

Matrix and Detection Summary:

Matrix Group ID	Protocol	Results reported in the range of:	Limit of Detection (LOD)*	Highest Approved Level*
FM MG1 - Corn	Base Range	0 - 10 ppm	0.10 ppm	10 ppm
	Dilution A	0 - >100 ppm	2.0 ppm	100 ppm
FM MG2 - DDGS	Base Range	0 - 30 ppm	0.20 ppm	30 ppm
	Dilution A	0 - >100 ppm	30 ppm	100 ppm
FM MG4 - Sorghum	Base Range	0 - 10 ppm	0.20 ppm	10 ppm
	Dilution A	0 - >100 ppm	10 ppm	100 ppm

**Do not assume accuracy for results reported below the protocol's LOD or above the protocol's highest approved level.*

Important Notes:

- Before testing, the enclosed Multi-Matrix Barcode Card (MMBC) must be scanned just once for each kit lot to upload information to the QuickScan
- QuickScan Software Version 5.1.1 Update 1 or later is required

A Summary Guide for testing is provided on Page 10-11. More details for each step in the process are described below and are important for achieving optimal, accurate results.

Contents of Kit:

- 50 TotalTox Strips packed in a moisture-resistant canister
- 50 reaction tubes
- 100 pipette tips (1-200 µL)
- DB5 Buffer
- Multi-Matrix Barcode Card - kit lot specific

Matrices

Note: Scanning the Multi-Matrix Barcode Card once per kit lot is required. The QuickScan software will prompt users to select a Matrix Group (MG) before proceeding to the result screen. **If you only plan to test matrices within the MG1 group (Corn), scan the side of the MMBC card that only has the MG1 barcode. This allows the software to skip the Matrix Group selection step.**

- | | | | |
|---|--------------------------------|--|--------------------------------|
| <ul style="list-style-type: none"> ▪ Corn ▪ Sorghum | SET A PROCEDURE: PAGE 5 | <ul style="list-style-type: none"> ▪ DDGS | SET B PROCEDURE: PAGE 6 |
|---|--------------------------------|--|--------------------------------|

Intended Use

TotalTox™ Fumonisin is designed to quickly provide quantitative results for the presence of total fumonisins. Matrix and Detection Summary on Page 1 lists the Limit of detection (LOD) and Assay range for each matrix.

How the Test Works

A composite sample is collected, ground, and extracted to solubilize any fumonisin present. The extract is further diluted into Buffer before being run on the test strip. Each strip has an absorbent pad at each end. The sample extract travels up the test strip and is absorbed into the larger pad at the top of the strip. At the end of the reaction time, the strip is cut at the top of the arrow tape, the bottom pads are discarded, and the strip is inserted into the QuickScan reader to obtain quantitative results.

Matrix specific extractions and analysis protocols are chosen for accuracy and precision. Each matrix is assigned to a Matrix Group (MG). Each MG has a common standard curve, Limit of Detection (LOD), and maximum reported value. When the user selects the MG during testing, the QuickScan System software reads the test strip, retrieves information encoded in the strip's barcode and on the Multi-Matrix Barcode Card (MMBC), and uses the appropriate curve to obtain a result for the matrix being tested.



Precautions – Read First!

SAFETY

1. **Disposal of fumonisin-contaminated materials.** Follow your facility's safety procedures for disposal of samples and extracts potentially containing or known to contain fumonisin.

