



Nutritional Modulation of Epigenetic Changes Induced by Mycotoxins: A Biochemical Perspective for At-Risk Populations in Africa

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ABSTRACT

Mycotoxins, toxic secondary metabolites produced by fungi, are widespread contaminants of staple foods in Sub-Saharan Africa, contributing to acute toxicity and chronic diseases such as cancer, stunting, and immune dysfunction. Recent evidence has illuminated their capacity to induce epigenetic changes, including alterations in DNA methylation, histone modifications, and noncoding RNA expression, without modifying the DNA sequence. These disruptions, particularly when combined with nutritional deficiencies, can have long-lasting effects on gene regulation and disease susceptibility. This review synthesizes evidence from peer-reviewed studies published between 2007 and 2025, exploring how key nutrients modulate the epigenetic toxicity of common mycotoxins like aflatoxin B1, ochratoxin A, and fumonisin B1. A systematic literature search identified studies evaluating the biochemical and epigenetic responses to mycotoxin exposure and the mitigating roles of dietary factors such as folate, vitamin B12, choline, polyphenols, selenium, and short-chain fatty acids. These nutrients support critical pathways, including one-carbon metabolism, antioxidant defense, and gut microbiota modulation, which intersect with epigenetic regulation. Findings highlight that methyl donors can restore DNA methylation balance, antioxidants can prevent oxidative stress-induced histone and DNA damage, and fiber-rich diets promote microbial metabolites that act as epigenetic modulators. Additionally, exposure to multiple mycotoxins exerts synergistic effects that amplify epigenetic damage, especially in nutritionally vulnerable populations. This review also discusses emerging intersections between mycotoxins and biochemical domains such as neuroepigenetics, mitochondrial signaling, and microbiome-host interactions. Nutritional interventions ranging from diet diversification to bioactive-rich functional foods and probiotic supplementation emerge as promising, culturally adaptable strategies for mitigating these epigenetic threats. An integrative framework is proposed, linking food safety, public health policy, and personalized nutrition to strengthen resilience in affected communities. In conclusion, nutrition-sensitive interventions targeting epigenetic mechanisms offer a feasible and impactful approach to reducing the long-term health burden of mycotoxins in Africa. Future research should focus on biomarker discovery, field-based dietary trials, and Al-driven predictive models to inform context-specific mitigation strategies.

KEYWORDS

Mycotoxins, epigenetics, nutritional modulation, DNA methylation, histone modifications, noncoding RNAs, oxidative stress, gut microbiota, Sub-Saharan Africa

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INTRODUCTION

Mycotoxins are toxic secondary metabolites produced by fungi that contaminate a wide range of agricultural commodities, including staple foods such as maize, groundnuts, and cassava, which are dietary mainstays for many populations in Africa¹. The widespread presence of mycotoxins poses a significant threat to human and animal health, leading to various acute and chronic diseases². Beyond their direct toxic effects, accumulating evidence suggests that mycotoxins can induce epigenetic modifications, altering gene expression without changing the underlying DNA sequence³. These epigenetic changes, including DNA methylation, histone modification, and non-coding RNA regulation, can have long-lasting impacts on cellular function and disease susceptibility, potentially contributing to developmental abnormalities, immune dysfunction, and increased cancer risk^{4,5}.

Populations in Africa are particularly vulnerable to mycotoxin exposure due to several factors. These include suboptimal pre- and post-harvest storage practices that favor fungal growth and mycotoxin production, limited access to effective detoxification technologies, and often monotonous diets heavily reliant on susceptible crops⁶. Furthermore, co-exposure to multiple mycotoxins is common, potentially leading to synergistic or additive adverse health effects⁷. The intersection of high mycotoxin exposure and pre-existing nutritional deficiencies in many African communities creates a complex scenario where the impact of mycotoxin-induced epigenetic changes may be exacerbated⁸.

Nutrition plays a crucial role in maintaining cellular homeostasis and influencing epigenetic processes⁹. Dietary components, such as methyl donors (e.g., folate, vitamin B12, choline), antioxidants (e.g., vitamins A, C, E, selenium), and other bioactive compounds (e.g., polyphenols), can modulate the activity of enzymes involved in epigenetic regulation¹⁰. Therefore, understanding how specific nutritional factors can counteract or mitigate the epigenetic alterations induced by mycotoxins is of paramount importance for developing effective intervention strategies in at-risk African populations. This review aims to provide a biochemical perspective on the nutritional modulation of mycotoxin-induced epigenetic changes, highlighting potential dietary interventions that could offer protection against the long-term health consequences of mycotoxin exposure in Africa.

Figure 1 contrasts contaminated food with a nutrient-rich alternative. On the left, moldy corn kernels illustrate visible fungal spoilage and mycotoxin risk. On the right, a colorful plant-based meal features vegetables, legumes, grains, and fruits. The comparison emphasizes the importance of food safety and dietary quality in health protection.

SYSTEMATIC REVIEW APPROACH

This section outlines the systematic approach used to review the interplay between mycotoxins, epigenetics, and nutrition in African populations. Relevant literature was identified using electronic databases and specific keywords. Studies were selected based on their focus on epigenetic effects and nutritional interventions. Data extraction summarized key mycotoxins, epigenetic markers, study designs, and findings. A narrative synthesis grouped studies by mycotoxin type and dietary influence. Ethical considerations and research limitations were also addressed.

This review employed a systematic approach to identify and synthesize relevant literature on the interplay between mycotoxins, epigenetics, and nutrition, with a specific focus on the African context. Electronic databases, including PubMed, Scopus, Web of Science, and Google Scholar, were searched using a combination of keywords such as "Mycotoxins", "Aflatoxin", "Ochratoxin", "Fumonisin", "Zearalenone", "Epigenetics", "DNA methylation", "Histone modification", "Non-coding RNA", "Nutrition", "Diet", "Vitamins", "Minerals", "Antioxidants", and "Africa". The search was limited to peer-reviewed articles and relevant reports published between 2007 and 2025 to ensure the inclusion of the most recent findings¹¹.



