



# QualiPlate™ Kit for Cry1C

Catalog Number AP 056 NW V50

## Highlights:

- Screen single cotton seed or leaf samples for the presence or absence of Cry1C

## Contents of Kit:

- 50 Cry1C antibody-coated solid plates
- Cry1C Positive Control
- Cry1C Enzyme Conjugate
- Substrate

**Note:** To handle bulk packaged Cry1C Enzyme Conjugate and Substrate, pour off 11.5 milliliters of each per plate to be run each day. Use a multiple-channel pipette to dispense. Do not pour excess Substrate back into the reagent bottle.

## Intended Use

The EnviroLogix QualiPlate Kit for Cry1C is designed for the non-quantitative laboratory detection of Cry1C protein in cotton leaf or seed samples.

## How the Test Works

This EnviroLogix QualiPlate Kit is a “sandwich” Enzyme-Linked ImmunoSorbent Assay (ELISA). In the test, plant leaf or seed sample extracts are added to test wells coated with antibodies raised against Cry1C toxin. Any residues present in the sample extract bind to the antibodies, and are then detected by addition of enzyme (horseradish peroxidase)-labeled Cry1C antibody.

After a simple wash step, the results of the assay are visualized with a color development step; color development is proportional to Cry1C concentration in the sample extract.

*Lighter color = Lower concentration  
Darker color = Higher concentration*

## Materials Not Provided

- Disposable Tissue Extractors, EnviroLogix Cat. # ACC 002
- PBS/0.05% Tween-20 Wash Buffer (may be purchased in 1L dry packets from Sigma Chemicals, Cat#P-3563, or prepared from salts on site). Store at controlled ambient temperature for up to one week, then discard.
- PBS/0.55% Tween-20 Extraction Buffer (may be prepared by adding 0.5 mL Tween-20 to 100 mL already prepared PBS/0.05% Tween-20 Wash Buffer). Store refrigerated when not in use; warm to room temperature prior to assay.
- 1 N Hydrochloric acid (HCl) Stop Solution. Prepare by adding 83 mL of concentrated HCl (36.5-38.0%) to 917 mL of distilled or deionized water; work in a fume hood and use proper protective gear. This reagent may be stored at room temperature for 2 years.
- disposable tip, adjustable air-displacement pipettes which will measure 100 microliters ( $\mu$ L), preferably of multi-channel style.
- marking pen (indelible)
- tape or Parafilm®
- timer
- microtiter plate reader
- wash bottle, or microtiter plate or strip washer
- orbital plate shaker (optional)

## Sample Preparation

### Sample Extraction:

Sample extraction protocols are to be designed and validated by the individual users of this kit. The following suggestions are guidelines, and define the manner in which the kit is performance tested by the manufacturer.

#### Green leaf samples:

1. Take 2 leaf punch samples (approximately 10 milligrams each) by snapping the tube cap of the Disposable Sample Extractor down on the leaf. Insert the pestle into the tube and grind the tissue by rotating the pestle against the sides of the tube with twisting motions. Continue this process for 20-30 seconds, or until the leaf tissue is well ground. Use a new extraction device for each sample. *Use extreme caution to prevent sample-to-sample cross-contamination with plant tissue or exudate.*
2. Add 0.5 mL to 1 mL of Extraction Buffer to the tube.
3. Repeat the grinding step to mix tissue with Extraction Buffer. Repeat this protocol for each sample to be tested, using a new tube and pestle for each. Allow the solids to settle in each tube for a few minutes.

#### Single seed samples:

1. Crush cotton seeds and extract each with 0.75 to 1 mL of Extraction Buffer. Mix thoroughly, then allow solids to settle before transferring extract to the assay plate.

## 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 reagents, sample extracts, and pipettes so that step 1 can be performed in 15 minutes or less. The use of a multichannel pipette is strongly recommended.
- Use a disposable-tip air-displacement pipette and a clean pipette tip to add Control and sample extracts 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 edge to guide you when adding the samples and reagents. It is recommended that at least two wells each of Blank (Extraction Buffer) and Cry1C Positive Control be run on each plate. Additional quality control samples may be added at the discretion of the user. Sample extracts may be run in either single or duplicate wells. See example of typical assay setup, Figure 1A, on page 4.

1. Add **100 µL** of Extraction Buffer **Blank**, **100 µL** of **Cry1C Positive Control**, and **100 µL** of each **sample extract** to their respective wells.