Items Not Provided:	*Available Accessories:		
<ul style="list-style-type: none"> • 50 EB17 dissolvable pouches (1 pkt per 25g sample) • QuickScan System* • Incubator (base + block)* • Bunn grinder or equivalent • 20-mesh screen (available through Seedburo or other vendor) • Digital scale for weighing samples • Extraction cups with lids* or other suitable vessels for sample extraction • Graduated cylinder* • Orbital/rotary shaker • Pipette to deliver 100 µL* • Pipette to deliver 50 µL (for Dilution A if desired)* • Pipette to deliver larger volumes (for Dilution A if desired)* • EB18 Extraction Buffer* for DDGS (see Notes, p.4, for more info) • Timer • Scissors • Distilled, deionized or bottled water 	<i>Item</i>	<i>Catalog No.</i>	<i>Part #</i>
	QuickScan™ System	ACC 331	12721
	EB17 Extraction Buffer pouches, 50	ACC 117	12938
	5 oz Sample cups/lids <i>Case of 500; for extracting samples up to 30g</i>	20-0047	10167
	10 oz Sample cups/lids <i>Case of 100; for extracting samples >30g</i>	20-0129	12383
	Graduated cylinder (100 mL)	ACC 068	11207
	MiniPet pipette 100 µL (one/location free)	ACC 041	11203
	EB18 Extraction Buffer 10X Concentrate <i>See instructions and SDS under 'Notes'</i>	KR 270-530	11930
	Coffee filters (100)	ACC 083	11434
	Centrifugation Set: <i>Disposables for 50 tests</i>	ACC 010	11214
	Microcentrifuge	ACC 064 E	11204
	50g Sample Extraction Set <i>Additional EB17 dissolvable pouches and sample cups (100)</i>	ACC 099	12409
	DB5 Buffer <i>Additional Buffer for DDGS, requires > 100 µL per Strip</i>	KR-266-7	11665
	Dilution Set: <i>Blue dilution tubes and EB17 dissolvable pouches for 50 tests</i>	ACC 103	12500
	Dilution Tubes: <i>Blue dilution tubes for non-EB17 dilution, 50</i>	ACC 098	12236
	MiniPet pipette 50 µL	ACC 051	11203
	1 mL adjustable pipette	ACC 1303-PRO-1000	11964
	Pipette tips for 1 mL pipette (50)	20-0127	12243
	Incubator	ACC BSH301	12458
*Available as Accessories			

Sample Preparation

1. Collect a composite sample according to your own sampling plan or USDA/GIPSA guidelines. Consult USDA/GIPSA reference documents to help design a plan that fits your needs. Contact Technical Support for more information.
2. Grind samples to provide a consistency such that 95% passes through a 20-mesh sieve.
3. Mix ground material thoroughly before sub-sampling, to minimize variability.
4. Weigh 25-50g samples into containers that will allow enough head room for the liquid to move forcefully when shaken vigorously.

Sample Clarification

Depending on the sample matrix, there may be multiple acceptable methods for removing particulate from the extract. Refer to Matrix Group instructions (pages 5-6) or Summary Table (pages 11-12).

Centrifugation	Filtration
1. Fill a microcentrifuge tube with extract.	1. Add an approved coffee filter (e.g. BUNN Part #BUNBCF100B) to a clean vessel.
2. Centrifuge for the specified time at 2000 x g (rcf, <i>not rpm</i>).	2. Pour extract into filter. Wait no more than 2 min.
3. Use the top layer of extract for all matrices except flour; there may be a white floating layer above that extract that should not be used for testing.	3. Pull back the filter to access the filtered extract.

Testing in Base Range

Refer to Matrix Group instructions (pages 5-6) or Summary Table (page 11-12) for base range testing.

Range with Dilution

If after running and reading the test, the initial result is greater than the upper end of the Base Range, samples can be diluted and retested to extend quantitation (see Summary on Page 1). Combine extract with the appropriate Dilution Reagent to create a diluted extract. Measure carefully and mix well.

Dilution Reagent

<i>Corn/MG1; Sorghum/MG4</i>	<i>DDGS/MG2</i>
EB17 Dilution Solution: Dissolve 1 EB17 pouch in 150 mL of water and mix well; Dilution Solution mixture will appear cloudy. Label, date, and document the preparation. Dilution Solution can be stored at ambient temperature for 30 days. Thoroughly mix before use.	EB18 Dilution Solution: Dissolve 1 EB17 pouch in 150 mL of EB18 and mix well; Dilution Solution mixture will appear cloudy. Label, date, and document the preparation. Dilution Solution can be stored at ambient temperature for 30 days. Thoroughly mix before use.

