

Metabolism, Toxicity, Detoxification, Occurrence, Intake and legislations of Fumonisin - A review

ABSTRACT

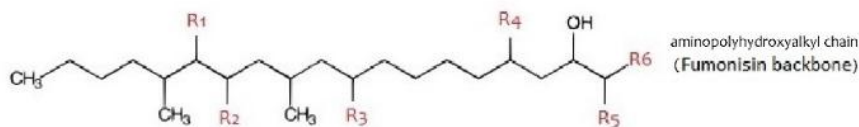
Fumonisin is a group of mycotoxins generated by the *Fusarium spp.* in foods and feeds. More than 15 isomers of fumonisin have been recognized, and the B series of fumonisins are the primary and referral isomers of fumonisins. Fumonisin B can cause, leukoencephalomalacia in rabbits and horses and porcine pulmonary edema in swine. Also, fumonisin B is nephrotoxic, hepatotoxic, immunotoxic and carcinogenic. Fumonisin B blocks sphingolipid biosynthesis (and hinders the synthesis of ceramide) by a noticeable resemblance to sphingosine and sphinganine. This paper gives a review of the toxicity, occurrence, and mechanism of carcinogenicity, hepatotoxicity, nephrotoxicity, and immunotoxicity of fumonisins. Fumonisin is primarily found on several foods and feed in Africa, America, Europe, Asia, and Oceania. In this paper, we summarize current information on the worldwide contamination of feeds and foods by fumonisins. Because of economic losses induced by fumonisins and their harmful effects on animal and human health, the various procedure for detoxifying infected feeds and foods have been illustrated in this review, including; biological, physical, and chemical processes. In addition, we discuss dietary intakes and maximum limits of fumonisins in some countries.

Keywords: Fumonisin; Toxicity; Detoxification; Mechanism; Occurrence; Intake

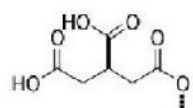
INTRODUCTION

Fumonisin is a group of more than ten mycotoxins created by *Fusarium* species like; *F. globosum*, *F. oxysporum*, *F. proliferatum*, *F. verticillioides* and other species of *Fusarium*, *Alternaria alternata f. sp. lycopersici*, and *Aspergillus niger* (Scott, 2012; Shimizu et al., 2015).

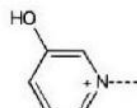
Fumonisin has a noncyclic structure (in contrast to most mycotoxins). In this structure, there is a chain with 19- or 20- carbon aminopolyhydroxyalkyl that diesterified by tricarballic acid groups (propane-1,2,3-tricarboxylic acid) Fig 1. Hitherto, various chemically associated series or groups of fumonisins were isolated. These series consist of A, B, C, and P. The main detected forms of fumonisins in foods, are the B series of fumonisins (Jackson and Jablonski, 2004). Fumonisin B₁, fumonisin B₂, and fumonisin B₃ are the broadest mycotoxins between more than 15 fumonisin forms that have been described until now (Humpf and Voss, 2004).



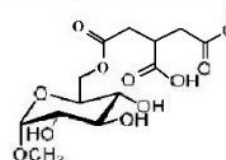
Fumonisins	Group					
	R1	R2	R3	R4	R5	R6
FA ₁	TCA	TCA	OH	OH	NHCOCH ₃	CH ₃
FA ₂	TCA	TCA	H	OH	NHCOCH ₃	CH ₃
FA ₃	TCA	TCA	OH	H	NHCOCH ₃	CH ₃
FAK ₁	=O	TCA	OH	OH	NHCOCH ₃	CH ₃
FB ₁	TCA	TCA	OH	OH	NH ₂	CH ₃
FB ₂	TCA	TCA	H	OH	NH ₂	CH ₃
FB ₃	TCA	TCA	OH	H	NH ₂	CH ₃
FB ₄	TCA	TCA	H	H	NH ₂	CH ₃
FC ₁	TCA	TCA	OH	OH	NH ₂	H
FP ₁	TCA	TCA	OH	OH	3HP	CH ₃
FP ₂	TCA	TCA	H	OH	3HP	CH ₃
FP ₃	TCA	TCA	OH	H	3HP	CH ₃
PH _{1a}	TCA	OH	OH	OH	NH ₂	CH ₃
PH _{1b}	OH	TCA	OH	OH	NH ₂	CH ₃
AP ₁ (Hydrolyzed FB ₁)	OH	OH	OH	OH	NH ₂	CH ₃
N-(carboxymethyl) FB ₁	TCA	TCA	OH	OH	NH(C ₂ H ₃ O ₂)	CH ₃
N-(deoxy-D-fructos-1-yl)B ₁	TCA	TCA	OH	OH	NH(C ₆ H ₁₁ O ₅)	CH ₃
Fumonisin B ₁ -di(methyl- α -D-glucopyranoside)	MG	MG	OH	OH	NH ₂	CH ₃



Tricarballic Acid (TCA)



3-Hydroxypyridinium (3HP)



Methyl- α -D-glucopyranoside (MG)

Fig. 1. Chemical structures of the fumonisins (Humpf and Voss, 2004; Jackson and Jablonski, 2004).

Fungi-producing fumonisin contaminated apple, barley, beef, breakfast cereals, black tea, corn, cornbread, corn flour, corn flakes, corn grits, corn snacks, basmati rice, crunchy nut, egg, milk, oats, polenta, popcorn, row corn, soybean, canned foods, tomato, tortilla, and wheat (Soriano and Dragacci, 2004b).

Intake of fumonisin B induced a different form of toxic effect in animals, including leukoencephalomalacia in horses (Giannitti et al., 2011), change in weight of the body and internal organ in broiler chicken (Ledoux et al., 1992; Weibking et al., 1993a), and pulmonary edema as well as hepatocellular necrosis in piglet (Colvin et al., 1993; Pósa et al., 2016). Moreover, renal and hepatic toxicity have been detected in different animals, such as rabbits, lambs, turkeys, mice, rats, and broilers (Edrington et al., 1995; Orsi et al., 2009; Voss et al., 1995; Weibking et al., 1993a; Weibking et al., 1993b).

In human, fumonisins increased risk of neural tube defects (NTD) and developing esophageal cancer (Missmer et al., 2006; Stockmann-Juvala and Savolainen, 2008).

METHODS

This is a narrative review on Fumonisin, and the databases including Science Direct, PubMed, and Google Scholar were used to collect the published articles from 1980 through 2018. The review was

46 conducted using keywords: [*Fumonisin* OR *mycotoxin* OR *fumonisin B* OR *Fusarium*] AND [toxicity
 47 OR detoxification OR degradation OR mechanism OR metabolism OR occurrence OR prevalence OR
 48 intake OR limitation]. The list of references of included articles was also searched to identify additional
 49 articles. After first screening by the title and abstract, the eligible articles were downloaded, and we
 50 chose easy and suitable phrases. Inclusion criteria in our study were included: (1) Full-text available.
 51 (2) Review, mini-review, original, narrative articles, and books. (3) Published paper in English
 52 language (to avoid avoid mistake in the translation process) among 1980 to 2018 year. (4) Detect
 53 concentration of fumonisin B₁, B₂, and B₃ in barley, wheat, oat, rice, corn and corn product such as
 54 corn grits, corn flakes, corn flour, corn meal, and corn kernel. The articles were excluded when they
 55 did not meet these criteria.

57 1. OCCURRENCE

58 Due to increases in global grain exchange, fungi spread has been transferred from one country to
 59 another (Placinta et al., 1999). In *Fusarium* fungi, this hazard expected to be minimum whereas these
 60 phytopathogens are field sooner than storage organisms. The global infection of animal feeds and
 61 foodstuffs with fumonisins is described in Table 1.

62 **Table 1. Occurrence of fumonisins from human foods, cereals, and crops in various countries.**

63 Nation-seed	Fumonisin B ₁ (mg/kg)	Fumonisin B ₂ (mg/kg)	Fumonisin B ₃ (mg/kg)	Reference
Barley				
Brazil	2.43			(Mallmann et al., 2001)
Korea	0 to 2667.3	0 to 1521.1		(Choi et al., 2018)
France	Not detected			(Malmauret et al., 2002)
Spain	0.2 to 11.6	0.5		(Castella et al., 1999a)
UK	Not Detected	Not Detected	Not Detected	(Patel et al., 1997)
Corn				
Argentina	Average of fumonisins in 2003: 10.2 and in 2004: 4.7 µg/kg			(Broggi et al., 2007)
Brazil	0.2 to 38.5	0.1 to 12		(Sydenham et al., 1992)
Brazil	5.45 to 10.59	3.62 to 10.31		(Hirooka et al., 1996)
Brazil	0.5 to 1.38	0.01 to 0.59		(Wild et al., 1998)
Brazil	0.2 to 6.1			(Vargas et al., 2001)
Brazil	78.92			(Mallmann et al., 2001)
Brazil	3.2	3.4	1.7	(Van Der Westhuizen et al., 2003)
Brazil	0.066 to 7.832	0.11 to 1.201		(Scussel et al., 2014)

Honduras	0.068 to 6.5			(Julian et al., 1995)
Uruguay	0.165 to 3.688			(Pineiro et al., 1997)
USA	0 to 1.614			(Tseng and Liu, 2001)
USA	0.058 to 1.976	0.054 to 0.890		(Li et al., 2001)
Venezuela	0.025 to 15.05			(Medina-Martínez and Martínez, 2000)
China	0.872 to 0.890	0.33 to 0.448		(Yoshizawa et al., 1994)
China	0.08 to 21	0.05 to 4.35	0.06 to 1.66	(GAO and YOSHIZAWA, 1997)
China	<0.05 to 25.97	<0.10 to 6.77	<0.10 to 4.13	(Ueno et al., 1997)
China	Total fumonisins <0.5 to 16.0			(ZHANG et al., 1997)
China	0.058 to 1.976	0.056 to 0.89	0.053 to 0.385	(Li et al., 2001)
China	0.003 to 71.121			(Gong et al., 2009)
China	0.268	0.537	0.472	(Li et al., 2015)
China	0.0165 to 0.3159			(Shi et al., 2018)
India	0.07 to 8			(Shetty and Bhat, 1997)
India	<1 to 100			(Jindal et al., 1999)
Iran	1.270 to 3.980	0.190 to 1.175	0.155 to 0.960	(Shephard et al., 2000)
Iran	223.64			(Alizadeh et al., 2012)
Japan	<0.05 to 4.1	<0.1 to 10.2		(Ueno et al., 1993)
Philippines	Total fumonisins 0.3 to 10			(Bryden et al., 1996)
Taiwan	0.63 to 18.8	0.05 to 1.4		(Yoshizawa et al., 1996)
Taiwan (Australia)	≤0.477			(Tseng and Liu, 2001)
Taiwan (USA)	≤1.614			(Tseng and Liu, 2001)
Taiwan (South)	≤0.865	≤0.12		(Rheeder et al.,

