10 deaths in Mutomo & 9 in Makueni	2006	[70, 103]
Humans	75	Machakos, Makueni, Kitui	Maize	Acute poisoning, 75 cases, 32 deaths	2005	[69, 70]
Humans	331	Eastern/Central Machakos, Kitui and Makueni areas	Contaminated maize	Acute poisoning, 125 deaths	2004	[238]
Humans	6	Thika	Mouldy maize	6 deaths	2003	[239]
Poultry/dogs	Large numbers	Coast	Contaminated feed	150 deaths	2002	[240]
Humans	3, 26	Meru North, Maua	Mouldy maize, contaminated maize	Severe liver damage, 16 deaths	2001	[39]
Humans	3	Meru North	Maize	Acute effects, 3 deaths	1998	[38]
Poultry	Large numbers	Kenya	Imported maize	Deaths	1984/1985	[241, 242]
Humans	12	Machakos	Poorly stored maize	Deaths	1981	[178]
Poultry/dogs	Large numbers	Nairobi, Mombasa, Eldoret	Poorly stored feed	Deaths	1977/1978	[242, 243]
Ducklings	16,000	Rift valley	Peanut ration	Deaths	1960	[82]

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Years are those in which the aflatoxicoses occurred rather than the years the data were published. Data are from [38, 92, 97]. A case report is also filed of a possible aflatoxicosis of a 17- year-old school boy [215].

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Kenya Agricultural and Livestock Research Organization in connection with International Institute of Tropical Agriculture (IITA) in 2018 developed a farmer-centered manual for management of AFs in maize and peanuts [234]. The manual gives a general overview of AFs (structures, health and economic effects), how to control AFs, drying, threshing, sorting and some of the farming practices that favours AF growth. It was particularly drafted to provide ample guidance on the best practices for limiting AF contamination of maize and peanuts, and to raise the value of these dietary staples.

Further, there are some projects running in the country to handle the plague of mycotoxins and these include the Aflacontrol project and Purchasing for Progress (P4P) Programme. The Aflacontrol project strives to minimize the ravage of AFs in maize and peanut value chains and is spearheaded by International Food Policy Research Institute (IFPRI). In addition, it seeks to increase the understanding of the economic and health impacts of AF contamination, identify and promote cost effective methods and technologies available to reduce contamination of foods and feeds. The project, funded by Bill and Melinda Gates Foundation, has partnership with International Maize and Wheat Improvement Center (CIMMYT), University of Pennsylvania (USA), United States Uniformed Health Services, Kenya Agricultural Research Institute (KARI) and Agricultural Cooperative Development Initiative (ACDI-VOCA). The project has been experimented in Mbere (Embu), Makueni, Homa Bay, Kisii and Rongo at the household level [97]. So far, it has released policy briefs, and held inception, and one-year national workshops to disseminate information on AFs. These are targeted at the Ministries of Agriculture and Public Health, who are the key players in mitigating AFs. On the other hand, the Purchasing for Progress Programme is led by World Food Programme that purchases maize from local farmers, usually ensuring strict adherence to AF limits in the grains. The grains are procured at fairer prices, encouraging the farmers to adhere to good pre-, peri-, and post-harvest practices [97]. Several partnerships are currently running in the country, some are with FAO and CDC to mitigate AFs in Kenya. These have been discussed in sufficient details a previous review by Mutegi et al. [38].

7.2 Scholarly Efforts

Earlier reports on the fate of AFs during processing of maize into *Muthokoi*-a traditional Kenyan food revealed that traditional maize preparation methods such as fermentation and dehulling in Eastern Kenya reduced AFs by up to 71% [83]. The findings of this study indicate that exposure to acute AF levels could be minimized during food processing and preparation. Generally, these processing techniques have been traditionally used for increasing the palatability of different food recipes but can should also be promoted as strategies capable of reducing AF contamination of grains [97].

Intermediate processes such as sorting and dehusking were shown to reduce AF in peanuts [244]. Soaking peanuts in water, *magadi*, sodium hypochlorite and ammonium persulphate significantly reduced AF levels by 27.7%, 18.4%, 18.3% and 1.6% respectively; while boiling in *magadi*, local ash, baking powder and water reduced AF levels by 43.8%, 41.8%, 28.9% and 11.7% respectively. Similarly, Kirui [184] while assessing the levels of residual AFs following various treatments using physicochemical and traditional cooking methods for maize and maize products reported that boiling maize reduced total AFs from 83.1 ± 0.3 to 7.0 ± 3.9 $\mu\text{g/kg}$, dry decortication reduced the level from 51.3 ± 15.3 to 9.6 ± 0.8 $\mu\text{g/kg}$, while boiling

with *magadi soda* (or maize wood ash) reduced the level from 59.5 ± 3.82 to 13.4 ± 0.42 $\mu\text{g/kg}$. Solar irradiation (for 18 hours) reduced the levels from 60.8 ± 1.8 to 13.7 ± 0.1 $\mu\text{g/kg}$ while ultraviolet irradiation (for 18 hours) reduced the levels from 81.7 ± 0.5 to 61.4 ± 4.5 $\mu\text{g/kg}$. The author reiterated that only dry decortication method and boiling with *magadi soda* followed by washing with water and boiling reduced AFs significantly to below the maximum advisory limit of 10 $\mu\text{g/kg}$.

In the same struggle, a probiotic yoghurt was formulated with AFB₁ binding *Streptococcus thermophilus*, *Lactobacillus rhamnosus* GR-1 and *Weissella cibaria* NN20 isolated from fermented *Kimere*, a dough food product made from millet [180, 183]. Forty primary school children, with maize being a regular part of their diet were randomly assigned to consume 200 ml yoghurt or control milk daily for 7 days, followed by a 7 day washout and another 7 day treatment. After both 7 day treatment periods, AF concentration in urine samples were significantly lower than baseline in the probiotic group ($p > 0.01$) but increased in the milk group. This suggested that locally produced probiotic yoghurt could reduce AF poisoning in Kenyan children, corroborating previous observations in our laboratory [245, 246]. Similarly, fermentation of milk into *Lala*-a traditional fermented drink and yogurt significantly reduced AFM₁ levels by 71.8% (in *Lala* after 15 hour room temperature incubation) and 73.6% in yogurt after incubation at 45 °C for 4 hours [135].

In another intervention survey, the use of a calcium montmorillonite clay (calcium silicate 100, popularized as ACCS100) in food reduced the bioavailability of AFs [182]. It was reported to be palatable, effective, and acceptable, though further evaluation in the AF-endemic parts of Eastern Kenya as well as its efficacy to ameliorate AFs to levels incapable of triggering poisoning yet remains to be established.

Another study [43] screened maize lines resistant to *A. flavus* infection, together with a biocontrol strategy. Two African maize lines (GAF4 and KDV1) were reported to have different fungal loads for the aflatoxigenic isolate (KSM014), fourteen days after infection, with no significant variation in *A. flavus* biomass between diseased and non-diseased maize tissues for GAF4. Meanwhile KDV1 had a significantly higher *A. flavus* biomass ($p < 0.05$) in infected shoots and roots compared to the control. The biocontrol strategy using an atoxigenic isolate (KSM012) against the toxigenic isolate (KSM014), showed aflatoxin production inhibition at the co-infection ratio, 50:50 for both maize lines (KDV1 > 99.7% and GAF 69.4%), as confirmed by bioanalytical techniques. It was indicated that the maize lines, which exhibited resistance to *A. flavus* with the appropriate biocontrol strategy could reduce aflatoxicosis outbreaks.

In a 2020 study [46], the possibility of using *Pseudomonas* and *Bacillus* bacterial spp was explored in soils from Eastern Kenya (Semi-arid) and Western Kenya (Sub-humid-Semi humid) [110]. *Pseudomonas* ($n = 7$) and *Bacillus* ($n = 5$) were identified in the two regions, though the latter recorded higher occurrence of *Bacillus*. Because these bacterial spp have been frequently associated with biological control of several plant pathogens including *Aspergillus* spp, a regression analysis was done to ascertain if there were any associations between the occurrence of *Aspergillus* spp and these bacterial spp in the studied regions. Weak relationships between occurrence of *A. flavus* and *Pseudomonas* spp in the Western region ($R^2 = 0.03693$) and the Eastern region ($R^2 = 0.06126$) as well as occurrence of *Bacillus* spp in the Western region ($R^2 = 0.196$) and in the Eastern region of Kenya ($R^2 = 0.03693$). The same observation

was made for the relationship between occurrence of *Trichoderma viride* in both Eastern ($R^2 = 0.03406$) and Western ($R^2 = 0.2266$) regions of the country. As a consequence, the authors deduced that the occurrence of the bacterial spp had little influence on occurrence of *A. flavus* in the two regions. To ascertain their assertion, an *in vitro* preliminary assay to determine the inhibitory potential of both *Pseudomonas* and *Bacillus* spp against *A. flavus* proliferation was done. Unfortunately, none of the bacterial strains from either spp had an inhibitory effect on *A. flavus* proliferation [46].

8. Suggested Management Strategies

8.1 Pre-Harvest Management

Staple food crop varieties that are disease, drought and pest tolerant or less susceptible to fungal growth should be bred and planted. This approach is so far the best for reduction of effects of mycotoxin-producing fungal species [247]. Valencia red (a peanut variety) was reported to be the least contaminated with AFs and had higher oil content than Uganda local, Homa Bay local and Local red [121]. Food oils and microorganisms are viable inhibitors of AF biosynthesis [6] through interference with the signal transduction regulatory networks involved in AF gene expression, blocking activities of AF biosynthetic cytosolic enzymes, downregulating fungal genes of the oxidative stress defence system that combats metabolic and environmental stressors [248]. The oils also inhibit fungal pathogenesis factors, disrupting mitochondrial respiration, a critical process that provides *acetyl*-CoA for AF biosynthesis and they are also associated with morphological alterations in the mycelium, such as vacuolation of cytoplasm and attenuation of cell wall [248]. Further, host and parasite macro- and micromolecular trafficking that suggests the possibility to circumvent the AF scourge through the utilization of cross species RNA interference have been attempted in maize and peanuts [249, 250]. This equips the plant with molecules that shuts down AF biosynthesis upon infection with aflatoxigenic fungi, thwarting AF accumulation. Particularly, UBI, COH, 26s, ATP, PPK, IMP, ABC and aflM were recommended as the suitable genes for RNAi silencing of *A. flavus in vivo* [249, 250]. This may, however, be impeded by the current policy on genetically modified organisms in the country.

Timely harvesting of grains with the husks upon maturity in dry conditions and early removal of any damaged maize kernels or cobs is a feasible AF reduction strategy [251]. Visual sorting, winnowing, washing, crushing and dehulling have been found to contribute up to a 40–80% reduction in AF levels in grains [151, 252]. Sorting is highly recommended for reducing AF content in foods, peculiarly in peanuts [251, 253–255] and cassava chips. Sorting can be done using clean water; the damaged seeds or grains are buoyant while good ones sink and can be cooked directly. Soaking and cooking in *magadi soda*, malting and roasting are other methods that have been used to reduce the levels of AFs [83, 252, 256, 257]. *Magadi soda* and wood ash is used by the Kalenjin of Rift Valley region, Nyanza and Western provinces to increase food palatability, offers convenience as it reduces cooking time but also reduces phytates and increases availability of niacin [258].