Corn/Sorghum Extended Range: Testing samples at levels greater than 10 ppm (> 10 ppm in Base Range)

- A1. Mix 700 µL Dilution Reagent + 50 µL clarified extract in a blue Dilution Tube or other suitable vessel.
- A2. Rerun assay as before (see Page 5). Example: for corn, pipette 100 µL DB5 + 100 µL of the diluted extract into a new reaction tube; place tube in 22°C incubator for 2 min[^], add a new test strip, and wait 4 minutes for test results. Note: for sorghum extended range, the diluted extract volume is 200 µL (see Summary Table page 10).
- A3. In the QuickScan Results Screen, use the dilution tab pull down menu to select Dilution A (1:A). The System will adjust and display the fumonisin level from diluted samples. Adjusted results are valid in the range of **2-100 ppm**.

DDGS Extended Range: Testing samples at levels greater than 30 ppm (> 30 ppm in Base Range)

- B1. Mix 400 µL Dilution Reagent + 100 µL clarified extract in a blue Dilution Tube or other suitable vessel.
- B2. Rerun assay as before (see Page 6). Example: for DDGS, pipette 200 µL DB5 + 100 µL of the diluted extract into a new reaction tube; place tube in 22°C incubator for 2 min[^], add a new test strip, and wait 5 minutes for test results.
- B3. In the QuickScan Results Screen, use the dilution tab pull down menu to select Dilution A (1:A). The System will adjust and display the fumonisin level from diluted samples. Adjusted results are valid in the range of **30-100 ppm**.

[^] The tube acclimation step is only required if the temperature of the testing environment is unknown or outside of 20 - 24°C (68 - 75°F).

Use of the QuickScan System

Detailed instructions for use of the QuickScan System are supplied with each unit and can also be found at www.envirologix.com/quickscan. The lot-specific Multi-Matrix Barcode Card (MMBC) must be scanned into the system prior to testing. In summary, a strip is inserted into the reader and the strips are read by touching or clicking on the “Read Test” area of the screen. The “Select Matrix Groups” screen will appear if more than one barcode was scanned into the system from the MMBC. Select the group that displays the matrix run. Results are then recorded in an electronic worksheet, allowing each user to report and track data easily.

Kit Storage

This Kit should be stored refrigerated. Note the shelf life on the kit box. Prolonged exposure to high temperatures may adversely affect the test results; protect all components from extreme hot or cold temperatures. Do not leave in direct sunlight or in a vehicle. Do not open the desiccated canister until ready to use the strips.

Cross-reactivity

The following mycotoxins have been tested with this kit and no false positive results occurred at the 100 ppm level: Aflatoxin, DON (deoxynivalenol), T2, Zearalenone.

Notes

- This product is currently not applicable for use in testing any other crops beyond those specified in this Product Insert.
- Pipettes lose calibration accuracy over time. Calibrate or replace pipettes at least annually.
- Immediate shaking of the sample after water addition is critical to ensure the EB17 packet does not cause clumps which may interfere with test results.
- This assay is calibrated against reference samples supplied by Trilogy Analytical Laboratory, Washington, MO, and other vendors and associated HPLC data.
- As with all screening tests, it is recommended that results be confirmed by an alternate method when necessary.
- The assay has been optimized for use with the protocols provided in the kit. Deviation from these protocols may invalidate the results of the test. Room temperature components, proper and thorough mixing, timing, and accurate pipetting are essential to accurate results.
- The results generated through the proper use of this diagnostic tool reflect the condition of the working sample directly tested. Extrapolation as to the condition of the originating lot, from which the working sample was derived, should be based on sound sampling procedures and statistical calculations which address random sampling effects, non-random sampling effects and assay system uncertainty. A negative result obtained when properly testing the working sample does not necessarily mean the originating lot is entirely negative for the analyte in question.
- **IMPORTANT:** If used, the 10X EB18 Extraction Buffer should be considered an irritant (SDS available at https://www.envirologix.com/?attachment_id=3004). Avoid contact with the skin, eyes, or clothing. Wear personal protective equipment including safety glasses, gloves, and a lab coat when handling.
 - **To prepare 1X EB18 Buffer Solution:** Mix 1 part 10X EB18 Extraction Buffer with 9 parts of water. 1X solution expires one week from date of mixing when stored at room temperature, or 4 weeks when stored at 2-8°C. Bring to room temperature before using.