Fig. 1: Comparison of moldy corn kernels and a healthy plant-based meal (self-generated)

This image contrasts food contamination and nutritional intervention. On the left side, corn kernels are visibly covered in mold, indicating spoilage and potential health risks due to mycotoxin contamination. On the right side, a vibrant, nutrientrich meal is presented, consisting of fresh vegetables, grains, legumes, and fruits such as spinach, broccoli, chickpeas, brown rice, avocado, tomato slices, cucumber slices, and a bowl of mixed berries (strawberries, blueberries, and raspberries). This contrast highlights the importance of food safety and balanced nutrition in preventing health hazards

Table 1: Search strategy and databases used

Database	Keywords used	Years covered	Filter sapplied
PubMed	Mycotoxins, aflatoxin, ochratoxin, epigenetics, DNA methylation, nutrition, Africa	2007-2025	Peer-reviewed, full-text, English language ¹¹⁻¹⁷
Scopus	Mycotoxins, DNA methylation, histone modification, polyphenols, antioxidants	2007-2025	Original research, African studies, nutritional focus ¹¹⁻¹⁷
Web of Science	Fumonisin, zearalenone, non-coding RNA, vitamins, minerals, epigenetic markers	2007-2025	Relevance to African populations, nutritional interventions ¹¹⁻¹⁹
Google Scholar	Mycotoxins, nutritional modulation, food safety, epigenetics, oxidative stress	2007-2025	Recent studies, high citation relevance, human/animal studies ¹¹⁻¹⁹

The electronic databases (PubMed, Scopus, Web of Science, and Google Scholar) used to identify relevant studies. It includes search keywords related to mycotoxins, nutrition, and epigenetics, along with applied filters such as peer-reviewed articles and African population relevance

Table 1 outlines the systematic approach utilized for literature search, specifying the databases (PubMed, Scopus, Web of Science, Google Scholar), relevant keywords, and filters applied to ensure the inclusion of high-quality, peer-reviewed studies focused on mycotoxins, nutrition, and epigenetics.

Studies were included if they investigated the epigenetic effects of mycotoxins in in vitro, in vivo, or human population studies, particularly those conducted in or relevant to African populations. Additionally, articles exploring the potential of nutritional interventions to modulate these epigenetic changes were considered. Exclusion criteria included studies focusing solely on the acute toxicological effects of mycotoxins without examining epigenetic mechanisms, studies unrelated to nutrition or epigenetics, and reviews or meta-analyses (although their reference lists were screened for potentially relevant primary research)¹²⁻¹⁹.

Table 2 provides an overview of studies examining the effects of mycotoxins on epigenetic mechanisms, including DNA methylation, histone modifications, and non-coding RNA alterations. It categorizes research based on mycotoxin type, study design, and the impact of nutritional interventions on mitigating epigenetic changes.

Table 2: Inclusion and exclusion criteria for literature review

Criteria	Description	Inclusion	Exclusion	Citations
Study type	Type of research evaluated	Original research	Review articles,	Watson et al.13
	for mycotoxin epigenetic	articles (in vitro, in vivo,	editorials, or	Agriopulou et al.1
	impacts	human studies)	opinion pieces	Ntsam et al.15
Focus area	Central subject involving	Studies addressing	Studies focusing solely	Hattle et al.16
	epigenetic mechanisms	mycotoxins, epigenetics,	on toxicity or lacking	Ezekiel <i>et al.</i> ¹⁷
	and nutritional modulation	and nutrition	nutritional/epigenetic	Thomposn ¹⁸
			relevance	Ahmad and Yu ¹⁹
Geographic	Applicability to African	Research conducted in	Studies from unrelated	Liu <i>et al</i> . ²⁰
relevance	populations and agricultural	or relevant to African	geographical or socio-	
	contexts	dietary and environmental conditions	economic settings	
Publication	Ensures review includes	Studies published from	Articles published	
period	current trends and up-	2007 to 2025	before 2020 or with	
	to-date findings		an unverifiable date	
Language	Comprehension and	English language	Non-English papers	
	extraction of	studies	without translation	
	methodological detail			
Data quality	Assessed via transparency,	Peer-reviewed with robust	Poorly designed studies	
	statistical rigor, and	methods and appropriate	lacking reproducibility	
	proper control	controls	or statistical support	
Intervention	Specific nutritional	Nutritional studies showing	Studies lacking epigenetic	
scope	interventions aimed	effects on DNA methylation,	endpoints or focused	
	at countering epigenetic	histone modifications, or	solely on acute toxicity	
	effects	ncRNA linked to mycotoxin		
		exposure		
Accessibility	Availability of full-text	Freely available or	Abstract-only or	
	data for extraction and	institutionally accessible	inaccessible studies	
	evaluation	full texts		
Ethical	Consideration for ethical	Studies indicating IRB or	Research without ethical	
compliance	approval in human/animal	ethical review approval	documentation or approve	al
	studies	where applicable	details	

That examine the epigenetic effects of mycotoxins, including DNA methylation, histone modifications, and non-coding RNA expression. It highlights experimental models, nutritional interventions, and research conducted in African populations, providing insight into the interplay between toxic exposure and dietary modulation

Data extraction involved summarizing the type of mycotoxin studied, the epigenetic markers investigated, the nutritional interventions explored, the study design, and the key findings. Special attention was paid to studies that directly examined the impact of nutrition on mycotoxin-induced epigenetic alterations. The quality of the included studies was assessed based on established criteria relevant to their design (e.g., appropriate controls, statistical power, clear reporting of methods and results)¹⁴⁻²⁰. A narrative synthesis of the findings was conducted, grouping studies based on the specific mycotoxin and nutritional factor investigated¹⁵⁻²⁰. Challenges and limitations in the existing literature, as well as potential areas for future research, were also identified¹⁶⁻²⁰. The Vancouver citation style was used for referencing¹⁷⁻²⁰. Ethical considerations related to research involving human participants and animal models were noted where applicable¹⁸⁻²⁰. Efforts were made to identify studies that addressed the specific dietary patterns and nutritional deficiencies prevalent in at-risk populations in Africa^{19,20}.