Africa)				1994)
Taiwan (South Africa)	≤0.05 to 0.9	<0.05 to 0.25		(Rheeder et al., 1994)
Taiwan (Thailand)	≤0.334			(Tseng and Liu, 2001)
Vietnam		Total fumonisins 0.3 to 9.1		(Bryden et al., 1996)
Australia		Total fumonisins 0.3 to 40.6		(Bryden et al., 1996)
Australia	≤0.477			(Tseng and Liu, 2001)
Austria	<15			(Lew et al., 1991)
Croatia	0.01 to 0.06	0.01		(Doko et al., 1995)
Croatia	The highest concentrations fumonisins 25.5, mean values of 4.509			(Pleadin et al., 2012)
Greece	0.1 to 0.56			(De Nijs et al., 1998a)
Portugal	0.09 to 2.3	0.25 to 4.45		(Doko et al., 1995)
Poland	0.01 to 0.02	<0.01		(Doko et al., 1995)
Romania	0.01 to 0.02	0.01		(Doko et al., 1995)
Spain	≤22	≤0.7		(Sanchis et al., 1995)
Spain	70 to 334	102 to 379		(Castellá et al., 1996)
Spain	0.2 to 19.2	0.2 to 5.9		(Castella et al., 1999b)
Spain	0.035 to 0.043	0.019 to 0.022		(Arino et al., 2007)
The Netherlands	Traces to 0.380			(De Nijs et al., 1998b)
The Netherlands	Traces to 3.35			(De Nijs et al., 1998a)
UK	0.2 to 6			(Preis and Vargas, 2000)
Benin	Total fumonisins: 6.1 to 12 in 1999-2003			(Fandohan et al., 2005)
Ethiopia	0.606	0.202	0.136	(Getachew et al., 2018)
Ghana	0.011 to 1.655	0.01 to 0.77	0.07 to 0.224	(Kpodo et al., 2000)

Malawi	0.02 to 0.115		0.03	(Doko et al., 1996)
Morocco	1.930			(Zinedine et al., 2006)
Nigeria	0.164 to 2.09 (0.852)	0.046 to 0.710 (0.262)	0.010 to 0.186 (0.069)	(Akinmusire et al., 2018)
South Africa	<10 to 83			(Sydenham et al., 1990a)
South Africa	≤0.63		≤0.25	(Rheeder et al., 1994)
South Africa	0.05 to 117.5		0.05 to 22.9	(Rheeder et al., 1992)
South Africa	0.2 to 46.9		0.15 to 16.3	(Sydenham et al., 1990b)
South Africa	<0.2 to 2			(Dutton and Kinsey, 1995)
South Africa	0.012 to 8.514			(Mngqawa et al., 2015)
South Africa (Argentina)	0.05 to 0.7	<0.05 to 0.5	<0.05 to 0.5	(Stockenström et al., 1998)
South Africa (USA)	0.9 to 3.9	0.3 to 1.2	0.08 to 0.6	(Stockenström et al., 1998)
Tanzania	0.025 to 0.165		0.06	(Doko et al., 1996)
Zimbabwe	0.125		0.04	(Doko et al., 1996)
Corn flakes				
Argentina	0.002 to 0.038	Not detected		(Solovey et al., 1999)
Brazil	0.66		0.03	(Mac Jr and Valente Soares, 2000)
Uruguay	0.218	Not detected		(Pineiro et al., 1997)
USA		Total fumonisins: <0.25		(Pohland, 1996)
USA	≤0.088	Not detected		(Castelo et al., 1998)
USA or Canada	0.012 to 0.155			(Kim et al., 2003)
Korea	0.018 to 0.143			(Kim et al., 2002)
China	0.001 to 0.171	<0.0002 to 0.025	0.0002 to 0.031	(Li et al., 2015)
Germany		Total fumonisins <0.01 to 1		(Usleber and Märtilbauer, 1998)

Italy	0.01	Not detected	(Doko and Visconti, 1994)
Italy	0.020 to 1.092	0.006 to 0.235	(Solfrizzo et al., 2001a)
Nordic countries	0.005 to 1.030	0.004 to 0.243	(Petersen and Thorup, 2001)
Serbia	0 to 0.434 (0.067)	0 to 0.145 (0.019)	(Torović, 2018)
Spain	0.02 to 0.1		(Sanchis et al., 1994)
Switzerland	0.055		(Pittet et al., 1992)
The Netherlands	1.43		(De Nijs et al., 1998b)
Turkey	Not detected	Not detected	(OMURTAG, 2001)
South Africa	Not detected	Not detected	(Sydenham et al., 1991)
Corn flour			
Argentina	0.038 to 1.86	0.02 to 0.768	(Hennigen et al., 2000)
Brazil	≤1.46	≤0.51	(Mac Jr and Valente Soares, 2000)
USA	Total fumonisins: <0.25 to 1		(Pohland, 1996)
China	0.06 to 0.2	<0.10	(Ueno et al., 1993)
Italy	3.54	0.84	(Doko and Visconti, 1994)
Nordic countries	0.017 to 0.86	0.007 to 0.024	(Petersen and Thorup, 2001)
Serbia	0 to 1.738 (0.162)	0 to 0.394 (0.042)	(Torović, 2018)
UK	Total fumonisins 0.218		(Patel et al., 1996)
The Netherland	0.04 to 0.09		(De Nijs et al., 1998b)
Corn grits			
Argentina	0.092 to 0.494	0.02 to 0.1	(Hennigen et al., 2000)
Argentina	1.1	0.425	(Torres et al., 2001)
Brazil	0.17 to 1.23	0.05 to 0.3	(Mac Jr and Valente Soares, 2000)

USA	Average 0.6	Average 0.4		(Sydenham et al., 1991)
USA	Total fumonisins: 0.251 to 1			(Pohland, 1996)
USA	Total fumonisins: <0.25			(Pohland, 1996)
Japan	0.2 to 2.6	0.3 to 2.8		(Ueno et al., 1993)
China	0.0002 to 2.238	0.0002 to 0.547	0.0002 to 0.402	(Li et al., 2015)
Germany	0.0139			(Usleber et al., 1994)
Italy	3.76	0.9		(Doko and Visconti, 1994)
Nordic countries	0.007			(Petersen and Thorup, 2001)
Spain	0.03 to 0.09	Not detected		(Sanchis et al., 1994)
Switzerland	0 to 0.79	0 to 0.16		(Pittet et al., 1992)
South Africa	<0.05 to 0.19	<0.05 to 0.12		(Sydenham et al., 1991)
Corn kernel				
Bahrain	0.025			(De Nijs et al., 1998a)
China	5.3 to 8.4	2.3 to 4.3		(Ueno et al., 1993)
China	1.878	0.853		(Guo et al., 2016)
Nepal	0.05 to 4.6	0.1 to 5.5		(Ueno et al., 1993)
Indonesia	0.051 to 2.44	<0.376		(Ali et al., 1998)
Egypt	69 to 4495			(Fadl, 1997)
Ghana	0.07 to 33.1	0.06 to 12.3		(Kpodo et al., 2000)
Kenya	0.11 to 12			(Kedera et al., 1999)
Corn meal				
Argentina	0.06 to 2.86	0.061 to 1.09	0.018 to 1.015	(Solovey et al., 1999)
Argentina	0.603 to 1.171	0.717		(Torres et al., 2001)
Brazil	0.56 to 4.93	0.21 to 1.38		(Mac Jr and Valente Soares, 2000)
Canada	0.05			(Sydenham et al., 1991)

Peru	0.66	0.13	(Sydenham et al., 1991)
USA	Average: 1	0.3	(Sydenham et al., 1991)
USA	Total fumonisins: <0.25 to >1		(Pohland, 1996)
China	<0.5 to 8.8	<0.5 to 2.8	<0.5 to 0.9 (Groves et al., 1999)
Turkey	0.25 to 2.66	0.55	(OMURTAG, 2001)
South Africa	Average: 0.14	Average: 0.08	(Sydenham et al., 1991)
Oat			
Brazil	0.17		(Mallmann et al., 2001)
UK	Total fumonisins not detected		(Patel et al., 1997)
Rice			
Iran	21.59		(Alizadeh et al., 2012)
China 1999	3.410 to 16.79		(Sun et al., 2017)
China 2010	0.0001 to 0.00164		(Sun et al., 2017)
China 2014	0 to 0.74		(Sun et al., 2017)
UK	Total fumonisins not detected		(Patel et al., 1997)
Wheat			
Brazil	24.35		(Mallmann et al., 2001)
Argentina_flour	0.0003	0.00124	(Cendoya et al., 2018)
France	Not detected		(Malmauret et al., 2002)
Spain	0.2 to 8.8	0.2	(Castella et al., 1999a)
UK	Total fumonisins not detected		(Patel et al., 1997)

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65

1.1 North and South America

66 Corn is the most prevalent source of Fumonisin (Table 1). The level of Fumonisin in South America
67 is higher than in North America maybe because of their different climate conditions. For instance,
68 concentration of fumonisin in corn in Brazil reached to 38.5 mg/kg (Sydenham et al., 1992), while
69 the percentage of Fumonisin in corn product of North America such as corn flour, corn grits, and corn

70 flakes rarely reached to 1 mg/kg. This decline probably prove that detoxification method was more
71 effective for the control of Fumonisin in North America than South America.

72 In Brazil, the incidence of fumonisins in corn was detected by (Scussel et al., 2014), (Sydenham et al.,
73 1992), (Hirooka et al., 1996), (Wild et al., 1998), (Vargas et al., 2001), (Mallmann et al., 2001) and
74 (Van Der Westhuizen et al., 2003), contamination of corn with fumonisins in Brazil usually decreased
75 over 1999 to 2014.

76 The average of fumonisins in corn of Argentina was 10200 µg/kg in 2003 and 4700 µg/kg in 2004
77 (Broggi et al., 2007).