Set A: Corn, Sorghum

- Review Sample Preparation on Page 3 for grinding consistency and notes.
- Turn on the incubator and set to 22°C for a minimum of 10 minutes before testing. Ensure that the temperature display has stabilized and indicates “OK” before starting the assay. All reagents should be at room temperature.
- Use distilled, deionized, or flat (non-carbonated) bottled water.
- If testing 50-gram samples, additional EB17 Buffer pouches and larger extraction vessels are required (50g Sample Set, order Catalog No. ACC-099).

Sample Extraction

Corn, Sorghum	25g	Add 1 EB17 pouch to sample Add 75 mL water	Wet sample immediately, by vigorously shaking for 10 seconds by hand. If needed, shake the sample against the palm of your other hand or a hard surface to loosen up any dry sample areas. Immediately proceed to next shaking step.
	50g	Add 2 EB17 pouches to sample Add 150 mL water	

Shake: choose mechanical shaker or hand shaking

Shaker Table: mix at highest speed (≥ 300 rpm) for 1 minute	By Hand: shake vigorously for 2 minutes
Centrifuge: 30 seconds at 2000 x g (ref, <u>not rpm</u>)	Filter: Pour through approved coffee filter (ACC 083); wait no more than 2 minutes

Clarify Extract: choose centrifuge or filter

Combine Buffer and Extract, then Run Test Strips

- Add DB5 to the Reaction Tube (discard tip)
- Add clarified extract to the Reaction Tube
- Mix thoroughly with extract pipette tip, discard tip
- Place the Reaction Tube in the 22°C incubator; equilibrate for 2 minutes. Note: tube acclimation step is only required if the temperature of the testing environment is unknown or outside of 20 - 24°C (68 - 75°F)
- Add test strip to tube, arrows down, wait 4 minutes (run time)
- Immediately cut strips at the top of the arrow tape (discard bottom pads)
- Insert strip into QuickScan Reader

TIPS!

Get Complete Extraction

- Fully wet samples before the next shaking step
- Avoid delay between water addition and shaking
- Assure liquid is moving forcefully though the sample while shaking

For Best Performance

- Pipette up and down while mixing
- Read strips promptly after run time

Avoid Contamination

- Use a new Reaction Tube per test
- Keep DB5 capped, when possible
- Use new pipette tips for each step

TABLE A: Set A Matrix Summary (base range)

Matrix	LOD (ppm)	Add to sample extraction vessel in order (choose 25 or 50g):		Shake	Clarify	Reaction Tube	Run
Corn --- Sorghum	0.1 --- 0.2	A1. 25g sample A2. 1 x EB17 A3. 75 mL water	B1. 50g sample B2. 2 x EB17 B3. 150 mL water	1 min – shaker <u>or</u> 2 min – by hand	Filter <u>or</u> Centri- fuge	100 µL DB5 + 100 µL extract	4 min
		(immediately shake by hand 10 sec)					

Set B: DDGS

- Review Sample Preparation on Page 3 for grinding consistency and notes.
- Turn on the incubator and set to 22°C for a minimum of 10 minutes before testing. Ensure that the temperature display has stabilized and indicates “OK” before starting the assay. All reagents should be at room temperature.
- Prepare EB18 Extraction Buffer from concentrate (see Notes, Page 4, for preparation instructions and SDS). Use distilled, deionized, or flat (non-carbonated) bottled water.
- If testing 50-gram samples, additional EB17 Buffer pouches and larger extraction vessels are required (50g Sample Set, order Catalog No. ACC-099).