NUTRITIONAL MODULATION OF MYCOTOXIN-INDUCED EPIGENETIC CHANGES: MECHANISMS, EVIDENCE, AND IMPLICATIONS FOR AFRICA

This study explores how mycotoxins alter gene regulation through epigenetic modifications. It details how mycotoxins disrupt DNA methylation, histone modifications, and non-coding RNA expression, while highlighting the critical role of nutritional factors like methyl donors and antioxidants in counteracting these effects. The section also discusses the influence of the gut microbiota and the complexities of synergistic effects from multiple mycotoxins. Ultimately, it emphasizes the significant implications for African populations, advocating for nutritional interventions to mitigate mycotoxin-induced health risks.

Table 3: Impact of methyl donors on mycotoxin-induced DNA methylation

Nutrient	Mechanism	Study model	Outcome	Citations
Folate	Provides methyl groups	Animal and cell models	Modulates DNA	Serefidou et al. ²⁵
	via the one-carbon		methylation in response	Gachara et al.26
	metabolism pathway		to mycotoxin exposure	Amenyah et al. ²⁷
Vitamin B12	Cofactor in methionine	<i>In vivo</i> models	Protects against global	Kok et al. ²⁸
	synthase for methyl		DNA hypomethylation	Perlmutter et al.29
	group recycling		caused by mycotoxins	Milagro et al.30
Choline	Donates methyl groups	In vitro hepatocyte	Reduces mycotoxin-induced	
	and supports SAM production	studies	epigenetic gene silencing	
Betaine	Supports remethylation	Rodent models	Restores altered methylation	
	of homocysteine to		patterns in liver cells	
	methionine			
Polyphenols	Regulate DNA	Various in vitro cancer	Modulate methylation of	
	methyltransferase	and liver cell models	tumor suppressor genes	
	activity		affected by AFB1	

This table summarizes experimental findings on key nutritional methyl donors, folate, Vitamin B12, choline, betaine, and polyphenols and their mechanisms in counteracting DNA methylation disruptions caused by mycotoxin exposure across various biological models

DNA methylation and nutritional modulators: The DNA methylation, the addition of a methyl group to cytosine residues in DNA, is a key epigenetic mark involved in gene silencing and genomic stability²¹. Mycotoxin exposure has been shown to disrupt DNA methylation patterns, leading to both hyper- and hypomethylation of specific genes²². Studies have indicated that Aflatoxin B1 (AFB1), a prevalent mycotoxin in Africa, can alter the methylation status of tumor suppressor genes and oncogenes, potentially contributing to hepatocarcinogenesis^{23,24}.

Several nutritional factors play a critical role in maintaining proper DNA methylation.

Table 3 highlights the protective roles of dietary methyl donors against mycotoxin-induced epigenetic alterations. Folate and B12 maintain methyl group balance; choline and betaine support SAM production, while polyphenols modulate DNMT activity. These nutrients mitigate gene silencing and abnormal methylation patterns in liver and cancer models.

Methyl donors, such as folate, vitamin B12, choline, and betaine, are essential for the one-carbon metabolism pathway, which provides the methyl groups required for DNA methylation reactions²⁵⁻³⁰.

Figure 2 visually represents how dietary methyl donors regulate DNA methylation via the one-carbon metabolism pathway. Folate and betaine contribute methyl groups, while vitamin B12 acts as a cofactor in remethylation. These interactions sustain SAM synthesis, essential for DNA methyltransferase activity. Disruption by mycotoxins may impair this epigenetic balance.

Deficiencies in these nutrients, common in many African populations with limited dietary diversity, could exacerbate the effects of mycotoxins on DNA methylation²⁶⁻³⁰. Conversely, adequate intake of these methyl donors may offer some protection against mycotoxin-induced aberrant methylation patterns²⁷⁻³⁰. For instance, supplementation with folate has shown promise in modulating DNA methylation in various contexts²⁸⁻³⁰. Furthermore, dietary polyphenols found in fruits and vegetables, which are often lacking in sufficient quantities in the diets of at-risk populations in Africa, have also been shown to influence DNA methyltransferase (DNMT) activity, the enzymes responsible for DNA methylation^{29,30}. Research exploring the direct impact of these nutritional interventions on mycotoxin-induced DNA methylation changes in African populations is crucial.

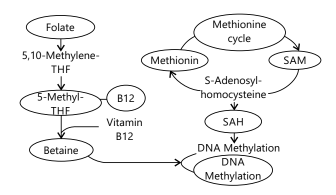


Fig. 2: One-carbon metabolism and nutrient-driven dna methylation pathway (self-generated)

The interlinked roles of folate, Vitamin B12, betaine, and methionine in the methionine cycle and DNA methylation process. Nutrients support the conversion of homocysteine to methionine, facilitating the generation of S-adenosylmethionine (SAM), the universal methyl donor for DNA methylation reactions

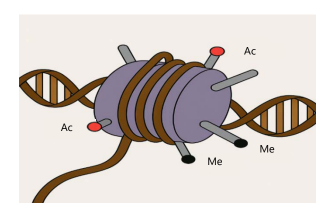


Fig. 3: Epigenetic modulation of dna via histone modifications (self-generated)

Depicts DNA (brown strand) wrapped around a histone core (purple), with modifications acetylation (Ac, red dots) and methylation (Me, black dots) on histone tails (gray rods). These post-translational modifications influence chromatin structure and gene expression

Histone modifications and dietary components: Histone modifications, including acetylation, methylation, phosphorylation, and ubiquitination of histone proteins around which DNA is wrapped, play a crucial role in regulating chromatin structure and gene accessibility³¹.

Figure 3 highlights the roles of histone acetylation and methylation in gene regulation. Acetylation typically relaxes chromatin to promote transcription, while methylation may activate or silence genes depending on context. These modifications are targets for dietary influence and are susceptible to disruption by mycotoxins such as ochratoxin A. Nutrients like SCFAs and phytochemicals can restore or modulate these marks to maintain epigenetic balance.