Protection of crops from pest attack is key in AF management. This can be done using ash while in storage for maize [259, 260] and plant essential oils with bioinsecticidal activity [261, 262]. Biocontrol strategies employing concoctions from plants have been investigated and

reported to inhibit *A. flavus* mycelial growth and proliferation. Essential oils of *Azadirachta indica* (neem) and *Morinda lucida* have been reported to retard aflatoxigenic *A. flavus* growth and its AF biosynthesis potential in inoculated maize grains [263]. Powder of *Aframomum danielli* (Zingiberaceae) can regulate moulds and insect infestation in maize and soybeans in storage for over a year under ambient conditions [264].

Competitive exclusion has been reported as a feasible AF control strategy. A shift of strain profile from toxigenic to atoxigenic is a viable biological control strategy. Kenya has approved a biocontrol product, a chemical that is introduced to the soil to help minimise the amounts of the toxic fungi that produces AFs. Studies have shown that the product (known as Aflasafe KE01) reduces AF contamination and helps improve the quality of food [8, 234]. A biopesticide, consisting of a rhizosphere-competent non-aflatoxigenic strain of *Aspergillus* with competitive saprophytic ability may competitively exclude toxigenic strains from infecting crops [265, 266]. For peanuts, a commercial non-toxigenic *A. flavus* strain, NRRL 21882 has been traded as Afla-Guard® in the United States [267]. Fluorescent pseudomonads and several strains of *Trichoderma* spp inhabit the rhizosphere of many crop plants and have been identified as potentially promising biocontrol agents against *A. flavus*. Since the beginning of the 21st century, many actinomycetes (*Streptomyces* spp) strains, *Trichoderma* (> 250), *Pseudomonas* (> 100) spp have been isolated, evaluated and validated to possess antagonism towards *A. flavus* [268]. Significant reduction of *A. flavus* populations and peanut kernel infection occurred in both greenhouse and field experiments. Two *Trichoderma* isolates (Tv 47 and Tv 23) and two bacterial isolates (*P. cepacia* B 33 & *P. fluorescens* Pf 2) were effective in reducing AF content in the peanut kernels. However, the efficacy of these agents warrant establishment under Kenyan conditions so that affordable, readily available and effective formulations can be developed for use. Promising biocontrol agents tested under greenhouse and field conditions in Africa and Asia has so far proved effective in reducing AFs up to 79% [269].

8.2 Post-Harvest Management

Proper drying of produce to moisture contents between 12-14%, preferably 12.5% or below is recommended. Harvested crops should be shelled and cleaned prior to storage to reduce incidences of pest infestation which may induce mould growth [270]. Further, storage facilities should be well ventilated to ensure temperatures between 25 °C and 32 °C and sustained relative humidity above 65% suitable for aflatoxin growth are not attained [11]. Moisture of 12–13% and temperatures below 18 °C does not favour the growth of *Aspergillus* fungi [271].

Following good agricultural, good storage and good manufacturing practices as well as use of advanced agricultural technologies can reduce AF contamination [152]. Novel food processing techniques such as use of ozone, pulsed light, electrolyzed water, electron beam, microwave, cold plasma, gamma and ultraviolet irradiation can reduce AF concentrations in agricultural foods. Food additives such as citric acid have been reported to sequester AFs in combination with moisture at high pressures and temperatures [272].

Clays (e.g. Novasil Plus) has been reported to bind AFs [273], reducing their available concentration. Compounds such as curcumin can alter the microsomal activation of AFB₁ and reduce the AFB₁ toxicity by increasing its detoxification. Chemoprotection against AFs ingested by animals has also been reported [274]. It utilizes compounds such as esterified

glucomanoses and other yeast extracts that increases the animal's detoxification process or otherwise prevent the production of AF-epoxide, thereby reducing or blocking AFB₁-induced hepatocarcinogenesis. Oltipraz and chlorophyll are used to reduce the biologically effective dose and acts by binding AFs, thereby rendering them biologically unavailable to humans and animals [274].

The management strategies suggested each have their advantages and limitations. Thus, biocontrol measures in synchrony with other physical and chemical methods along with improved packaging materials should be implemented to manage the plague of AFs in Kenya.

9. Conclusion

Aflatoxin exposure is ubiquitous in Kenya and the different commodities have relatively high levels of AFs, usually above statutory compliance limits by several folds. Maize, peanuts and their products are the most contaminated food crops in Kenya. Variations in AF exposure are evident between the different regions of the country and is fundamentally a function of diet and economic status. Large-scale, evidence-based interventions are required to reduce exposure. More exposure assessments, including co-exposure with other mycotoxins alongside routine monitoring of AFs should be adopted.

Data Availability

This article is a review article and no raw data were collected. Any data used and/or analyzed are within this article.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

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References

1. K. R. N. Reddy, B. Salleh, B. Saad, et al., "An overview of mycotoxin contamination in foods and its implications for human health," *Toxin Reviews*, vol. 29, pp. 3-26, 2010.

- 1351 2. J. W. Bennett, and M. Klich, "Mycotoxins," *Clinical Microbiology Reviews*, vol. 16, no. 3, pp. 497–
1352 516, 2003.
- 1353 3. J. M. Wagacha, and J. W. Muthomi, "Mycotoxin problem in Africa: current status, implications to food
1354 safety and health and possible management strategies," *International Journal of Food Microbiology*,
1355 vol. 124, no. 1, pp. 1–12, 2008.
- 1356 4. D. C. Kemboi, G. Antonissen, P. E. Ochieng et al., "A review of the impact of mycotoxins on dairy
1357 cattle health: challenges for food safety and dairy production in Sub-Saharan Africa," *Toxins*, vol. 12,
1358 no. 222, 2020.
- 1359 5. V. Kumar, M. S. Basu, and T. P. Rajendran, "Mycotoxin research and mycoflora in some
1360 commercially important agricultural commodities," *Crop Protection*, vol. 27, pp. 891-905, 2008.
- 1361 6. J. H. Williams, T. D. Phillips, P. E. Jolly et al., "Human aflatoxicosis in developing countries: A
1362 review of toxicology, exposure, potential health consequences, and interventions," *The American*
1363 *Journal of Clinical Nutrition*, vol. 80, no. 5, pp. 1106–1122, 2004.
- 1364 7. Y. Gebrehiwet, S. Ngqangweni, and J. F. Kirsten, "Quantifying the trade effect of sanitary and
1365 phytosanitary regulations of OECD countries on South African food exports," *Agrekon*, vol. 46, pp.
1366 23-38, 2007.
- 1367 8. B. Mutahi, "Kenya's ugali scare: How safe is your maize flour?" [https://www.bbc.com/news/world-](https://www.bbc.com/news/world-africa-50407159)
1368 [africa-50407159](https://www.bbc.com/news/world-africa-50407159)
- 1369 9. A. Sirma, J. F. Lindahl, K. Makita et al., "The impacts of aflatoxin standards on health and nutrition in
1370 sub-Saharan Africa: the case of Kenya," *Global Food Security*, vol. 18, pp. 57-61, 2018.
- 1371 10. N. Baranyi, S. Kocsube, C. Vagvolgyi, and J. Varga, "Current trends in aflatoxin research," *Acta*
1372 *Biologica Szegediensis*, vol. 57, pp. 95–107, 2013.
- 1373 11. P. Villers, "Aflatoxins and safe storage," *Frontiers in Microbiology*, vol. 5, no. 158, pp. 1–6, 2014.
- 1374 12. W. P. Blount, "Turkey X disease," *Journal of British Turkey Federation*, vol. 9, no. 52, pp. 52–61,
1375 1961.
- 1376 13. J. L. Richard, "Discovery of aflatoxins and significant historical features," *Toxin Reviews*, vol. 27, no.
1377 3-4, pp. 171–210, 2008.
- 1378 14. K. Sargeant, A. Sheridan, J. O'Kelly, and R. B. A. Carnaghan, "Toxicity associated with certain
1379 samples of groundnuts," *Nature*, vol. 192, no. 4807, pp. 1096-1097, 1961.
- 1380 15. F. D. Asplin, and R. B. A. Carnaghan, "The toxicity of certain groundnut meals for poultry with special
1381 reference to their effect on ducklings and chickens," *Veterinary Records*, vol. 73, pp. 1215–1219, 1961.
- 1382 16. T. Omara, W. Nassazi, T. Omute et al., "Aflatoxins in Uganda: an encyclopedic review of the etiology,
1383 epidemiology, detection, quantification, exposure assessment, reduction, and control," *International*
1384 *Journal of Microbiology*, vol. 2020, Article ID 4723612, 18 p., 2020.
- 1385 17. Q. Wu, A. Jezkova, Z. Yuan et al., "Biological degradation of aflatoxins," *Drug Metabolism Reviews*,
1386 vol. 41, no. 1, pp. 1–7, 2009.
- 1387 18. INCHEM, Principles of Evaluating Chemical Effects on the Aged Population: International
1388 Programme on Chemical Safety-Environmental Health Criteria 144, World Health Organization,
1389 Geneva, Switzerland, 1993.
- 1390 19. H. S. Chun, H. J. Kim, H. E. Ok, J.-B. Hwang, and D.-H. Chung, "Determination of aflatoxin levels in
1391 nuts and their products consumed in South Korea," *Food Chemistry*, vol. 102, no. 1, pp. 385–391,
1392 2007.
- 1393 20. A. W. Yunus, E. Razzazi-Fazeli, and J. Bohm, "Aflatoxin B1 in affecting broiler's performance,
1394 immunity, and gastrointestinal tract: a review of history and contemporary issues," *Toxins*, vol. 3, pp.
1395 566-590, 2011.
- 1396 21. H. Strosnider, E. Azziz-Baumgartner, M. Banziger et al., "Workgroup report: Public health strategies
1397 for reducing AF exposure in developing countries," *Environmental Health Perspectives*, vol. 114, pp.
1398 1898-03, 2006.
- 1399 22. K. D. Raney, D. J. Meyer, B. Ketterer, T. M. Harris, and F. P. Guengerich, "Glutathione conjugation of
1400 aflatoxin B1 exo- and endo-epoxides by rat and human glutathione S-transferases," *Chemical Research*
1401 *in Toxicology*, vol. 5, no. 4, pp. 470–478, 1992.

