



# Mycotoxin Handbook



**NEOGEN'S MYCOTOXIN TESTS & APPROVALS**
**GIPSA AOAC**

Aflatoxin			
8015	<b>Reveal for Aflatoxin</b> – screens at 20 ppb, 25 samples.	✓	
8085	<b>Reveal Q+ for Aflatoxin</b> – range of 2–100 ppb, up to 25 samples.	✓	
8086	<b>Reveal Q+ for Aflatoxin Green</b> – (water-based extraction) range of 2–150 ppb, up to 25 samples.		
8088	<b>Reveal Q+ MAX for Aflatoxin</b> – range 2–300ppb, up to 25 samples.	✓	
8030	<b>Veratox for Aflatoxin</b> – range 5–50 ppb, up to 40 samples.	✓	✓
8031	<b>Veratox for Aflatoxin HS (High Sensitivity)</b> – range of 1–8 ppb, up to 38 samples.		
8032	<b>Veratox MAX for Total Aflatoxin HS (High Sensitivity)</b> – range of 1–10ppb, up to 38 samples.	✓	
8035	<b>Veratox MAX for Total Aflatoxin</b> – range of 5–50 ppb, up to 38 samples		
8019	<b>Veratox for Aflatoxin M1</b> – range of 5–100 ppt, up to 38 samples		
8043	<b>NeoColumn for Aflatoxin</b> – wide bore, clean-up column, 50 columns.		
DON (vomitoxin)			
8385	<b>Reveal Q+ for DON</b> – range of 0.3–6 ppm, up to 25 samples.	✓	
8388	<b>Reveal Q+ MAX for DON</b> – range 0.3–30 ppm, up to 25 samples.		
8335	<b>Veratox for DON 2/3</b> – range of 0.5–5.0 ppm, up to 38 samples.	✓	✓
8331	<b>Veratox for DON 5/5</b> – range of 0.5–5.0 ppm, up to 38 samples.	✓	
8332	<b>Veratox for DON HS</b> – range of 25–250 ppb, up to 38 samples.		
8340	<b>NeoColumn for DON</b> – Wide Bore, Clean-Up Column, 50 columns.		
Fumonisin			
8885	<b>Reveal Q+ for Fumonisin</b> – range of 0.3–6 ppm, up to 25 samples.	✓	
8888	<b>Reveal Q+ MAX for Fumonisin</b> – range 0.25–50ppm, up to 25 samples.		
8830	<b>Veratox for Fumonisin</b> – range of 1–6 ppm, up to 38 samples.		✓
8835	<b>Veratox for Fumonisin 5/10</b> – range of 0.5–6 ppm, up to 38 samples.		
8840	<b>Veratox for Fumonisin 5/5</b> – range of 0.25–6 ppm, up to 38 samples.		
8832	<b>Veratox for Fumonisin HS (High Sensitivity)</b> – range of 50–600 ppb, up to 38 samples.		
Ochratoxin			
8685	<b>Reveal Q+ for Ochratoxin</b> – range of 2–20 ppb, 25 samples.		
8688	<b>Reveal Q+ MAX for Ochratoxin</b> – range 2–100 ppb, up to 25 samples.		
8610	<b>Veratox for Ochratoxin</b> – range of 2–25 ppb, up to 38 samples.		
8630	<b>Veratox for Ochratoxin Grain</b> – range of 2–25 ppb, up to 38 samples.		
8640	<b>NeoColumn for Ochratoxin A</b> – wide bore, clean-up column, 50 columns.		
T-2/HT-2 Toxins			
8285	<b>Reveal Q+ for T-2/HT-2 Toxins</b> – range of 50–600 ppb, up to 25 samples.		
8288	<b>Reveal Q+ MAX for T2/HT2</b> – range 50–3000 ppb, up to 25 samples.		
8230	<b>Veratox for T-2/HT-2 Toxins</b> – range of 25–250 ppb, up to 38 samples.		
8240	<b>NeoColumn for T-2/HT-2</b> – wide bore, clean-up columns, 50 columns.		
Zearalenone			
8185	<b>Reveal Q+ for Zearalenone</b> – range of 50–1200 ppb, up to 25 samples.	✓	
8188	<b>Reveal Q+ MAX for Zearalenone</b> – range 25–1500 ppb, up to 25 samples.		
8110	<b>Veratox for Zearalenone</b> – range of 25–500 ppb, up to 38 samples.		
8140	<b>NeoColumn for Zearalenone</b> – wide bore, clean-up column, 50 columns.		

Updated approvals and validated commodities list available on request.

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## FREQUENTLY ASKED QUESTIONS

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### WHAT ARE MYCOTOXINS?

Mycotoxins are toxins produced by organisms categorized as fungi, including mushrooms, yeasts and molds. Fungi of one species or another, or their spores, can be found virtually everywhere. When the growth conditions are right for specific fungi, they will grow very rapidly into colonies, and produce toxins specific to that fungus as a by-product. Growth conditions, which include temperature, humidity, and available organic food sources, can not only affect whether or not a specific fungus will grow, but also the characteristics of the mycotoxin that it may produce.

Mycotoxins can be produced wherever fungi growth conditions exist, for example, in grains preharvest in the field and postharvest in storage. In either case, damage from insects, mishandling and environmental stress can enable the fungi to invade the grains' seeds.

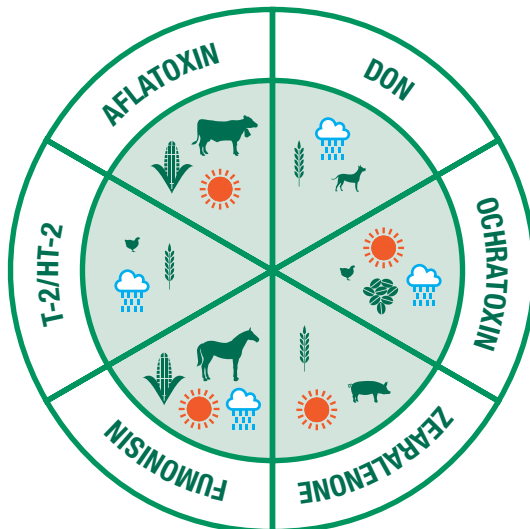
### ARE MYCOTOXINS HARMFUL?

As their name implies, mycotoxins are generally considered toxic, although not all mycotoxins are demonstrably toxic to every animal that may ingest them. Some mycotoxins, such as aflatoxin, have been shown to be dangerous to both humans and animals, others dangerous to only specific animal species, and still others, such as penicillin, only lethal to other fungi and bacteria.

Extensive mold growth in grains can have other obvious negative effects, such as producing changes in the grains' color, consistency, and smell—which may make the grains undesirable to livestock and as a human food. Mold growth can also rob grains of their fat, protein, and vitamin content, and lead to nutritional deficiencies in livestock.

### HOW MANY MYCOTOXINS ARE THERE?

Researchers have identified thousands of mycotoxins thus far, and continually identify new mycotoxins. Subtypes of numerous mycotoxins have also been identified. Within the identified mycotoxins and their subtypes, a relative few have been determined to pose a significant threat to the health of humans and animals. Those that have been proven to threaten health include aflatoxin, deoxynivalenol (a.k.a., DON or vomitoxin), fumonisin, ochratoxin, T-2/HT-2 toxins, and zearalenone.



### CAN MYCOTOXINS BE “KILLED” OR OTHERWISE NEUTRALIZED?

Unlike the fungi that produces them, mycotoxins are chemical substances that are not alive, and cannot be “killed”. The only known treatment to reduce aflatoxin levels, for example, is ammoniation, which leaves the kernels black and smelling like ammonia. There are no proven treatments to both neutralize a mycotoxin and preserve the integrity of the contaminated commodity.

