

Catalog Number AP 013

Highlights:

- *High Sensitivity Protocol detects the PAT enzyme from the bar gene (PAT/bar) found in 0.1% StarLink corn*
- *Rapid Protocol screens individual seeds or leaf samples for the presence of PAT/bar*

Contents of Kit:

- *1 antibody-coated 96-well plate*
- *PAT/bar Enzyme Conjugate*
- *1 packet of Buffer Salts*
- *Substrate*
- *Stop Solution*

Intended Use

The QualiPlate Kit for LibertyLink PAT/*bar* is designed for the qualitative laboratory detection of phosphinothricin acetyl transferase enzyme (PAT) coded for by the *bar* gene in grain, leaf, or seed. Two assay protocols are presented: The High Sensitivity Protocol will detect the PAT enzyme from the *bar* gene (PAT/*bar*) found in 0.1% StarLink® corn (by weight) and requires 2.5 hours to run. The Rapid Protocol (one hour total) is intended for use in screening individual seeds or leaf samples for the presence of PAT/*bar*. LibertyLink PAT from the *pat* gene, at concentrations present in T25 corn, is not detected in either format.

How the Test Works

This QualiPlate Kit is a “sandwich” Enzyme-Linked ImmunoSorbent Assay (ELISA). In the test, **corn** or **cotton** sample extracts are added to test wells coated with antibodies raised against PAT from the *bar* gene. Any residues present in the sample extract bind to the antibodies, and are then detected by addition of enzyme (horseradish peroxidase)-labeled PAT/*bar* antibody.

After a simple wash step, the results of the assay are visualized with a color development step; color development is proportional to PAT/*bar* concentration in the sample extract.

Lighter color = Lower concentration

Darker color = Higher concentration

How the Kit Performs

The High Sensitivity Protocol will detect 0.1% StarLink corn grain. Samples are judged to contain more or less PAT/*bar* enzyme than 0.1% StarLink standard by comparing sample absorbances to the absorbance of a 0.1% StarLink sample extract (prepared by the user).

The Rapid Protocol is a strictly qualitative test, in which seed and leaf samples are screened for the presence or absence of the PAT/*bar* enzyme at the levels commonly seen in commercially modified crops. It is recommended that the user prepare extracts from known conventional and known PAT/*bar* expressing plant tissues to run in each assay as negative and positive controls.

Materials not Provided

- distilled or deionized water for preparing Wash/Extraction Buffer
- glass bottles or flask plus graduated cylinder with 1 liter capacity for preparation and storage of Wash/Extraction Buffer
- test or centrifuge tubes for extraction of grain
- Tissue Extraction Kit: snap-cap tubes and pestles for extraction of leaf samples (EnviroLogix Cat. No. ACC 002, 100/package)
- centrifuge capable of 5000 x g
- disposable tip, adjustable air-displacement pipettes which will measure 50 and 100 microliters (µL)
- marking pen (indelible)
- tape or Parafilm®



Prepare
Wash/Extraction Buffer

Preparation of Solutions

Wash/Extraction Buffer: Add the contents of the packet of **Buffer Salts** (phosphate buffered saline, pH 7.4 - Tween 20) to 1 liter of distilled or deionized water, and stir to dissolve. Store refrigerated when not in use; warm to room temperature prior to assay. If more extraction buffer is needed, order item # P-3563 from Sigma Chemical Co. (St. Louis, MO), or prepare the equivalent.

- timer
- microtiter ELISA plate reader
- wash bottle, or microtiter plate or strip washer
- multi-channel pipette that will measure 50 and 100 μ L
- racked dilution tubes for loading samples into the plate with a multi-channel pipette, or the equivalent
- orbital plate shaker (optional)

Sample Preparation

Note: PAT/*bar* protein is not stable in solution. Extraction should be performed in 30 minutes or less. Do not extract overnight.