- 1402 23. J. K. Jewers, "Mycotoxins and their Effect on Poultry Production," Tropical Development and
1403 Research Institute (TDRI), London, UK, 2015.
- 1404 24. D. M. Dereszynski, S. A. Center, J. F. Randolph et al., "Clinical and clinicopathologic features of dogs
1405 that consumed foodborne hepatotoxic aflatoxins: 72 cases (2005-2006)," *Journal of the American*
1406 *Veterinary Medical Association*, vol. 232, no. 9, pp. 1329–1337, 2008.
- 1407 25. G. C. Dors, S. S. Caldas, V. Feddern et al., "Aflatoxins: contamination, analysis and control," In:
1408 *Aflatoxins-Biochemistry and Molecular Biology*, InTechOpen, Shanghai, China, pp. 415–438, 2011.
- 1409 26. N. Benkerroum, "Retrospective and prospective look at aflatoxin research and development from a
1410 practical standpoint," *International Journal of Environmental Research and Public Health*, vol. 16, no.
1411 3633, 2019.
- 1412 27. M. Klvana, and U. Bren, "Aflatoxin B1-Formamidopyrimidine DNA adducts: relationships between
1413 structures, free energies, and melting temperatures," *Molecules*, vol. 24, no. 150, 2019.
- 1414 28. A. P. Wacoo, D. Wendi, P. C. Vuzi, and J. F. Hawumba, "Methods for detection of aflatoxins in
1415 agricultural food crops," *Journal of Applied Chemistry*, vol. 2014, Article ID 706291, 15p., 2014.
- 1416 29. J. Mehrzad, A. M. Malvandi, M. Alipour, and S. Hosseinkhani, "Environmentally relevant level of
1417 aflatoxin B1 elicits toxic pro-inflammatory response in murine CNS-derived cells," *Toxicology*
1418 *Letters*, vol. 279, pp. 96-106, 2017.
- 1419 30. M. E. Smela, S. S. Currier, E. A. Bailey, and J. M. Essigmann, "The chemistry and biology of aflatoxin
1420 B(1): From mutational spectrometry to carcinogenesis," *Carcinogenesis*, vol. 22, no. 4, pp. 535-545,
1421 2001.
- 1422 31. L. Wojnowski, P. C. Turner, B. Pedersen et al., "Increased levels of aflatoxin-albumin adducts are
1423 associated with CYP3A5 polymorphisms in The Gambia, West Africa," *Pharmacogenetics*, vol. 14,
1424 pp. 691-700, 2004.
- 1425 32. G. E. Neal, D. L. Eaton, D. J. Judah, and A. Verma, "Metabolism and toxicity of aflatoxins M1 and B1
1426 in human-derived in vitro systems," *Toxicology and Applied Pharmacology*, vol. 151, pp. 152–158,
1427 1998.
- 1428 33. L. L. Bedard, and T.E. Massey, "Aflatoxin B1-induced DNA damage and its repair," *Cancer Letters*,
1429 vol. 241, pp. 174-183, 2006.
- 1430 34. Z. Zhuang, Y. Huang, Y. Yang, and S. Wang, "Identification of AFB1-interacting proteins and
1431 interactions between RPSA and AFB1," *Journal of Hazardous Materials*, vol. 301, pp. 297-303, 2016.
- 1432 35. C. de Oliveira, and P. Germano, "Aflatoxins: current concepts on mechanisms of toxicity and their
1433 involvement in the etiology of hepatocellular carcinoma," *Revista de Sa'ude P'ublica*, vol. 31, pp. 417-
1434 424, 1997.
- 1435 36. G. S. Bbosa, D. Kitya, A. Lubega et al., "Review of the biological and health effects of aflatoxins on
1436 body organs and body systems, in *Aflatoxins-Recent advances and future prospects*," InTechOpen, pp.
1437 239-265, 2013.
- 1438 37. A. P. Wacoo, S. Alier, T. Okoth et al., "The roles of novel proteins in the reduction of aflatoxin-
1439 induced liver cancer," (in press), 2020.
- 1440 38. C. K. Mutegi, P.J. Cotty, and R. Bandyopadhyay, "Prevalence and mitigation of aflatoxins in Kenya
1441 (1960-to date)," *World Mycotoxin Journal*, vol. 11, no. 3, pp. 341-357, 2018.
- 1442 39. C. Probst, H. Njapau, and P. J. Cotty, "Outbreak of an acute aflatoxicosis in Kenya in 2004:
1443 Identification of the causal agent," *Applied and Environmental Microbiology*, vol. 8, no. 73, pp. 2762-
1444 2764, 2007.
- 1445 40. J. W. Muthomi, L. N. Njenga, J. K. Gathumbi, and G. N. Chemining'wa, "The occurrence of aflatoxins
1446 in maize and distribution of mycotoxin-producing fungi in Eastern Kenya," *Plant Pathology Journal*,
1447 vol. 8, no. 3, pp. 113-119, 2009.
- 1448 41. S. Okoth, B. Nyongesa, V. Ayugi et al., "Toxigenic potential of *Aspergillus* species occurring on maize
1449 kernels from two agroecological zones in Kenya," *Toxins*, vol. 4, no. 11, pp. 991–1007, 2012.
- 1450 42. D. R. Oloo, S. Okoth, P. Wachira et al., "Genetic profiling of *Aspergillus* Isolates with Varying
1451 Aflatoxin production potential from different maize-growing regions of Kenya," *Toxins*, vol. 11, no. 8,
1452 pp. 467, 2019.

- 1453 43. A. Mitema, S. Okoth, and S. M. Rafudeen, "The development of a qPCR assay to measure *Aspergillus*
1454 *flavus* biomass in maize and the use of a biocontrol strategy to limit aflatoxin production," *Toxins*, vol.
1455 11, no. 3, pp. 179, 2019.
- 1456 44. C. K. Mutegei, H. K. Ngugi, S. L. Hendriks, and R. B. Jones, "Factors associated with the incidence of
1457 *Aspergillus* section *Flavi* and aflatoxin contamination of peanuts in the Busia and Homa bay districts of
1458 western Kenya," *Plant Pathology Journal*, vol. 61, no. 6, pp. 1143-1153, 2012.
- 1459 45. G. K. Muriuki, and G. M. Siboe, "Maize flour contaminated with toxigenic fungi and mycotoxins in
1460 Kenya," *African Journal of Health Sciences*, vol. 2, no. 1, pp. 236-241, 1995.
- 1461 46. E. Monda, J. Masanga, and A. Alakonya, "Variation in occurrence and aflatoxigenicity of *Aspergillus*
1462 *flavus* from two climatically varied regions in Kenya," *Toxins*, vol. 13, no. 34, 2020.
- 1463 47. Md.-S. Islam, K. A. Callicott, C. Mutegei, R. Bandyopadhyay, and P. J. Cotty, "*Aspergillus flavus*
1464 resident in Kenya: High genetic diversity in an ancient population primarily shaped by clonal
1465 reproduction and mutation-driven evolution," *Fungal Ecology*, vol. 35, pp. 20-33, 2018.
- 1466 48. S. Okoth, M. Boevre, A. de Vidal et al., "Genetic and toxigenic variability within *Aspergillus flavus*
1467 population isolated from maize in two diverse environments in Kenya," *Frontiers in Microbiology*, vol.
1468 9, pp.57, 2018.
- 1469 49. C. N. Menza, and W. M. Muturi, "Occurrence of Aflatoxigenic *Aspergillus* species in peanut varieties
1470 in Busia and Kisii Central Districts, Kenya," *Open Journal of Medical Microbiology*, vol. 8, pp. 98-
1471 108, 2018.
- 1472 50. F. J. Varga, and R. A. Samson, "A reappraisal of fungi producing AFs," *World Mycotoxin Journal*, vol.
1473 2, pp. 263–277, 2009.
- 1474 51. C. Soares, P. Rodrigues, S. W. Peterson, N. Lima, and A. Venancio, "Three new species of *Aspergillus*
1475 section *flavi* isolated from almonds and maize in Portugal," *Mycologia*, vol. 104, no. 3, pp. 682–697,
1476 2012.
- 1477 52. A. C. Baquião, M. M. de Oliveira, T. A. Reis et al., "Polyphasic approach to the identification of
1478 *Aspergillus* section *Flavi* isolated from Brazil nuts," *Food Chemistry*, vol. 139, pp. 1127–1132, 2013.
- 1479 53. J. W. Ndung'u, A. O. Makokha, C. A. Onyango et al., "Prevalence and potential for aflatoxin
1480 contamination in groundnuts and peanut butter from farmers and traders in Nairobi and Nyanza
1481 provinces of Kenya," *Journal of Applied Biosciences*, vol. 65, pp. 4922-4934, 2013.
- 1482 54. M. E. Murithi, "Prevalence of *Fusarium* and *Aspergillus* species in maize grain from Kitui, Machakos
1483 and Meru and use of near infra-red-light sorting to remove fumonisins and aflatoxin contaminated
1484 grain in Kenya," MSc Thesis, University of Nairobi, Kenya, 112 p., 2014.
- 1485 55. C. Probst, F. Schulthess, and P. J. Cotty, "Impact of *Aspergillus* section *Flavi* community structure on
1486 the development of lethal levels of aflatoxins in Kenyan maize (*Zea mays*)," *Journal of Applied*
1487 *Microbiology*, vol. 108, pp. 600-610, 2010.
- 1488 56. C. Probst, K. A. Callicott, and P. J. Cotty, "Deadly strains of Kenyan *Aspergillus* are distinct from
1489 other aflatoxin producers," *European Journal of Plant Pathology*, vol. 132, pp. 419-429, 2012.
- 1490 57. E. G. Lizárraga-Paulin, E. Moreno-Martinez, and S. P. Miranda-Castro, "Aflatoxins and their impact
1491 on human and animal health: an emerging problem," In: Guevara-González RG, editor. *Aflatoxins -*
1492 *biochemistry and molecular biology*, Rijeka (Croatia), InTech, pp. 255-282, 2011.
- 1493 58. P. Ochungo, J. F. Lindahl, T. Kayano et al., "Mapping aflatoxin risk from milk consumption using
1494 biophysical and socioeconomic data: A case study of Kenya," *African Journal of Food, Agriculture,*
1495 *Nutrition and Development*, vol. 16, no. 3, pp. 11066-11085, 2016.
- 1496 59. H. S. Nzioki, "Review of aflatoxin research in Kenya: NARS (KARI) perspective," Nairobi, Kenya,
1497 2008.
- 1498 60. J. L. Leroy, J. S. Wang, and K. Jones, "Serum aflatoxin B 1-lysine adduct level in adult women from
1499 Eastern Province in Kenya depends on household socio-economic status: A cross sectional study,"
1500 *Social Science & Medicine*, vol. 146, pp. 104-110, 2015.
- 1501 61. C. K. Mutegei, H. K. Ngugi, S. L. Hendriks, and R. B. Jones, "Prevalence and factors associated with
1502 aflatoxin contamination of peanuts from Western Kenya," *International Journal of Food Microbiology*,
1503 vol. 130, pp. 27-34, 2009.