Likewise, extreme heat and freezing do not destroy mycotoxins. Mycotoxins have also been shown to be resistant to breakdown in an animal’s digestive system—meaning that they can be passed along in meat and dairy products.

Commodities known to contain a harmful level of a certain mycotoxin are diverted away from use in products destined to be consumed by animals known to be especially sensitive to that mycotoxin. For example, corn products known to contain harmful amounts of fumonisin, a mycotoxin of special concern to horses and rabbits, would be diverted away from use in horse and rabbit feed.

### DO BLACK LIGHTS WORK TO DETECT AFLATOXIN IN CORN?

Studies have shown that using black light to detect aflatoxin in corn produces unreliable results. The bright yellow-green fluorescence that a black light can produce detects the presence of kojic acid, not aflatoxin. Kojic acid is one of many by-products of *Aspergillus flavus*, one of the two major producers of aflatoxin. But, *Aspergillus flavus* can produce aflatoxin without producing kojic acid, and it can produce kojic acid without producing aflatoxin. In addition, kojic acid can dissipate over time, thus a sample that once “glowed” may not at a later time.

Additionally, the other major producer of aflatoxin, *Aspergillus parasiticus*, does not produce kojic acid at all. So, while a black light procedure can seem to detect corn contaminated with aflatoxin at times, the procedure is an unreliable indicator for the presence of aflatoxin.

### HOW DO YOU OBTAIN A REPRESENTATIVE SAMPLE FROM A VERY LARGE QUANTITY, SUCH AS A HOPPER CAR?

Not even the best, most accurate test systems, such as Neogen’s, can detect the accurate level of possible mycotoxin contamination in a large load of a commodity if the sample tested is not representative of the entire load. Representative samples for mycotoxin testing are much more difficult to achieve because mycotoxin contamination tends not to be as evenly distributed in a load as other testing targets, such as protein, moisture, and fiber.

Carrier	Probe Length	Probes Per Compartment
Barges	12 feet	1
Hopper car	10–12 feet	1
Boxcar	6 feet	5
Truck	5–6 feet	7
Hopper-bottom truck	6–10 feet	2

Taken from the *The USDA Grain Inspection, Packers & Stockyards Administration (GIPSA) recommended sampling guidelines*.

For more information go to <http://www.gipsa.usda.gov/GIPSA/webapp?area=home&subject=lr&topic=hb-hb-1>

## MAJOR MYCOTOXINS

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### HOW FINE MUST A SAMPLE BE GROUND BEFORE TESTING?

The Federal Grain Inspection Service (FGIS) recommends grinding corn so that 95% passes through a 20 mesh screen, which is about the consistency of fine ground coffee. The sample's particle size is extremely important to subsequent test results. One kernel of corn can hold a very high amount of toxin. Unless kernels are ground and distributed evenly throughout the sample to be tested, variable and inaccurate results can occur. Proper cleaning of equipment between sampling is recommended to prevent cross contamination.

### WHAT IS A PPM?

"One part per million" is a lot to think about. Here are some facts that put 1 ppm into perspective.

- There are approximately 13,960 kernels of wheat in 1 pound. One kernel in 71 pounds is equal to 1 ppm.
- There are approximately 3,500,000–4,000,000 grains of sand per pound. If you take 4 grains out of the pound you have removed 1 ppm.
- "One part per billion" is 1,000 times smaller than 1 ppm. For example, one second in 32 years is 1 ppb.

## MAJOR MYCOTOXINS

The amount of mycotoxin required to produce adverse effects in humans and animals varies by the mycotoxin, and can even vary from animal to animal of the same species. The amount of risk posed by mycotoxins is a combination of the level of contamination of a given commodity, and the total amount of mycotoxins ingested by a specific animal.

### Aflatoxin

Aflatoxin is a toxic and carcinogenic substance produced by certain strains of the molds *Aspergillus flavus* and *A. parasiticus*. There are four principle types of aflatoxin: B<sub>1</sub>, B<sub>2</sub>, G<sub>1</sub> and G<sub>2</sub> in grains. Aflatoxin B<sub>1</sub> is the most frequently encountered of the group and the most toxic. The commodities most affected by aflatoxin are corn, peanuts, cottonseed, milo, and the majority of tree nuts.

The effects in animals of ingesting excessive amounts of the toxin range from chronic health and performance problems to death. Aflatoxin has been shown to cause liver damage or cancer, decreased milk and egg production, immune suppression and interference with reproductive efficiency.



Regulatory limits for aflatoxin are issued by regional authorities:

## Food

Region	Commodity	Maximum Level		
		B1	Total	M1
<b>US</b>	All foods except milk		20 ppb	
	Milk			0.5 ppb
<b>EU</b>	Cereals, processed	2 ppb	4 ppb	
	Dried fruits to be processed	5 ppb	10 ppb	
	Groundnuts and nuts to be processed	8 ppb	15 ppb	
	Groundnuts, nuts, and dried fruit for direct human consumption	2 ppb	4 ppb	
	Maize to be processed	5 ppb	10 ppb	
	Spices	5 ppb	10 ppb	
	Almonds, pistachios, and apricots for direct human consumption	8 ppb	10 ppb	
	Almonds, pistachios, and apricots to be processed	12 ppb	15 ppb	
	Dietary foods for special medical purposes intended specifically for infants	0.10 ppb		0.025 ppb
	Dried fruit for direct human consumption	2 ppb	4 ppb	
	Hazelnuts and Brazil nuts for direct human consumption	5 ppb	10 ppb	
	Hazelnuts and Brazil nuts to be processed	8 ppb	15 ppb	
	Infant formula and follow-on formula, including infant milk and follow-on milk			0.025 ppb
	Processed cereal-based foods and baby foods for infants and young children	0.10 ppb		
	Raw milk, heat-treated milk and milk for the manufacture of milk-based products	2 ppb	4 ppb	0.05 ppb
	Tree nuts for direct human consumption	5 ppb	10 ppb	
<b>Japan</b>	All food		10 ppb	
	Milk			0.5 ppb
<b>Indonesia</b>	Corn and its products	15 ppb	20 ppb	
	Spices	15 ppb	20 ppb	
	Dairy products			0.5 ppb
	Dried milk and related products			5 ppb

*Continued overleaf*

## MAJOR MYCOTOXINS

<b>Korea</b>	Grain, beans, peanut, nuts and their processed food (grinding, cutting, etc)	10 ppb	15 ppb	
	Processed cereal products and processed bean product	10 ppb	15 ppb	
	Nutmeg, turmeric, dried red pepper, dried paprika and spice products containing these	10 ppb	15 ppb	
	Wheat flour, dried fruits	10 ppb	15 ppb	
	Confectioneries (peanut or nut-containing food)	10 ppb	15 ppb	
	Processed corn products for popcorn	10 ppb	15 ppb	
	Soybean paste, red pepper paste, curry powder	10 ppb	15 ppb	
	Meju	10 ppb	15 ppb	
	Steamed rice	10 ppb	15 ppb	
	Baby foods for infants and young children	0.1 ppb		
	Raw milk and milk before processing			0.5 ppb
<b>Malaysia</b>	Groundnuts, almonds, hazelnuts, pistachios, Brazil nuts (shelled, for further processing)		15 ppb	
	Groundnuts, almonds, hazelnuts, pistachios, Brazil nuts (shelled, ready to eat)		10 ppb	
	Cereal based food for infants and children	0.1 ppb		
	Milk			0.5 ppb
	Infant formula and follow-up formula (ready to drink)			0.025 ppb