High Sensitivity Protocol for Ground Grain/Seed

This protocol requires that a small sample be analyzed. It is essential that this sample be well mixed and representative of the larger bulk. The test will detect 0.1% StarLink corn, containing PAT/*bar* (one kernel in a sample of 1000 kernels).

Note: Thorough mixing of the bulk grain sample and determination of an appropriate sampling plan are critical to the results of this testing, and are the responsibility of the user of this test kit. The USDA/GIPSA has prepared several guidance documents to address the issues involved in obtaining representative grain samples from static lots - such as trucks, barges, and railcars - and for taking samples from grain streams. Sampling plans should be chosen that best meet the needs of both the buyer and seller in terms of acceptable risks. Increasing the number of kernels in the sample and taking multiple samples will increase the likelihood of obtaining representative samples, and maximize the probability of detecting any contamination in the grain lot. For further information on USDA/GIPSA guidelines for obtaining representative samples and assessing detection probabilities for biotech grain, see the websites listed on page 3.

It is the responsibility of the user to ensure proper sampling and thorough mixing prior to analysis. Once representative samples have been obtained from the truck or container, they can be reduced in size using a splitter and uniformly ground and mixed. The finer the grind, the faster and more efficient the extraction.

1. Weigh 5 grams of ground corn sample into a 15 mL capacity vial or tube.
2. Add 12.5 mL of Wash/Extraction Buffer to each 5 gram sample. For all other grain sample sizes, add Wash/Extraction Buffer at the rate of 2.5 mL per gram of grain. Cap and shake vigorously by hand or vortex for 20-30 seconds. Let stand at room temperature to extract.
3. The extracted samples/controls must be clarified by centrifuging the extract at 5000 x g for 5 minutes. Insert a pipette tip below any floating lipid layer and above the pellet to remove the clarified sample.

Rapid Protocol for Screening Single Seeds or Leaf Punches:

Individual seeds:

1. Crush seeds: Seeds may be placed in a baggie and smashed with a hammer, then transferred to a tube; or, a seed crusher/48-well plate combination may be used (for example Hypure #HSC-100, PerkinElmer, Norton, OH, with Costar plate #3548, Corning Life Sciences, Acton, MA, or equivalent). Check to be sure that all seeds have been crushed. Take extreme care not to cross-contaminate between seed samples.
2. Add 0.5 mL of Wash/Extraction Buffer to each crushed corn seed, or 1 mL to each crushed cotton seed. Mix for at least 30 seconds, then allow particles to settle.

Reference Websites:

www.gipsa.usda.gov/fgis/handbook/gihbk1_inspec.aspx
 USDA *Grain Inspection Handbook, Book 1, Grain Sampling*. This document provides a comprehensive overview of recommended sampling guidelines for static lots and grain streams. It reviews the various types of equipment and strategies that can be used to obtain a representative grain sample from different types of containers.

www.gipsa.usda.gov/fgis/biotech/sample2.htm
 Guidance document entitled *Sampling for the Detection of Biotech Grains*, which provides important statistical sampling considerations when testing for the presence of biotech grains. It covers the basis for making probability determinations in accepting lots based upon different assumptions with respect to sample size, number of samples, sample preparation, etc.

www.gipsa.usda.gov/fgis/biotech/sample1.htm
Practical Application of Sampling for the Detection of Biotech Grains. This one-page application guide provides a table that gives sample sizes for selected lot concentrations and probability of rejecting the specified concentrations. It also provides a formula for making the calculation for other combinations.

www.gipsa.usda.gov/fgis/biotech/samplingplan1.xls This website provides a simple to use Sample Planner (29k Excel Spreadsheet). The planner allows you to enter different assumptions in terms of sample size, number of samples, acceptable quality level and to determine the probability of accepting lots with given concentration levels. It also plots the probabilities in graph form for easy interpretation. Specific data can be saved for documentation and future analyses.