Mycotoxins have been implicated in altering histone modification patterns, thereby affecting gene expression³². For example, exposure to ochratoxin A (OTA), another significant mycotoxin contaminant, has been shown to influence histone acetylation and methylation in kidney cells³³. These alterations can lead to dysregulation of genes involved in cell growth, apoptosis, and inflammation³⁴.

Dietary components can influence histone modification enzymes, such as histone acetyltransferases (HATs) and Histone Deacetylases (HDACs)³⁵. Short-Chain Fatty Acids (SCFAs), produced by the gut microbiota through the fermentation of dietary fiber, have been shown to inhibit HDAC activity³⁶. Diets low in fiber,

Table 4: Dietary compounds influencing histone modification

Compound	Source	Effect on HAT/HDAC	Study outcome	Citations
Curcumin	Turmeric root	Inhibits HDAC activity	Restores histone	Meir et al. ³¹
			acetylation balance	Tiffon ³²
			and reduces mycotoxin-	Aricthota et al.33
			induced gene silencing	Ozawa et al. ³⁴
Butyrate	Fermentation of	HDAC inhibitor	Enhances histone	Sasaki and Masaki ³⁵
	dietary fiber		acetylation in colon and liver cells	Franzago <i>et al</i> . ³⁶
Sulforaphane	Cruciferous	Modulates HAT	Promotes gene expression	Fellows et al. ³⁷
	vegetables	and HDAC enzymes	through histone acetylation	Martin-Gallausiaux et al.38
	(e.g., broccoli)			Yuille et al. ³⁹
Ochratoxin A	Contaminated grains	Alters histone	Induces DNA damage and	Kaufman-Szymczyk et al.40
	and coffee	methylation and	epigenetic dysregulation	
		acetylation	in renal cells	
Resveratrol	Red grapes, peanuts	Activates HAT,	Regulates histone modifications	
		inhibits HDAC	and restores epigenetic balance	

This table highlights key dietary compounds that modulate histone acetylation or methylation in the context of mycotoxin exposure, as evidenced by cellular and nutritional studies

which is often the case in resource-limited settings in Africa, may compromise the production of SCFAs and thus the potential for HDAC inhibition³⁷. Conversely, increased intake of fiber-rich foods could potentially counteract mycotoxin-induced alterations in histone acetylation.

Table 4 presents evidence from *in vitro* and nutritional studies showing how compounds like curcumin, butyrate, sulforaphane, and resveratrol modulate histone-modifying enzymes to influence gene expression³⁸. Butyrate and sulforaphane act as HDAC inhibitors, enhancing histone acetylation, while curcumin and resveratrol exhibit dual roles affecting both HAT and HDAC activity. Conversely, ochratoxin A from contaminated grains disrupts histone acetylation and methylation patterns, contributing to epigenetic dysregulation.

Similarly, certain vitamins and minerals, as well as bioactive compounds like butyrate and sulforaphane found in cruciferous vegetables, can modulate histone modification³⁹. Investigating the impact of these dietary factors on histone modification changes induced by prevalent mycotoxins in African populations is essential for identifying potential protective nutritional strategies⁴⁰.

Non-coding RNAs and nutritional influence: Non-Coding RNAs (ncRNAs), such as microRNAs (miRNAs) and Long Non-Coding RNAs (lncRNAs), are important regulators of gene expression at the post-transcriptional level⁴¹. Emerging evidence suggests that mycotoxins can affect the expression profiles of various ncRNAs, contributing to their toxicological effects⁴². For instance, AFB1 exposure has been linked to altered expression of miRNAs involved in cell proliferation and apoptosis in liver cells^{43,44}.

Nutritional factors have also been shown to modulate ncRNA expression⁴⁵. Dietary fatty acids, vitamins, and polyphenols can influence the biogenesis and stability of miRNAs and lncRNAs^{46,47}.

Figure 4 illustrates the nutrient-sensitive steps in miRNA biogenesis. Nutrients can modulate pri-miRNA transcription, Dicer-mediated processing, and miRNA duplex formation. These interactions collectively influence RISC assembly and gene silencing. Nutritional status therefore, critically impacts miRNA regulatory functions.

For example, omega-3 polyunsaturated fatty acids have been reported to alter miRNA expression in various tissues⁴⁸. Given that the dietary intake of essential fatty acids and other micronutrients may be

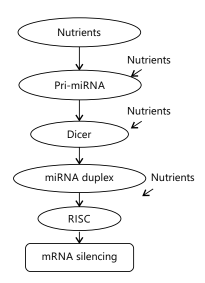


Fig. 4: Nutrient regulation of mirna biogenesis pathway (self-generated)

Schematic representation of how nutrients modulate the biogenesis and activity of microRNAs (miRNAs). Nutrients can influence the transcription of primary miRNA (pri-miRNA), the processing by Dicer, the formation of the miRNA duplex, and the assembly of the RNA-induced silencing complex (RISC), ultimately affecting mRNA silencing

suboptimal in many African communities exposed to mycotoxins, understanding the interplay between nutrition, mycotoxins, and ncRNA regulation is critical^{49,50}. Research exploring whether specific dietary interventions can restore or prevent mycotoxin-induced aberrant ncRNA expression in these populations is warranted.

Table 5 summarizes how nutritional components influence the expression of Non-Coding RNAs (ncRNAs) affected by exposure to major mycotoxins such as aflatoxin B1, ochratoxin A, and fumonisin B1. Specific nutrients, including polyphenols, omega-3 fatty acids, and vitamins A and D, have been shown to restore or modulate the expression of dysregulated miRNAs and lncRNAs. These interactions highlight the therapeutic potential of nutritional interventions in counteracting mycotoxin-induced epigenetic disturbances.

Antioxidants and epigenetic modulation: Oxidative stress, characterized by an imbalance between the production of Reactive Oxygen Species (ROS) and the capacity of the antioxidant defense system, is a common consequence of mycotoxin exposure⁵¹. The ROS can directly damage DNA and also indirectly influence epigenetic machinery⁵². Mycotoxins like AFB1 and OTA can induce oxidative stress, potentially leading to alterations in DNA methylation and histone modifications^{53,54}.