**Feed**

Region	Commodity	Maximum Level	
		B1	Total
<b>US</b>	Corn and peanut products intended for finishing beef cattle		300 ppb
	Cottonseed meal intended for beef cattle, swine, or poultry		300 ppb
	Corn and peanut products intended for finishing swine of 100 lbs or greater		200 ppb
	Corn and peanut products intended for breeding beef cattle, breeding swine, or mature poultry		100 ppb
	Corn, peanut products and other animal feeds and feed ingredients, excluding cottonseed meal, intended for immature animals		20 ppb
	Corn, corn products, cottonseed meal, and other animal feed and feed ingredients		20 ppb
<b>EU</b>	All feed materials	20 ppb	
	Complementary and complete feed	10 ppb	
	Compound feed for dairy cattle and calves, dairy sheep and lambs, dairy goats and kids and young poultry animals	5 ppb	
	Compound feed for cattle, sheep, goats, pigs and poultry not listed above	20 ppb	
<b>Japan</b>	Corn	20 ppb	
	Formula feed for cattle, pig, domestic fowl, quails	20 ppb	
	Formula feed for suckling piglet	20 ppb	
	Formula feed for dairy cattle	10 ppb	
<b>China</b>	Corn, peanut meal, cottonseed meal, rapeseed meal	50 ppb	
	Soybean meal	30 ppb	
	Complementary, complete and concentrated feeding stuff for piglets	10 ppb	
	Complementary, complete and concentrated feeding stuff for fattening pigs	20 ppb	
	Complementary, complete and concentrated feeding stuff for young broilers, chicks	10 ppb	
	Complementary, complete and concentrated feeding stuff for broilers, layers	20 ppb	
	Complementary, complete and concentrated feeding stuff for young ducks, ducklings	10 ppb	
	Complementary, complete and concentrated feeding stuff for ducks, layers	15 ppb	
	Complementary, complete and concentrated feeding stuff for quails	20 ppb	
	Supplementary feeding stuffs for dairy cattle	10 ppb	
	Supplementary feeding stuffs for beef cattle	50 ppb	
<b>Indonesia</b>	Feed and corn (final products)		50 ppb
	Feed for layer, broiler, and pigs		50 ppb
	Feed for quails		40 ppb
	Feed for ducks		20 ppb

## MAJOR MYCOTOXINS

### DON

Deoxynivalenol (DON) is most commonly produced by the pink mold *Fusarium graminearum*. DON, a member of the trichothecene family, is produced by fungi living on cereal commodities such as wheat, corn, barley and ensilages. The toxicological effects attributed to DON include: nausea (vomiting), feed refusal, gastroenteritis, diarrhea, immunosuppression and blood disorders.



Regulatory limits for DON are issued by regional authorities:

### Food

Region	Commodity	Maximum Level
		<b>Total</b>
<b>US</b>	Finished wheat products for consumption by humans	1,000 ppb
<b>Canada</b>	Uncleaned soft wheat for use in non-staple foods	2.0 mg/kg
	Uncleaned soft wheat for use in baby foods	1.0 mg/kg
<b>EU</b>	Bread, pastries, biscuits, cereal snacks and breakfast cereals	500 ppb
	Cereals for human consumption - cereal flour, bran, and germ	750 ppb
	Pasta (dried)	750 ppb
	Milling fractions of maize with particle size > 500 micron	750 ppb
	Milling fractions of maize with particle size < 500 micron	1,250 ppb
	Processed cereal-based food for babies	200 ppb
	Unprocessed cereals other than wheat durum, oats and maize	1,250 ppb
	Unprocessed durum wheat, oats, and maize not intended for wet milling	1,750 ppb
<b>Japan</b>	Wheat and wheat products	1,100 ppb (tentative)
<b>China</b>	Wheat and wheat products	1,000 ppb
	Corn and corn products	1,000 ppb
	Barley and barley products	1,000 ppb
<b>Indonesia</b>	Corn and its products	1,000 ppb
	Wheat and its products	1,000 ppb
	Wheat flour and its products (pastry, bakery, biscuits, snacks)	500 ppb
	Pasta and noodles	750 ppb
	Wheat based breast-milk substitute products	200 ppb

**Feed**

Region	Commodity	Maximum Level
<b>US</b>	Grains and grain by-products for ruminating beef and feedlot cattle older than 4 months, chickens	10 ppm
	Grain and grain by-products for swine	5 ppm
	Grain and grain by-products for other animals	5 ppm
<b>EU</b>	Cereals and cereal products with the exception of maize by-products	8 ppm
	Maize by-products	12 ppm
	Complementary and complete feeding stuffs	5 ppm
	Complementary and complete feeding stuffs for pigs	0.9 ppm
	Complementary and complete feeding stuffs for calves (<4 months), lambs, and kids	2 ppm
<b>Japan</b>	Formula feed (cows over 3 months after birth)	4,000 ppm
	Formula feed (except for cows over 3 months after birth)	1,000 ppm
<b>China</b>	Complementary and complete feeding stuffs for swine	<1 ppm
	Complementary and complete feeding stuffs for calves	<1 ppm
	Complementary and complete feeding stuffs for lactating animals	<1 ppm
	Complementary and complete feeding stuffs for cattle	< 5 ppm
	Complementary and complete feeding stuffs for poultry	< 5 ppm

## MAJOR MYCOTOXINS

### Fumonisin

Discovered in 1989, fumonisins are a family of mycotoxins produced by different species of the mold *Fusarium*. These molds commonly infect corn (in fact, they are considered ubiquitous in corn) and rice, hence the potential for fumonisins to be found in feed and foodstuffs is high. Fumonisin affects various animals differently and have been linked to esophageal cancer in humans. The Environmental Protection Agency classifies fumonisins as Category II-B carcinogens.



Horses are extremely sensitive to low amounts of fumonisin, which can cause leukoencephalomalacia (liquefaction of the brain). In swine, research has shown fumonisin attacks the cardiopulmonary system causing pulmonary edema, as well as liver and pancreatic lesions.

Regulatory limits for fumonisin are issued by regional authorities:

### Food

Region	Commodity	Guidance Level (B <sup>1</sup> , B <sup>2</sup> , B <sup>3</sup> )	Maximum Level (B <sup>1</sup> + B <sup>2</sup> )
		<b>Total</b>	
<b>US</b>	Degermed dry milled corn products (i.e. flaking grits, corn meal, corn flour)	2,000 ppb	
	Cleaned corn used for popcorn	3,000 ppb	
	Whole or partially degermed dry milled corn products; dry milled corn bran; cleaned corn used for mass production	4,000 ppb	
<b>EU</b>	Milling fractions of maize with particle size > 500 micron		1,400 ppb
	Milling fractions of maize with particle size < 500 micron		2,000 ppb
	Maize intended for human consumption		1,000 ppb
	Maize snacks, maize based breakfast cereals		800 ppb
	Processed maize-based foods for babies		200 ppb
	Unprocessed maize not intended for wet milling		4,000 ppb
<b>Indonesia</b>	Corn and corn as a raw material	2,000 ppb	
	Corn food products (flakes, popcorn, corn chips)	1,000 ppb	
<b>Korea</b>	Grain products & cereals (containing >50% corn, corn processed products, corn powder)	1,000 ppb	
	Processed corn products for popcorn	1,000 ppb	
	Confectioneries (containing > 50% corn)	1,000 ppb	
	Corn	4,000 ppb	
	Corn processed food (grinding, cutting etc) corn powder	2000 ppb	