78 The infection of wheat, oat and barely by fumonisins were also detected by (Mallmann et al., 2001).

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81 1.2 Asia and Oceania

82 In China, the contamination of corn with fumonisins was reported by (Yoshizawa et al., 1994); (Ueno
83 et al., 1997), (GAO and YOSHIZAWA, 1997), (Li et al., 2001), (Li et al., 2015), (Gong et al., 2009) and
84 (Shi et al., 2018). Based on these studies the most extreme concentration of fumonisin B₁, B₂ and B₃
85 were 25.97 mg/kg, 6.77 mg/kg and 4.13 mg/kg respectively. Also, (ZHANG et al., 1997) reported that
86 in China total fumonisins concentration was 0.5 to 16 mg/kg. In Iran (Shephard et al., 2000)
87 investigated infection of corn with fumonisin B₁, B₂, and B₃. Also, (Alizadeh et al., 2012) reported the
88 corn's contamination with fumonisin B₁. The high concentration of Fumonisin in corn of Iran and
89 China, justify the high prevalence of esophageal cancer in Iranian and Chinese people.

90 The contamination of corn with fumonisin B₁ and B₂ were detected by (Ueno et al., 1993) in Japan.
91 (Bryden et al., 1996) declared that the measure of total fumonisins in corn of Philippines and Vietnam
92 was 0.3 to 10 mg/kg and 0.3 to 9.1 mg/kg, respectively.

93 Contamination of Taiwan's corn with fumonisins was investigated by (Yoshizawa et al., 1996), (Tseng
94 and Liu, 2001) and (Rheeder et al., 1994). Increasing in level of Fumonisin in Taiwan's corn declared
95 that legislation and control program in this country was not efficient and it is necessary to change their
96 programmes.

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98 1.3 Europe

99 (Doko et al., 1995) published a review article on information about the occurrence of fumonisins from
100 some European nations (Croatia, Poland, Portugal, and Romania). The highest concentration of
101 fumonisins in Croatia was 25.2 mg/kg, and the mean value was 4.509 mg/kg (Pleadin et al., 2012).

102 In Spain, contamination of corn with fumonisins investigated by (Sanchis et al., 1995), (Arino et al.,
103 2007), (Castellá et al., 1996), and (Castella et al., 1999b). Also, (Castella et al., 1999a) reported the
104 concentration of fumonisin B₁ and B₂ in wheat and barley.

105 Fumonisin B₁ was not found in wheat and barley of France (Malmauret et al., 2002).

106 (Lew et al., 1991) reported the corn contamination with fumonisin B₁ in Austria.

107 In oat, barley and wheat of United Kingdom (Patel et al., 1997) have not detected fumonisins but
108 (Preis and Vargas, 2000) declared the concentration of fumonisin B₁ in corn of UK (0.2 to 6 mg/kg).

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110 1.4 Africa

111 Albeit majority African territory has weather distinguished by high temperature and high humidity that
112 is suitable for the development of molds, little data is accessible on the occurrence of toxins of
113 *Fusarium*. High infection of the basic material is a developing problem in Africa. Regulative problems
114 are not accessible in the territory of food retailing and exhibition, and mycotoxin issues now have
115 been combined with some food infection in some parts in Africa (Zinedine et al., 2007).

116 The maximum level of fumonisin B₁ in researches on corn of South Africa is very high and achieved
117 to 117.5 mg/kg in (Rheeder et al., 1992) and 8.514 in new literature by (Mngqawa et al., 2015).

118 (Getachew et al., 2018) detected the fumonisin B₁, B₂, and B₃ in corn of Ethiopia.

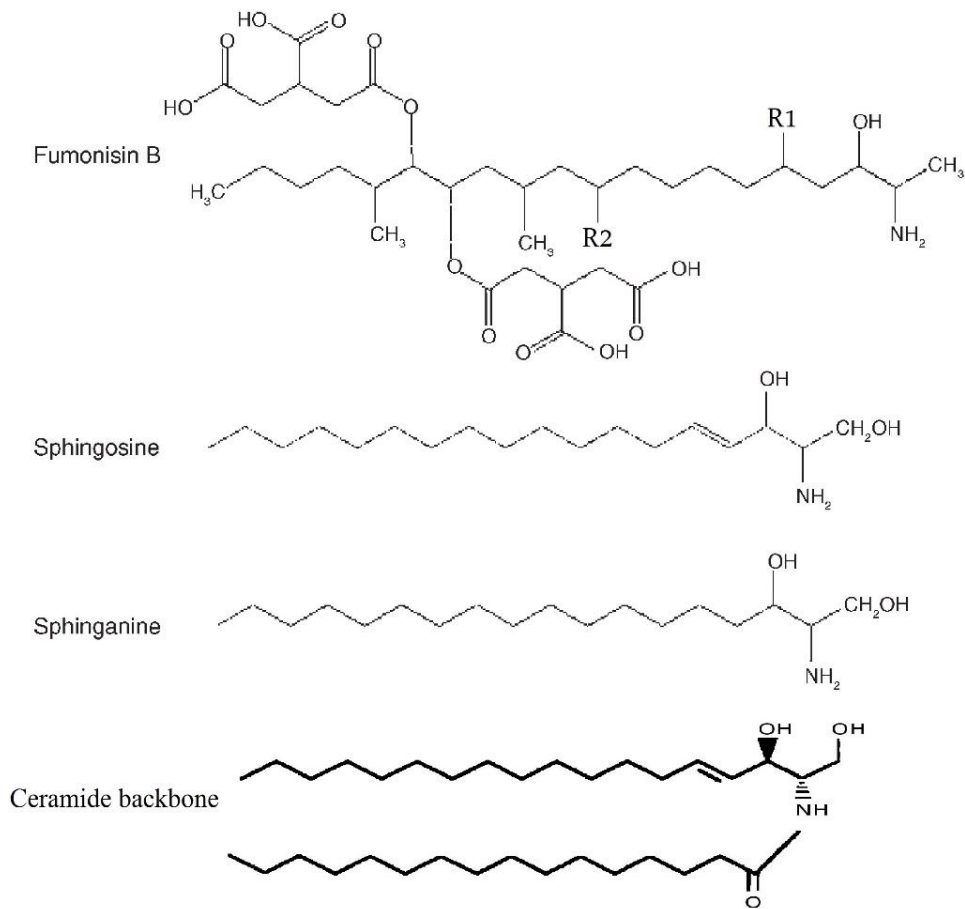
119 Evaluation of fumonisins on corn products of Africa is low, and these investigation consisted of corn
120 meal (Sydenham et al., 1991), corn kernel (Kedera et al., 1999), and corn grits (Sydenham et al.,
121 1991).

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123 2. METABOLISM AND MECHANISM OF FUMONISINS

124 Structure of fumonisin B has a noticeable similarity to sphinganine and sphingosine. In Fig. 2 both
125 sphingosine and sphinganine are intermediates in the degradation and biosynthesis of sphingolipids.
126 Furthermore, (D'mello et al., 1999) reported that fumonisin B obstruct sphingolipid biosynthesis by
127 specifically inhibiting sphingosine (sphinganine) N-acyltransferase, *in vitro* and *in situ*.

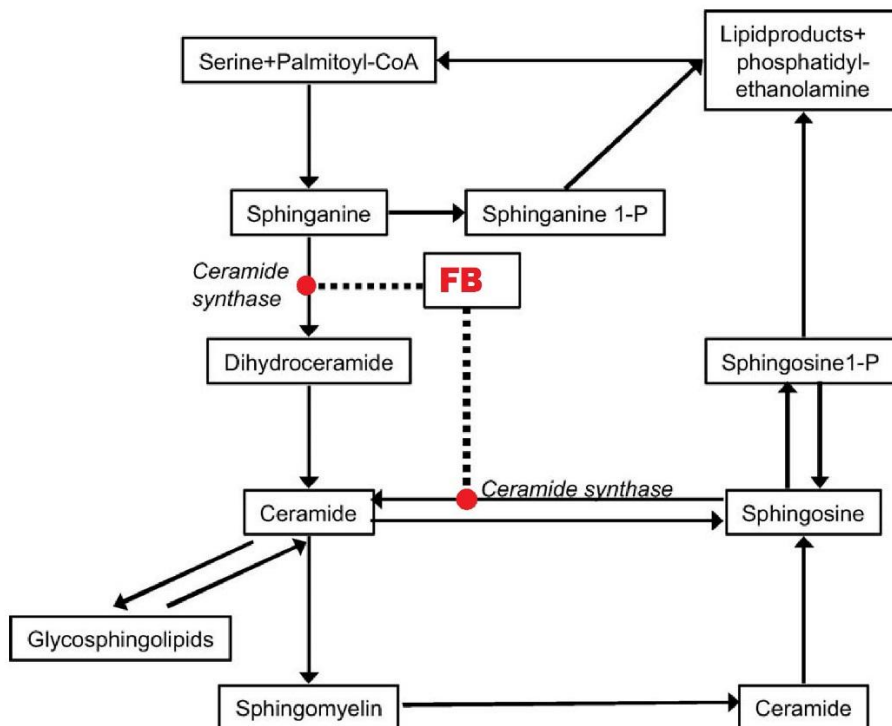
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Fig. 2. Structures of fumonisin B, sphingosine, sphinganine and ceramide backbone(Jackson and Jablonski, 2004); (Merrill Jr et al., 2001)

Sphingolipids are a group of lipids that can be detected in the whole of eukaryotic cells. All of the sphingolipids include a sphingoid (long-chain base backbone). Sphingolipids are critical basic molecules and rule as regulators of a numeral of cell act (Merrill et al., 1997). In Fig. 3 location of working of fumonisin B-induced inhibition of the enzyme CER synthase, is presented.