Sample Extraction

DDGS	25g	Add 1 EB17 pouch to sample Add 100 mL EB18 Extraction Buffer	Wet sample immediately, by vigorously shaking for 10 seconds by hand. If needed, shake the sample against the palm of your other hand or a hard surface to loosen up any dry sample areas. Immediately proceed to next shaking step.
	50g	Add 2 EB17 pouches to sample Add 200 mL EB18 Extraction Buffer	

Shake: choose mechanical shaker or hand shaking

Shaker Table: mix at highest speed (≥ 300 rpm) for 1 minute	By Hand: shake vigorously for 2 minutes
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Clarify Extract: Centrifuge:

Centrifuge: 30 seconds at 2000 x g (rcf, <u>not rpm</u>)
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Combine Buffer and Extract, then Run Test Strips

- Add DB5 to the Reaction Tube (discard tip)
- Add clarified extract to the Reaction Tube
- Mix thoroughly with extract pipette tip, discard tip
- Place the Reaction Tube in the 22°C incubator; equilibrate for 2 minutes. Note: tube acclimation step is only required if the temperature of the testing environment is unknown or outside of 20 - 24°C (68 - 75°F)
- Add test strip to tube, arrows down, wait 5 minutes (run time)
- Immediately cut strips at the top of the arrow tape (discard bottom pads)
- Insert strip into QuickScan Reader

TIPS!

Get Complete Extraction

- Fully wet samples before the next shaking step
- Avoid delay between water addition and shaking
- Assure liquid is moving forcefully though the sample while shaking

For Best Performance

- Pipette up and down while mixing
- Read strips promptly after run time

Avoid Contamination

- Use a new Reaction Tube per test
- Keep DB5 capped, when possible
- Use new pipette tips for each step

TABLE B: Set B Matrix Summary (base range)

Matrix	LOD (ppm)	Add to sample extraction vessel in order (choose 25 or 50g):		Shake	Clarify	Reaction Tube	Run
DDGS	0.2	A1. 25g sample	B1. 50g sample	1 min – shaker <u>or</u> 2 min – by hand	Centri-fuge	200 μ L DB5 + 100 μ L extract	5 min
		A2. 1 x EB17	B2. 2 x EB17				
		A3. 100 mL EB18	B3. 200 mL EB18				
		(immediately shake by hand 10 sec)					



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Safety data sheet

SECTION 1. Identification of the substance/mixture and of the company/undertaking

1.1 Product identifier
Trade name: **DB 5 Dilution Buffer**
Part number: 11150, 11665, 12495 (KR-266)

1.2 Relevant identified uses of the substance or mixture and uses advised against application of the substance / the preparation :
Laboratory chemicals; kit component. Not to be used for purposes other than those specified in product literature.

1.3 Details of the supplier of the safety data sheet
Manufacturer/Supplier: EnviroLogix Inc., 500 Riverside Industrial Pkwy., Portland ME 04103, USA
Phone: (207) 797-0300

1.4 Emergency telephone number: (207) 797-0300 Technical Service

SECTION 2. Hazards identification.

2.1 Classification of the substance or mixture
Classification according to 29CFR 1910.1200: Eye Damage Category 1
Aquatic Toxic, Chronic Category 2

2.2 Label elements
Labeling according to 29CFR 1910.1200:

Pictogram:	
Signal word:	Warning
Hazard Statements:	H318 Causes serious eye damage H411 Toxic to aquatic life with long lasting effects
Precautionary Statements:	P264 Wash hands thoroughly after handling P280 Wear protective gloves/protective clothing/eye Protection/face protection P305+P351+P338 IF IN EYES: Rinse cautiously with Water for several minutes. Remove contact lenses If present and easy to do. Continue rinsing. P337+P313 IF eye irritation persists: Get medical attention/advice
2.3 Other Statements	Restricted to professional users

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SECTION 3. Composition/information on ingredients.