62. C. Mutegi, M. Wagacha, J. Kimani et al., "Incidence of aflatoxin in peanuts (*Arachis hypogaea* Linnaeus) from markets in Western, Nyanza and Nairobi Provinces of Kenya and related market traits," *Journal of Stored Products Research*, vol. 52, pp. 118–127, 2013.
63. M. A. Obonyo, and E. N. Salano, "Perennial and seasonal contamination of maize by aflatoxins in Eastern Kenya," *International Journal of Food Contamination*, vol. 5, no. 6, 2018.
64. C. K. Mutegi, J. M. Wagacha, M. E. Christie, J. Kimani, and L. Karanja, "Effect of storage conditions on quality and aflatoxin contamination of peanuts (*Arachis hypogaea* L.)," *International Journal of AgriScience*, vol. 3, no. 10, pp. 746-758, 2013.
65. C. K. Mutegi, S. L. Hendricks, R. B. Jones, J. J. Okello, and H. K. Ngugi, "Role of collective action and handling practices on aflatoxin contamination of groundnuts," Paper presented at the African Crop Science Conference Proceedings, 2007.
66. G. Kiarie, P. Dominguez-Salas, S. Kang'ethe, D. Grace, and J. Lindahl, "Aflatoxin exposure among young children in urban low-income areas of Nairobi and association with child growth," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 16, pp. 10967-10990, 2016.
67. W. K. N. Moturi, "Factors likely to enhance mycotoxin introduction into the human diet through maize in Kenya," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 8, pp. 291-303, 2008.
68. P. Koskei, C. C. Bii, P. Musotsi, and S. M. Karanja, "Postharvest Storage Practices of Maize in Rift Valley and Lower Eastern Regions of Kenya: A cross-sectional study," *International Journal of Microbiology*, vol. 2020, Article ID 6109214, 2020.
69. E. Azziz-Baumgartner, K. Lindblade, K. Gieseke et al., "Case-control study of an acute aflatoxicosis outbreak, Kenya, 2004," *Environmental Health Perspectives*, vol. 113, no. 12, pp. 1779-1783, 2005.
70. J. H. Daniel, L. W. Lewis, Y. A. Redwood et al. Comprehensive assessment of maize aflatoxin levels in eastern Kenya," *Environmental Health Perspectives*, vol. 119, no. 12, pp. 1795, 2011.
71. L. Lewis, M. Onsongo, H. Njapau et al., "Aflatoxin contamination of commercial maize products during an outbreak of acute aflatoxicosis in Eastern and Central Kenya," *Environmental Health Perspectives*, vol. 113, no. 12, pp. 1763-1767, 2005.
72. S. K. Mutiga, V. Hoffmann, J. Harvey et al., "Assessment of aflatoxin and fumonisin contamination of maize in western Kenya," *Phytopathology*, vol. 105, no. 9, pp. 1250–1261, 2015.
73. E. W. Mwihia, P. G. Mbuthia, G.S. Eriksen et al., "Occurrence and levels of aflatoxins in fish feeds and their potential effects on fish in Nyeri, Kenya," *Toxins*, vol. 10, no. 543, 16p., 2018.
74. T. N. Kiama, J. Lindahl, A. J. Sirma et al., "Kenya dairy farmer perception of moulds and mycotoxins and implications for exposure to aflatoxins: A gendered analysis," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 16, no. 3, pp. 11106-11125, 2016.
75. S. K. Mutiga, V. Were, V. Hoffmann et al., "Extent and drivers of mycotoxin contamination: inferences from a survey of Kenyan maize mills," *Phytopathology*, vol. 104, pp. 1221-1231, 2014.
76. A. J. Sirma, E. O. Ouko, G. Murithi et al., "Prevalence of aflatoxin contamination in cereals from Nandi county, Kenya," *International Journal of Agricultural Sciences and Veterinary Medicine*, vol. 3, no. 3, pp. 55-63, 2015.
77. W. R. Nabwire, J. Ombaka, C. P. Dick et al., "Aflatoxin in household maize for human consumption in Kenya, East Africa," *Food Additives & Contaminants: Part B Surveillance*, vol. 13, no. 1, pp. 45-51, 2020.
78. S. A. Okoth, and M. A. Kola, "Market samples as a source of chronic aflatoxin exposure in Kenya," *African Journal of Health Sciences*, vol. 20, no. 1, pp. 56–61, 2012.
79. J. K. Kibugu, D. Mburu, L. K. Munga et al., "Food-borne mycotoxin hazards in the Kenyan market-a retrospective study," *bioRxiv preprint*, 2019, <https://doi.org/10.1101/773747>
80. M. I. Obade, P. Andang'O, C. Obonyo, and F. Lusweti, "Exposure of children 4 to 6 months of age to aflatoxin in Kisumu County, Kenya," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 15, no. 2, pp. 9949-9963, 2015.
81. G. M. Kenji, A. M. Mvula, H. Koaze, N. Baba, "Aflatoxin contamination of Kenyan maize flour and malted Kenyan and Malawian grains," *Scientific Reports of the Faculty of Agriculture, Okayama University, Okayama, Japan*, vol. 89, pp. 5–7, 2000.

- 1556 82. F. G. Peers, and C. A. Linsell, "Dietary aflatoxins and liver cancer-a population based study in Kenya,"
1557 British Journal of Cancer, vol. 27, no. 6, pp. 473-487, 1973.
- 1558 83. C. Mutungi, P. Lamuka, S. Arimi, J. Gathumbi, and C. Onyango, "The fate of aflatoxins during
1559 processing of maize into muthokoi—a traditional Kenyan food," Food Control, vol. 19, no. 7, pp. 714–
1560 721, 2008.
- 1561 84. E. W. Gachomo, E. W. Mutitu, and O. S. Kotchoni, "Diversity of fungal species associated with
1562 peanuts in storage and the levels of aflatoxins in infected samples," International Journal of
1563 Agricultural Biology, vol. 6, pp. 955-959, 2004.
- 1564 85. D. N. A. Orony, J. O. Lalah, and I. O. Jondiko, "Carcinogenic polycyclic aromatic hydrocarbons
1565 (PAHs), aflatoxins and nitrosamines in processed fish from the Winam Gulf Area of Kenya,"
1566 Polycyclic Aromatic Compounds, vol. 36, pp. 295-317, 2015.
- 1567 86. D. K. Nyangaga, "Traders' awareness and level of aflatoxins in human foods and cattle feeds in
1568 selected markets and stores in Nairobi county, Kenya," MSc Thesis, Kenyatta University, Nairobi,
1569 Kenya, 2014.
- 1570 87. E. K. Kangeth'e, G. M. M'ibui, T. F. Randolph, and A. K. Lang'a, "Prevalence of aflatoxin M1 and B1
1571 in milk and animal feeds from urban smallholder dairy production in Dagoretti division, Nairobi,
1572 Kenya," East African Medical Journal, vol. 84, no. 11, pp. S83-86, 2007.
- 1573 88. T. P. Lanyasunya, L.W. Wamae, H. H. Musa, O. Olowofeso, and I. K. Lokwaleput, "The risk of
1574 mycotoxins contamination of dairy feed and milk on smallholder dairy farms in Kenya," Pakistan
1575 Journal of Nutrition, vol. 4, no. 3, pp. 162–169, 2005.
- 1576 89. E. K. Kang'ethe, and K. A. Lang'a, "Aflatoxin B1 and M1 contamination of animal feeds and milk
1577 from urban centers in Kenya," African Health Sciences, vol. 9, no. 4, pp. 218–226, 2009.
- 1578 90. J. F. Lindah, I. N. Kagera, and D. Grace, "Aflatoxin M1 levels in different marketed milk products in
1579 Nairobi, Kenya," Mycotoxin Research, vol. 34, pp. 289-295, 2018.
- 1580 91. L. Keter, R. Too, N. Mwikwabe et al., "Risk of fungi associated with aflatoxin and fumonisin in
1581 medicinal herbal products in the Kenyan market," Scientific World Journal, vol. 2017, pp. 1-6, 2017.
- 1582 92. S. Okoth, "Improving the Evidence Base on Aflatoxin Contamination and Exposure," Series:
1583 Agriculture and Nutrition, The Technical Centre for Agricultural and Rural Cooperation, Wageningen,
1584 The Netherlands, CTA Working Paper 16/13, 2016.
- 1585 93. A. Medina, M. K. Gilbert, B. M. Mack et al., "Interactions between water activity and temperature on
1586 the Aspergillus flavus transcriptome and aflatoxin B1 production," International Journal of Food
1587 Microbiology, vol. 256, pp. 36-44, 2017.
- 1588 94. G. Mahukua, H. S. Nzioki, C. Mutegi et al., "Pre-harvest management is a critical practice for
1589 minimizing aflatoxin contamination of maize," Food Control, vol. 96, pp. 219-226, 2019.
- 1590 95. N. N. Nduti, P. N. Njeru, M. Mwaniki, and G. Reid, "Aflatoxin variations in maize flour and grains
1591 collected from various regions of Kenya," African Journal of Food, Agriculture, Nutrition and
1592 Development, vol. 17, no. 1, pp. 11743-11756, 2017.
- 1593 96. K. Hell, and C. Mutegi, "Aflatoxin control and prevention strategies in key crops of Sub-Saharan
1594 Africa," African Journal of Microbiology Research, vol. 5, no. 5, pp. 459-466, 2011.
- 1595 97. E. Kang'ethe, "Situational analysis: improving food safety in the maize value chain in Kenya,"
1596 University of Nairobi, Nairobi, Kenya, 12p., 2011.
- 1597 98. S. Egal, A. Hounsa, Y. Y. Gong et al., "Dietary exposure to aflatoxin from maize and groundnut in
1598 young children from Benin and Togo, West Africa," International Journal of Food Microbiology, vol.
1599 104, no. 2, pp. 215–224, 2005.
- 1600 99. G. S. Shephard, "Risk assessment of aflatoxins in food in Africa," Food Additives and
1601 Contamination Part A, vol. 25, pp. 1246–1256, 2008.
- 1602 100. A. R. Abizari, H. A. Garti, G. Gajate-Garrido et al., "Technological and Market Interventions for
1603 Aflatoxin Control in Ghana: Final Report," Global Alliance for Improved Nutrition, 2016.
- 1604 101. ACIDI/VOCA, Kenya Maize Development Program (KMDP). ACIDI/VOCA, Washington, DC, USA,
1605 2015.
- 1606 102. G. Kenji, "Aflatoxins in traditional beer (Busaa) in Nairobi, Kenya," African Journal of Food &
1607 Nutritional Sciences, vol. 3, pp. 7-8, 2003.