140
141 **Fig. 3. A summarized scheme of the sites of action of fumonisin B-induced inhibition of the**
142 **enzyme ceramide synthase on the pathway of de novo sphingolipid synthesis and turnover in**
143 **mammalian cells and (Stockmann-Juvala and Savolainen, 2008).**
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145 2.1 Mechanism of Fumonisin in Apoptosis and Cancer

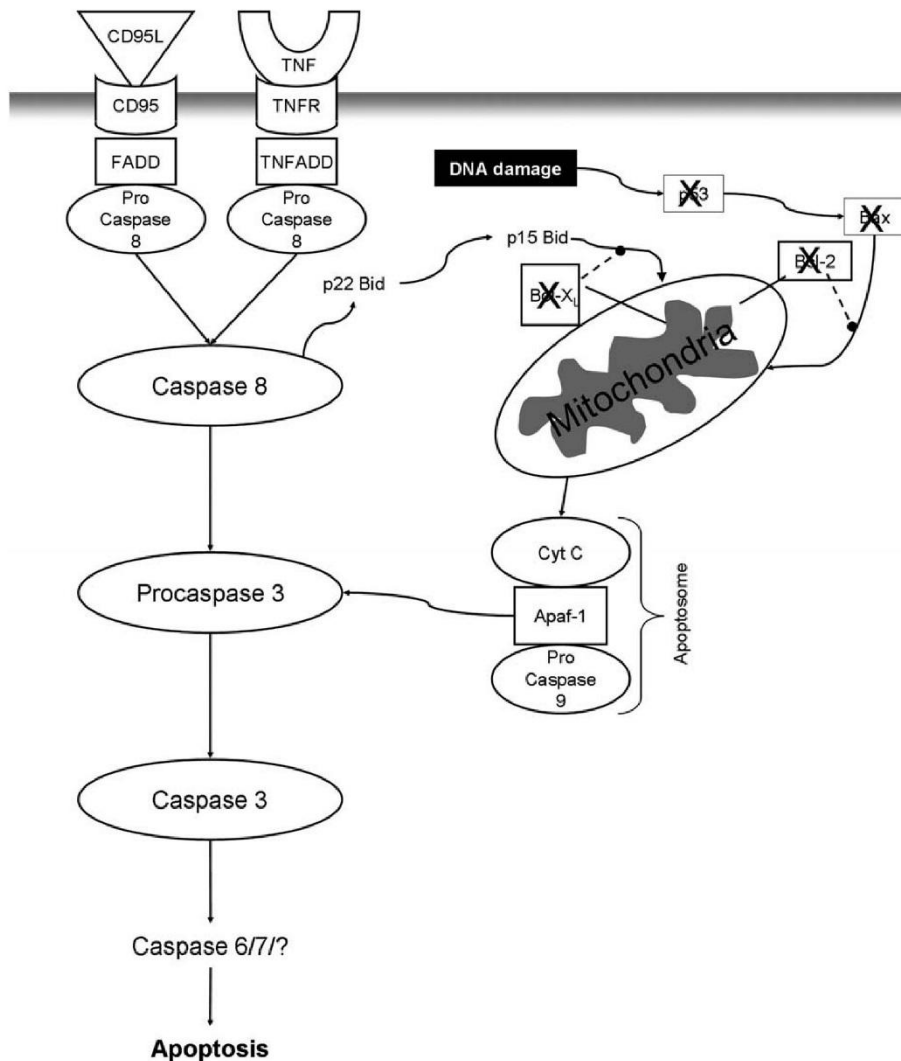
146 Interruption of sphingolipid metabolism can cause the increase in; available sphingoid backbone and
147 their 1-phosphates, **may change the** compound sphingolipids, and **can** decrease the biosynthesis of
148 ceramide (CER). Available sphingoid backbone induced cell death, but fumonisins by inhibition of
149 CER synthase can restrain cell death (Riley et al., 2001).

150 Feedback of the apoptosis and carcinogenicity effects induced by fumonisin B₁ can be some
151 mechanisms including oxidative damage, lipid peroxidation and maybe induction of hepatic, and renal
152 tumors can happen (Stockmann-Juvala and Savolainen, 2008). Also, (Yin et al., 1998) discovered
153 that FB₁ was able to promote the production of free radicals (by increasing the rate of oxidation) and
154 by lipid peroxidation in membranes can accelerate chain reactions.

155 Increasing in sphinganine of tissue by FB was able to elevate beginning a cascade of cellular
156 changes that probably product the carcinogenicity and toxicity by an unknown mechanism(s).
157 However, in the following of sphinganine-induced cell proliferation, apoptosis and cancer incidence
158 might be elevated (Jackson and Jablonski, 2004).

159 In some studies following fumonisin B₁ treatment in different cells of human and animals, has been
160 shown that apoptosis caused by fumonisin B₁ does not entail p53 or Bcl-2 group proteins and protect
161 cells from the apoptosis by baculovirus gene (CpIAP). Baculovirus gene obstructs induced apoptosis
162 by the tumor necrosis factor (TNF) pathway that cleaved caspase-8. The mitochondrial pathway
163 perhaps is consisted of induced apoptosis by fumonisin B₁ by the actuation of Bid and release
164 cytochrome c (Stockmann-Juvala and Savolainen, 2008).

165 (Wang et al., 2014) reported that fumonisin B₁ in human normal esophageal epithelial cells (HEECs)
166 stimulated the proliferation. Mechanism of the proliferation of HEECs is, decreasing in protein
167 expression of cyclin E, p21, and p27 and increase in protein expression of cyclin D1.



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Fig. 4. A schematic landscape of the pathways conduct to apoptosis and the mechanisms probably consisted of fumonisin B₁ -induced activation of caspase-3 resulted in apoptosis. X mark showed the mechanisms that do not consisted of the apoptosis caused by fumonisin B₁ (Stockmann-Juvala and Savolainen, 2008).

2.2 Mechanism of Fumonisins in Hepatotoxicity

Accumulation of sphingoid base because of induced fumonisin B₁ can induce TNF- α and make the hepatotoxicity in mice. Also, TNF- α receptor 1b is important mediating in the hepatotoxic responses by a rise in the circulation of liver enzymes (Sharma et al., 2000).

2.3 Mechanism of Fumonisins in Immunotoxicity

Exposure to FB₁ in human dendritic cells; increases the exhibition of IFN- γ and the associated chemokine CXCL9. Nevertheless, fumonisin B₁ may decline the lipopolysaccharide-induced liver and brain expression of IL-1 β and IFN- γ in addition to the induced lipopolysaccharide expression of IL-1 β , IL-6, and the chemokines CCL3 and CCL5 in human dendritic cells (Stockmann-Juvala and Savolainen, 2008).

In piglets, fumonisin B₁ exposure can increase expression of IL-18, IL-8, and IFN- γ mRNA. But mRNA measure of TNF- α , IL-1 β in piglet alveolar macrophages and levels of IL-4 may decrease (Halloy et al., 2005); (Taranu et al., 2005).

After exposure to fumonisin B₁ in mouse, a raise expression of TNF- α and interleukin-1 β (IL-1 β) has been observed in kidney and the liver. Also, FB₁ can raise expression of IFN- γ , IL-1 α , IL-18, IL-12, IL-10, and IL-6 in the liver of mouse (Stockmann-Juvala and Savolainen, 2008).

194 **2.4 Mechanism of Fumonisin in Some Disorder**
 195 (Smith et al., 2002) recommended that the induced fumonisin B₁ by destruction of cardiovascular
 196 action can be one of the major elements that triggers happening of equine leukoencephalomalacia
 197 through increase in serum and sphingosine concentrations and myocardial sphinganine.
 198 Interruption of sphingolipid metabolism resulted in FB₁ before the pregnancy and during the first
 199 trimester may affect folate uptake and cause development risk of NTD (Marasas et al., 2004); (Cornell
 200 et al., 1983).
 201 FB₁ increases sphingosine and/or sphinganine concentrations, reduces the mechanical potency of the
 202 left ventricle and blocks L-type Ca channels. Pulmonary edema could generally caused by acute left-
 203 sided heart failure (Constable et al., 2000); (Smith et al., 2000).
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206 3. TOXICITY OF FUMONISINS

207 In the human and different animal, fumonisins beget some toxic effects such as carcinogenic,
 208 hepatotoxic, and nephrotoxic. Moreover, sensitivity to fumonisins is a different in human and varies
 209 animals. For example; based on (Bondy et al., 1997), rats are more sensitive to fumonisin B₁ than
 210 mice. We summarized disorder effects, dosage, duration and source of fumonisin in Table 2.
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Table 2. In vivo disorder effects induced by fumonisins

	Dosage and source	Fumonisin	Duration	Effects	References
Human	Both FB1 and FB2 High corn intake higher risk than low corn intake		case-control study	Developing esophageal cancer	(Stockmann-Juvala and Savolainen, 2008)
Human	FB1 in corn of three area of China, average of contamination was; 2.84, 1.27, and 0.65 mg/kg		1 year	Esophageal- and hepato-carcinogenesis	(Sun et al., 2007)
The human colon cell line HT-29	5 mg of FB1 was dissolved in PBS at a concentration of 1,380 uM		24 to 72 h	Main and early induction of Lipid peroxidation, assessing IL-8 secretion, increase in membrane microviscosity	(Minervini et al., 2014)
Human cells	Medicine with FB1 for 24, 48, 72 and 96 h			The proliferation of human esophageal epithelial cells (HEECs)	(Wang et al., 2014)
Women	Exposure to FB1 corn tortilla intake during the first trimester and before the pregnancy.		case-control study	Raise the risk of NTD	(Missmer et al., 2006)
Lamb	Intraruminally; 11.1, 22.2, 45.5 mg fumonisin B1, B2, B3/kg b.w		9 days	Tubular nephrosis, mild hepatopathy, diarrhea, lethargy, death	(Edrington et al., 1995)
Cattle	Feeding; 15, 31, 148 µg fumonisins /kg b.w		31 days	Increase in the AST, GGT, LDH, bilirubin, cholesterol and lymphocyte blastogenesis Mild microscopic liver lesions	(Osweiler et al., 1993)
Cattle	Intravenous; 1 mg fumonisin B1/kg b.w		7 days	Lethargy, the decrease in appetite Increase in Sa/So, proliferation and hepatocyte apoptosis, the proliferation of bile ductular cells, vacuolar change, proliferation of	(Mathur et al., 2001)