Leaf testing:

1. Take a single leaf punch of approximately 5 millimeters diameter, using a micro-tube cap or a paper punch. Mash the leaf tissue with a pestle matched to the micro-tube, or with a disposable pipette tip, or a Hypure cutter (HCT-200, PerkinElmer, Norton, OH) in a 96-well plate (Costar #3370, Corning Life Sciences, Acton, MA, or equivalent).
2. Add 0.25 mL of Wash/Extraction Buffer per leaf punch. Mix for at least 30 seconds, then allow particles to settle. Take extreme care not to cross-contaminate between leaf samples.

NOTE: It is recommended that the kit user extract known conventional and known PAT/*bar*-containing samples of the matrix being tested, and run these as Negative and Positive Controls (NC and PC) in each assay.

How to Run the Assay

- Read all of these instructions before running the kit.
- Allow all reagents to reach room temperature before beginning (at least 30 minutes with un-boxed plates and reagents at room temperature - do not remove plates from bag with desiccant until they have warmed up).
- Organize all Control and sample extracts, and pipettes so that Step 1 can be performed in 15 minutes or less. The use of a multi-channel pipette is strongly recommended.
- Use a disposable-tip air-displacement pipette and a clean pipette tip to add each Calibrator and sample extract to the wells. Conjugate, Substrate, and Stop Solution may be added in the same manner; alternatively, use a repeating pipette with a disposable tip on the end of the Combitip for these three reagents.
- Use the well identification markings on the plate frame to guide you when adding the samples and reagents. In a qualitative assay, the Blank (BL), Negative Control (NC), and the Positive Control (PC) and 90 sample extracts (S) may be run in single wells on one plate. (See the Qualitative Assay Example Plate Layout - Figure 1A).

HIGH SENSITIVITY PROTOCOL: The High Sensitivity Protocol will detect 0.1% StarLink corn (by weight) in ground grain/seed, and requires 2.5 hours of total assay incubation time.

Procedure

1. Add 50 μ L of PAT/*bar*-enzyme Conjugate to each well, followed immediately by 50 μ L of Wash/Extraction Buffer Blank (BL), 50 μ L of the centrifuged 0.1% StarLink ground corn and Negative ground corn extracts (PC and NC), and 50 μ L of each centrifuged sample extract (S) to their respective wells, as shown in the Example Plate Layout (Figure 1A)

NOTE: In order to minimize setup time it is recommended that a multi-channel pipette be used in steps 1, 5, and 7.

2. Thoroughly mix the contents of the wells by moving the plate in a rapid circular motion on the benchtop for a full 20-30 seconds. Be careful not to spill the contents!
3. Cover the wells with tape or Parafilm to prevent evaporation and incubate at ambient temperature for 2 hours. If an orbital plate shaker is available, shake plate at 200 rpm.
4. After incubation, carefully remove the covering and vigorously shake the contents of the wells into a sink or other suitable container. Flood the wells



Allow all reagents to reach room temperature before beginning



Leaf punch



Extract sample



Centrifuge to clarify sample extract
(High Sensitivity protocol only)

completely with Wash/Extraction Buffer, then shake to empty. Repeat this wash step three times. Alternatively, perform these four washes (300 μ L/well) with a microtiter plate or strip washer. Slap the inverted plate on a paper towel to remove as much liquid as possible.

5. Add 100 μ L of Substrate to each well.
6. Thoroughly mix the contents of the wells, as in step 2. Cover the wells with new tape or Parafilm and incubate for 30 minutes at ambient temperature. Use orbital shaker if available.
7. Caution: Stop Solution is 1.0N Hydrochloric acid. Handle carefully.
8. Add 100 μ L of Stop Solution to each well and mix thoroughly. This will turn the well contents yellow.

NOTE: Read the plate within 30 minutes of the addition of Stop Solution.

RAPID PROTOCOL: The Rapid Protocol is less sensitive (able to detect 1% StarLink corn) but only requires one hour of total assay incubation time.