**NOTE:** It is strongly recommended that a multi-channel pipette be used in steps 1, 4, 7, and 9.



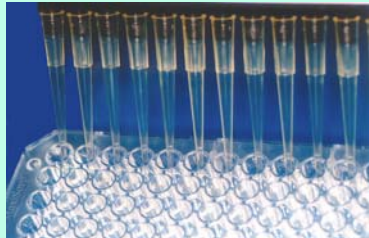
Prepare wash buffer and extraction solutions



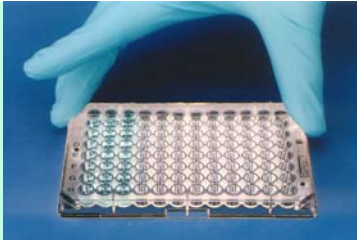
Punch leaf sample



Crush single seed, extract



Add Conjugate, Control, and sample extract



Mix plate



Incubate

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

2. Thoroughly mix the contents of the wells by moving the plate in a rapid circular motion on the bench top 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 15 minutes. If an orbital plate shaker is available shake plate at 200 rpm.

4. Add **100 µL Cry1C Enzyme Conjugate** to each well of the plate. Thoroughly mix the contents of the wells by moving the plate in a rapid circular motion on the bench top for a full 20-30 seconds.

5. Cover the wells with tape or Parafilm to prevent evaporation and **incubate at ambient temperature for 1 hour**. If an orbital plate shaker is available shake plate at 200 rpm.

**NOTE:** Users shall determine appropriate incubation times to give the best results with the tissue disruption/extraction methods in use.

6. 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 Buffer, then shake to empty. Repeat this wash step three times. Alternatively, perform these four washes (300 µL/well) with a microtiter plate or strip washer. Slap the plate on a paper towel to remove as much water as possible.

7. Add **100 µL of Substrate** to each well.

8. Thoroughly mix the contents of the wells, as in step 2. Cover the wells with new tape or Parafilm and **incubate for 15 to 30 minutes at ambient temperature**. Use orbital shaker if available.

**NOTE:** Users shall determine appropriate incubation times to give the best results with the tissue disruption/extraction methods in use.

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

9. Add **100 µL 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.

## How to Interpret the Results

### Spectrophotometric Measurement

1. Set the wavelength of your 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 Extraction Buffer **Blank** wells. If the reader cannot do this, measure and record the optical density (OD) of each well's contents, then subtract the average OD of the **Blank** wells from each of the readings.

### Interpreting the Results

Compare the OD's of the sample extracts to those of the Positive Control to determine presence or absence of Cry1C endotoxin in your sample extract. Samples with absorbances close to that of the Blank wells (and less than that of the Positive Control wells) are presumed to be free of Bt endotoxin. Samples with absorbances significantly higher than those of the Blank wells are positive for Bt endotoxin content.

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

	1	2	3	4	5	6	7	8	9	10	11	12
A	BL	S7	S15	S23	S31	S39	S47	S55	S63	S71	S79	S87
B	PC	S8	S16	S24	S32	S40	S48	S56	S64	S72	S80	S88
C	S1	S9	S17	S25	S33	S41	S49	S57	S65	S73	S81	S89
D	S2	S10	S18	S26	S34	S42	S50	S58	S66	S74	S82	S90
E	S3	S11	S19	S27	S35	S43	S51	S59	S67	S75	S83	S91
F	S4	S12	S20	S28	S36	S44	S52	S60	S68	S76	S84	S92
G	S5	S13	S21	S29	S37	S45	S53	S61	S69	S77	S85	BL
H	S6	S14	S22	S30	S38	S46	S54	S62	S70	S78	S86	PC

“BL” = Blank wells (Extraction Buffer)

“PC” = Cry1Ab Positive Control Wells

“S..” = sample extracts

## Precautions and Notes

- Store all QualiPlate components at 4°C to 8°C (39°F to 46°F) when not in use.
- Do not expose QualiPlate 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 test plates from one QualiPlate with reagents or test plates from a different QualiPlate.
- **Do not expose Substrate to sunlight** during pipetting or while incubating in the test wells.
- Do not dilute or adulterate test reagents or use samples not called for in the test procedure.
- Cry1 proteins can be degraded by heat and sunlight. Take samples from green, actively growing leaves. Leaf samples that cannot be extracted immediately may be stored frozen for up to 1 week prior to analysis. Seeds may be stored for at least 6 months under cool, dry conditions.
- 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.

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