**Feed**

Region	Commodity	Guidance Level (B <sup>1</sup> ,B <sup>2</sup> ,B <sup>3</sup> )	Maximum Level (B <sup>1</sup> +B <sup>2</sup> )
		<b>Total</b>	
<b>US</b>	Corn and corn by-products for equids and rabbits	5 ppm	
	Corn and corn by-products for swine and catfish	20 ppm	
	Corn and corn by-products for breeding ruminants, breeding poultry and breeding mink	30 ppm	
	Poultry raised for slaughter	100 ppm	
	All other species of livestock and pet animals	10 ppm	
<b>EU</b>	Maize and maize based products		60 ppm
	Complementary and complete feeding stuffs for pigs, horses, rabbits, and pet animals		5 ppm
	Complementary and complete feeding stuffs for fish		10 ppm
	Complementary and complete feeding stuffs for poultry, calves (<4 months), lambs and kits		20 ppm
	Complementary and complete feeding stuffs for adult ruminants (>4 months) and mink		50 ppm

### Ochratoxin

Ochratoxin, commonly produced by the molds *Aspergillus ochraceus* and *Penicillium viridicatum*, can be found in corn, barley, green coffee and various dried fruits. Ochratoxin may be present in conjunction with aflatoxin, one of the most potent naturally-occurring carcinogens. In fact, ochratoxin is a suspected carcinogen.

Ochratoxin affects kidneys in animals exposed to naturally-occurring levels of this mycotoxin. Turkeys and other poultry exhibited lower productivity levels during field outbreaks of ochratoxicosis. Symptoms included retarded growth and decreased feed conversion. It has also been known to affect egg production in laying hens.

Although there has been no advisory or regulatory level for ochratoxin issued by the Food and Drug Administration, many agree that levels between 10–20 parts per billion (ppb) for commodities destined for human or animal consumption may cause health problems and economic losses. Some international markets have set regulation limits ranging from 5 to 50 ppb.



Regulatory limits for ochratoxin are issued by regional authorities:

## Food

Region	Commodity	Maximum Level
<b>EU</b>	Dried vine fruits	10 ppb
	Processed cereals and cereal products	3 ppb
	Roasted coffee beans	5 ppb
	Soluble coffee	10 ppb
	Unprocessed cereals	5 ppb
	Wine, grape juice and grape must	2 ppb
	Dietary foods for special medical purposes intended specifically for infants	0.50 ppb
	Liquorice	20 ppb
	Liquorice for use in food	80 ppb
	Nutmeg, ginger, turmeric	15 ppb
	Processed cereal-based foods and baby foods for infants and young children	0.50 ppb
	Spices - capsicum, piper	30 ppb
	<b>Malaysia</b>	Cereal-based food for infants and children
<b>Korea</b>	Coffee or ground coffee or coffee powder	5 ppb
	Instant coffee or soluble coffee, decaffeinated coffee	10 ppb
<b>China</b>	Grains, beans, and their products	5 ppb
<b>Indonesia</b>	Cereal (rice, corn, sorghum, wheat) and their products	5 ppb
	Spices	20 ppb
	Coffee	5 ppb
	Instant coffee	10 ppb
	Dried raisins	10 ppb
	Grape juice	2 ppb
	Beer	0.2 ppb
	Cereal grain based breast-milk substitute products	0.5 ppb

## Feed

Region	Commodity	Maximum Level
<b>EU</b>	Cereal and cereal products	0.25 ppm
	Complementary and complete feeding stuffs for pigs	0.05 ppm
	Complementary and complete feeding stuffs for poultry	0.1 ppm
<b>China</b>	Complementary and complete feeding stuffs, corn	< 100 ppb

## MAJOR MYCOTOXINS

### T-2/HT-2 Toxins

T-2/HT-2 toxins are trichothecene mycotoxins produced by several species of *Fusarium* molds. As T-2 toxin is readily metabolized to HT-2 toxin, and the toxins have been shown to produce numerous adverse effects on many animals, these two mycotoxins are frequently evaluated together.

Animals affected by the toxins include swine, dairy cattle, poultry, dogs, cats and horses. Effects of the toxins include digestive disorders, hemorrhaging, edema, oral lesions, dermatitis, and blood disorders. Damage caused by the toxins to the digestive track is irreversible. In the most severe cases, these toxins will cause death. T-2 toxin is the principal causal toxin in the human disease alimentary toxic aleukia.

Poultry studies have shown T-2 intoxication has led to a reduction in weight gain and other problems such as beak lesions, poor feathering, motor function impairment and increased susceptibility to *Salmonella* spp.

The best protection against these mycotoxins is monitoring for their presence in feeds and foods. That means testing all along the pathway from initial harvest of grains to the finished product.

Regulatory limits for T-2/HT-2 are issued by regional authorities:



### Food

Region	Commodity	Maximum Level
EU	Unprocessed cereals: Barley (including malting barley)	200 ppb
	Oats ( with husk)	1,000 ppb
	Wheat, rye and other cereals	100 ppb
	Cereal grains for direct human consumption: Oats	200 ppb
	Maize	100 ppb
	Other cereals	50 ppb
	Cereal grains for human consumption: Oat bran and flaked oats	200 ppb
	Cereal bran except oat bran, oat milling products other than oat bran and flaked oats, and maize milling products	100 ppb
	Other cereal milling products	50 ppb
	Breakfast cereals including formed cereal flakes	75 ppb
	Bread, pastries, biscuits, cereal snacks, pasta	25 ppb
	Cereal based foods for infants and young children	15 ppb

### Feed

Region	Commodity	Maximum Level
EU	Oat milling products (husks)	2,000 ppb
	Other cereal products	500 ppb
	Compound feed, with the exception of feed for cats	250 ppb
	Compound feed for cats	50 ppb
China	Complementary and complete feeding stuff for swine	<1 ppm
	Complementary and complete feeding stuff for poultry	<1 ppm



### Zearalenone

Zearalenone is primarily produced by the mold *Fusarium graminearum*, which also commonly produces DON. Hence, there is evidence that if zearalenone is detected, there is a high probability that other fusarial mycotoxins may be present. Zearalenone is classified as an estrogenic mycotoxin because it frequently causes estrogenic responses in animals.

When zearalenone-contaminated feed or grain is eaten by livestock, it can cause a wide variety of reproductive problems. In swine, it causes vulvovaginitis, low birth weights, fetal reabsorption, aborted pregnancies, reduced litter sizes, abnormal estrus and feminization of immature males. The FDA has issued advisory levels for zearalenone at <500 ppb.



Regulatory limits for zearalenone are issued by regional authorities:

#### Food

Region	Commodity	Maximum Level
EU	Bread, pastries and biscuits	50 ppb
	Cereals for human consumption - cereal flour, bran and germ	75 ppb
	Refined maize oil	400 ppb
	Milling fractions of maize with particle size > 500 micron	200 ppb
	Milling fractions of maize with particle size < 500 micron	300 ppb
	Processed cereal and maize-based food for babies	20 ppb
	Unprocessed cereals other than maize	100 ppb
	Unprocessed maize not intended for wet milling	350 ppb
	Maize intended for human consumption, maize snacks, maize based breakfast cereals	100 ppb
Korea	Grain and their process foods (grinding, cutting etc)	200 ppb
	Confectioneries	50 ppb
	Baby foods for infants and young children	20 ppb
China	Wheat corn and their products	60 ppb

#### Feed

Region	Commodity	Maximum Level
EU	Cereal and cereal products with the exception of maize by-products	2 ppm
	Maize by-products	3 ppm
	Complementary and complete feeding stuffs for piglets, young sows, puppies, kittens, dogs and cats for reproduction	0.1 ppm
	Complementary and complete feeding stuffs for adult dogs and cats other than for reproduction	0.2 ppm
	Complementary and complete feeding stuffs for sows and fattening pigs	0.25 ppm
	Complementary and complete feeding stuffs for calves, dairy cattle, sheep and goats	0.5 ppm
Japan	Formula feed	1000 ppb
China	Complementary and complete feeding stuffs, corn	< 500 ppb

## CONFIRMATION METHODS

High performance liquid chromatography (HPLC) is the preferred instrument based confirmation method for mycotoxins. Testing requires a skilled technician, a validated test method, and appropriate equipment.