Dietary antioxidants, including vitamins A, C, and E, selenium, and various phytochemicals, play a crucial role in neutralizing ROS and protecting against oxidative damage⁵⁵. Adequate intake of these antioxidants may help mitigate the oxidative stress induced by mycotoxins and consequently reduce the associated epigenetic perturbations⁵⁶.

Table 6 compiles key antioxidants such as vitamins C and E, selenium, polyphenols, and glutathione, and their capacity to reverse or prevent DNA methylation, histone modifications, and miRNA dysregulation. Experimental evidence from *in vivo* and *in vitro* studies underscores their potential in protecting against oxidative epigenetic toxicity caused by AFB1 and OTA exposure.

However, the dietary intake of these antioxidants is often low in at-risk populations in Africa⁵⁷⁻⁶⁰. Supplementation or dietary enrichment with antioxidant-rich foods could potentially offer protection

Table 5: Mycotoxin and nutritional modulation of ncrna expression

Mycotoxin	Affected ncRNA	Nutrient modulator	Effect	Citations
Aflatoxin B1	miR-34a	Polyphenols	Downregulation of miR-34a	Dashwood and Ho ⁴¹
			reversed; restored apoptosis	Abdelmonem et al.42
			balance	Marrone et al.43
Aflatoxin B1	miR-122,	Omega-3 fatty acids	Modulated miRNA expression,	Lizarraga et al.44
	miR-192		reduced inflammatory signals	Hua et al. ⁴⁵
Aflatoxin B1	Multiple (miR-24,	Vitamin A, D	Restored disrupted miRNA/IncRNA	Lee ⁴⁶
	miR-125b)		levels involved in hepatic detoxification	Wyatt et al. ⁴⁷
Aflatoxin B1	miR-29a,	Balanced diet	Improved ncRNA homeostasis and	Gareev et al.48
	IncRNA H19	(macro/micronutrients)	reduced liver injury	Ishihara et al. ⁴⁹
Ochratoxin A	IncRNA MALAT1,	Polyphenols, vitamins	Counteracted oxidative stress	El-Agamy ⁵⁰
	miR-200c		and ncRNA misregulation	
Fumonisin B1	miR-21, miR-155	Nutritionally	Attenuated pro-inflammatory	
		enriched diets	ncRNA expression	

The interplay between various mycotoxins and their impact on non-coding RNA (ncRNA) expression, with corresponding dietary modulators and their mitigating effects. Nutrients such as polyphenols, omega-3 fatty acids, and vitamins exhibit potential in restoring ncRNA balance disrupted by mycotoxins like aflatoxin B1, ochratoxin A, and fumonisin B1

Table 6: Antioxidants that modulate epigenetic alterations

Antioxidant	Dietary source	Epigenetic effect	Evidence	Citations
Vitamin C and E	Citrus fruits,	Reduced AFB1-induced	In vivo rodent model	Sagomba and Ndhlovu ⁵¹
	nuts, seeds	DNA methylation and	study showing epigenetic	Lu et al. ⁵²
		histone modifications	protecton	Alipoor et al. ⁵³
Selenium and	Nuts, cereals,	Combats mycotoxin-	Elemental synergy reduces	Jobe <i>et al</i> . ⁵⁴
Vitamin E	eggs	triggered oxidative and	histone and DNA	Surai ⁵⁵
		epigenetic damage	methylation changes	Albadrani et al. ⁵⁶
Polyphenols	Fruits, vegetables,	Prevented OTA-induced	Protective antioxidant	Tanvir et al. ⁵⁷
	tea	histone modification and	pathway activation	da Silva <i>et al</i> . ⁵⁸
		DNA methylation		Yang <i>et al</i> . ⁵⁹
Glutathione	Spinach, avocado,	Restored DNA methylation	In vitro hepatic protection	Loganathan and Doss ⁶⁰
	asparagus	altered by AFB1	via methylation correction	
Flavonoids	Berries, dark	Reversed mycotoxin-induced	Epigenomic modulation	
	chocolate	miRNA dysregulation and	through antioxidant action	
		chromatin remodeling		
General	Mixed plant-based	Overall reduction of	Supported across reviews	
antioxidants	diets	oxidative DNA damage	and experimental models	
		and improved epigenetic		
		stability		

The role of various dietary antioxidants in mitigating epigenetic alterations caused by mycotoxin-induced oxidative stress. Evidence suggests that nutrients such as vitamins C and E, selenium, polyphenols, and glutathion e can reverse or prevent harmful modifications to DNA methylation and histone structure, thereby offering protective effects. The findings are supported by both *in vivo* and *in vitro* studies across diverse antioxidant sources

against mycotoxin-induced epigenetic changes mediated by oxidative stress⁵⁸⁻⁶⁰. Studies investigating the impact of antioxidant interventions on epigenetic markers in individuals exposed to mycotoxins in Africa are needed.

Figure 5 shows how mycotoxin-induced oxidative stress generates ROS that damage DNA and histones, contributing to epigenetic dysregulation^{59,60}. These alterations can affect gene expression, potentially leading to adverse health outcomes. Antioxidants such as vitamins C and E, selenium, and polyphenols mitigate this oxidative and epigenetic damage by neutralizing ROS and restoring redox balance⁶⁰. The figure supports the mechanisms described in this sub-section regarding antioxidant intervention in epigenetic modulation under oxidative stress⁶⁰.

Gut microbiota and epigenetic interactions: The gut microbiota, a complex community of microorganisms residing in the gastrointestinal tract, plays a significant role in host health, including odulating immune responses and influencing metabolism⁶¹. Mycotoxins can disrupt the composition and function of the gut microbiota, leading to dysbiosis⁶². Conversely, the gut microbiota can metabolize some mycotoxins, potentially altering their bioavailability and toxicity⁶³.