Procedure

1. Add 50 μ L of PAT/*bar*-enzyme Conjugate to each well, followed immediately by 50 μ L of Wash/Extraction Buffer Blank (BL), 50 μ L of the user-prepared Positive and Negative Control extracts (PC and NC), and 50 μ L of each sample extract (S) to their respective wells, as shown in the Example Plate Layout (Figure 1A).

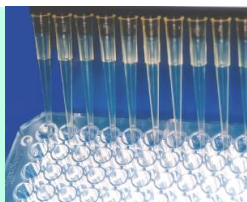
NOTE: In order to minimize setup time it is recommended that a multi-channel pipette be used in steps 1, 5, and 7.

2. Thoroughly mix the contents of the wells by moving the plate in a rapid circular motion on the benchtop for a full 20-30 seconds. Be careful not to spill the contents!
3. Cover the wells with tape or Parafilm to prevent evaporation and incubate at ambient temperature for 45 minutes. If an orbital plate shaker is available, shake plate at 200 rpm.
4. After incubation, carefully remove the covering and vigorously shake the contents of the wells into a sink or other suitable container. Flood the wells completely with Wash/Extraction Buffer, then shake to empty. Repeat this wash step three times. Alternatively, perform these four washes (300 μ L/well) with a microtiter plate or strip washer. Slap the inverted plate on a paper towel to remove as much liquid as possible.
5. Add 100 μ L of Substrate to each well.
6. Thoroughly mix the contents of the wells, as in step 2. Cover the wells with new tape or Parafilm and incubate for 15 minutes at ambient temperature. Use orbital shaker if available.

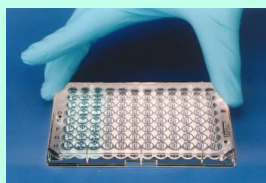
Caution: Stop Solution is 1.0N Hydrochloric acid. Handle carefully.

7. Add 100 μ L of Stop Solution to each well and mix thoroughly. This will turn the well contents yellow.

NOTE: Read the plate within 30 minutes of the addition of Stop Solution.



Add Conjugate, controls and samples



Mix plate



Incubate



Wash plate



Read plates in a Plate Reader within 30 minutes of the addition of Stop Solution

How to Interpret the Results

Spectrophotometric Measurement

1. Set the wavelength of the microtiter plate reader to 450 nanometers (nm). (If it has dual wavelength capability, use 600, 630 or 650 nm as the reference wavelength.)
2. Set the plate reader to blank on the Wash/Extraction Buffer Blank wells (this should automatically subtract the mean optical density (OD) of the Blank wells from each control and sample OD). If the reader cannot do this, it must be done manually.

Note: The mean OD of the BLANK wells should not exceed 0.2.

Interpret the Results

For leaf and seed screening, compare the OD's of sample extracts to OD's of extracts from known leaf or seed samples (conventional and PAT/*bar*-expressing varieties). Note: very low positive results in a seed-screening assay may be due to cross-contamination during crushing. Truly positive seeds will result in OD readings typically greater than 0.500.

Figure 1A. Example of a typical Qualitative assay setup.

	1	2	3	4	5	6	7	8	9	10	11	12
A	BL	S6	S14	S22	S30	S38	S46	S54	S62	S70	S78	S86
B	NC	S7	S15	S23	S31	S39	S47	S55	S63	S71	S79	S87
C	PC	S8	S16	S24	S32	S40	S48	S56	S64	S72	S80	S88
D	S1	S9	S17	S25	S33	S41	S49	S57	S65	S73	S81	S89
E	S2	S10	S18	S26	S34	S42	S50	S58	S66	S74	S82	S90
F	S3	S11	S19	S27	S35	S43	S51	S59	S67	S75	S83	BL
G	S4	S12	S20	S28	S36	S44	S52	S60	S68	S76	S84	NC
H	S5	S13	S21	S29	S37	S45	S53	S61	S69	S77	S85	PC