Gas chromatography (GC) and thin layer chromatography (TLC) are also popular confirmation methods.

## COMMODITY VALIDATION LISTS

Neogen's test kits have been validated on a variety of commodities from the most susceptible to the most obscure based on customer requests. Please contact a Neogen representative for the most up-to-date list.

## pH ADJUSTMENT PROCEDURE

Commodities to be tested should have a pH of 6.0–8.0. Most raw or unprocessed grains, such as corn or wheat, have a pH between 6.0–8.0 and will not need to be adjusted. To ensure the accuracy of subsequent testing, excessively acidic or alkaline samples should be adjusted using this method:

1. Grind and extract sample per the test kit's written instructions.
2. Filter 5 mL into a clean test tube.
3. Check pH with pH paper or meter.

**If acidic (pH is below 6):** Adjust the pH with 1N NaOH (sodium hydroxide) to 6.0–8.0. Add one drop of 1N NaOH to the sample extract, vortex or swirl to mix and re-check the pH. If still acidic add another drop and check pH. Continue until the pH is 6.0–8.0

**If alkaline (pH is above 8):** Adjust the pH with 1N HCl (hydrochloric acid) to 6.0–8.0. Add one drop of 1N HCl to the sample extract, vortex or swirl to mix and re-check the pH. If still alkaline add another drop and check pH. Continue until the pH is 6.0–8.0

4. The sample extract is now ready to test.

## SCREENING VS. QUANTIFYING RESULTS

Neogen's rapid tests for the detection of mycotoxins are available in multiple formats. Neogen's Reveal tests are the easiest available for those who require only a simple yes/no result, providing screening results in as little as 2 minutes. Neogen's Reveal Q+, Veratox and NeoColumn formats can provide screening results, or results in exact parts per million or billion, in just minutes. Each requires only a minimal amount of training and equipment.

### A. Screening tests

1. **Reveal** – Designed for ease of use, Reveal test kits are extremely easy to use and interpret test strips that screen samples against set thresholds. The AccuScan Pro lateral flow test reader provides an easy method to objectively read, store, and analyze results from Neogen's Reveal product line.

### B. Quantitative tests

1. **Reveal Q+ MAX** – A quantitative lateral flow device that utilizes a common water based extraction which enables users to test for up to six mycotoxins from one sample.
2. **Reveal Q+** – These quantitative lateral flow devices provide an easy-to-use rapid test with unparalleled accuracy.
3. **Veratox** – These quantitative tests compare up to 19 samples at a time against test controls. Through the use of a microwell reader, the tests provide accurate sample results in parts per million or billion.
4. **NeoColumn** – These immunoaffinity columns efficiently clean and concentrate the toxins prior to analysis by HPLC, fluorometric reader, or Neogen's Veratox test kits.

## TECHNICAL INFORMATION

### Sample Extraction:

Sample extraction is always performed at a specific ratio of solid sample to liquid extraction solution. Sample sizes can vary from the written instruction as long as the sample to extraction solution ratio remains the same.

### Example: 1:5 extraction ratio

Ground Sample	Extraction Solution
5 g	25 mL
10 g	50 mL
50 g	250 mL

More representative results are achieved with a greater sample size. For example the USDA-FGIS recommends using a 50 g sample for aflatoxin testing and blending to extract (50 g + 250 mL). However, using a blender to extract is not always feasible and by using a smaller sample, the process can be sped up and simplified by utilizing disposable extraction cups and supplies.

### Water-Based Extraction

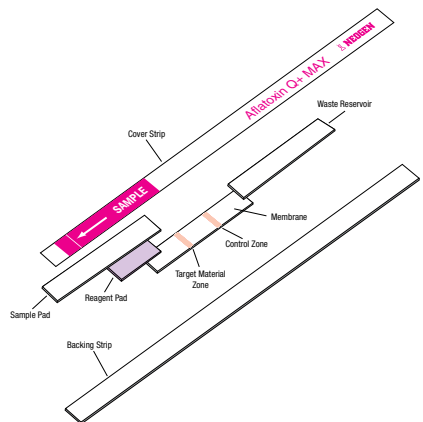
Neogen has test kits available that use a water-based extraction method. These kits simply require distilled or deionised water to be added to the extraction packets, offering a user and environmentally friendly extraction method. Tests include the Reveal Q+ MAX range, Veratox MAX for Total Aflatoxin and Veratox MAX HS for Total Aflatoxin. Please contact a Neogen representative for a complete list.

Please contact us or refer to the kit inserts for information on different sample methods.

## HOW DO NEOGEN'S MYCOTOXIN TESTS WORK?

### A. Reveal, Reveal Q+ and Reveal Q+ MAX

Neogen's Reveal tests for the detection of mycotoxins are lateral flow assays based on a competitive immunoassay format. The extract is wicked through a reagent zone, which contains antibodies specific for the target mycotoxin conjugated to colloidal gold particles. If the target mycotoxin is present, it will be captured by the particle-antibody complex. The mycotoxin-labeled antibody complex is then wicked onto a membrane, which contains a zone of mycotoxin. This zone captures any unbound mycotoxin antibody, allowing the particles to concentrate and form a visible line. As the level of mycotoxin in a sample increases, free mycotoxin will bind with the antibody-gold particles. This allows less antibody-gold to be captured in the test zone. Therefore, as the concentration of target mycotoxin in the sample increases, the test line density decreases. The membrane also contains a control line which will always form regardless of the presence of mycotoxin, ensuring the strip is functioning properly. For the Reveal Q+ and Reveal Q+ MAX tests, an AccuScan reader is utilized to convert the line densities into a quantitative result displayed in ppm or ppb.



## EXAMPLE PROCEDURE

### Example: Reveal Q+ MAX for Aflatoxin Test Procedure



1. Prepare by entering values into the Reveal AccuScan® Gold Reader



2. Obtain a representative sample. Grind and weigh out a 10 gram sample



3. Add contents of 1 MAX 1 packet to the sample



4. Add 50 mL distilled or deionized water to the sample



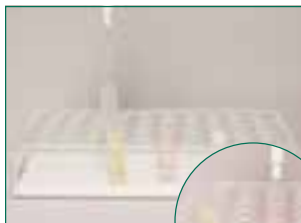
5. Shake vigorously for 3 minutes, or blend for 1 minute



6. Allow to settle, then filter



7. Add 100 µL of sample diluent to the red dilution cup



8. Add 100 µL sample extract to the red dilution cup and mix up and down 5 times



9. Transfer 100 µL diluted sample extract to sample cup



10. Place a new Reveal Q+MAX for Aflatoxin strip into the sample cup. Set a timer for 6 minutes



11. Remove promptly at 6 minutes and interpret results using the Reveal AccuScan® Gold reader.



12. Results shown on the AccuScan Gold

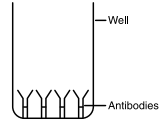
Please refer to kit insert for instructions and proper ratios for certain commodities.