103. B. N. Muture, and G. Ogana, "Aflatoxin levels in maize and maize products during the 2004 food poisoning outbreak in Eastern Province of Kenya," *East African Medical Journal*, vol. 82, no. 6, pp. 275-279, 2005.
104. A. Pietri, M. Zanetti, and T. Bertuzzi, "Distribution of aflatoxins and fumonisins in dry-milled maize fractions," *Food Additives Contaminants*, vol. 26, pp. 372-380, 2009.
105. Aflacontrol Project Note 3, Prevalence of aflatoxin in Kenya: Summary of Findings January - June 2010. November 2010, Washington, USA, 2010.
106. S. Collins, G. Mahuku, H. S. Nzioki, C. Narrod, and P. Trench, "Aflatoxin in Kenya: An overview," Aflacontrol Project Note 1. July 2010, Washington, DC, 2010.
107. J. W. Muthomi, B. K. Mureithi, G. N. Chemining'wa, J. K. Gathumbi, and E. W. Mutitu, "Aspergillus and Aflatoxin B1 Contamination of Maize and Maize Products from Semi-Arid and Humid Regions of Kenya," *International Journal Agricultural Sciences*, vol. 2, pp. 22-34, 2012.
108. Building an Aflatoxin Safe East African Community, Technical Policy Paper 8, Aflatoxin Standards for Food Knowledge Platform 2015, Situational Analysis East Africa Region, 30 pages, <http://aflasafe.com/wp-content/uploads/pdf/TPP-8-Aflatoxin-Standards-for-Food.pdf>
109. R. M. Kilonzo, J. K. Imungi, W. M. Muiru, P. O. Lamuka, and P. M. K. Njage, "Household dietary exposure to aflatoxins from maize and maize products in Kenya," *Food Additives & Contaminants Part A*, vol. 31, no. 12, pp. 2055-2062, 2014.
110. G. W. Gachara, A. K. Nyamache, J. Harvey et al., "Genetic diversity of *Aspergillus flavus* and occurrence of aflatoxin contamination in stored maize across three agro-ecological zones in Kenya," *Agriculture & Food Security*, vol. 7, p.52, 2018.
111. A. K. Kipkoech, M. A. Okiror, J. R. Okalebo, and H. K. Maritim, "Production efficiency and economic potential of different soil fertility management strategies among groundnut farmers of Kenya," *Science World Journal*, vol. 2, pp. 15-21, 2007.
112. T. L. Bucheyeki, E. M. Shenkalwa, T. X. Mapunda, and L. W. Matata, "On-farm evaluation of promising groundnut varieties for adaptation and adoption in Tanzania," *African Journal of Agricultural Research*, vol. 3, pp. 531-536, 2008.
113. N. Okoko, N. Kidula, L. Wasilwa et al., "Participatory evaluation and dissemination of improved groundnut varieties and technologies for processing and utilization," In: *Transforming Agriculture for Improved Livelihoods through Agricultural Product Value Chains. The Proceedings of the 12th KARI Biennial Scientific Conference, 8-12th November 2010, Nairobi, Kenya*, pp.1403-1408, 2010.
114. E. N. K. Okoko, D. J. Rees, J. K. Kwach, and P. Ochieng, "Participatory evaluation of groundnut production in Southwest Kenya," In: *Towards Increased Use of Demand Driven Technology KARI/DFID NARP II PROJECT, End of Project Conference 23rd-26th March 1999, Nairobi, Kenya*, pp. 305-307, 1999.
115. H. Nyirahakizimana, L. Mwamburi, J. Wakhisi et al., "Occurrence of *Aspergillus* Species and Aflatoxin Contamination in Raw and Roasted Peanuts from Formal and Informal Markets in Eldoret and Kericho Towns, Kenya," *Advances in Microbiology*, vol. 3, pp. 333-342, 2013.
116. FAOSTAT, Production: Crops, Food and Agriculture Organization of the United Nations, 2012.
117. D. Nabuuma, D. Nakimbugwe, Y. B. Byaruhanga et al., "Formulation of a drinkable peanut-based therapeutic food for malnourished children using plant sources," *International Journal of Food Sciences and Nutrition*, vol. 64, pp. 467-475, 2012.
118. S. A. Okoth, and M. Ohingo, "Dietary aflatoxin exposure and impaired growth in young children from Kisumu district, Kenya: Cross sectional study," *African Journal of Health Sciences*, vol. 11, no. 1, pp. 43-54, 2004.
119. M. C. Nelson, M. W. Margaret, and K. M. Lucy, "Oil contents and aflatoxin levels in peanut varieties produced in Busia and Kisii Central Districts, Kenya," *Tropical Medicine & Surgery*, vol. 4, no. 204, 2p., 2016.
120. R. Nyaguthie, 6 foods that can be contaminated with aflatoxins if not well stored, 2020, <https://www.tuko.co.ke/324051-6-foods-contaminated-aflatoxins-stored.html>

121. C. N. Menza, W. M. Muturi, and M. L. Kamau, "Oil contents and aflatoxin levels in peanut varieties produced in Busia and Kisii Central Districts, Kenya," *Tropical Medicine & Surgery*, vol. 4, no. 204, pp. 1-6, 2016.
122. A. Seetha, W. Munthali, H. W. Msere et al., "Occurrence of aflatoxins and its management in diverse cropping systems of Central Tanzania," *Mycotoxin Research*, vol. 33, no. 4, pp. 323-331, 2017.
123. A. U. Achidi, O. A. Ajayi, M. Bokanga, B. Maziya-Dixon, "The use of cassava leaves as food in Africa," *Ecology and Food Nutrition*, vol. 44, pp. 423-435, 2005.
124. J. A. Montagnac, C. R. Davis, and S. A. Tanumihardjo, "Nutritional value of cassava for use as a staple food and recent advances for improvement," *Comprehensive Reviews in Food Science and Food Safety*, vol. 8, pp. 181-194, 2009.
125. A. P. Cardoso, E. Mirione, M. Ernesto et al., "Processing of cassava roots to remove cyanogens," *Journal of Food Composition Analysis*, vol. 18, pp. 451-460, 2005.
126. P. K. Gacheru, G. O. Abong, M. W. Okoth et al., "Cyanogenic content, aflatoxin level and quality of dried cassava chips and flour sold in Nairobi and Coastal regions of Kenya," *Current Research in Nutrition and Food Science*, vol. 3, no. 3, pp. 197-206, 2015.
127. M. F. Abdallah, R. Krska, and M. Sulyok, "Mycotoxin contamination in sugarcane grass and juice: first report on detection of multiple mycotoxins and exposure assessment for aflatoxins B1 and G1 in humans," *Toxins*, vol. 8, pp. 343, 2016.
128. M. E. Alpert, M. S. R. Hutt, G. N. Wogan, and C. S. Davidson, "Association between aflatoxin content of food and hepatoma frequency in Uganda," *Cancer*, vol. 28, no. 1, pp. 253-260, 1971.
129. I. Kagera, P. Kahenya, F. Mutua et al., "Status of aflatoxin contamination in cow milk produced in smallholder dairy farms in urban and peri-urban areas of Nairobi County: a case study of Kasarani sub county, Kenya," *Infection Ecology & Epidemiology*, vol. 9, no. 1, pp.1547095, 2019.
130. A. Sirma, D. Senerwa, J. Lindahl et al., "Aflatoxin M1 survey in dairy households in Kenya," Poster prepared for the Food Africa Midterm Seminar, Helsinki, Finland, 16p., 2014.
131. Y. Kirino, K. Makita, D. Grace et al., "Survey of informal milk retailers in Nairobi, Kenya and prevalence of aflatoxin M1 in marketed milk," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 16, no. 3, pp. 11022-11038, 2016.
132. G. Langat, M. Tetsuhiro, T. Gono, V. Matiru, and C. Bii, "Aflatoxin M1 contamination of milk and its products in Bomet County, Kenya," *Advances in Microbiology*, vol. 6, pp. 528-536, 2016.
133. D. Senerwa, A. J. Sirma, N. Mtimet et al., "Prevalence of aflatoxin in feeds and cow milk from five counties in Kenya," *African Journal of Food, Agriculture, Nutrition and Development*, vol. 16, p. 11004-11021, 2016.
134. G. Anyango, F. Mutua, I. Kagera et al., "A survey of aflatoxin M1 contamination in raw milk produced in urban and peri-urban areas of Kisumu County, Kenya," *Infection Ecology & Epidemiology*, vol. 8, no. 1547094, 2018.
135. M. M. Kuboka, J. K. Imungia, L. Njue et al., "Occurrence of aflatoxin M1 in raw milk traded in peri-urban Nairobi, and the effect of boiling and fermentation," *Infection Ecology & Epidemiology*, vol. 9, no. 1625703, 2019.
136. G. J. A. Speijers and M. H. M. Speijers, "Combined toxic effects of mycotoxins," *Toxicology Letters*, vol. 153, no. 1, pp. 91-98, 2004.
137. C. J. Kedera, R. D. Plattner, and A. E. Desjardins, "Incidence of *Fusarium* spp. and levels of Fumonisin B1 in maize in Western Kenya," *Applied Environmental Microbiology*, vol. 65, pp. 41-44, 1998.
138. S. K. Mbugua, and J. K. Gathumbi, "The Contamination of Kenyan Lager Beers with *Fusarium* Mycotoxins" *Journal of Institute of Brewing*, vol. 110, pp. 227-229, 2004.
139. A. E. Alakonya, E. O. Monda, and S. Ajanga, "Fumonisin B1 and aflatoxin B1 levels in Kenyan maize," *Journal of Plant Pathology*, vol. 91, no. 2, pp. 459-464, 2009.
140. I. Rodrigues, J. Handl, and E. M. Binder, "Mycotoxin occurrence in commodities, feeds and feed ingredients sourced in the Middle East and Africa," *Food Additives & Contaminants Part B: Surveillance*, vol. 4, pp. 168-179, 2011.

- 1708 141. M. C. Kirui, A. E. Alakonya, K. K. Talam, G. Tohru, and C. C. Bii, "Total aflatoxin, fumonisin and
1709 deoxynivalenol contamination of busaa in Bomet County, Kenya," *African Journal of Biotechnology*,
1710 vol. 13, pp. 2675-2678, 2014.
- 1711 142. C. M. Makau, J. W. Matofari, P. S. Muliuro, and B. O. Bebe, "Aflatoxin B1 and Deoxynivalenol
1712 contamination of dairy feeds and presence of aflatoxin M1 contamination in milk from smallholder
1713 dairy systems in Nakuru, Kenya," *International Journal of Food Contamination*, vol. 3, no. 6, 2016.
- 1714 143. N. K. Njeru, C. A. O. Midega, J. W. Muthomi, J. M. Wagacha, and Z. R. Khan, "Influence of socio-
1715 economic and agronomic factors on aflatoxin and fumonisin contamination of maize in Western
1716 Kenya," *Food Science and Nutrition*, vol. 7, pp. 2291-2301, 2019.
- 1717 144. A. Stevens, C. Saunders, J. Spence et al., "Investigations into "diseases" of Turkey poult," *Veterinary*
1718 *Record*, vol. 72, pp. 627-628, 1960.
- 1719 145. M. Ozturk, "P53 mutation in hepatocellular carcinoma after aflatoxin exposure," *Lancet*, vol. 338, pp.
1720 1356-1359, 1991.
- 1721 146. R. Jaetzold, and H. Schmidt, "Farm Management Handbook of Kenya. Natural Condition and Farm
1722 Management Information," Ministry of Agriculture and German Agricultural Team (GTZ), vol. II/C,
1723 Nairobi, Kenya, 2009.
- 1724 147. L. N. Kirimi, T. S. Sitko, F. Jayne et al., "A Farm Gate to Consumer Value Chain Analysis of Kenya's
1725 Maize Marketing System," Egerton University Working Paper Series, 44p., 2011.
- 1726 148. R. Bandyopadhyay, A. Akande, C. Mutegi et al., "Biological control of aflatoxins in Africa: Current
1727 status and potential challenges in the face of climate change," *World Mycotoxin Journal*, vol. 9, pp.
1728 771-789, 2016.
- 1729 149. J. M. Milani, "Ecological conditions affecting mycotoxin production in cereals: A review," *Veterinary*
1730 *Medicine*, vol. 58, no. 8, pp. 405-411, 2013.
- 1731 150. N. W. Turner, S. Subrahmanyam, and S. A. Piletsky, "Analytical methods for determination of
1732 mycotoxins: a review," *Analytica Chimica Acta*, vol. 632, no. 2, pp. 168-180, 2009.
- 1733 151. T. B. Whitaker, "Detecting mycotoxins in agricultural commodities," *Molecular Biotechnology*, vol.
1734 23, no. 1, pp. 61-72, 2003.
- 1735 152. M. Kamle, D. K. Mahato, S. Devi et al., "Fumonisin: impact on agriculture, food, and human health
1736 and their management strategies," *Toxins*, vol. 11, no. 328, 2019.
- 1737 153. F. Berthiller, C. Crews, C. Dall'Asta et al., "Masked mycotoxins: a review," *Molecular Nutrition and*
1738 *Food Research*, vol. 57, pp. 165-186, 2013.
- 1739 154. V. Nagl, B. Woechtl, H. E. Schwartz-Zimmermann et al., "Metabolism of the masked mycotoxin
1740 deoxynivalenol-3-glucoside in pigs," *Toxicology Letters*, vol. 229, pp. 190-197, 2014.
- 1741 155. M. Gareis, J. Bauer, J. Thiem et al., "Cleavage of zearalenone-glycoside, a "masked" mycotoxin,
1742 during digestion in swine," *Journal of Veterinary Medicine*, vol. 37, pp. 236-240, 1990.
- 1743 156. N. Broekaert, M. Devreese, T. De Mil et al., "Oral bioavailability, hydrolysis, and comparative
1744 toxicokinetics of 3-acetyldeoxynivalenol and 15-acetyldeoxynivalenol in broiler chickens and pigs,"
1745 *Journal of Agricultural Food Chemistry*, vol. 63, pp. 8734-8742, 2015.
- 1746 157. C. Dall'Asta, M. Mangia, F. Berthiller et al., "Difficulties in fumonisin determination: the issue of hidden
1747 fumonisins," *Analytical and Bioanalytical Chemistry*, vol. 395, pp.1335-1345, 2009.
- 1748 158. A. Vidal, S. Marín, V. Sanchis et al., "Hydrolisers of modified mycotoxins in maize: α-Amylase and
1749 cellulase induce an underestimation of the total aflatoxin content," *Food Chemistry*, vol. 248, pp. 86-
1750 92, 2018.
- 1751 159. N. V. Beloglazova, M. De Boevre, I. Y. Goryacheva et al., "Immunochemical approach for
1752 zearalenone-4-glucoside determination," *Talanta*, vol. 106, pp. 422-430, 2013.
- 1753 160. C. Dall'Asta, G. Galaverna, G. Aureli, A. Dossena, and R. Marchelli, "A LC/MS/MS method for the
1754 simultaneous quantification of free and masked fumonisins in maize and maize-based products,"
1755 *World Mycotoxin Journal*, vol. 1, pp. 237-246, 2008.
- 1756 161. F. Tao, H. Yao, Z. Hruska et al., "Recent development of optical methods in rapid and non-destructive
1757 detection of aflatoxin and fungal contamination in agricultural products," *Trends in Analytical*
1758 *Chemistry*, vol. 100, pp. 65-81, 2018.