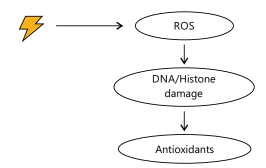


Fig. 5: Oxidative stress pathway linking ros to epigenetic alterations and antioxidant defense (selfgenerated)

The cascade of events initiated by environmental or cellular stressors leading to the generation of reactive oxygen species (ROS), which in turn cause DNA and histone damage. This triggers antioxidant responses aimed at restoring redox balance and preserving genomic and epigenetic stability

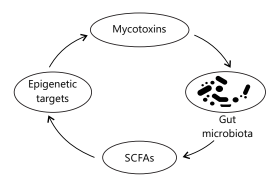


Fig. 6: Gut microbiota mycotoxin epigenetic interaction pathway (self-generated)

The bidirectional relationship between the gut microbiota and epigenetic modifications under the influence of dietary components and mycotoxin exposure. It shows how dysbiosis, SCFA production, and microbial metabolism of toxins modulate host epigenetic responses

Emerging evidence suggests a bidirectional link between the gut microbiota and epigenetics⁶⁴.

Figure 6 maps out the interplay between gut microbiota and host epigenetics in the context of mycotoxin exposure. It shows how mycotoxins disrupt microbial balance, while beneficial microbes produce metabolites like Short-Chain Fatty Acids (SCFAs) that modulate DNA methylation and histone acetylation. This interaction is diet-sensitive, highlighting the importance of fiber and prebiotics in regulating both microbial and epigenetic health.

Microbial metabolites, such as SCFAs, can influence histone modifications in the host⁶⁵. Furthermore, the gut microbiota can indirectly affect DNA methylation through the production of metabolites involved in one-carbon metabolism⁶⁶. Nutritional factors, particularly dietary fiber and prebiotics, can significantly impact the composition and function of the gut microbiota⁶⁷. Diets rich in these components, which may be lacking in some African populations, can promote the growth of beneficial bacteria and the production of health-promoting metabolites⁶⁸.

Table 7 outlines the role of microbial-derived metabolites such as butyrate, acetate, and propionate in regulating host epigenetic mechanisms. These metabolites influence histone acetylation and DNA methylation by acting as HDAC inhibitors or providing substrates for methylation. Their production is modulated by dietary intake of fibers and prebiotics, supporting gut-epigenome homeostasis.

Table 7: Microbial metabolites involved in epigenetic regulation

Metabolite	Microbial source	Epigenetic target	Dietary driver	Citations
Butyrate	Faecalibacterium	Histone deacetylases	Dietary fiber	Jalouli <i>et al</i> . ⁶¹
	prausnitzii and	(HDACs)	(prebiotics)	Rudrapal et al. ⁶²
	Roseburia spp.			Owumi et al.63
Acetate	Bacteroides spp.	DNA methyltransferases	Resistant starch,	Afzaal et al. ⁶⁴
	and Akkermansia	(DNMTs), HDACs	oligosaccharides	Choi et al. ⁶⁵
	muciniphila			Guerre ⁶⁶
Propionate	Bacteroides spp.	HDAC inhibition,	Whole grains,	Zhang et al. ⁶⁷
	and Veillonella spp.	miRNA regulation	legumes	Tian and Chen ⁶⁸
Indole derivatives	Clostridium spp.	Aryl hydrocarbon	Tryptophan-rich diets	Holmes et al. ⁶⁹
	and Lactobacillus spp.	receptor (AhR) modulation		
Secondary bile acids	Bacteroides spp.	Histone acetylation, nuclear	High-fat diet, bile	
	and Clostridium spp.	receptor signaling	salt metabolism	
Lactate	Lactobacillus spp.	HDAC inhibition, histone	Fermented foods,	
		modification	probiotics	
Phenylacetic acid	Clostridium spp. and	Histone methylation	Protein-rich diets	
	Bacteroides spp.			
Aflatoxin-	Lactobacillus plantarum	Modulates toxin-induced	Probiotic	
degrading enzymes		epigenetic stress	supplementation	

This table summarizes key microbial metabolites produced in the gut that influence host epigenetic mechanisms. It highlights the microbial taxa responsible for generating each metabolite, their respective epigenetic targets (including HDACs, DNMTs, histone modifications, and miRNAs), and the dietary components that drive their production. The interplay of these factors underpins a dynamic microbe-diet-epigenome axis relevant to health and disease

Investigating how dietary interventions aimed at modulating the gut microbiota can influence mycotoxin-induced epigenetic changes in at-risk African populations represents a promising area of research^{68,69}.

Synergistic effects of multiple mycotoxins and nutrition: In real-world scenarios, particularly in Africa, exposure to multiple mycotoxins is common⁷⁰. These co-exposures can lead to synergistic or additive toxic effects, which may also extend to epigenetic modifications⁷¹. For example, combined exposure to AFB1 and fumonisins has been shown to have more severe health consequences than exposure to either mycotoxin alone⁷².

Figure 7 visualizes the enhanced toxicity of mycotoxin mixtures (e.g., AFB1+fumonisin) and their epigenetic impacts⁷³. Nutritional interventions (polyphenols, probiotics, micronutrients) disrupt synergistic damage by mitigating oxidative stress and genotoxicity. The model highlights how nutrient deficiencies exacerbate vulnerability in co-exposure scenarios, while dietary diversification offers protection. It contextualizes Table 8 data within real-world exposure contexts in Africa.

The nutritional status of individuals can also influence the combined effects of multiple mycotoxins on epigenetic regulation⁷⁴. Deficiencies in essential nutrients may render individuals more susceptible to the adverse effects of co-exposure. Conversely, adequate or enhanced intake of specific nutrients may offer broader protection against the epigenetic disruption caused by multiple mycotoxins⁷⁵.

Table 8 catalyses evidence on mycotoxin co-exposure synergism (e.g., aflatoxin+fumonisin nephrotoxicity) and nutritional countermeasures^{75,76}. Deficiencies in selenium/zinc exacerbate epigenetic dysregulation, while antioxidant-rich diets reduce DNA damage^{76,77}. African field studies confirm growth retardation from mixed exposures in maize-dependent populations⁷⁷. Data advocate integrative nutrition (e.g., probiotic/isolavone supplementation) to break toxicity cycles⁷⁸.