			proximal renal tubular cells, apoptosis, and karyomegaly.	
Bovine granulosa cell	0, 0.3, 1, 3, 10 µM of FB1	48 h	No effect on; cell proliferation, progesterone production, CYP11A1 and CYP19A1 gene expression, Slightly inhibited estradiol production Decreasing in serum-induced granulosa cell (GC) proliferation	(Albonico et al., 2017)
Broiler chicken	Feeding; 0, 100, 200, 300 or 400 mg fumonisin B1/kg b.w	21 days	The decline in body weight Increase in the liver-, proventriculus-, and gizzard-weights, Serum calcium, cholesterol, and AST	(Ledoux et al., 1992)
Broiler chicken	Feeding; 0, 75, 150, 225, 300, 375, 450, 525 mg fumonisin B1/kg b.w	21 days	Increase in liver and kidney weights, MCV, MCHC, Sa/So Histological lesions in the liver	(Weibking et al., 1993a)
Broiler chicken	Dietary; 0, 20, 40, 80 mg fumonisin B1/kg b.w	21 days	Increase in the Sa/So, GGT, AST, the weights of liver, proventriculus, spleen, kidney, and bursa of Fabricius.	(Henry et al., 2000)
Broiler chicken	Dietary; 0, 50, 100 or 200 mg fumonisin B1/kg b.w	21 days	Cell proliferation in response to mitogens, immunosuppress	(Li et al., 1999)
Broiler chicken	Dietary; 300 mg fumonisin B1/kg b.w	21 days	Increase activities of AST, LDH, GGT	(Kubena et al., 1997)
Broiler chicken Cobb 500	Orally and postnatal; 100 mg fumonisin B1/kg b.w	21 days	Increase in the liver weight, Sa/So, hepatic TBARS, Vit C, catalase	(Poersch et al., 2014)
Broiler chicken	diet (18.6 mg FB1+FB2/kg)	More than 17 days	Reduce in villus height and crypt depth of the ileum, Shift in the microbiota composition in the ileum Decreasing in abundance of Candidatus Savagella and Lactobacillus spp Increase in Clostridium perfringens content caused to higher percentage of birds developed subclinical necrotic enteritis	(Antonissen et al., 2015)
Broiler chicken	Purified FB (0 or 10 ppm)	34 days	Higher feed-to-gain ratio than control, Serum nitric oxide (NO) levels were elevated	(Lee et al., 2017)
Chicken Embryos	Injection in air cell of chicken eggs; 0, 2, 4, 8, 16, 32, and 64 µg fumonisin/egg	In 72h of incubation	Not microscopic abnormalities but haemorrhages of the neck, thoracic area, and	(Henry and Wyatt, 2001)

				head of the dead embryos	
Turkey	Dietary; 0, 100, 200 mg fumonisin B1/kg b.w	21 days		Increase in AST, alkaline phosphatase, MCV, MCH, liver-, kidney-, and pancreas-weights Biliary hyperplasia, thymic cortical atrophy, hypertrophy of Kupffer's cells, and moderate broaden out of the proliferating hypertrophied zones of tibial physes The decrease in spleen and heart weights, body weight gains, cholesterol	(Weibking et al., 1993b)
Duck	Orally; 0, 5, 15, 45 mg fumonisin B1/kg b.w	12 days		Body weight gain was slightly retarded, liver hyperplasia Increase in liver weight, total protein, AST, Sa/So, LDH, GGT, cholesterol	(Bailey et al., 2001)
Mouse embryos	Exposure of FB1	Long term Short-term		NTD; 65% in continuing experimentation and by almost 50% in temporary experimentation	(Sadler et al., 2002)
Mice	Subcutaneous; 2.25 mg fumonisin B1/kg b.w	5 days		Hepatotoxic effects, increase in AST and liver enzymes in circulation	(Sharma et al., 2000)
Mice	Dietary; 0, 14, 70, and 140µmol fumonisin B1, B2, B3, hydrolyzed fumonisin B1, fumonisin P1, N-(carboxymethyl)fumonisin B1 or N-(acetyl)fumonisin B1/kg	28 days		Increase in whole bile acids, alkaline phosphatase, cholesterol, hepatocellular apoptosis, macrophage pigmentation, Kupffer cell hyperplasia, and hepatocellular hypertrophy.	(Howard et al., 2002)
Mice	Gavage; 1-75 mg fumonisin B1/kg	14 days		In the liver, mitosis, anisokaryosis, and hepatocellular single cell necrosis Increase in ALT, serum cholesterol, blood urea nitrogen in male, vacuolated lymphocytes and myeloid cells Mild decreases in ion transport of kidney	(Bondy et al., 1997)
Mice	Dietary; 0, 1, 3, 9, 27, or 81 ppm FB1	13 weeks		Hepatopathy	(Voss et al., 1995)
Mice	150 mg/kg diet of FB1	16 weeks		Decreasing in number of parietal cells, gastric mucosa height and mitotic index in the gastric glands, Mild to moderate gastric atrophy, proliferative activity of	(Alizadeh et al., 2015)

				gastric glands lower than the control	
Female B6C3F1 mice	Fed 50 or 80 ppm FB1	2-year feeding		Hepatocellular adenomas and carcinomas	(Howard et al., 2001)
Mice	8 mg/kg, i.p. for	4 days		No changes in the; Indirect nitric oxide (NOx) content, TBARS, ascorbic acid, organ-to-body weight ratio, organ-to-adrenal gland weight ratio or organ-to-brain weight	(Dassi et al., 2018)
				Increasing in non-protein thiols (NPSH) levels in liver and lungs	
				decreaseing in Ferric reducing antioxidant power (FRAP) content in liver and kidneys	
Rat	Dietary; 0, 1, 3, 9, 27, or 81 ppm FB1	13 weeks		Nephrosis	(Voss et al., 1995)
Male BD IX rats	Intake of 50 ppm FB1	Up to 2 years	2	Culminated in the appearance of hepatocellular carcinomas and cholangiocarcinomas	(Gelderblom et al., 2001a)
Male F344 rats	FB1	2-year feeding		No hepatocarcinogenic effects ,but FB1 caused renal tubule tumors	(Howard et al., 2001)
Male BD IX rats	0.08 and 0.16 mg FB/100 g of (bw)/day over	2 years		Induce cancer, mild toxic, and preneoplastic lesions	(Gelderblom et al., 2001b)
Rabbit	Gavage; 0, 31.5, 630 mg fumonisin B1/kg b.w	Single dose		Increase in AP, ALT, AST, GGT, urea, total protein, and creatinine	(Orsi et al., 2009)
Rabbit	Gavage; 1.75 mg fumonisin B1/kg b.w	9,13 days		Focal small bilateral hemorrhages in the white matter cerebral, malacia, apoptosis in kidney and liver	(Bucci et al., 1996)
Rabbit	10 mg/kg fumonisin B1	4 weeks		Increasing in liver weight dramatically, change in active monovalent cation	(Szabó et al., 2014)
Horse	Intravenously; 1.25-4 , 1-4 mg fumonisin B1/kg b.w	33-35 days		Lesions of LEM Apathy, incoordination, walking into objects, changes in temperament, paralysis of the lips and tongue,	(Kellerman et al., 1990)
Horse	Intravenously; 0.125 mg fumonisin B1/kg b.w	0-9 days		Apathy, trembling, paresis of the lower lip and tongue, reluctance to move, a wide-based stance, ataxia, tetanic convulsion, inability to drink or eat Focal necrosis in the medulla oblongata and severe edema in brains, bilaterally symmetrical.	(Thiel et al., 1991)

Horse	Feeding; 160-3800 µg fumonisin B1/kg b.w 20-950 µg fumonisin B1/kg b.w			FB1 is the major fumonisin in LEM in horses	(Thiel et al., 1991)
Arabian horse	Dietary; 12.490 µg fumonisin B1/kg b.w, 5.251 µg fumonisin B2/kg b.w			Blindness, hyperexcitability, four leg ataxia, circling, aimless walking, death Focal areas of hemorrhage, softening of the sub-cortical white matter and brown-yellow discoloration Microscopic brain lesions; wide areas of malacia within the white matter of the brainstem, cerebral hemispheres, and cerebellum	(Giannitti et al., 2011)
Pony	Feeding; 1-88 ppm fumonisin B1, B2, B3	120 days		Leukoencephalomalacia and hepatic necrosis	(Ross et al., 1993)
Horses hoof cells	FB1 0.125–10 µg/mL		24 to 48 h	No effect on dermal or epidermal cells, increasing in supernatants of explants, reducing in lamellar integrity at noncytotoxic concentrations	(Reisinger et al., 2016)
Pigs	Intravenously; 4.6-7.9 mg fumonisin B1/kg b.w Orally; 48-166 ppm FB1	15 days		Pulmonary edema and hepatic necrosis	(Haschek et al., 1992)
Pigs	Dietary; 16 mg fumonisin B1/kg b.w			Hydrothorax, variably severe pulmonary edema, icterus and hepatocellular necrosis	(Colvin et al., 1993)
Pigs	Dietary; 20 ppm fumonisin B1	42 days		Strong edema in the lung, mild degenerative changes in the kidneys, slight edema in the different interior organs	(Pósa et al., 2016)
Pigs	Feeding; 10 mg/kg fumonisin B ₁		4 weeks	Higher sphinganine/sphingosine ratio and gained less weight	(Régnier et al., 2017)
Gilt	Dietary; 0.1 g fumonisin B1/kg b.w	7, 27-80 days		Nodular hyperplasia in liver, hyperkeratosis, parakeratosis, formation of papillary, hyperplastic plaques in esophageal mucosa	(Stan et al., 1993)
Weaned piglets	Orally; 5 mg fumonisin B1/kg b.w	Single dose		Increase in cholesterol, alkaline phosphatase and highest Sa and Sa/So ratios in plasma and urine	(Dilkin et al., 2010)

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3.1 Carcinogenicity

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Contamination of wheat, corn and rice with Fumonisin B can increase the risk of esophageal cancer in human (Stockmann-Juvala and Savolainen, 2008), (Alizadeh et al., 2012), (Sun et al., 2007) by stimulating the proliferation of human esophageal epithelial cells (HEECs) (Wang et al., 2014). Also,

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218 Mathur in 2001 observed some different effects of stimulation of the proliferation in liver cells
219 consisted of a proliferation of bile ductular cells and hepatocyte proliferation in cattle (Mathur et al.,
220 2001).

221 In rats, continuing intake of FB₁ (up to 2 years) consequences the introduction of renal tubule tumors,
222 hepatocellular adenomas, cholangiocarcinomas, and carcinomas (Gelderblom et al., 2001a; Howard
223 et al., 2001).

224

225 3.2 Hepatotoxic Effect

226 Fumonisin could create a mild hepatopathy in lambs (Edrington et al., 1995). Nonetheless
227 hepatotoxic effects of Fumonisin in cattle is more extensive than lamb, and consisted of increases in
228 gamma-glutamyl transpeptidase (GGT), lactate dehydrogenase (LDH), serum aspartate
229 aminotransferase (AST), cholesterol and bilirubin, and mild microscopic liver lesions (Osweiler et al.,
230 1993) hepatic lesions were distinguished by the different severity of disorganized hepatic cords and
231 hepatocyte apoptosis (Mathur et al., 2001). Therefore, there is a possibility that cattle are more
232 sensitive to Fumonisin than lamb.