- 1759 162. J. M. Castelino, M. N. Routledge, S. Wilson et al., "Aflatoxin exposure is inversely associated with
1760 IGF1 and IGFBP3 levels in vitro and in Kenyan school children," *Molecular Nutrition and Food*
1761 *Research*, vol. 59, pp. 574-581, 2015.
- 1762 163. O. Guguyu, American companies jostle for aflatoxin test kits deal, 2017,
1763 [https://www.standardmedia.co.ke/business/article/2001243799/american-companies-jostle-for-](https://www.standardmedia.co.ke/business/article/2001243799/american-companies-jostle-for-aflatoxin-test-kits-deal)
1764 [aflatoxin-test-kits-deal](https://www.standardmedia.co.ke/business/article/2001243799/american-companies-jostle-for-aflatoxin-test-kits-deal)
- 1765 164. H. Yao, Z. Hruska, R. Kincaid et al., "Correlation and classification of single fluorescence
1766 hyperspectral data with AF concentration in corn kernels inoculated with *A. flavus* spores," *Food*
1767 *Additives & Contamination Part A*, vol. 27, no. 5, pp. 701-709, 2010.
- 1768 165. J. Boonena, S. V. Malyshev, L. Taevernier et al., "Human skin penetration of selected model
1769 mycotoxins," *Toxicology*, vol. 301, pp. 21- 32, 2012.
- 1770 166. P. Larsson, and H. Tjalve, "Intranasal instillation of Aflatoxin B1 in rats: Bioactivation in the nasal
1771 mucosa and neuronal transport to the olfactory bulb," *Toxicological Science*, vol. 55, pp. 383-391,
1772 2000.
- 1773 167. L. K. Kamdem, I. Meineke, U. Gödtel-Armbrust, J. Brockmöller, and L. Wojnowski, "Dominant
1774 Contribution of P450 3A4 to the Hepatic Carcinogenic Activation of Aflatoxin B1," *Chemical*
1775 *Research in Toxicology*, vol. 19, no. 4, pp. 577-586, 2006.
- 1776 168. T. Kamataki, H. Hashimoto, M. Shimoji et al., "Expression of CYP3A7, a human fetus-specific
1777 cytochrome P450, in cultured cells and in the hepatocytes of p53-knockout mice," *Toxicology Letters*,
1778 vol. 82-83, pp. 879-882, 1995.
- 1779 169. E. K. Kang'ethe, M. Gatwiri, A. J. Sirma et al., "Exposure of Kenyan population to aflatoxins in foods
1780 with special reference to Nandi and Makueni counties," *Food Quality and Safety*, vol. 1, no. 2, pp.
1781 131-137, 2017.
- 1782 170. M. N. Routledge, and Y. Y. Gong, "Developing biomarkers of human exposure to mycotoxins," in
1783 *Determining Mycotoxins and Mycotoxigenic Fungi in Food and Feed*, S. De Saeger, Ed., pp. 225-244,
1784 Woodhead Publishing, Cambridge, UK, 2011.
- 1785 171. P. L. Skipper, D. H. Hutchins, R. J. Turesky et al., "Carcinogen binding to serum-albumin," in
1786 *Proceedings of the American Association for Cancer Research*, American Association for Cancer
1787 Research, Houston, TX, USA, February 1985.
- 1788 172. S. Egal, A. Hounsa, Y. Y. Gong et al., "Dietary exposure to aflatoxin from maize and groundnut in
1789 young children from Benin and Togo, West Africa," *International Journal of Food Microbiology*, vol.
1790 104, no. 2, pp. 215-224, 2005.
- 1791 173. B. I. Agag, "Mycotoxins in foods and feeds: aflatoxins," *Assiut University Bulletin for Environmental*
1792 *Researches*, vol. 7, pp. 173-206, 2004.
- 1793 174. B. Chapot and C. P. Wild, "ELISA for quantification of AFalb adducts and their application to human
1794 exposure assessment," in *Techniques in Diagnostic Pathology*, M. Warhol, D. van Velzen, and G. R.
1795 Bullock, Eds., pp. 1-13, Academic Press, San Diego CA, USA, 1991.
- 1796 175. G. Sabbioni, S. Ambs, G. N. Wogan, and J. D. Groopman, "The aflatoxin-lysine adduct quantified by
1797 high-performance liquid chromatography from human serum albumin samples," *Carcinogenesis*, vol.
1798 11, no. 11, pp. 2063-2066, 1990.
- 1799 176. P. F. Scholl, L. McCoy, T. W. Kensler, and J. D. Groopman, "Quantitative analysis and chronic
1800 dosimetry of the aflatoxin B1 Plasma albumin adduct lys-AFB1 in rats by isotope dilution mass
1801 spectrometry," *Chemical Research in Toxicology*, vol. 19, no. 1, pp. 44-49, 2006.
- 1802 177. C. A. Linsell, "Cancer Incidence in Kenya, 1957-1963," *British Journal of Cancer*, vol. 21, pp. 465-
1803 473, 1967.
- 1804 178. A. Ngindu, P. Kenya, D. Ocheng et al., "Outbreak of acute hepatitis caused by aflatoxin poisoning in
1805 Kenya," *The Lancet*, vol. 319, no. 8285, pp. 1346-1348, 1982.
- 1806 179. G. N. Marete, L. W. Kanja, J. M. Mbaria et al., "Effects of the use of good agricultural practices on
1807 aflatoxin levels in maize grown in Nandi County, Kenya," *Sci*, vol. 2, 26p., 2020.
- 1808 180. N. N. Nduti, G. Reid, M. Sumarah et al., "Weissella cibaria Nn20 isolated from fermented kimere
1809 shows ability to sequester AFB1 in vitro and ferment milk with good viscosity and Phin comparison to
1810 yogurt," *Food Science & Nutrition Technology*, vol. 3, pp. 000137, 2018.

- 1811 181. V. Hoffmann, K. Jones, and J. L. Leroy, "The impact of reducing dietary aflatoxin exposure on child
1812 linear growth: a cluster randomised controlled trial in Kenya," *BMJ Global Health*, vol. 3, pp. e000983,
1813 2018.
- 1814 182. A. O. Awuor, E. Yard, J. H. Daniel et al., "Evaluation of the efficacy, acceptability and palatability of
1815 calcium montmorillonite clay used to reduce aflatoxin B1 dietary exposure in a crossover study in
1816 Kenya," *Food Additives & Contaminants: Part A*, vol. 34, pp. 93-102, 2017.
- 1817 183. N. Nduti, A. McMillan, S. Seney et al., "Investigating probiotic yoghurt to reduce an aflatoxin B1
1818 biomarker among school children in Eastern Kenya: Preliminary study," *International Dairy Journal*,
1819 vol. 63, pp. 124-129, 2016.
- 1820 184. C. C. Kirui, "Assessment of the traditional and improved processing methods in the reduction of
1821 aflatoxin levels in maize and maize products," MSc Thesis, University of Nairobi, Nairobi, Kenya,
1822 60p., 2016.
- 1823 185. E. E. Yard, J. H. Daniel, L. S. Lewis et al., "Human aflatoxin exposure in Kenya, 2007: a cross-
1824 sectional study," *Food Additives & Contaminants: Part A*, vol. 30, no. 7, pp. 1322-1331, 2013.
- 1825 186. Y. Y. Gong, S. Wilson, J. K. Mwatha et al., "Aflatoxin exposure may contribute to chronic hepatomegaly
1826 in Kenyan school children," *Environmental Health Perspectives*, vol. 120, pp. 893-896, 2012.
- 1827 187. C. P. Wild, Y. Z. Jiang, G. Sabbioni, B. Chapot, and R. Montesano, "Evaluation of methods for
1828 quantitation of aflatoxin-albumin adducts and their application to human exposure assessment," *Cancer*
1829 *Research*, vol. 50, no. 2, pp. 245-51, 1990.
- 1830 188. S. M. Maxwell, F. Apeagyei, H. R. De Vries, D. D. Mwanmut, and R. G. Hendrickse, "Aflatoxins in
1831 breast milk, neonatal cord blood and sera of pregnant women," *Journal of Toxicology-Toxins Reviews*,
1832 vol. 8, no. 1-2, pp. 19-29, 1989.
- 1833 189. H. Autrup, T. Seremet, J. Wakhisi, and A. Wasunna, "Aflatoxin exposure measured by urinary excretion
1834 of aflatoxin B1-guanine adduct and hepatitis B virus infection in areas with different liver cancer
1835 incidence in Kenya," *Cancer Research*, vol. 47, no. 13, pp. 3430-3433, 1987.
- 1836 190. S. H. Henry, T. Whitaker, I. Rabbani et al., "Aflatoxin M1," JECFA 47, Food and Drug
1837 Administration, Washington DC, USA, 2001.
- 1838 191. A. W. Onyango, O. Receveur, and S. A. Esrey, "The contribution of breastmilk to toddlers diet in
1839 Western Kenya," *Bulletin of the World Health Organization*, vol. 80, pp. 292-299, 2002.
- 1840 192. D. Githang'a, and A. Awuor, "Acute Aflatoxin exposure and impacts: The Kenyan example, and
1841 response towards outbreaks," Presentation at the PACA Meeting on Engaging The Health And
1842 Nutrition Sector in Aflatoxin Control in Africa, African Union, Addis Ababa, Ethiopia, pp. 23-24,
1843 March 2016.
- 1844 193. R. Echodu, H. Edema, G. M. Malinga et al., "Is nodding syndrome in Northern Uganda linked to
1845 consumption of mycotoxin contaminated food grains?" *BMC Research Notes*, vol. 11, no. 1, pp. 678,
1846 2018.
- 1847 194. R. Echodu, G. M. Malinga, K. J. Moriku, E. Ovuga, and G. Haesaert, "Prevalence of aflatoxin,
1848 ochratoxin and deoxynivalenol in cereal grains in Northern Uganda: Implication for food safety and
1849 health," *Toxicology Reports*, vol. 6, pp. 1012-1017, 2019.
- 1850 195. A. W. Kigotho, "Fungal problems in Kenya," *The Lancet*, vol. 350, no. 9079, pp. 718, 1997.
- 1851 196. G. Njag, "Isolation of fungal agents from formulated and commercial feeds in three fish farms in humid
1852 tropical environments of Kenya," In: *Proceedings of the 5th Annual National Biosafety Conference*,
1853 National Biosafety Authority, Nairobi, Kenya, 2016.
- 1854 197. A. Atukwase, A. Kaaya, and C. Muyanja, "Factors associated with Fumonisin contamination of maize
1855 in Uganda," *Journal of the Science of Food and Agriculture*, vol. 89, no.14, pp. 2393-2398, 2009.
- 1856 198. A. Nafi'u, and K. T. K. Anandapandian, "The occurrence of waterborne diseases in drinking water in
1857 Nakaloke Sub-county, Mbale district, Uganda," *International Journal of Scientific Research*, vol. 5, no.
1858 10, pp. 1416-1421, 2016.
- 1859 199. Y. Y. Gong, L. Torres-Sanchez, L. Lopez-Carrillo et al., "Association between tortilla consumption and
1860 human urinary fumonisin B1 levels in a Mexican population," *Cancer Epidemiology Biomarkers &*
1861 *Prevention*, vol. 17, no. 3, pp. 688-694, 2008

