Probe qPCR Detecting GMO Marker MON87411 from Maize

DESCRIPTION

Many countries in the world regulate the cultivation and trade with genetically modified organisms (GMO). The enforcement of such regulations depends on the ability to detect and quantify the presence of GMOs in food, feed, and seed products. Real-time qPCR method is sensitive and robust, and is regarded as the gold standard for GMO analysis.

The Maize GMO MON87411 qPCR Kit is designed for identifying GMO marker MON87411 presence or absence in food, feed and seed products using real-time quantitative polymerase chain reaction(qPCR), primers, and labeled probe. The Kit includes maize GMO MON87411 Positive and Negative controls, and PCR internal controls, qPCR Super Mix, maize GMO MON87411 Prime-Probe Mix, in which the probe is labeled with Fam, and Hex is labeled for PCR internal control. These aids in the straightforward interpretation of the results.

PRINCIPLE

GMO marker identification is based on qPCR amplification of a specific region of the relevant target genome. The amplified product is detected using target-specific fluorescent probes that bind to the amplified product. As the PCR product accumulates, there is an increased fluorescent signal from the bound probes. Monitoring the fluorescence intensities during the PCR run allows the detection of the accumulating PCR product in real time.

KEY FEATURES

- Highly sensitivity and specificity for identification of GMO MON87411.
- High efficiency: the optimal systemic conditions for PCR amplifications.
- Streamlined protocol: Just add DNA Template, and water.
- ♦ No cross reactivity with others.

APPLICATIONS

Detect GMO MON87411 in food, and animal feed.

KIT CONTENTS

Name	100RXN
qPCP Super Mix	0.8 mL
Primer-probe Mix	0.6 mL
Positive Control DNA	60 µL
Negative Control DNA	60 µL

MON87411 gene is labeled with FAM; PCR internal control is labeled with Hex.

STORAGE CONDITION

The kit is shipped on ice and stored at -20°C for long-term storage. Shelf life of 12 months after receipt.

PCR PROTOCOL

1. DNA extraction: The methods for DNA extraction can be used for any suitable preparation of DNA purification from food samples. We recommend that the Fast genomic DNA extraction method is used for this purpose (catalog: TBS6008).

2. Set up PCR reaction for each sample in 20 µL

Reaction Component	Volume (µL)	
qPCR Super Mix	7.0	
Primer-probe Mix	5.0	
Nuclease-free Water	3.0	
DNA sample	5.0	
Final Volume	20 μL	

Internal control should be included as below: Positive Control or Negative control (5 μ L /reaction).

3. Suggested PCR conditions

	Amplification	PCR	
Step	HOLD	CYCLE (40 cycles)	
		Denature	Anneal/ Extend
Temperature	95 °C	95 °C	60 °C
Time	1 min	15 sec	60 sec

DATA ANALYSIS

Positive Reaction: Sample Ct < or = 37, and Positive, Negative and Blank controls are normal.

Negative Reaction: Sample $Ct \ge 38$, and Positive, Negative and Blank controls are normal.

PCR internal control is positive in all samples, positive and negative controls. The positive response indicates a normal PCR amplification. Otherwise, the PCR reaction may be inhibited. Repeat Reaction: If one of the control reactions is not normal, PCR reaction is failed, and should be repeated.

RELATIVE PRODUCTS

TBS6008: Fast Genomic DNA Extraction TBS6025: Microbial DNA Magnetic Extraction TBS43001: Maize GMO 98140 qPCR Detection TBS43003: Maize GMO DBT418 qPCR Detection TBS43004: Maize GMO LY038 qPCR Detection TBS43005: Maize GMO Mon863 qPCR Detection TBS43007: DAS-59132 qPCR Detection TBS42025: 4-In-1 Aspergillus Species qPCR TBS42026: O157H7 E. Coli qPCR TBS42028: Salmonella qPCR TBS42029: Salmonella qPCR TBS42031: Listeria Monocytogen qPCR TBS42032: Listeria Species qPCR TBS42033: Bacillus Cereus qPCR

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