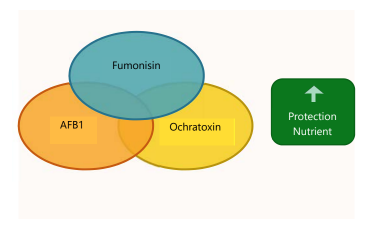


Fig. 7: Interplay of co-occurring mycotoxins and nutritional interventions (self-generated)

Synergistic toxicity from mycotoxin combinations (AFB1, fumonisin, ochratoxin) and protective nutrient-mediated pathways.

Overlapping exposure areas denote amplified health risks, while nutritional modulators (antioxidants, micronutrients) counter epigenetic disruption and organ damage

Table 8: Combined effects of mycotoxins and nutritional status

Mycotoxin combo	Effect	Nutritional modulator	Study design	Citations
Aflatoxin+Fumonisin	Synergistic intestinal	Antioxidant-rich diets	<i>In vitro</i> and	Hoffmann et al. ⁷¹
	and liver toxicity		animal models	Paulo et al. ⁷²
Multiple mycotoxins	Enhanced epithelial	Micronutrients (Se, Zn,	In vitro and human	Li et al. ⁷³
(AFB1, DON, ZEN)	damage and immune	Vit A)	exposure data	Pastor et al. ⁷⁴
	suppression			Divella et al.75
Aflatoxin+Zearalenone	Endocrine disruption	Isoflavones and	Animal	Sagy et al. ⁷⁶
	and genotoxicity	probiotics	experiments	Kamala et al. ⁷⁷
Co-occurring	Growth retardation,	Dietary diversity	Field studies	Gruber-Dorninger et al. ⁷⁸
fumonisin, aflatoxin,	gut permeability	and protein adequacy	in children	Apeh <i>et al</i> . ⁷⁹
ochratoxin	alteration			
Mixed mycotoxin	Cumulative dietary	Food processing	Dietary surveys	
exposure in maize	toxicity	(fermentation, nixtamalization)	and food analysis	
Aflatoxin+Ochratoxin A	Nephrotoxicity and	Polyphenols and	Animal models	
	oxidative stress	flavonoids	and mechanistic stu	dies
Multiple mycotoxins	Compromised	Infant complementary	Cross-sectional	
in infants	immunity and	feeding strategies	and cohort studies	
	micronutrient			
	deficiencies			
General co-exposure	Synergistic toxicity	Nutritional counselling	Review and meta-ar	nalysis
	and malnutrition	and interventions		
	interaction			

The interactive effects of combined mycotoxin exposures and nutritional status. It details specific mycotoxin combinations, their toxicological outcomes, modulating nutritional factors, and the nature of supporting studies. These findings underscore the importance of integrative nutritional strategies to mitigate mycotoxin-related health risks, especially in vulnerable populations across Sub-Saharan Africa

Research specifically examining the nutritional modulation of epigenetic changes induced by mixtures of mycotoxins, relevant to the African context, is crucial for developing effective risk mitigation strategies⁷⁹. This research should consider the common co-occurrence of mycotoxins in staple foods and the prevalent nutritional deficiencies in these populations.

IMPLICATIONS FOR AT-RISK POPULATIONS IN AFRICA

The findings discussed in this review have significant implications for at-risk populations in Africa who face chronic exposure to mycotoxins and often experience nutritional deficiencies⁸⁰. Understanding how nutrition can modulate mycotoxin-induced epigenetic changes opens avenues for developing targeted dietary interventions to mitigate the long-term health risks associated with mycotoxin exposure⁸¹.



Fig. 8: Multidimensional approach to mycotoxin risk mitigation (self-generated)

Four-pillar framework for reducing mycotoxin-related health risks: Food Safety (contamination control), Nutrition Education (dietary awareness), Community Action (local implementation), and Research (evidence generation). Arrows denote interdependencies between pillars

Table 9: Recommendations for nutritional interventions

Strategy	Nutrient focus	Implementation approach	Target population	Citations
Dietary	Vitamins, minerals,	Promote varied diets	Rural and urban	Xu et al. ⁸¹
diversification	antioxidants	including cereals, legumes,	low-income	Chen et al.82
		fruits, and vegetables	households	Cheng et al.83
Functional food	Anti-epigenetic	Encourage consumption	Populations at risk	Turner et al.84
promotion	bioactives	of biofortified and	of aflatoxin exposure	Nji et al. ⁸⁵
		functional foods		Visser et al.86
Nutrition-sensitive	Mycotoxin awareness,	Integrate food safety	Women, caregivers, and	Kimanya et al. ⁸⁷
food safety	micronutrients	into nutrition education	school-age children	Ingenbleek et al.88
education		programs		Wild and Gong ⁸⁹
Community-based	Sanitation-linked	Local programs	Underserved rural	
hygiene interventions	nutrient retention	addressing water,	communities	
		sanitation, and		
		hygiene (WASH)		
Policy-driven	Nutrient-enabling	National aflatoxin	Policymakers, educators,	
public health	environments	mitigation strategies and	and food vendors	
campaigns		food system reforms		
Risk communication	Environmental	Holistic assessment of	General African population	ns
and systems approach	nutrition interaction	environmental and	in hotspot regions	
		dietary toxins		

This table presents strategic nutritional interventions to mitigate mycotoxin exposure risks across African communities. Dietary diversification, functional food intake, food safety education and hygiene-centered programs are recommended for vulnerable populations. Policy campaigns and systems-based communication target broader public health infrastructure and environmental factors

Figure 8 proposes an integrated strategy to address mycotoxin exposure in African populations. Food safety practices reduce contamination in staples⁸¹, while nutrition education promotes protective diets rich in methyl donors and antioxidants⁸². Community-driven hygiene interventions enhance nutrient retention^{82,83}, and research identifies epigenetic biomarkers for intervention efficacy⁸³. The model aligns with Table 9 evidence-based recommendations for vulnerable groups.