3.2 Mixture				
Chemical name	CAS No	EC No	Classification According to 29CFR 1910.1200	Amount (%)
Sodium Tetraborate Decahydrate	1303-96-4	215-540-4	H560 Rep 1B	<3 %
p-tertiary Octylphenoxy polyethyl alcohol (Triton X-100)	9002-93-1		H302 Acute Tox. Oral 4 H315 Skin Irrit. 2 H318 Eye Dam. 1 H411 Aquatic Chronic 2	1 %
Surfynol	9014-85-1		H315 Skin irritation 2 H318 Eye damage 1 H335 STOT SE 3	2 %
1,2-Benzisothiazolin-3-one (Proxel-GXL)	2634-33-5	220-120-9	H302 Acute Tox. 4; H315 Skin Irrit. 2 H317 Skin Sens. 1 (C _≥ 0.05%) H318 Eye Dam. 1; H400 Aquatic Acute 1	0.048 %

SECTION 4. First aid measures.

4.1 Description of first aid measures

After inhalation : In case of inhalation. Remove to fresh air. If not breathing give artificial respiration. Get medical attention immediately.

After skin contact : In case of skin contact. Remove contaminated clothing and shoes immediately. Wash affected areas with mild soap or detergent for at least 10 minutes or until no evidence of chemical remains.

After eye contact : In case of eye contact, immediately flush eyes with plenty of water for at least 15 minutes. Lifting eyelids occasionally, until no evidence of chemical remains. Get medical attention immediately.

After swallowing : In case of ingestion. DO NOT induce vomiting unless directed to do so by medical personnel. Never give anything by mouth to an unconscious person. Call a physician immediately.

4.2 Most important symptoms and effects, both acute and delayed: None

4.3 Indication of any immediate medical attention and special treatment needed: None

SECTION 5. Firefighting measures.

5.1 Extinguishing media: CO₂, extinguishing powder or water spray. Fight larger fires with water spray or alcohol resistant foam.

5.2 Special hazards arising from the substance or mixture: None

5.3 Advice for firefighters: Wear protective gear appropriate for fire conditions including respiratory protective gear.

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SECTION 6. Accidental release measures.

6.1 Personal precautions, protective equipment and emergency procedures: In the case of spilled mixture wear gloves to prevent skin contact. In the case of a large spill, additional protection is recommended.

6.2 Environmental precautions: Do not discharge mixture to sewer system or waterways.

6.3 Methods and material for containment and clean-up: Absorb in paper towel and discard in appropriate waste. Clean with water afterwards. Large spills may be neutralized with dilute solutions of sodium carbonate or calcium oxide.

6.4 References to other sections: For safe handling refer to Section 7. For information on PPE refer to Section 8. For disposal refer to Section 13

SECTION 7. Handling and storage.

7.1 Precautions for safe handling: Practice good chemical hygiene when handling. Avoid contact with eyes, skin, and clothing.

7.2 Conditions for safe storage, including any incompatibilities: Store in tightly closed, non-metal container, in a corrosive compatible area. Prevent direct sunlight and heat. Store in well aired storage rooms.

7.3 Specific end uses: Apart from the uses mentioned in section 1.2, no other specific uses are stipulated

SECTION 8. Exposure controls/personal protection.

8.1 Exposure limits: Components with limit values that require monitoring at the workplace:

	EH40/2005	OSHA
Sodium Tetraborate Decahydrate	8 Hr TWA = 5mg/m ³	8 Hr TWA = 10 mg/m ³

8.2 Exposure Controls:

8.2.1 Engineering controls
Facilities using this mixture should be equipped with an eyewash and safety shower. Use general or local exhaust ventilation to keep airborne concentrations below permissible exposure limits.

8.2.2 General protective and hygienic measures:
The usual precautionary measures should be adhered to when handling chemicals.

Eye Protection:
Safety glasses with side shields, goggles. Use equipment for eye protection tested and approved under appropriate government standards such as NIOSH (US) or EN 166 (EU). Eye and face protection regulations are described by OSHA (US) in 29CFR1910.133. Do not wear contact lenses when working with chemicals

Hand Protection:
Handle with gloves. Gloves must be inspected prior to use. Use proper glove removal technique (without touching glove's outer surface) to avoid skin contact with this product. Dispose of contaminated gloves after use in accordance with applicable laws and good laboratory practices. Wash and dry hands. The selected protective gloves have to satisfy the specifications of EU Directive 89/686/EEC and the standard EN 374 derived from it.