233 Increasing dietary fumonisin B₁ caused the increase in liver weights, serum calcium, cholesterol, and
234 AST levels. Also, biliary hyperplasia and multifocal hepatic necrosis were present in broiler chicken
235 (Ledoux et al., 1992). In researches of (Kubena et al., 1997; Weibking et al., 1993a), chickens fed
236 with fumonisin B₁, sphinganine: sphingosine (Sa: So) ratio, serum glutamate oxaloacetate
237 aminotransaminase (SGOT), levels of free sphinganine in the serum, AST ratios, LDH, and GGT were
238 increased. Nonetheless, total liver lipids of chicks were decreased significantly. Subacute treatment of
239 broiler chicks with fumonisin B₁ bring about hepatic oxidative stress simultaneously by SA/SO
240 gathering. Also, TBARS (Thiobarbituric acid reactive substance) levels, catalase activity, and Vit C
241 content were increased (Poersch et al., 2014). Therefore, (according to the measure of treatment with
242 fumonisin) sensitivity to Fumonisin in broiler chicken is increased in comparison with the latest
243 research. In addition, hepatotoxic effects of Fumonisin besides of change in the level of liver
244 enzymes can influence other factors like Vit C content, TBARS, and even liver weight of broiler
245 chicken.

246 Feeding the turkey with fumonisin B₁ caused increases in liver weights and serum AST levels.
247 However, serum cholesterol, alkaline phosphatase, MCH (mean cell hemoglobin) and MCV (mean
248 cell volume) were declined. Likewise, hypertrophy of Kupffer's cells and biliary hyperplasia were
249 present in these turkeys (Weibking et al., 1993b).

250 In ducks FB₁ increased the level of cholesterol, total protein, alanine aminotransferase (ALT), LDH,
251 GGT and SA/SO (sphinganine to sphingosine ratio) in the plasma. Also, FB₁ caused to growth in liver
252 weight by liver hyperplasia (Bailly et al., 2001). These effect of fumonisin in ducks probably created
253 by SA to SO ratio and oxidative damages.

254 FB₁ in mice decreased Ferric reducing antioxidant power (FRAP) content in liver and increased; non-
255 protein thiols (NPSH) levels (Dassi et al., 2018) and liver enzymes like AST and ALT in circulation
256 (Sharma et al., 2000). Also, FB₁ increased serum levels of the total bile acids, alkaline phosphatase,
257 and cholesterol, and created microscopic effect such as hepatocellular hypertrophy, hepatocellular
258 apoptosis, Kupffer cell hyperplasia, hepatocellular single cell necrosis, mitosis, anisokaryosis, and
259 macrophage pigmentation (Bondy et al., 1997; Howard et al., 2002)

260 Effects of FB₁ in rabbits are significant increase in liver weight (Szabó et al., 2014), alkaline
261 phosphatase (AP), total protein, AST, ALT, and GGT. Forethermore, degeneration of hepatocytes and
262 apoptosis were the prominent degenerative changes in liver of rabbits (Bucci et al., 1996; Orsi et al.,
263 2009).

264 Because of fumonisin B₁, B₂, and B₃, hepatic necrosis in ponies occurred (Ross et al., 1993).

265 Effect of fumonisin in the liver of piglet was apoptosis, necrosis, hepatocyte proliferation, hyperplastic
266 hepatic nodules (in chronic studies), icterus, and hepatocellular necrosis. Besides serum cholesterol,
267 alkaline phosphatase, AST activities, and sphinganine and sphingosine concentrations in kidney,
268 heart, lung, and liver were elevated. But there were no detectable portal triads or central veins,
269 adjacent parenchyma, and the perilobular connective tissue was compressed (Colvin et al., 1993;
270 Dilkin et al., 2010; Haschek et al., 2001; Stan et al., 1993). The hepatic changes especially
271 disorganization in piglet by fumonisin probably is because of an acute pathway of this mycotoxin.

272

273 3.3 Kidney Toxicity

274 Fumonisin in the kidney of lambs revealed with tubular nephrosis (Edrington et al., 1995).

275 Accumulation of sphingosine and sphinganine in the kidney of calves created renal lesion like
276 vacuolar change, karyomegaly, apoptosis, dilatation of proximal renal tubules (that included protein
277 and cellular debris) and the proliferation of proximal renal tubular cells (Mathur et al., 2001).

278 Effect of fumonisin in the kidney of turkeys and broiler chicken was increasing in kidney weight (Henry
279 et al., 2000; Weibking et al., 1993a; Weibking et al., 1993b).
280 In both sexes of rats, fumonisins caused decreasing in the weight of kidney, nephrosis in outer
281 medulla (especially in female rats) (Voss et al., 1995). Ferric reducing antioxidant power (FRAP)
282 content in kidney of mice was decreased (Dassi et al., 2018).
283 (Bucci et al., 1996; Orsi et al., 2009) reported that the effect of fumonisin in the kidney of the rabbit
284 was apoptosis and degeneration of renal tubule epithelium, also level of urea and creatinine were
285 increased.
286 Fumonisin in the kidney of pigs created a mild degenerative change, and in the urine of pigs, the
287 highest Sa/So ratio and Sa ratio were produced in the 48th h (Dilkin et al., 2010; Pósa et al., 2016).
288 According to these studies, toxic effects of fumonisins in the kidney is not extensive such as liver and
289 sensitivity of kidney of rodents and chicken to fumonisins is lesser than other animals.

3.4 Leukoencephalomalacia

291 Fumonisin (especially fumonisin B₁) are the causal factor in the development of LEM in horses (Thiel
292 et al., 1991). The lethality rates, mortality, and morbidity in horses were 85.7%, 10%, and 11.6%
293 respectively (Giannitti et al., 2011).

294 In horses with LEM because of fumonisins, the brain lesions were observed such as; severe to early
295 bilaterally symmetrical edema of the brain, brown-yellow discoloration, focal necrosis in the medulla
296 oblongata, focal or multifocal areas of hemorrhage, sporadically pyknotic nucleus all over the areas of
297 rarefaction hemorrhage, softening of the sub-cortical white matter, cavitations crowded with
298 proteinaceous edema with rarefaction of the white matter, mild percolation by infrequent eosinophils
299 and neutrophils, intracytoplasmic eosinophilic globules, inflamed glial cells with plentiful eosinophilic
300 cytoplasm, inflamed glial cells with plentiful eosinophilic cytoplasm, cell edges were separated,
301 hyperchromatic, edema, necrosis, wide parts of malacia in the white matter of the cerebral
302 hemispheres, cerebellum, and brainstem (Giannitti et al., 2011; Kellerman et al., 1990; Thiel et al.,
303 1991). Perhaps these brain lesions (that were emerged by fumonisin in horses) leads horses to show
304 nervous signs, consisted mainly of; apathy, incoordination, walking into objects, changes in
305 temperament, just in one horse paralysis of the tongue and lips, paresis of tongue and the lower lip,
306 inability to drink or eat, a wide-based stance, reluctance to move, trembling, hyperexcitability, four leg
307 ataxia, blindness, tetanic convulsion, aimless walking and circling developed by death (Giannitti et al.,
308 2011; Kellerman et al., 1990; Thiel et al., 1991).

309 Fumonisin created leukoencephalomalacia in rabbits, and the bilateral brain microscopic lesions
310 consisted of small focal bleeding in the malacia, cerebral white matter, and bleeding in the
311 hippocampus (Bucci et al., 1996). However, brain lesions and nervous signs because of
312 leukoencephalomalacia in rabbits, is not as extensive and prevalent as horses. Therefore the brain of
313 horses is more sensitive than rabbits, to fumonisins.

3.5 Porcine Pulmonary Edema (PPE)

314 Usual damage of Fumonisin B in pigs was severe edema in the lung by inhibiting sphingolipid
315 biosynthesis, phagocytosis in pulmonary macrophages, and gathering of substance material in
316 pulmonary capillary endothelial cells (Haschek et al., 2001; Pósa et al., 2016).

317 The clinical signs in pigs because of pulmonary edema (induced by fumonisins) consisted of;
318 hydrothorax and respiratory distress (revealed by increasing respiratory rate and effort with open
319 mouth and abdominal breathing). Lethal pulmonary edema appears during 4 to 7 days after the daily
320 feed or intravenous treatment of FB₁ (Colvin et al., 1993; Haschek et al., 2001).

3.6 Other Toxic Effects

321 Exposure to FB₁ during the first trimester and before the pregnancy emerged to developed the hazard
322 of neural tube defects (NTD; because of the defeat of the neural tube to close, embryonic defects of
323 the spinal cord and brain happened) (Haschek et al., 1992; Missmer et al., 2006). Also, fumonisins in
324 human colon cells caused to main and early induction of lipid peroxidation, assessing IL-8 secretion,
325 and increasing in membrane microviscosity (Minervini et al., 2014).

326 Feeding by fumonisin in calves has some effects such as; impairing the lymphocyte blastogenesis
327 (Osweiler et al., 1993), decreasing in serum-induced granulosa cell (GC) proliferation (Albonico et al.,
328 2017), lethargy, increasing of sphingosine and sphinganine concentrations in the heart, lung, and
329 skeletal muscle. Raise in the concentration of sphinganine, but not sphingosine, in brains of managed
330 calves (Mathur et al., 2001) and have not effect on cell proliferation, progesterone production,
331 CYP11A1 and CYP19A1 gene expression of bovine granulosa cell (Albonico et al., 2017).

338 Diarrhea and lethargy detected in fumonisin administrated lambs (Edrington et al., 1995).
 339
 340 In broiler chicks, FB₁ had a bad effect on weight, water consumption, feed efficiency, and body (Henry
 341 et al., 2000). Also, fumonisins reduced villus height and crypt depth of the ileum, abundance of
 342 *Candidatus Savagella* and *Lactobacillus* spp (Antonissen et al., 2015), and body weight, but
 343 *Clostridium perfringens* content (reason of subclinical necrotic enteritis), the weight of bursa of
 344 Fabricius, gizzard, and proventriculus was increased. Other effects of FB₁ were diarrhea, thymic
 345 cortical atrophy, rickets (Henry et al., 2000; Ledoux et al., 1992), slightly inhibition in estradiol
 346 production (Antonissen et al., 2015), and elevation in level of serum nitric oxide (NO) (Lee et al.,
 347 2017).
 348 (Henry and Wyatt, 2001) reported that fumonisin in the egg could cause extreme hemorrhages of the
 349 thoracic area, head, neck of the dead embryos.

350
 351 Fumonisin B₁ in turkey appeared thymic cortical atrophy, and moderate enlarging of the proliferating
 352 and degenerating hypertrophied zones of tibial physis (Weibking et al., 1993b).