Promoting dietary diversity, increasing the intake of fruits, vegetables, and whole grains rich in methyl donors, antioxidants, and fiber, and addressing micronutrient deficiencies are crucial steps⁸³.

Table 9 synthesizes intervention strategies to disrupt mycotoxin-induced epigenetic damage in Africa. Dietary diversification combats micronutrient deficiencies linked to susceptibility^{83,84}, while functional foods

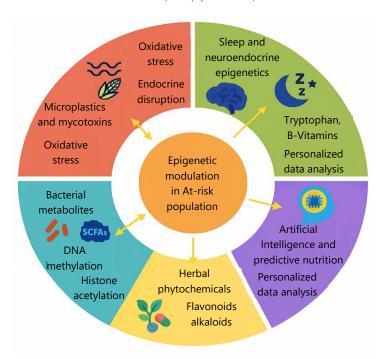


Fig. 9: Integrated nutritional-epigenetic response model to environmental pollutants (self-generated)

Five major domains influencing epigenetic responses to environmental toxins. Nutritional components and emerging technologies offer protective modulation. This integrative model underscores the complexity of diet-environment-gene interactions in vulnerable populations

(e.g., biofortified crops) deliver anti-epigenetic bioactives⁸⁴. Policy-driven food safety campaigns and WASH programs address environmental determinants⁸⁵, and systems-based risk communication enables holistic responses⁸⁶.

Furthermore, research is needed to identify specific nutritional biomarkers that can indicate susceptibility to mycotoxin-induced epigenetic damage and to monitor the effectiveness of dietary interventions⁸⁷.

Culturally appropriate and sustainable food-based strategies should be prioritized⁸⁸. Public health initiatives aimed at improving food safety practices, reducing mycotoxin contamination in staple crops, and promoting nutrition education are essential components of a comprehensive approach to protect the health of these vulnerable populations⁸⁹. Future research should focus on conducting well-designed intervention studies in African settings to evaluate the efficacy of specific nutritional strategies in modulating mycotoxin-related epigenetic alterations and improving health outcomes.

INTEGRATIVE PERSPECTIVES ON EMERGING BIOCHEMICAL INTERACTIONS AND NUTRITIONAL EPIGENETICS

Recent advances in nutritional epigenetics have expanded our understanding of how biochemical signals interact with environmental toxins to modulate gene expression. Mycotoxins are increasingly being considered alongside other emerging pollutants such as microplastics, which may similarly disrupt epigenetic homeostasis through oxidative stress, mitochondrial dysfunction, and interference with hormone signaling pathways⁹⁰. These pollutants, including mycotoxins, can influence systemic biochemical networks, with cumulative effects on metabolic, immune, and neuroendocrine systems.

A growing body of evidence indicates that neurobiological rhythms, particularly those governing sleep and circadian cycles, are under both nutritional and e pigenetic regulation⁹¹. Mycotoxins have been associated with altered sleep architecture and endocrine function, potentially via modulation of melatonin pathways and DNA methylation enzymes. Dietary modulation through polyphenols, methyl donors, and specific amino acids may provide a corrective mechanism to restore disrupted neuroepigenetic circuits.

Moreover, bacterial-derived metabolites are emerging as potent biochemical mediators that influence host immunity, neurotransmitter regulation, and gene expression through epigenetic remodeling⁹². These findings emphasize the potential of microbiome-targeted nutritional strategies such as the use of prebiotics, probiotics, and postbiotics to counteract mycotoxin-induced epigenetic alterations.

Parallel insights from herbal biochemistry offer another promising intervention pathway. Certain phytochemicals from traditional herbal medicines possess epigenetic activity, including inhibition of Histone Deacetylases (HDACs) and DNA Methyltransferases (DNMTs), thereby modulating gene expression in a manner that can mitigate mycotoxin toxicity⁹³. The integration of nutraceuticals with known molecular targets could strengthen community-based dietary interventions.

Finally, Artificial Intelligence (AI) is beginning to play a pivotal role in biochemical research, offering computational models to predict nutrient toxin epigenome interactions⁹⁴. The AI-based platforms can help identify population-specific dietary strategies that optimally counteract the epigenetic burden of mycotoxins, accounting for genetic polymorphisms, metabolic profiles, and environmental exposures. Future research in African populations should incorporate these novel perspectives to enhance the effectiveness of nutrition-sensitive interventions.

Figure 9 illustrates how diverse biochemical and environmental factors influence epigenetic modulation in at-risk populations. Key contributors include microplastics, gut microbiota, sleep regulation, herbal phytochemicals, and Al-guided nutrition. Arrows represent directional influences on DNA methylation, histone acetylation, and overall epigenetic stability.

CONCLUSION

Mycotoxin exposure poses a significant epigenetic and public health challenge in Sub-Saharan Africa, especially among nutritionally vulnerable populations. This review highlights the pivotal role of nutrition in modulating mycotoxin-induced alterations in DNA methylation, histone modifications, and noncoding RNA expression. Key dietary components, including methyl donors, antioxidants, and fiber, offer promising protective mechanisms against these epigenetic disruptions. Integrating nutrition-sensitive strategies with food safety interventions can mitigate long-term health risks. Culturally appropriate dietary interventions and public health education are essential for sustainable impact. Future research should prioritize biomarker discovery and targeted trials to inform evidence-based policies.

SIGNIFICANCE STATEMENT

This manuscript highlights a critical yet often overlooked dimension of public health in Sub-Saharan Africa, the intersection between chronic mycotoxin exposure and nutrition-driven epigenetic regulation. By synthesizing current evidence on how key dietary components such as methyl donors, antioxidants, and microbiota modulators influence gene expression and mitigate toxin-induced epigenetic disruptions, the study presents a compelling framework for preventive health. It emphasizes the potential of nutrition as a cost-effective, culturally appropriate strategy to reduce the long-term health burden associated with mycotoxins. The findings offer valuable insights for integrating nutritional interventions into food safety policies and population health programs.

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