Precautions and Notes

- Store all Plate Kit components at 4°C to 8°C (39°F to 46°F) when not in use.
- Do not expose Plate Kit components to temperatures greater than 37°C (99°F) or less than 2°C (36°F).
- Allow all reagents to reach ambient temperature (18°C to 27°C or 64°F to 81°F) before use.
- Do not use kit components after the expiration date.
- Do not use reagents or plates from one Kit with reagents or plates from a different Kit.
- Do not expose Substrate to sunlight during pipetting or while incubating in test wells.
- The assay has been optimized to be used with the protocol provided in the kit. Deviation from this protocol may invalidate the results of the test.
- As with all tests, it is recommended that results be confirmed by an alternate method when necessary.
- Observe any applicable regulations when disposing of samples and kit reagents.
- Use extreme caution to prevent sample-to-sample cross-contamination with samples, fluids, or disposables.



**For Technical Support
Contact Us At:**

EnviroLogix
500 Riverside Industrial
Parkway
Portland, ME 04103-1486
USA

Tel: (207) 797-0300
Toll Free: 866-408-4597
Fax: (207) 797-7533

e-mail:
info@envirologix.com

website:
www.envirologix.com



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Safety Data Sheet
According to OSHA 29CFR 1910.1200

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

1.1 Product identifier
Trade name: **Wash Buffer Salts**
Part number: 50-0091, 10099

1.2 Relevant identified uses of the substance or mixture and uses advised against application of the substance or the preparation:
Laboratory chemicals

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

1.4 Emergency telephone number:

SECTION 2. Hazards identification

2.1 Classification of the Substance or Mixture:
Classification according to OSHA 29CFR 1910.1200 (Hazard Communication): Not a hazardous substance or mixture

2.2 Label Elements:
None required according to 29CFR 1910.1200
Other indications: None

2.3 Additional Information:
No other information

SECTION 3. Composition/information on ingredients

3.1 Mixture: Powdery solid
Synonyms: PBS

Hazardous Components

Chemical name	CAS No	EC No	Amount (%)	Classification
Potassium Chloride	7447-40-7	231-211-8	1-5 %	Aquatic Acute 3, Aquatic Chronic 3, H412

Based on the amount of hazardous ingredients in this product, it is not considered hazardous according to 29CFR 1910.1200

SECTION 4. First aid measures

4.1 Description of first aid measures:
After inhalation: Supply fresh air; consult doctor in case of breathing difficulties.
After skin contact: Flush skin with plenty of water for at least 15 minutes. Remove contaminated clothing. Seek medical attention if irritation develops.
After eye contact: Rinse opened eye for several minutes under running water. Seek medical attention if irritation develops.
After swallowing: If swallowed, consult with medical staff or poison control center to determine if any immediate response or follow up actions are recommended. Never give anything by mouth to an unconscious person.

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

4.3 Indication of any immediate medical attention and special treatment needed:
No special treatment is required

SECTION 5. Firefighting measures

5.1 Extinguishing media:
Suitable extinguishing agents: 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:
Carbon oxides, Oxides of Phosphorus, Potassium, Sodium, Hydrogen Chloride gas

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

SECTION 6. Accidental release measures

6.1 Personal precautions, protective equipment and emergency procedures:
Use PPE, avoid dust formation, ensure adequate ventilation, avoid breathing dust

6.2 Environmental precautions:
Prevent further leakage or spillage if safe to do so. Do not let product enter drains. Discharge to the environment must be avoided.

6.3 Methods and material for containment and clean up:
Pick up and arrange disposal without creating dust. Sweep up and shovel. Keep in suitable closed containers for disposal

6.4 Reference 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. Prevent formation of dust.

7.2 Conditions for safe storage, including any incompatibilities:
Keep containers closed, store in a dry, well ventilated space.