Breathing Equipment:
Appropriate respiratory protection should be determined according to local conditions using risk analysis protocols. An approved disposable air purifying particulate respirator may be used as a backup to engineering controls. Always use respirators and components tested and approved under appropriate government standards such as NIOSH (US) or CEN (EU).

8.2.3 Environmental exposure controls:
Contain spills, do not allow into environment

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SECTION 9. Physical and chemical properties.

9.1 Information on basic physical and chemical properties:

a) Appearance:	Clear liquid, colorless to slight yellow.
b) Odor:	None
c) Odor Threshold:	No Data Available
d) pH:	8.6
e) Melting point/freezing point:	No Data Available
f) Boiling point/Boiling range:	No Data Available.
g) Flash point:	Not applicable.
h) Evaporation rate:	No Data Available
i) Flammability (solid, gaseous):	No Data Available
j) Upper/lower flammability or explosive limits:	No Data Available
k) Vapor pressure:	No Data Available
l) Vapor density:	No Data Available
m) Relative density:	No Data Available
n) Solubility(ies):	Fully miscible, water.
o) Partition Coefficient: n-Octanol/water:	No Data Available
p) Auto-ignition temperature:	No Data Available
q) Decomposition temperature:	No Data Available
r) Viscosity:	No Data Available
s) Explosive properties:	No Data Available.
t) Oxidizing properties:	No Data Available

9.2 Other information: No further relevant information available.

SECTION 10. Stability and reactivity.

10.1 Reactivity: No data available

10.2 Chemical Stability: Stable under normal temperatures and pressures.

10.3 Possibility of hazardous reactions: Under normal conditions of storage and use, hazardous reactions will not occur.

10.4 Conditions to avoid: No specific data

10.5 Incompatible materials: No Data Available.

10.6 Hazardous decomposition products: Under normal conditions of storage and use, hazardous decomposition products should not be produced.

SECTION 11. Toxicological information.

Information on Toxicological Effects


Triton X-100
Acute toxicity: Oral LD50 –Rat- 1800mg/kg
Dermal LD50- Rabbit- 8000 mg/kg


Sensitization: No sensitizing effects known

CMR (carcinogenicity, mutagenicity and toxicity for reproduction) effects: No CMR effects.

Additional toxicological information: No Additional Information

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SECTION 12. Ecological information.		
12.1 Toxicity:	Fish: LC50 Pimephales promelas (fathead minnow) – 8.9mg/l – 96.0 hr Daphnia: EC50 – Daphnia – 26 mg/l – 48 hr	
12.2 Persistence and degradability :	No Data Available	
12.3 Bio accumulative potential:	No Data Available	
12.4 Mobility in soil :	No Data Available	
12.5 Results of PBT and vPvB assessment:	Not available as a chemical safety assessment, not required/not conducted.	
12.6 Other adverse effects:	No Data Available	
SECTION 13. Disposal considerations.		
Waste treatment methods:	Contact a licensed professional waste disposal service to dispose of this material. Disposal of surplus or waste solutions must be in accordance with applicable local, state, and national laws and regulations.	
SECTION 14. Transport information.		
14.1 UN-Number DOT, ADR, ADN, IMDG, IATA :	Not Hazardous for Transport	
14.2 UN proper shipping name DOT, ADR, ADN, IMDG, IATA :	Not Hazardous for Transport	
14.3 Transport hazard class(es) DOT, ADR, ADN, IMDG, IATA):	Not Hazardous for Transport	
14.4 Packing group (DOT, ADR, IMDG, IATA):	Not Hazardous for Transport	
14.5 Environmental hazards	No environmental hazard.	
14.6 Special precautions for user :	None	
14.7 Transport in bulk according to Annex II of MARPOL 73/78 and the IBC code:	No information available.	
SECTION 15. Regulatory information.		
15.1 Safety, health, and environmental regulations		
US Federal Regulations		
OSHA	Not a hazardous material	
SARA 313	Not listed	
US State Regulations		
European/International Regulations		
European labeling in accordance with EC Directives	Not hazardous according to European directives	
15.2 Chemical Safety Assessment	Not carried out	
SDS DB5 Dilution Buffer		