353
 354 In mice, fumonisins can cause adrenal cortical cell vacuolation and mild to moderate gastric atrophy,
 355 and may cause increases in serum cholesterol. Vacuolated lymphocytes and myeloid cells were also
 356 detected in mice due to fumonisins (Bondy et al., 1997). Also, fumonisins decreased number of
 357 parietal cells, proliferative activity of gastric glands, gastric mucosa height and mitotic index in the
 358 gastric glands (Alizadeh et al., 2015). In contrast, (Dassi et al., 2018) did not detect changes in the;
 359 indirect nitric oxide (NO_x) content, TBARS, ascorbic acid, organ-to-body weight ratio, organ-to-
 360 adrenal gland weight ratio or organ-to-brain weigh.

361 Fumonisin in pigs had some effects such as; decrease in left ventricular dP/dT (max) (an indicator of
 362 heart contractility). But mean pulmonary artery pressure, heart rate, mean systemic arterial pressure,
 363 cardiac output, and pulmonary artery wedge pressure (by obstruction of L-type Ca channels) were
 364 increased, by get up sphinganine and/or sphingosine mass. Also, in previous studies, parakeratosis,
 365 postpone in the pattern of papillary of the distal esophageal mucosa (part of stratum basale),
 366 hyperkeratosis, and hyperplastic nodules in the liver cell, esophageal plaques, and right ventricular
 367 hypertrophy were detected (Haschek et al., 2001; Régnier et al., 2017; Stan et al., 1993).
 368 Effects of fumonisins on hoof cells of horses was increasing in supernatants of explants, reducing in
 369 lamellar integrity at noncytotoxic concentrations, but fumonisins have not effect on dermal or
 370 epidermal cells (Reisinger et al., 2016).

371 372 4. DIETARY INTAKE

373 In the European diet, the total intake of FB₁ has been evaluated at 1.4 µg/kg of body weight/week
 374 (Soriano and Dragacci, 2004a). Daily intake of fumonisins in varies countries and foods, were
 375 summarized in Table 3.

376 In (Soriano and Dragacci, 2004a); (Creppy, 2002) articles, tolerable daily intake (TDI) of FB₁ was
 377 reported 800 ng/kg. Also, provisional-maximum-tolerable-daily-intake (PMTDI) of fumonisin was noted
 378 2 µg/kg of body weight per day by the no-observed-effect-level (NOEL) of 0.2 mg/kg of body
 379 weight/day and a safety aspect of one hundred.

380 Using the simulation model, mean concentrations of fumonisin B₁ in milk evaluated 0.36 µg/kg.
 381 Whenas the pretended TDI from milk for females and males fell lesser European Union guidelines
 382 (Coffey et al., 2009).

383 (Orsi et al., 2009) demonstrated that feces are the major way of excretion of fumonisin B₁ in rabbits,
 384 by comparing the concentration of FB₁ in urine, liver, and feces.

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 386 **Table 3. Daily intake of fumonisins for different countries and foods**

Food	Nation	Intake (ng/kg bw/day)	Explantion of	Reference
Beer	USA	20 to 54	Computed on the base of the 60 kg body weight	(Hlywka and Bullerman, 1999)
Cereal commodities	France	22.8	All children in france	(Soriano and Dragacci, 2004a)
Cereal commodities	France	4.6	All female adults in france	(Soriano and Dragacci, 2004a)
Cereal commodities	France	3.2	All male adults in france	(Soriano and Dragacci, 2004a)

Cereal commodities	France	9.96	All people in france	(Soriano and Dragacci, 2004a)
Cereal commodities	Germany	31.8	Users >14 years	(Soriano and Dragacci, 2004a)
Cereal commodities	Norway	430	6 month babies	(Soriano and Dragacci, 2004a)
Corn	Brazil	392	Camputed on the base of the 70 kg body weight from urban area	(Mac Jr and Valente Soares, 2000)
Corn	Brazil	1276	Camputed on the base of the 70 kg body weight from rural area people	(Mac Jr and Valente Soares, 2000)
Corn	Brazil	4.1	Conventional corn	(Ariño et al., 2007)
		3.4	Organic corn	
		3.8	Total	
Corn	France	45.6	All children in france	(Soriano and Dragacci, 2004a)
Corn	France	12.4	All female adults in france	(Soriano and Dragacci, 2004a)
Corn	France	7.4	All male adults in france	(Soriano and Dragacci, 2004a)
Corn	France	9.96	All people in france	(Soriano and Dragacci, 2004a)
Corn	Germany	8.7	Users >14 years	(Soriano and Dragacci, 2004a)
Corn	Switzerland	30		(Zoller et al., 1994)
Corn	The Netherlands	3.1	Adults	(Soriano and Dragacci, 2004a)
Corn	USA	80		(Humphreys et al., 2001)
Corn	USA	600000 to 2100000	Natural outbreak of LEM in horses	(Thiel et al., 1992)
Corn	Zimbabwe	140 and 5760	Shamva district	(Murashiki et al., 2017)
Corn	Zimbabwe	180 and 8092	Makoni district	(Murashiki et al., 2017)
Corn commodity	Brazil	63.3	São Paulo population	(Bordin et al., 2014)
Food with corn based	Argentina	0.73 to 2.29	Camputed on the base of the 70 kg body weight	(Torres et al., 2001)
Food with corn based	Brazil	maximum probable daily intake (MPDI): 256.07 average probable daily intake (APDI): 120.58		(Martins et al., 2012)
Food with corn based	Canada	89	All children	(Kuiper-Goodman et al., 1996)
Food with corn based	Canada	190	Child users	(Kuiper-Goodman et al., 1996)
Food with corn based	Denmark	400		(Petersen and Thorup, 2001)
Food with corn based	South Africa	14,000 to 440,000	A group of people exhibiting a high prevalence of human esophageal	(Thiel et al., 1992)
Food with corn based	South Africa	5,000 to 59,000	A group of people exhibiting a less prevalence of human esophageal	(Thiel et al., 1992)
Food with	UK	30		(Gregory et al., 1990)

corn based				
Corn inferred commodities	Belgium	16.7		(Soriano and Dragacci, 2004a)
Corn inferred commodities	China	450 to 15,810 (Mean=3020)	Computed on the base of the 50 kg body weight	(Li et al., 2001)
Corn inferred commodities	Germany	10.4	Users >14 years	(Soriano and Dragacci, 2004a)
Corn inferred commodities	Italy	185.6	Italian users	(Soriano and Dragacci, 2004a)
Corn inferred commodities	Italy	24.6	All people in Italy	(Soriano and Dragacci, 2004a)
Corn inferred commodities	Norway	0.24	Adult male and female population	(Soriano and Dragacci, 2004a)
Corn inferred commodities	Norway	0.50	Adult male and female users	(Soriano and Dragacci, 2004a)
Corn powder	Argentina	79 to 198	For samples during 1996/1997 and January 1998	(Hennigen et al., 2000)
Corn pieces	Germany	69.8	Users >14 years	(Soriano and Dragacci, 2004a)
Corn pieces	Italy	283.6	Italian users	(Soriano and Dragacci, 2004a)
Corn pieces	Italy	15.9	All people in Italy	(Soriano and Dragacci, 2004a)
Food	Mexico	0.4 (0-23.2)	User in state of Morelos	(Wild and Gong, 2009)
Food	Burkina Faso	0.8 (0-2.4)	All users	(Wild and Gong, 2009)
Food	South Africa	3.8	User in Transkei	(Wild and Gong, 2009)
Food	South Africa	0	User in KwaZulu-Natal	(Wild and Gong, 2009)
Food	Guatemala	3.5	Urban area	(Wild and Gong, 2009)
Food	Guatemala	15.6	Rural area	(Wild and Gong, 2009)
Food	Guatemala	0.2-23	All users	(Torres et al., 2013)
Rice	France	12.1	All children in france	(Soriano and Dragacci, 2004a)
Rice	France	5.6	All female adults in france	(Soriano and Dragacci, 2004a)
Rice	France	5.6	All male adults in france	(Soriano and Dragacci, 2004a)
Rice	France	5.7	All people in france	(Soriano and Dragacci, 2004a)
Rice	Germany	0.6	Users >14 years	(Soriano and Dragacci, 2004a)
Wheat commodities	France	345.1	All children in france	(Soriano and Dragacci, 2004a)
Wheat commodities	France	230.8	All female adults in france	(Soriano and Dragacci, 2004a)
Wheat commodities	France	256	All male adults in france	(Soriano and Dragacci, 2004a)
Wheat commodities	France	240.08	All people in france	(Soriano and Dragacci, 2004a)
Wheat commodities	Italy	62.1	Italian users	(Soriano and Dragacci, 2004a)
Wheat commodities	Italy	10.6	All people in Italy	(Soriano and Dragacci, 2004a)
Food and feeds	Germany	bad case scenario: 21,000 mean case scenario: 1,100	German users	(Zimmer et al., 2008)

387 **5. MAXIMUM LIMITATION**

388 There are different variables that may affect the foundation of tolerances for specific mycotoxins, such
 389 as the delivery of mycotoxins through products, regulations of trade contact in different countries,
 390 availability data of toxicological or dietary exposure, and the accessibility of techniques for analysis
 391 (Van Egmond, 1993).

392 Deadline level for fumonisins in maize and other cereals, at the moment change from 5 to 100000
 393 µg/kg. Present laws of fumonisins in feeds and foods set by nations from America, Africa, Europe,
 394 and Asia and described by (AC04318739, 2004); (Abdallah et al., 2015) are provided in Table 4.

395 **Table 4. Maximum limits for Fumonisin in feeds and foods in different countries (AC04318739,**
 396 **2004); (Abdallah et al., 2015)**

Country	Maximum limit (µg/kg)	Commodity
Bulgaria (FB1, FB2)	1000	Maize and processed products thereof
Cuba (FB1)	1000	Maize, rice
France (FB1)	3000	Cereals & cereal products
Iran (FB1, FB2)	1000	Maize
Singapore (FB1, FB2)	Not given	Corn & corn products
Switzerland (FB1, FB2)	1000	Maize
Taiwan (FB1)	Based on the result of risk evaluation	Maize commodities
USA (FB1, FB2, FB3)	2000	Disinfected dry milled corn commodities (e.g. corn grits, flaking grits, corn meal, corn flour with fat content of <2.25%, dry weight basis)
	3000	purified corn purpose of popcorn
	4000	Total of partially disinfected dry milled corn commodities (e.g. corn grits, flaking grits, corn meal, corn flour with fat content of <2.25%, dry weight basis); dehydrated milled corn bran; purified corn purpose of masa production
	5000	
	20000	Corn and corn derived purpose of rabbits and equids
	30000	Corn and corn derived purpose of catfish and swine
	60000	Corn and corn derived purpose of breeding mink, breeding poultry, and breeding ruminants (contains hens laying eggs and lactating dairy cattle for human use)
	100000	
	10000	Mink upbringing for pelt output and Ruminants >3 months old upbringing for slaughter
		Poultry upbringing for slaughter
	Pet animals and all other species or classes of livestock	
European Union fumonisins	2000 1000	Unprocessed maize Maize products for human
European Union (FB1, FB2)	50 5	Animal feeds except Equines Feeds of Equines
Food and Drug	30	Animal feeds except Equines

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6. DETOXIFICATION OF FUMONISINS

Strategies of detoxification for infected feeds and foods to diminish or remove the toxic effects of fumonisins by biological, physical, and chemical processes are essential to boost food safety, hinder financial damage, and recover infected commodities. Data detected on biological, physical, and chemical processes for detoxification of fumonisins were abbreviated in Table 5.