7.3 Specific end use(s):
Apart from the uses mentioned in section 1.2, no other end uses are stipulated.

SECTION 8. Exposure controls/personal protection

8.1 Control parameters:
Components with workplace control Parameters: Contains no substances with occupational exposure limit values

8.2 Exposure controls
8.2.1 Appropriate engineering controls: Ensure eyewash and safety shower are nearby; provide ventilation if necessary

8.2.2 Personal Protective Equipment:
Eyes: 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

Hands: 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.

Respiratory protection: 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 CEV (EU).

Body: Use body protection relative to its type and amount of material being handled

8.2.3 Environmental controls: Sweep or wipe up spills, do not allow into sewers or drains

SECTION 9. Physical and chemical properties

9.1 Information on basic physical and chemical properties:

a) Appearance:	White powder.
b) Odor:	None
c) Odor Threshold:	No data available
d) pH:	7.4
e) Melting point/freezing point:	No data available
f) Boiling point/boiling range:	No data available
g) Flash point:	Not data available
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(s):	Water soluble
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 recommended storage conditions.

10.3 Possibility of hazardous reactions:
No data available

10.4 Conditions to avoid:
No data available

10.5 Incompatible materials:
Strong oxidizing agents and strong acids.

10.6 Hazardous decomposition products:
No data available

SECTION 11. Toxicological information

Acute toxicity:
Inhalation: No data available
Dermal: No data available
Skin corrosion/irritation: No data available
Serious eye damage: No data available
Respiratory or skin sensitization: No data available
Mutagenicity and toxicity for reproduction: No data available
Carcinogenicity: No component of this product at levels greater than 0.1 % is identified as probable, possible, or confirmed human carcinogen by IARC, ACGIH, NTP, or OSHA.

SECTION 12. Ecological information

12.1 Toxicity:
No data available

12.2 Persistence and degradability:
No data available

12.3 Bioaccumulative 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

Dispose of excess or unused product in accordance with Local, State and Federal regulations. Contact a licensed professional waste disposal service to dispose of this material.

SECTION 14. Transport information

14.1 UN Number (DOT, ADR, ADN, IMDG, IATA):
Not dangerous goods

14.2 UN proper shipping name (DOT, ADR, ADN, IMDG, IATA):
Not dangerous goods

14.3 Transport hazard class(es) (DOT, ADR, ADN, IMDG, IATA):
Not applicable

14.4 Packing group (DOT, ADR, IMDG, IATA):
Not applicable

14.5 Environmental hazards:
Not applicable

14.6 Special precautions for user:
Not applicable

14.7 Transport in bulk according to Annex II of MARPOL/73/78:
Not applicable

SECTION 15. Regulatory information

15.1 Safety, health and environmental regulations/legislation specific for the substance or mixture

US Federal Regulations
SARA Section 302 (Extremely Hazardous Substances): Not listed
Clean Air Act: Not listed
Clean Water Act: Not listed
OSHA: Not listed

US State Regulations
Massachusetts Right to Know: Disodium Hydrogenorthophosphate CAS No 7558-79-4 Rev Date: 2007-03-01
California Prop. 65 Components: Contains no chemicals known to cause cancer, birth defects, or reproductive harm

15.2 Chemical Safety Assessment
Not carried out

SECTION 16. Other information

Hazard Code
H412 Harmful to aquatic life with long lasting effects

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.

EHS Department
EnviroLogix Inc.