1862 200. P. C. Turner, V. J. Burley, J. A. Rothwell et al., "Dietary wheat reduction decreases the level of urinary
1863 deoxynivalenol in UK adults," *Journal of Exposure Science & Environmental Epidemiology*, vol. 18,
1864 no. 4, pp. 392–399, 2008.

1865 201. C. P. Shirima, M. E. Kimanya, M. N. Routledge et al., "A prospective study of growth and biomarkers
1866 of exposure to aflatoxin and fumonisin during early childhood in Tanzania," *Environmental Health*
1867 *Perspectives*, vol. 123, pp. 173-178, 2015.

1868 202. C. Srey, M. E. Kimanya, M. N. Routledge, C. P. Shirima, and Y. Y. Gong, "Deoxynivalenol exposure
1869 assessment in young children in Tanzania," *Molecular Nutrition & Food Research*, vol. 58, pp. 1574-
1870 1580, 2014.

1871 203. S. Piekkola, P. C. Turner, M. Abdel-Hamid et al., "Characterisation of aflatoxin and deoxynivalenol
1872 exposure among pregnant Egyptian women," *Food Additives & Contaminants: Part A*, vol. 29, pp.
1873 962-971, 2012.

1874 204. Y. Y. Gong, A. Hounsa, S. Egal et al., "Postweaning exposure to aflatoxin results in impaired child
1875 growth: a longitudinal study in Benin, West Africa," *Environmental Health Perspectives*, vol. 112, pp.
1876 1334-1338, 2004.

1877 205. K. F. Cardwell, and S. H. Henry, "Risk of exposure to and mitigation of effect of aflatoxin on human
1878 health: a West African example," *Toxin Reviews*, vol. 23, pp. 217-247, 2004.

1879 206. E. O. Farombi, "Aflatoxin contamination of foods in developing countries: implications for
1880 hepatocellular carcinoma and chemopreventive strategies," *African Journal of Biotechnology*, vol. 5,
1881 pp. 1-14, 2006.

1882 207. WHO, WHO Child Growth Standards: Length/height-for-age, weight-for-age, weight-for-length,
1883 weight-for-height and body mass index-for-age: methods and development. Geneva, Switzerland,
1884 2006.

1885 208. Y. Y. Gong, P. C. Turner, A. J. Hall, and C. P. Wild, "Aflatoxin exposure and impaired child growth in
1886 West Africa: An unexplored international public health burden?" In: Leslie JF, Banerjee R, Visconti A
1887 Eds. *Mycotoxins Detection Methods, Management, Public Health and Agricultural Trade*, pp. 53-66,
1888 2008.

1889 209. L. E. Smith, R. J. Stoltzfus, and A. Prendergast, "Food chain mycotoxin exposure, gut health, and
1890 impaired growth: a conceptual framework," *Advances in Nutrition*, vol. 3, pp. 526-531, 2012.

1891 210. International Agency for Research on Cancer. IARC monographs on the evaluation of carcinogenic
1892 risks to humans; vol. 56, Chemical agents and related occupations. IARC, Lyon, France, 100F: 1–599,
1893 2012.

1894 211. R. G. Hendrickse, "Of sick turkeys, kwashiorkor, malaria, perinatal mortality, heroin addicts and food
1895 poisoning: Research on the Influence of aflatoxins on child health in the tropics," *Annals of Tropical*
1896 *Medicine and Parasitology*, vol. 91, pp. 787-793, 1997.

1897 212. H. L. Mehl, and P. J. Cotty, "Variation in competitive ability among isolates of *Aspergillus flavus* from
1898 different vegetative compatibility groups during maize infection," *Phytopathology*, vol. 100, no. 2, pp.
1899 150-159, 2010.

1900 213. M. N. Okioma, "The 2004 and 2005 aflatoxin tragedies in Kenya-a case study," In: Leslie JF,
1901 Bandyopadhyay R, Visconti A, eds. *Mycotoxins: Detection Methods, Management, Public Health and*
1902 *Agricultural Trade*. Trowbridge: Cromwell Press, pp. 127-31, 2008.

1903 214. J. D. Groopman, L. G. Cain, and C. C. Krusler, "Aflatoxin exposure in human populations:
1904 measurements and relationship to cancer," *CRC Critical Reviews in Toxicology*, vol. 19, pp. 113-145,
1905 1988.

1906 215. O. W. Mwanda, C. F. Otieno, and E. Omonge, "Acute aflatoxicosis: case report," *East African Medical*
1907 *Journal*, vol. 82, pp. 320-324, 2005.

1908 216. K. Krishnamachari, R. V. Nagarajan, R. Bhat, and T. B. G. Tilak, "Hepatitis due to aflatoxicosis. An
1909 outbreak in Western India," *Lancet*, vol. 305, no. 7915, pp. 1061-1063, 1975.

1910 217. B. N. Tandon, L. Krishnamurthy, A. Koshy, et al., "Study of an epidemic of jaundice, presumably due
1911 to toxic hepatitis, in Northwest India," *Gastroenterology*, vol. 72, pp. 488-494, 1977.

1912 218. J. I. Pitt, "An Introduction to Mycotoxins, Food and Agriculture Organization, Rome Italy," In:
1913 Mycotoxin Prevention and Control in Food Grains: Eds. Semple RL, Frio AS, Hicks PA, Lozare JV,
1914 1989.

1915 219. A. Serck-Hanssen, "Aflatoxin-induced fatal hepatitis?," Archives of Environmental Health, vol. 20, no.
1916 6, pp. 729-731, 1970.

1917 220. A. Kamala, C. Shirima, B. Jani et al., "Outbreak of an acute aflatoxicosis in Tanzania during 2016,"
1918 World Mycotoxin Journal, vol. 11, no. 3, pp. 311-320, 2018.

1919 221. E. K. Kang'ethe, H. Korhonen, K. A. Marimba et al., "Management and mitigation of health risks
1920 associated with the occurrence of mycotoxins along the maize value chain in two counties in Kenya,"
1921 Food Quality & Safety, vol. 1, no. 4, pp. 268-274, 2017.

1922 222. Centers for Disease Control and Prevention (CDC). Outbreak of Aflatoxin Poisoning: Eastern and
1923 Central Provinces, Kenya, January-July 2004. Morbidity and Mortality Weekly Report. September 3,
1924 vol. 53, no.34, pp.790-793, 2004.

1925 223. R. C. Shank, J. E. Gordon, C. N. Wogan, A. Nondasuta, and B. Subhamani, "Dietary aflatoxins and
1926 human liver cancer. III. Field survey of rural Thai families for ingested aflatoxins," Food and
1927 Cosmetics Toxicology, vol. 10, pp. 171-179, 1972.

1928 224. F. G. Peers, and C. A. Linsell, "Dietary aflatoxins and human primary liver cancer," Annales de la
1929 Nutrition et de L'alimentation, vol. 31, no. 4-6, pp. 1005-1017, 1977.

1930 225. G.N. Wogan, "Dietary factors and special epidemiological situations of liver cancer in Thailand and
1931 Africa," Cancer Research, vol. 35, pp. 3499-502, 1975.

1932 226. N. D. McGlashan, "Primary liver cancer and food-based toxins. A Swaziland review," Ecology of
1933 Disease, vol. 1, no. 1, pp. 37-44, 1982.

1934 227. J. R. Barrett, "Liver cancer and aflatoxin: new information from the Kenyan outbreak," Environmental
1935 Health Perspectives, vol. 113, pp. A837-A838, 2005.

1936 228. B. Kucukcakan, and Z. Hayrulai-Musliu, "Challenging role of dietary aflatoxin B1 exposure and
1937 Hepatitis B infection on risk of hepatocellular carcinoma," Open Access Macedonian Journal of
1938 Medical Sciences, vol. 3, no. 2, pp. 363-369, 2015.

1939 229. S. Marin, A. J. Ramos, G. Cano-Sancho, and V. Sanchis, "Mycotoxins: occurrence, toxicology, and
1940 exposure assessment," Food and Chemical Toxicology, vol. 60, pp. 218-237, 2013.

1941 230. A. Petruzzello, "Epidemiology of Hepatitis B Virus (HBV) and Hepatitis C Virus (HCV) related
1942 hepatocellular carcinoma," The Open Virology Journal, vol. 12, pp. 26-32, 2018.

1943 231. C. P. Wild, and Y.Y. Gong, "Mycotoxins and human disease: a largely ignored global health issue,"
1944 Carcinogenesis, vol. 31, pp. 71-82, 2010.

1945 232. A. Magnussen, and M. A. Parsi, "Aflatoxins, hepatocellular carcinoma and public health," World
1946 Journal of Gastroenterology, vol. 19, no. 10, pp. 1508-1512, 2013.

1947 233. International Food Policy Research Institute. Aflatoxins Finding Solutions for Improved Food Safety.
1948 Unnevehr, L. and Grace, D. (Eds). Focus 20, p.1-62, November 2013.

1949 234. IITA, J. Atehnkeng, C. Mutegi et al., Management of aflatoxins in maize and groundnuts in Kenya: A
1950 Farmers' Training Manual, pp. 1-25, 2018.

1951 235. Kenya Bureau of Standards (KEBS). Kenya Standard KSEAS 2: 2017. Maize grains specifications.
1952 KEBS, Nairobi, Kenya, 2018.

1953 236. E Stronger, EAGC, http://eagc.org/wp-content/uploads/2018/01/EAGC_@10_Milestone.pdf.

1954 237. J. W. Muthomi, B. K. Mureithi, G. N. Chemining, J. K. Gathumbi, and E. W. Mutitu, "Aspergillus and
1955 Aflatoxin B1 contamination of Maize and Maize Products from Eastern and North Rift Regions of
1956 Kenya," Proceedings of the 12th KARI Biennial Conference, 8th November 2010, Nairobi, Kenya, p.
1957 344-352, 2010.