Table 5. Biological, physical, and chemical processes of fumonisins

Process	Observation	Reference
Biological process		
Lactic acid bacteria (Micrococcus luteus, acillus subtilis)	Binding to FB1 and FB2	(Scott, 2012)
Sphingopyxis sp.	Hydrolysis of FB1 to HFB1	(Scott, 2012)
Saccharomyces	Decrease in FB1 and FB2	(Scott et al., 1995).
Lactobacillus strains (L. plantarum B7 and L. pentosus X8)	Removing fumonisins (FB1 and FB2)	(Zhao et al., 2016)
Black yeasts Rhinoclodiella atrovirensa and Exophiala spinifera	Ester bonds was hydrolyzed of FB1	(Volcani Center, 2004)
Candida parapsilosis	Mycelial growth inhibition	(Fallah et al., 2016)
Physical process		
150–200 °C	87–100 % destruction of fumonisin B1 in corn cultures	(Volcani Center, 2004)
Extrusion and roasting	60–70 % loss of FB1 and FB2	(Scudamore, 2004)
Extrusion	30 % loss of FB1 and FB2	(Scudamore, 2004)
Extrusion	92 % loss of fumonisin B1	(Scudamore, 2004)
Extrusion	34-95% reduction of fumonisins	(Shanakhat et al., 2018)
Extrusion of drymilled products	Decrease in fumonisin accumulation by 30–90 % for mixing-type extruders and 20–50 % for non-mixing extruders	(Meister, 2001)
Baking corn	16 and 28 % loss of FB1	(Shapira and Paster, 2004)
Frying corn chips	loss of 67 % of the fumonisin	(Shapira and Paster, 2004)
Cooking and canning	Small influence on fumonisin measure (23%)	(Shephard et al., 2002)
Ethanol–water extraction solvent at 80 °C	The most environmentally friendly, least toxic, and cheapest	(Lawrence et al., 2000)
Cholestyramine	Adsorption 85% of FB1	(Solfrizzo et al., 2001b)
Activated carbon	Adsorption 62% of FB1	(Solfrizzo et al., 2001b)
Ammonia process	Reduce FB1 levels 30-45% No mutagenic potentials were apparent	(Norred et al., 1991)
Fructose	Obstruct the amine group of FB1, that is important for its toxicity	(Lu et al., 1997)
Chlorophorin	Reduced FB1 levels by 90–91%	(Beekrum et al., 2003)
Oxidizing agents	Little effects in FB1, but applicable because of the minimal cost and the minimal destruction of important	(Leibetseder, 2006)

	nutrients	
Bentonite	Adsorbed only 12% of the toxin FB1	(Solfrizzo et al., 2001b)
Celite	Not effective	(Solfrizzo et al., 2001b)
Chemical process		
Solution of SO ₂ at 60 °C for 6 h	Most impressive treatment to decline the measure of fumonisin B1	(Pujol et al., 1999)
Acidic aqueous solution by the addition of NaNO ₂ NaCl solution	Fumonisin B1 was significantly deaminated Fumonisin B1 had a little mass and that 86 % of the toxin could be eliminated	(Lemke et al., 2001) (Shetty and Bhat, 1999)
Ozone (O ₃)	No significant difference in FB1	(McKenzie et al., 1997)
Single (nixtamalization) or with Na-HCO ₃ + H ₂ O ₂ (modified nixtamalization)	Ca(OH) ₂ reduction of 100% FB1 and 40% decreased toxicity of brine shrimp by Ca	(Leibetseder, 2006)

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5.1 Biological Methods

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An enzymatic detoxification process is recombinant enzymes from the bacterium *Sphingopyxis* sp. resulted in hydrolysis of fumonisin B₁ to HFB₁; deamination of HFB₁ by aminotransferase (a miss of the two tricarballylic side-chains via carboxylesterase) in the existence of pyridoxal phosphate and pyruvate. Lactic acid bacteria such as *Micrococcus luteus* and *Bacillus subtilis* bind to fumonisin B₁ and fumonisin B₂. **Peptidoglycan** bind to atleast one tricarballylic acid part in the structure of FB₁ and especially FB₂ (Scott, 2012).

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***L. plantarum* MYS6 is having potent probiotic attributes and antifungal activity against fumonisin producing *F. proliferatum* MYS9.**(Deepthi et al., 2016)

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52.9% of FB₁ and 85.2% of FB₂ were removed by two *Lactobacillus* strains (*L. pentosus* X8 and *L. plantarum* B7), in the aqueous medium (Zhao et al., 2016).

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Fermentation using three different yeast strains (*Saccharomyces*) is a method for detoxification of fumonisins, thus a maximal decrease was observed in 28% and 17% for fumonisin B₁ and fumonisin B₂, respectively (Scott et al., 1995).

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Hydrolyzing ester bonds of fumonisin B₁ by black yeasts (*Exophiala spinifera* and *Rhinochloidiella atrovirensa*) reported by (Volcani Center, 2004).

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***Candida parapsilosis* could inhibit mycelial growth of *Fusarium* species from 74.54% and 56.36%, and the maximum and minimum decrease in total created fumonisin was 78% and 12%, respectively (Fallah et al., 2016). Therefore we can remove 17 to 85 % of Fumonisin using the biological process, and *Lactobacillus* knew as the most effective strains for detoxification of Fumonisin.**

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5.2 Physical and Chemical Methods

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Fumonisin B₁ needs a massive temperature (150–200 °C) to gain 87–100 % demolition in corn cultivation (Volcani Center, 2004).

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During extrusion of dry-milled products, decreasing in the measure of fumonisins was 20–50 % for non-mixing extruders and 30–90 % for mixing-type extruders (Saunders et al., 2001). For the production of cornflakes through the extrusion and roasting of raw corn, 60–70 % of fumonisins B₁ and B₂ were loosened. But removing of fumonisins only in the extrusion step was less than 30 % (Scudamore, 2004). Destroying of fumonisin B₁ in extrusion processing of grits, was 92 % (Scudamore, 2004). The economical, lowest toxic and most biodegradable solvent for fumonisin extraction is ethanol-water (Lawrence et al., 2000).

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In baking corn muffins, removing of fumonisin during **baking** for 20 minutes were amidst 16 and 28 % at 175 °C and 200 °C respectively. **In addition**, flotation the corn in water reduced the amount of fumonisin B₁, and frying corn chips for 15 minutes at 190 °C bring about a remove of 67 % of the fumonisin. **However**, spiked corn masa fried at 140–170 °C (while degradation begins to take placed above 180 °C) has not significant loss of fumonisin B₁ (Jackson et al., 1997),(Shapira and Paster, 2004).

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One of the most impressive management to decline the measure of fumonisin B1 is a 0.2 % solution of SO₂ at 60 °C for six hours (Pujol et al., 1999). **Nevertheless**, canning and cooking had a small influence on fumonisin measure (Shephard et al., 2002).

447 In (Solfrizzo et al., 2001b) studies, the adsorption capacity of cholestyramine for fumonisin B₁ was
448 85% from a solution including 200 µg/ml FB₁.
449 Detoxification of corn with ammonia process reduced fumonisin levels (30 to 45 %), and no mutagenic
450 potentials were found in the managed corn (Norred et al., 1991).
451 Obstruction the amine group of fumonisin B₁ by reaction with fructose is another way to the
452 detoxification of fumonisin B₁ (Lu et al., 1997).
453 The percentage of reduction of FB₁ in corn by single Ca(OH)₂ (nixtamalization) or with Na-HCO₃ +
454 H₂O₂ (modified nixtamalization), was 100% (Leibetseder, 2006).
455 Chlorophorin gets from vanillic acid, ferulic acid, caffeic acid, and iroko decreased FB₁ levels by 90–
456 91% (Beekrum et al., 2003).
457 Treatment with oxidizing agents is an economical method for detoxification of fumonisin B₁, but this
458 method isn't demonstrated in bioassays (Leibetseder, 2006).
459 The acidic aqueous solution such as NaNO₂ can create deamination in fumonisin B₁, significantly
460 (Lemke et al., 2001).
461 In the floating section after treatment with NaCl solution, 86% of FB₁ were removed (Shetty and Bhat,
462 1999).
463 Celite and O₃ couldn't make a significant difference in the level of FB₁, but bentonite adsorbed only
464 12% of the FB₁ (McKenzie et al., 1997; Solfrizzo et al., 2001b).
465 According to these reports, physical and chemical methods are the most effective way for
466 detoxification of Fumonisin (in comparison with the biological method), so intervention is necessary to
467 for remove the Fumonisin from feeds and foods.

468 CONCLUSION

469 Fumonisin can cause fatal diseases in animals and are classified as a potential human carcinogen.
470 In this paper, we review the aspects of studies concerning the ability of Fumonisin to cause varies
471 toxicity effects in different part of body in human and animal. We evaluate and compare the
472 occurrence of Fumonisin in many countries. We also evaluate the effects of different detoxification
473 method for removing the Fumonisin, mechanism of toxicity in cells of human and animals, the intake
474 of Fumonisin in various consumers, and compare the limitation of Fumonisin in several countries.
475 The authors suggestion for future investigation about Fumonisin are estimating the reproductive
476 effects of fumonisin, improving the information about the occurrence of fumonisin in different parts
477 of the world, extending masked Fumonisin in detoxification researches, improving the legislation
478 about Fumonisin to change daily intake of these mycotoxins, increasing attention to mechanisms of
479 Fumonisin on different types of animals and cells, cell-cell interactions, exposure pathway, and
480 exposure measures.

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