Reveal, Reveal Q+ and Reveal Q+ MAX Quick Reference Guide					
Product Number	Test	Extraction	Ratio	Incubations (minutes)	Other
8015	Reveal for Aflatoxin	10 g sample, 20 mL of 70% MeOH. Shake vigorously or blend for 1 minute	1:2	3	Dilute 200 µL of sample diluent and 200 µL of sample extract.
8085	Reveal Q+ for Aflatoxin	10 g sample, 50 mL of 65% ethanol. Shake vigorously for 3 minutes or blend for 1 minute	1:5	6	Dilute 500 µL of sample diluent and 100 µL of sample extract.
8086	Reveal Q+ for Aflatoxin Green	10 g sample, 50 mL Green Extraction Solution. Shake vigorously for 3 minutes or blend for 1 minute	1:5	6	Dilute 500 µL of sample diluent and 100 µL of sample extract.
8088	Reveal Q+ Max Aflatoxin	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	6	For 2-50 ppb, dilute 100 µL sample diluent and 100 µL sample extract
8385	Reveal Q+ for DON	10 g sample in 100 mL of DI or distilled water. Shake 3 minutes, settle and filter	1:10	3	Dilute 1 mL (1,000 µL) of sample diluent and 100 µL of sample extract.
8388	Reveal Q+ Max DON	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	5	dilute 1000 µL sample diluent and 100 µL sample extract
8885	Reveal Q+ for Fumonisin	10 g sample in 50 mL of 65% ethanol. Shake 3 minutes, settle and filter	1:5	6	Dilute 200 µL of sample diluent and 100 µL of sample extract.
8888	Reveal Q+ Max Fumonisin	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	5	Dilute 900 µL of dilution buffer and 200 µL sample diluent. Then add 200 µL sample sample diluent and 100 µL diluted sample extract
8685	Reveal Q+ for Ochratoxin	10 g sample in 40mL of 70% methanol. Shake 3 minutes, settle and filter	1:5	9	Dilute 200 µL of sample diluent and 100 µL of sample extract.
8688	Reveal Q+ Max Ochratoxin	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL* deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	5	dilute 100 µL sample diluent and 200 µL of sample extract*
8285	Reveal Q+ for T-2/HT-2	10 g sample in 100 mL of distilled water. Shake 3 minutes, settle and filter	1:10	6	Dilute 500 µL of sample diluent and 100 µL of sample extract.
8288	Reveal Q+ Max T2/HT2	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	5	Dilute 1500 µL sample diluent and 100 µL sample extract
8185	Reveal Q+ for Zearalenone	Corn: 10 g in 30 mL of 65% ethanol Wheat: 10 g in 50 mL of 65% ethanol. Shake vigorously for 3 minutes or blend for 1 minute	Corn 1:3 Wheat 1:5	6	Dilute 200 µL of sample diluent and 100 µL of sample extract.
8188	Reveal Q+ Max Zearalenone	10 g sample, (1) MAX 1 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes or blend for 1 minute	1:5	5	dilute 100 µL sample diluent and 100 µL sample extract

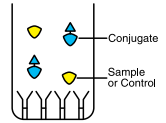
**B. Veratox**

Neogen's microwell mycotoxin tests are competitive direct enzyme-linked immunosorbent assays (CD-ELISAs). Each test kit contains antibody-coated microwells with antibodies specific to the kit's target mycotoxin. First, samples and controls are added to their respective test wells. Next, an enzyme conjugate (the target mycotoxin chemically linked with an enzyme) is added. The samples/controls and conjugate are mixed and transferred to antibody wells where they compete for the antibody binding sites. The more target substance in the sample, the less conjugate that binds in the wells. After an incubation, the wells are washed to remove all unbound materials. A substrate, which changes color in the presence of the conjugate, is then added to the wells. During an incubation, blue color develops in proportion to the amount of conjugate versus target mycotoxin in the wells. The more conjugate bound, the more blue color that develops, indicating less mycotoxin present. In the Veratox quantitative format, results are obtained by measuring the wells' color change in a microwell reader and comparing the readings against a standard curve.

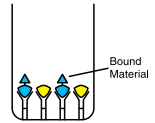
1. Microwells are coated with antibodies specific to the target substance



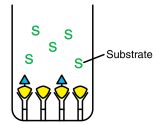
2. Conjugate competes with target substance/controls for antibody binding sites



3. Conjugate and target substance/controls remain bound in wells



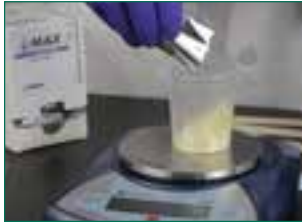
4. Substrate is added to produce a color change



5. Results are read visually or in a reader—the less blue color, or more red, the more target substance detected



**Example: Veratox MAX HS for Total Aflatoxin**



1. Weigh out 10 g sample, add one MAX 2 packet. Add 50 mL distilled or deionized water.



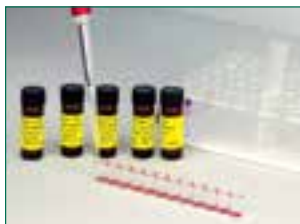
2. Shake for **3 minutes**. Allow to settle.



3. Filter using syringe.



4. Add 100  $\mu$ L conjugate to each red marked mixing well.



5. Add 100  $\mu$ L controls and samples to their respective wells.



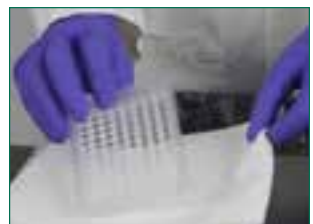
6. Mix. Transfer 100  $\mu$ L to antibody wells. Incubate at room temperature for **15 minutes**, sliding microwell holder back and forth gently for first **30 seconds**.



7. Dump liquid from antibody wells.



8. Wash wells thoroughly with deionized water. Repeat wash step five times.



9. Tap out water on absorbent paper towel.



10. Transfer 100  $\mu$ L substrate from the reagent boat to the antibody wells. Incubate at room temperature for **15 minutes**, sliding microwell holder back and forth gently for first **30 seconds**.



11. Transfer 100  $\mu$ L Red Stop from reagent boat to antibody wells.



12. Read results using a microwell reader with a 650 nm filter. Results should be read within 20 minutes of adding Red Stop.

## TECHNICAL INFORMATION

Please refer to kit insert for instructions and proper ratios for certain commodities.