1958 238. J. H. Lewis, L. W. Redwood, Y. A. Kieszak et al., "Comprehensive assessment of maize aflatoxin
1959 levels in eastern Kenya," Environmental Health Perspectives, vol. 119, no. 12, pp. 1795, 2011.

1960 239. J. Onsongo, "Outbreak of aflatoxin poisoning in Kenya," IDS Bulletin, vol. 5, pp. 3-4, 2004.

1961 240. H. Njapau, and P. J. C. Probst, "Outbreak of an acute aflatoxicosis in Kenya in 2004: Identification of
1962 the causal agent," Applied Environmental Microbiology, vol. 73, pp. 2762-2764, 2007.

- 1963 241. International Maize and Wheat Improvement Center (CIMMYT), United Nations Development Programme (UNDP), and the United States Agency for International Development (USAID). 1986. Aflatoxins in Maize. A Proceedings of the Workshop El Batan, Mexico, 7–11 April. CIMMYT, Mexico City, Mexico.
- 1964
- 1965
- 1966
- 1967 242. Food and Agriculture Organization of the United Nations (FAO). Aflatoxin in Kenya. FAO, Rome, Italy, 1988, <http://agris.fao.org/aos/records/QY870004888>
- 1968
- 1969 243. N. Muraguri, L. C. Omulkoola, G. M. Kenji, and G. A. Condier, “A survey of mycotoxins in human and animal foods: Part 1,” *East African Medical Journal*, vol. 58, no. 7, pp. 484-488, 1981.
- 1970
- 1971 244. C. K. Mutegi, “The extent of aflatoxin and *Aspergillus* section *Flavi*, *Penicillium* spp and *Rhizopus* spp contamination of peanuts from households in Western Kenya and the causative factors of contamination,” PhD Thesis, University of KwaZulu, Natal, South Africa, 120p., 2010.
- 1972
- 1973
- 1974 245. A. P. Wacoo, I. Mukisa, R. Meeme et al., “Probiotic enrichment and reduction of aflatoxins in a traditional African maizebased fermented food,” *Nutrients*, vol. 11, no. 2, pp. 265, 2019.
- 1975
- 1976 246. S. Byakika, I. M. Mukisa, A. P. Wacoo et al., “Potential application of lactic acid starters in the reduction of aflatoxin contamination in fermented sorghum-millet beverages,” *International Journal of Food Contamination*, vol. 6, no. 4, 2019.
- 1977
- 1978
- 1979 247. R. L. Brown, D. Bhatnagar, T. E. Cleveland, Z. Chen, and A. Menkir, “Development of maize host resistance to aflatoxigenic fungi,” *In: Mehdi MR (Eds). Aflatoxins: Recent Advances and Future Prospects*, InTechOpen, Rijeka, Croatia, 2013.
- 1980
- 1981
- 1982 248. L. Ranasinghe, B. Jayawardena, and K. Abeywickrama, “Fungicidal activity of essential oils of *Cinnamomum zeylanicum* (L.) and *Syzygium aromaticum* (L.) Merret L. M. Perry against crown rot and anthracnose pathogens isolated from banana,” *Letters in Applied Microbiology*, vol. 35, pp. 208-211, 2002.
- 1983
- 1984
- 1985
- 1986 249. J. Ssekandi, “Identification of Suitable Genes for RNAi Silencing of Aflatoxigenic *Aspergillus flavus* Isolated from Groundnuts in Uganda,” MSc Thesis, Makerere University, Kampala, Uganda, 66p., 2018.
- 1987
- 1988
- 1989 250. Y. Raruang, O. Omolehin, D. Hu et al., “Host induced gene silencing targeting *Aspergillus flavus* aflM reduced aflatoxin contamination in transgenic maize under field conditions,” *Frontiers in Microbiology*, vol. 11, no. 754, 2020.
- 1990
- 1991
- 1992 251. K. Hell, P. Fandohan, R. Bandyopadhyay et al., “Pre- and post-harvest management of AF in maize,” *In: Leslie JF, Bandyopadhyay R & Visconti A (Eds). Mycotoxins: Detection Methods, Management, Public Health and Agricultural Trade*, CABI Publishing, Wallingford, UK, 2008.
- 1993
- 1994
- 1995 252. P. Fandohan, D. Zoumenou, D. J. Hounhouigan et al., “Fate of aflatoxins and fumonisins during the processing of maize into food products in Benin,” *International Journal of Food Microbiology*, vol. 98, no. 3, pp. 249–259, 2005.
- 1996
- 1997
- 1998 253. D. L. Park, “Effect of processing on aflatoxin,” *In: Mycotoxins and Food Safety. Advances in Experimental Medicine and Biology*, DeVries JW & Jackson LS (Eds), Springer, Boston, MA, USA, vol. 504, 2002.
- 1999
- 2000
- 2001 254. P. Turner, A. Sylla, Y. Gong et al., “Reduction in exposure to carcinogenic aflatoxins by postharvest intervention measures in west Africa: a community-based intervention study,” *The Lancet*, vol. 365, no. 9475, pp. 1950–1956, 2005.
- 2002
- 2003
- 2004 255. C. B. N’dede, C. M. Jolly, S. D. Vodouhe, and P. E. Jolly, “Economic risks of AF contamination in marketing of peanut in Benin,” *Economics Research International*, vol. 2012, Article ID 230638, 12 pages, 2012.
- 2005
- 2006
- 2007 256. L. M. Martins, A. S. Sant’ana, B. T. Iamanaka et al., “Kinetics of aflatoxin degradation during peanut roasting,” *Food Research International*, vol. 97, pp. 1785-183, 2017.
- 2008
- 2009 257. A. O. Makokha, R. Oniang’O, S. M. Njoroge, and O. K. Kamar, “Effect of traditional fermentation and malting on phytic acid and mineral availability from *Sorghum bicolor* and finger millet *Eleusine caracana* grain varieties grown in Kenya,” *Food and Nutritional Bulletin*, vol. 23, no. 3, pp. 241–250, 2002.
- 2010
- 2011
- 2012 258. P. J. Muindi, S. Thomke, and R. Ekman, “Effect of magadi soda treatment on the tannin content and in vitro nutritive value of sorghum grains,” *Journal of Science of Food and Agriculture*, vol. 32, pp. 25-34, 2006.
- 2013
- 2014

- 2015 259. G. Avantaggiato, F. Quaranta, E. Desiderio, and A. Visconti, "Fumonisin contamination of maize
2016 hybrids visibly damaged by *Sesamia*," *Journal of the Science of Food and Agriculture*, vol. 83, no. 1,
2017 pp. 13–18, 2003.
- 2018 260. G. P. Munkvold, "Cultural and genetic approaches to managing mycotoxins in maize," *Annual Review*
2019 *of Phytopathology*, vol. 41, no. 1, pp. 99–116, 2003.
- 2020 261. T. Omara, K. F. Kateeba, B. Musau et al., "Bioinsecticidal activity of eucalyptol and 1R-alpha-pinene
2021 rich acetonc oils of *Eucalyptus saligna* on *Sitophilus zeamais* Motschulsky, 1855 (Coleoptera:
2022 Curculionidae)," *Journal of Health and Environmental Research*, vol. 4, no. 4, pp. 153–160, 2018.
- 2023 262. J. M. Kirima, M. Okuta, and T. Omara, "Chemical composition of essential oils from *Pinus caribaea*
2024 Morelet needles," *French-Ukrainian Journal of Chemistry*, vol. 8, no. 1, pp. 1-7, 2020.
- 2025 263. S. A. Bankole, "Effect of essential oils from two Nigerian medicinal plants (*Azadirachta indica* and
2026 *Morinda lucida*) on growth and aflatoxin B 1 production in maize grain by atoxigenic *Aspergillus*
2027 *flavus*," *Letters in Applied Microbiology*, vol. 24, no. 3, pp. 190–192, 1997.
- 2028 264. G. O. Adegoke, H. Iwahashi, Y. Komatsu, K. Obuchi, and Y. Iwahashi, "Inhibition of food spoilage
2029 yeasts and aflatoxigenic moulds by monoterpenes of the spice *Aframomum danielli*," *Flavour and*
2030 *Fragrance Journal*, vol. 15, no. 3, pp. 147–150, 2000.
- 2031 265. H. K. Abbas, C. Accinelli, and W. T. Shier, "Biological control of aflatoxin contamination in U.S.
2032 crops and the use of bioplastic formulations of *Aspergillus flavus* biocontrol strains to optimize
2033 application strategies," *Journal of Agricultural and Food Chemistry*, vol. 65, no. 33, pp. 7081–7087,
2034 2017.
- 2035 266. J. W. Dorner, "Biological control of aflatoxin contamination in corn using a nontoxigenic strain of
2036 *Aspergillus flavus*," *Journal of Food Protection*, vol. 72, pp. 801–804, 2009.
- 2037 267. J. W. Dorner, "Biological Control of Aflatoxin Contamination," *In: Aflatoxin and Food Safety*, H. K.
2038 Abbas (Eds), New York: Taylor and Francis, pp. 333-352, 2005.
- 2039 268. V. Anjaiah, R. P. Thakur, and N. Koedam, "Evaluation of bacteria and *Trichoderma* for biocontrol of
2040 pre-harvest seed infection by *Aspergillus flavus* in groundnut," *Biocontrol Science and Technology*,
2041 vol. 16, pp. 431-436, 2006.
- 2042 269. G. N. Harini, N. Kumar, B. Hameeda et al., " Biological management of *Aspergillus flavus* infection
2043 and aflatoxin contamination in groundnut by *Streptomyces* sp. CDA 19," *In: Proceedings of the 64th*
2044 *Indian Phytopathological Society Annual Meeting and National Symposium*, Hyderabad, India,
2045 December 2-4, pp. 133, 2011.
- 2046 270. A. N. Kaaya, and W. Kyamuhangire, "The effect of storage time and agroecological zone on mould
2047 incidence and aflatoxin contamination of maize from traders in Uganda," *International Journal of Food*
2048 *Microbiology*, vol. 110, no. 3, pp. 217–223, 2006.
- 2049 271. P. E. Sumner, and D. Lee, "Reducing Aflatoxin in Corn During Harvest and Storage," Atlanta, GA:
2050 The University of Georgia, Georgia College of Agriculture and Environmental Sciences, 2012.
- 2051 272. A. Méndez-Albores, J. Velez-Medina, E. Urbina-Álvarez, F. Martínez-Bustos, and E. Moreno-
2052 Martínez, "Effect of citric acid on aflatoxin degradation and on functional and textural properties of
2053 extruded sorghum," *Animal Feed Science and Technology*, vol. 150, pp. 316-329, 2009.
- 2054 273. B. Kabak and A. D. W. Dobson, "Biological strategies to counteract the effects of mycotoxins,"
2055 *Journal of Food Protection*, vol. 72, no. 9, pp. 2006–2016, 2009.
- 2056 274. T. W. Kensler, N. E. Davidson, P. A. Egner et al., "Mechanisms of Chemoprotection against Aflatoxin-
2057 Induced Hepatocarcinogenesis by Oltipraz," *In: Nygaard O.F., Upton A.C. (eds), Anticarcinogenesis*
2058 *and Radiation Protection 2*. Springer, Boston, MA, vol. 1, pp. 315-322, 1991.
- 2059