

### Intended use

Cry1F ELISA test kit is intended to be used for the qualitative detection of Cry1F toxin in cotton / corn leaf or seed.

The total incubation time of the assay is 75 minutes.

### Principle of the test

An antibody specific to the Cry1F protein molecule is immobilized on the micro well plates and second antibody specific for Cry1F molecule is conjugated with horse radish peroxidase ( HRP).

When the sample extracts are added to the micro titer wells, the Cry1F from sample binds to the antibody immobilized in the well. This binding is subsequently detected by addition of enzyme labeled antibody. After a washing step, substrate is added. The enzymatic reaction and development of color is proportional to the amount of Cry1F present in the sample. The reaction is terminated by addition of stopping solution. Absorbance is measured on a plate reader.

Light color indicates lower concentration while dark color indicates higher concentration of Cry1F protein in the sample.

### Cross-Reactivity

The Cry1F ELISA kit does not recognize Cry1AC, Cry2A, CP4EPSPS, PAT/pat, Vip3A or PAT/Bar genes.

### Material and equipment required but not provided

- ❖ Pipette with disposable plastic tips
- ❖ Multichannel pipette with disposable pipette tips
- ❖ Deionized or distilled water
- ❖ Graduated cylinders with one liter capacity
- ❖ Reagent troughs
- ❖ Plate shaker
- ❖ Plate ELISA washer or wash bottle
- ❖ Plate ELISA reader with 450/620 nm filter
- ❖ Table top centrifuge
- ❖ Marking pen, Parafilm, Pipette tips, Timer and Paper towels.

### Precautions

The Cry1F ELISA test is intended for in vitro use only. The reagents contain thimerosal as preservative. Prevent direct skin and eye contact with kit components. Seek medical attention in case of accidental ingestion of kit components.

### Contents of the kit:

**Kits are provided in 10 plates or 50 plates format.**

#### **Contents of 10-plate kit:**

1. Ten plates of 96 wells coated with Anti-Cry1F antibody, individually packed in a laminate bag with silica gel
2. Cry1F enzyme conjugate: One Ready- to- use bottle of 55 ml
3. Extraction buffer concentrate: One packet of powder with a 25 ml vial of Tween-20
4. Wash solution concentrate: One packet of powder with a 2.5 ml vial of Tween-20
5. TMB Substrate: One ready-to-use bottle of 110 ml
6. Cry1F Negative control: One ready-to-use vial of 10 ml



### **Contents of 50-plate kit:**

1. Fifty plates of 96 wells coated with Anti-Cry1F antibody, each packed individually in a laminate bag with silica gel
2. Cry1F enzyme conjugate: One ready to use bottle with 275 ml.
3. Extraction buffer concentrate: One packet of powder with a 25 ml vial of Tween-20
4. Wash solution concentrate: One packet of powder with a 2.5 ml vial of Tween-20
5. TMB Substrate: One ready-to-use bottle of 500 ml
6. Cry1F Negative control: One ready-to-use vial of 10 ml

### **Storage of The Kit**

The kit should be stored at 2 - 8° C. The unopened kit is stable till the expiry date printed on the kit label. The expiry date of each unopened component is printed on the label of the component.

### **Reagent preparation**

**The procedure for preparation of reagent is as follows**

#### **Preparation of working extraction buffer:**

Dissolve the packet of extraction buffer concentrate powder in 500 ml of distilled/deionised water and add 25 ml of Tween-20 which is provided along with the powder. Mix it well before use. The resultant extraction solution is 10X concentrate.

**This has to be further diluted with 4500 ml of water to have a working extraction buffer.**

Store unused portion of working extraction buffer at 2 - 8° C. Thaw to room temperature before using the same again.

#### **Preparation of Working wash buffer:**

Dissolve the packet of wash buffer concentrate powder in 500 ml of distilled/deionised water and add 2.5 ml of Tween-20 which is provided along with the powder. Mix it well before use. The resultant wash solution is 10X concentrate.

**This needs to be further diluted with 4500 ml of water to have working wash buffer**

Store unused portion of working wash buffer at 2 - 8° C. Thaw to room temperature before using the same again.

#### **Preparation of stop solution:**

Add 27.0 ml of 98 % H<sub>2</sub>SO<sub>4</sub> in 973 ml of deionized or distilled water. Work in fume hood while preparing stop solution. Store at room temperature for up to one year.

#### **Preparation of user positive control:**

The user is advised to prepare in-house positive control. Crush or grind one known positive seed into a uniform powder and add 1.0 ml of extraction buffer. Wait for 10 minutes before use. Prepare fresh before use.

### **Sample Preparation**

#### **Individual seed**

Crush or grind seed thoroughly into a uniform powder using mortar and grinder or seed crusher. Add 1.0ml of extraction buffer to each sample. Allow the mixture to stand for 30 minutes at room temperature (RT). Allow particle to settle and use only the supernatant to do the test. Wash and rinse grinding equipment carefully between samples to avoid cross contamination.

#### **Individual leaf:**

Weigh (two leaf punch, about 20 mg) and add it to a microtube. Crush leaf with pestle in 0.5 ml extraction buffer. Mix well and incubate for 30 minutes at RT. Allow particle to settle and use only the supernatant to do the test.

Avoid cross contamination between samples.

## Assay Procedure

Test protocol for individual seed or leaf samples

1. Allow all reagents to reach room temperature before use.
2. Add 50  $\mu$ l enzyme conjugate to each well.  
Add 50  $\mu$ l extraction buffer to blank well.
3. Add 50  $\mu$ l of positive control and 50  $\mu$ l of negative control in duplicate wells.
4. Add 50  $\mu$ l of sample extracts to microtiter wells. Mix contents of the plate carefully avoiding cross contamination.
5. Cover the plate and incubate it for 60 minutes at room temperature.
6. Aspirate and wash the wells 4 times with 300  $\mu$ l of washing solution.
7. Add 100  $\mu$ l substrate per well.
8. Cover the plate and incubate for 15 minutes at room temperature.
9. Stop the reaction by adding 100  $\mu$ l stop solution.
10. Measure the absorbance at 450 nm (primary filter) and 620/630 nm (secondary filter). Read the plate within 15 minutes after addition of stop solution.



### **Notes on Technique:**

1. Protect the plates from draught, strong light or direct sunlight during the test procedure.
2. Careful aspiration of the washing solution is essential for good assay precision.
3. Since the timing of the incubation steps is important to performance of the assay, use multi channel pipettes to dispense samples/reagents.
4. Plate readers measure absorbance vertically. Do not touch the bottom of the wells.

## Interpretation of Results

### **General test criteria:**

- ❖ Absorbance of the blank well should not exceed 0.1
- ❖ Absorbance value of blank should be subtracted from positive control, negative control and samples.
- ❖ The coefficient of variance between the duplicate positive control should be less than 15 %
- ❖ Mean positive control should have absorbance of at least 0.5
- ❖ Mean negative control should have absorbance below 0.1

If above criteria are not met, the test is invalid and should be repeated.

## Qualitative Test

For single corn/cotton leaf and seed samples:

**CUTOFF CALCULATION** : for single corn/cotton leaf or seed sample

Cutoff value: (Mean absorbance of negative control +0.3)

**Positive sample:** If sample absorbance is more than the cutoff value.

**Negative sample:** If sample absorbance is less than the cutoff value.

Individual seed or leaf samples are either positive or negative. Low level results normally indicate improper washing, cross contamination or technique. In such cases, retesting is recommended.

### **Performance characteristics**

Sensitivity: 100 %

Specificity: 100 %



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