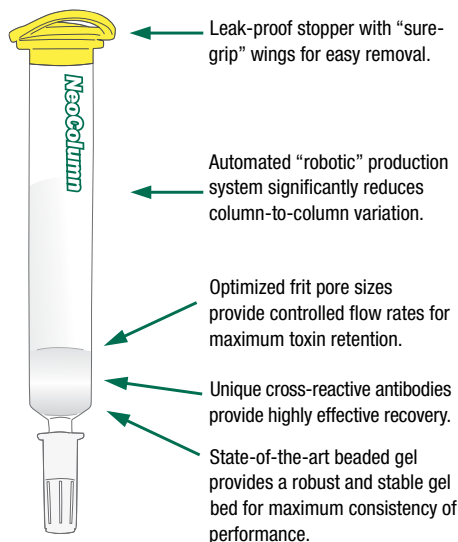
Veratox Quick Reference Guide						
Product Number	Test	Extraction	Ratio	Controls	Incubations (minutes)	Other
8030	Veratox for Aflatoxin	10 g sample, 50 mL of 70% MeOH. Shake vigorously for 3 minutes.	1:5	0, 5, 15, 50 (ppb)	2/3	None.
8031	Veratox HS for Aflatoxin	25 g sample, 125 mL of 70% MeOH. Blend for 2 minutes.	1:5	0, 1, 2, 4, 8 (ppb)	10/10	None.
8032	Veratox MAX for Total Aflatoxin HS	10 g sample, (1) MAX 2 aqueous extraction packet, 50 mL deionised water. Shake vigorously for 3 minutes.	1:5	0, 1, 3, 5, 10 (ppb)	15/15	None.
8035	Veratox MAX for Total Aflatoxin	10 g sample, (1) MAX 2 aqueous extraction packet, 50 mL deionised water, shake vigorously for 3 minutes.	1:5	0, 5, 15, 50 ppb	5/5	None.
8019	Veratox for Aflatoxin M1	Dried milk powders: 10g + 100 mL deionised water, extract on shaker for 30 minutes. Centrifuge for 10 minutes, and collect supernatant.	1:10	0, 5, 15, 30, 60, 100 ppt	45	None.
8331	Veratox for DON 5/5	10 g sample, 100 mL of DI or distilled water. Shake vigorously for 3 minutes.	1:10	0, 0.5, 1, 2, 6 (ppm)	5/5	None.
8335	Veratox for DON 2/3	10 g sample, 100 mL of DI or distilled water. Shake vigorously for 3 minutes.	1:10	0, 0.5, 1, 2, 6 (ppm)	2/3	None.
8830	Veratox for Fumonisin 10/10	10 g sample, 50 mL of 70% MeOH. Shake vigorously for 3 minutes.	1:5	0, 1, 2, 4, 6 (ppm)	10/10	100 µL of extract into dilution bottle.
8835	Veratox for Fumonisin 5/10	10 g sample, 50 mL of 70% MeOH. Shake vigorously for 3 minutes.	1:5	0, 0.5, 1, 3, 6 (ppm)	5/10	100 µL of extract into dilution bottle.
8840	Veratox for Fumonisin 5/5	10 g sample, 50 mL of DI water. Shake vigorously for 3 minutes.	1:5	0, 0.25, 1, 3, 6 (ppm)	5/5	100 µL of extract into 900 µL of diluent.
8610	Veratox for Ochratoxin	10 g sample, 40 mL of 50% MeOH. Shake for 3 minutes (wheat, barley and rye samples must be extracted in 70% MeOH).	1:4	0, 2, 5, 10, 25 (ppb)	10/10	None.
8630	Veratox for Ochratoxin Grain	10 g sample, 40 mL of 50% MeOH. Shake for 3 minutes.	1:4	0, 2, 5, 10, 25 (ppb)	10/10	None.
8230	Veratox for T-2/HT-2 Toxins	10 g sample, 50 mL of 70% MeOH. Shake vigorously for 3 minutes.	1:5	0, 25, 50, 100, 250 (ppb)	5/5	Dilute extract 1:1 in DI or distilled water.
8110	Veratox for Zearalenone	10 g sample, 50 mL of 70% MeOH. Shake vigorously for 3 minutes.	1:5	0, 25, 75, 150, 500 (ppb)	5/5	Dilute extract 1:5 in DI or distilled water.



### C. NeoColumn

The NeoColumn test format is a high performance immunoaffinity column designed for the clean-up and concentration of a sample prior to HPLC, GC-MS, ELISA and other analytical methods. Clean-up columns are available for aflatoxin in both narrow and wide bore columns and in the wide bore column for DON, ochratoxin A and zearalenone. These columns deliver highly accurate results and recoveries on a range of validated matrices.

NeoColumn for Aflatoxin DR is an affinity column immunoassay. Aflatoxin is extracted from a ground sample by blending and filtering. Extracted toxin in the filtrate is sampled and diluted with water. The diluted extract is filtered and applied to the column. Positive pressure is used to induce flow through the column allowing the antibody to capture any aflatoxin present. Then the column is washed to remove any non-bound materials. Bound aflatoxin is eluted using 100% methanol and collected in a test tube. Aflatoxin fluorescence is enhanced by the addition of a developer (bromine solution) and read in a calibrated fluorometer, which displays the concentration of aflatoxin.



NeoColumn Quick Reference Guide				
Product Number	Test	Limit of Detection	Recovery	Testing Time
8040 Narrow Bore	NeoColumn for Aflatoxin (for HPLC clean-up)	0.1 ppb	>90% each of B <sub>1</sub> , B <sub>2</sub> , G <sub>1</sub> , G <sub>2</sub> for cereal grains and nuts; conditions may vary depending on commodity	25 minutes
8043 Wide Bore	NeoColumn for Aflatoxin (for HPLC clean-up)	0.1 ppb	>90% each of B <sub>1</sub> , B <sub>2</sub> , G <sub>1</sub> , G <sub>2</sub> for cereal grains and nuts; conditions may vary depending on commodity	25 minutes
8047	NeoColumn for Aflatoxin DR (Direct-read & HPLC clean-up)	1 ppb	>90% B <sub>1</sub> >80% B <sub>2</sub> , G <sub>1</sub> , G <sub>2</sub>	5 minutes
8140	NeoColumn for Zearalenone	5 ppb	>90% for maize, wheat, animal feed and breakfast cereals; conditions may vary depending on commodity	20 minutes
8240	NeoColumn for T-2/HT-2 Toxins	125 ppb	≥95%	100 minutes
8340	NeoColumn for DON	0.1 ppm	>85%	25 minutes
8640	NeoColumn for Ochratoxin A	<0.1 ppb	>95% for cereals; conditions may vary depending on commodity	30 minutes

## EXAMPLE PROCEDURE

### Example: NeoColumn for Aflatoxin WB Test Procedure



1. Add 20 mL of prepared extract to the reservoir



2. Remove the bottom cap of the column to initiate the flow dropwise, allowing entire sample to pass through to top of frit. Replace the bottom cap on the column.



3. Add 2-3 mL of 25% methanol, and then reattach the reservoir/adaptor.



4. Add 20 mL of 25% methanol.



5. Remove the bottom cap and allow the wash to flow through the column.



6. Ensure all liquid is removed from the column.



7. Slowly elute the bound aflatoxin from the column by passing 2 mL of 50:50 acetonitrile/HPLC methanol through the column dropwise. Collect eluant into a clean glass vial.



8. Add 2 mL of HPLC grade water to the column, pushing through dropwise. The total elution volume now will be 4 mL.

Product No.	Product	Reveal	Reveal Q+	Reveal Q+ MAX	Veratox	NeoColumn
9401	Grinder	•	•	•	•	•
9427	Scale	•	•	•	•	
9428	Extraction container	•	•	•	•	•
9447 / 9368	Graduated cylinder	•	•	•	•	•
NA	DI or distilled water	•	•	•	•	•
9420	Filter syringe or equivalent	•	•	•	•	•
9421	Sample collection tube		•	•	•	•
9475	Sample cup rack	•	•	•		•
9402	Well holder				•	
9400	Wash bottle				•	
9426	Timer	•	•	•	•	•
9278 / 9272	100 µl pipettor	•	•	•	•	
9273	12-channel pipettor				•	
9407 / 9410	Pipette tips and rack	•	•	•	•	
9595	AccuScan Gold Reader	optional	•	•		
9303	Neogen 4700 Microwell Reader				•	
N/A	Plate rotary shaker				•*	
N/A	Vortex				•*	
N/A	Centrifuge				•*	
8089	MAX1 Extraction Packets			•		
8036	MAX2 Extraction Packets				•**	

\* Veratox Aflatoxin M1 only

\*\* Veratox MAX for Total Aflatoxin only

### Is the particle size of a ground sample important?

Yes it is. Mycotoxins are embedded in different grain particles, and need to be released so that they are properly extracted. USDA set a grind size so that 95% of the material goes through a mesh sieve and this should have the consistency of instant coffee.