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SECTION 16. Other information.		
<i>This information is true based on our present knowledge. However, EnviroLogix makes no representation of its accuracy or completeness. Persons receiving this information must exercise their independent judgment in determining the product's safety and suitability for its intended use. This document shall not constitute a guarantee for any specific product features and shall not establish a legally valid contractual relationship</i>		
EHS Department EnviroLogix Inc.		
Codes:		
H302 Harmful if swallowed	H315 Causes skin irritation	H317 May cause an allergic skin reaction
H318 Causes Serious Eye Damage	H335 May cause respiratory irritation	H411 Toxic to Aquatic Life with Long Lasting Effects
SDS DB5 Dilution Buffer		

Summary Guide for Approved Matrices

Approved Matrix	Add to Sample Extraction Vessel (in this order)	Then shake immediately	Clarify	Run the Base Range Protocol First, followed by Dilution Protocol, if necessary	Transfer to Reaction Tube	Add Reaction Tube to Incubator Set at 22°C	Add Strip for	Read in QuickScan: Dilution Tab on the Result Page Should Display
Corn (MG1); Sorghum (MG4)	1. 25g sample 2. 1 EB17 pouch 3. 75 mL water* 4. Immediately shake vigorously for 10 seconds by hand -----OR----- 1. 50g sample 2. 2 EB17 pouches 3. 150 mL water* 4. Immediately shake vigorously for 10 seconds by hand	1 min highest speed on shaker table or 2 min by hand	Filter or Centrifuge 30 sec. at 2000 x g	Base Range 0 – 10 ppm	100 µL DB5 buffer + 100 µL clarified extract in Reaction Tube	Acclimate tube for 2 min [^]	4 min.	1:1 (this is software default)
	Dilution A Corn: 2 – 100 ppm			<u>Pre-Mix</u> 700 µL EB17 Dilution Solution† + 50 µL clarified extract <u>Transfer</u> 100 µL of this Pre-Mix and 100 µL DB5	Acclimate tube for 2 min [^]	4 min.	1:A (this must be selected)	
	Dilution A Sorghum: 10 – 100 ppm			<u>Pre-Mix</u> 700 µL EB17 Dilution Solution† + 50 µL clarified extract <u>Transfer</u> 200 µL of this Pre-Mix and 100 µL DB5				

Notes:

*Use distilled, deionized, or flat (non-carbonated) bottled water

[^]The tube acclimation step is only required if the temperature of the testing environment is unknown or outside of 20 - 24°C (68 - 75°F)

†Refer to page 3 for Dilution Reagent instructions

Approved Matrix	Add to Sample Extraction Vessel (in this order)	Then shake immediately	Clarify	Run the Base Range Protocol First, followed by Dilution Protocol, if necessary	Transfer to Reaction Tube	Add Reaction Tube to Incubator Set at 22°C	Add Strip for	Read in QuickScan: Dilution Tab on the Result Page Should Display
DDGS (MG2)	1. 25g sample 2. 1 EB17 pouch 3. 100 mL EB18 4. Immediately shake vigorously for 10 seconds by hand -----OR-----	1 min highest speed on shaker table or 2 min by hand	Centrifuge 30 sec. at 2000 x g	Base Range 0 – 30 ppm	200 µL DB5 buffer + 100 µL clarified extract in Reaction Tube	Acclimate tube for 2 min [^]	5 min.	1:1 (this is software default)
	Dilution A 30 – 100 ppm			Pre-Mix 400 µL EB18 Dilution Solution [†] + 100 µL clarified extract Transfer 200 µL of this Pre-Mix and 100 µL DB5	Acclimate tube for 2 min [^]	5 min.	1:A (this must be selected)	

Notes:

*Use distilled, deionized, or flat (non-carbonated) bottled water

[^]The tube acclimation step is only required if the temperature of the testing environment is unknown or outside of 20 - 24°C (68 - 75°F)

[†]Refer to page 3 for Dilution Reagent instructions