Material Safety Data Sheet
OSHA 29CFR 1910.1200

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

1.1 Product identifier	Stop Solution
Trade name:	L.O.N HCl
Synonyms:	10825, 10827, 10828, 11193, 11776 (XGID007)
Part number:	Laboratory chemicals
1.2 Relevant identified uses of the substance or mixture and uses advised against application of the substance / the preparation :	
1.3 Details of the supplier of the safety data sheet	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	Hazard Classes	
Classification according to OSHA 29 CFR 1910.1200	Metal Corrosive (Cat. 1)	H290
	Skin Irritation (Cat 2)	H315
	Serious Eye damage (Cat. 1)	H318
2.2 Label elements		
Labeling according to OSHA 29CFR 1910.1200		
Hazard pictograms :		
Signal word :	Warning	
Hazard statements:	H290 May be corrosive to metals H315 Causes skin irritation H318 Causes serious eye damage	
Precautionary statements:	P281 Use personal protective equipment as required P302 + P352 IF ON SKIN: Wash with plenty of soap and water. P305 + P351 + P338 IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses if present and easy to do. Continue rinsing.	
2.3 Other Statements	None	

SECTION 3. Composition/information on ingredients

3.2 Mixture				
Aqueous solution 1N Hydrochloric Acid (1N HCl, 3% HCl)				
Chemical name	Amount (%)	CAS No	Classification According to OSHA 29CFR 1910.1200	
			EC No	Hazard Code
Hydrochloric acid	1-4%	7647-01-0	Hazard Classification:	
			May be Corrosive to Metals	
			H290	
			Causes Skin Irritation	
			H315	
			Causes Serious Eye Damage	
			H318	

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 area 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:	May cause skin irritation and eye damage
4.3 Indication of any immediate medical attention and special treatment needed:	DO NOT use sodium bicarbonate in an attempt to neutralize the acid.

SECTION 5. Firefighting measures

5.1 Extinguishing media:	CO ₂ , 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:	Hydrogen Chloride gas
5.3 Advice for firefighters:	Wear protective gear appropriate for fire conditions including respiratory protective gear.

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 cleanup:	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 use(s):	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:									
	<table border="1"> <thead> <tr> <th>Hydrogen Chloride</th> <th>European (Commission directive 96/94)</th> <th>USA (OSHA)</th> </tr> </thead> <tbody> <tr> <td></td> <td>8hr TWA = 5 ppm (7.5 mg/m³)</td> <td>Ceiling Limit = 5 ppm (7.5 mg/m³)</td> </tr> <tr> <td></td> <td>STEL = 10 ppm (15 mg/m³)</td> <td></td> </tr> </tbody> </table>	Hydrogen Chloride	European (Commission directive 96/94)	USA (OSHA)		8hr TWA = 5 ppm (7.5 mg/m ³)	Ceiling Limit = 5 ppm (7.5 mg/m ³)		STEL = 10 ppm (15 mg/m ³)	
Hydrogen Chloride	European (Commission directive 96/94)	USA (OSHA)								
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	STEL = 10 ppm (15 mg/m ³)									
8.2 Exposure 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.									
8.2.1 Engineering controls:										
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									

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:	Pungent (slight)
c) Color/Threshold:	No Data Available
d) pH:	1
e) Melting point/freezing point:	No Data Available
f) Boiling point/Boiling range:	No Data Available
g) Flash point:	Not applicable
h) Evaporation rate:	0.36 (Water) compared with n-Butyl Acetate = 1
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 but should be similar to that of water
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:	Metals, Alkali metals, bases, Amines
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			
Acute effects (toxicity tests):	7647-01-0 HCl	Effect Dose	Species
	Acute oral toxicity	LD50=920mg/kg	rabbit
	Acute dermal toxicity	No data	
	Acute inhalative toxicity	LC50 = 3124 mg/L	rat
Sensitization:	No sensitizing effects known		
CMR (carcinogenicity, mutagenicity and toxicity for reproduction) effects:	No CMR effects		
Additional toxicological information:	No Additional Information		

SECTION 12. Ecological information

12.1 Toxicity:	Aquatic toxicity (1N HCl)	Effect dose	Exposure time	Species
	Acute fish toxicity	LC50=826 mg/L	96h	Leuciscus idus
	Acute daphnia toxicity	No data		
	Acute algae toxicity	No data		