### What does MRM stand for and why should I use them?

MRM: Mycotoxin Reference Material. This is a naturally contaminated sample which has been verified by multiple methods, usually HPLC/LCMS. Technicians can then use that sample as if it were an unknown and then determine if they are getting the correct result. These types of samples are offered by Neogen.

### What's the difference between mycotoxin advisory limits and regulatory limits?

Advisory limits are just that; advisory levels set by experts within the regulatory community like the FDA, advising on limits for mycotoxins that would be agricultural best practices. However, being above those limits is not cause for recall. This is different from a regulatory limit which the FDA etc can set in order to institute a recall if that particular foodstuff is over that limit. Users should be aware of regulatory limits where they are operating but also where they are exporting to.



### What is the typical percentage CV of mycotoxin levels from rapid test kits?

10–15% CV or relative standard deviation is the typical %CV. But remember that test kits are only a portion of the total error contributed during analysis. Method variation also takes into account the technician, instrumentation or equipment itself, as well as sampling variability. These all come together as the total error for the method.

### How do we know if an analyst is proficient?

As a check on their proficiency, analysts should participate in a proficiency program. Neogen offers programs that provide proficiency testing samples on an annual or quarterly basis. These samples are tested as normal, and their results are reported through an online portal. The analysts' proficiency is then gauged against other users of that test method by determining how closely the analyst came to the correct answer, and the overall variability of the analyst's test results.



### How long is a sample valid for before running a test?

Because the mycotoxins in the sample aren't soluble for an infinite period of time, we typically recommend within 4 hours of extraction the analysis should be performed. Generally this means that if it's extracted in the morning, analysis should be carried out before lunch, likewise if the sample is extracted in the afternoon perform the analysis before going home. For that same reason, the solubility issue, we generally recommend not saving those extracts for extended periods of time to be analysed later, because the value can slowly decline overtime and not be indicative of the actual result.



**How large a sample should I grind?**

Refer to the GIPSA or your regional grain inspection handbook. These are very specific guidelines depending on the size of the vessel that specify exactly what size of a sample to collect. For trucks and container 2 lbs is typical, railcars is 3 lbs, and barges etc is 10 lbs. recommend that the entire sample be ground together so that it is representative of the whole lot.

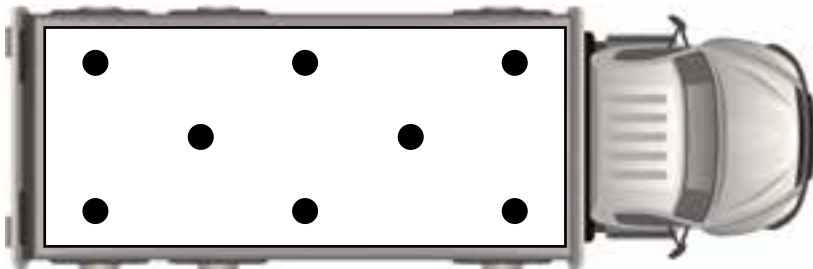
Commodity	Official Lot Type Minimum Sample Size (lbs. / grams)			Submitted Samples lbs. / grams
	Trucks or Containers	Railcars	Barges, Sublots & Composite Samples	
Aflatoxin	2 lbs./908 grams	3 lbs./1,362 grams	10 lbs./4,540 grams	10 lbs./4,450 grams (Recommended)
Deoxynivalenol (DON)	200 grams 2 lbs./908 grams (corn only)	200 grams 1000 grams (corn only)	200 grams 1000 grams (corn only)	200 grams 1000 grams (corn only) (Recommended)
Zearalenone	2 lbs./908 grams	3 lbs./1,362 grams	3 lbs./1,362 grams	10 lbs./4,540 grams (Recommended)
Fumonisin	2 lbs./908 grams	3 lbs./1,362 grams	3 lbs./1,362 grams	10 lbs./4,540 grams (Recommended)

**Why do I get different results on samples taken from the same truckload?**

A sample that is tested may not be fully representative of the total. Always encourage that a standard GIPSA or regional authority sampling plan be adhered to, so that the testing that happens from one lab to the next can be consistent. Typically, also, when comparing labs, its good practice to test from the same ground sample to ensure consistency in analysis.

**How many times should I probe a truck of grain to get a representative sample?**

GIPSA recommends that a multi-tier probe be used, and each of those probes have individual compartments within them. Depending on the size of the vessel the number of compartments can vary.



To obtain a representative sample a truck should be probed according to this pattern. Following this will ensure you have a representative sample and results generated are reliable.

**Why is there a wide range of acceptance for reference material?**

There are different sources of variance for particular testing – method itself, technician, equipment, sample. Samples are very well characterised by HPLC methods, but to account for other sources of variance a wider margin of error is given to the individual technician to be able to demonstrate proficiency when using that reference material.

### **What does LOQ stand for and why is it important?**

Limit of quantification (LOQ) is the lowest concentration of analyte in a sample that can be determined with acceptable precision and accuracy following the kit insert protocol. The limit of detection (LOD) is the lowest quantity of a substance that can be distinguished from the absence of that substance (blank value) within a stated confidence limit.

### **Helpful hints and pipetting techniques**

1. Swirl, don't shake, all reagents before using. Otherwise, the reagents will foam.
2. Always change pipette tips when there is a change in the reagent.
3. Veratox test kits should be stored at 2–8°C (35–46°F) but allowed to warm to ambient temperature, 18–30°C (64–86°C) before use to ensure optimum performance.
4. Prime pipette tips prior to dispensing all reagents. To prime the tip, draw up the reagent and discharge it back into the same container. Priming the tips coats the inside of the pipette tip so that the volume dispensed will be identical regardless of tip wetting properties.
5. When drawing or dispensing reagents, always drag the pipette tip against the container rim to remove liquid on the outside of the tip.
6. When dispensing reagents into the microwells, place the tip point against the inside wall of the micro well. This helps draw all of the liquid out of the tip and eliminates drops that form on the end of the pipette tip. In addition, placing the tip against the microwell holds the tip in place as the liquid is dispensed.
7. Always check the fluid levels in your tips prior to dispensing to be sure that the same amount is being collected each time (100 µL). If the proper amount was not collected or bubbles are present, refill the tip.
8. Most pipettors should be lubricated and calibrated at least every 12 months.
9. If a sample result is greater than the kit's stated range of quantitation (many times the kit's highest control), it is not considered a valid result. For accurate results you must dilute and rerun the sample as per kit insert.

**RESOURCES**

Canadian Grain Commission; [www.grainscanada.gc.ca](http://www.grainscanada.gc.ca)  
North American Miller’s Association (NAMA) 202/484-2200; [www.namamillers.org](http://www.namamillers.org)  
USDA GIPSA; [www.gipsa.usda.gov](http://www.gipsa.usda.gov) (Grain Inspection, Packers and Stockyards Administration)  
FAPAS® Central Science Laboratory, Sand Hutton, York, UK; Tel: (+44) 1904 462100; [www.fapas.com](http://www.fapas.com)  
North Dakota State University, Veterinary Diagnostic Laboratory; 701/231-8307; [www.vdl.ndsu.edu](http://www.vdl.ndsu.edu)  
AOAC International; [www.aoac.org](http://www.aoac.org)  
European Commission - Legislation on mycotoxins - <https://ec.europa.eu>  
European Food Safety Authority - Mycotoxins - <https://www.efsa.europa.eu>

**NOTES**

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