BÍONEER

[Cat. No.] K-2962

Introduction

Innovation • Value

AccuPower[®] MON87769 PCR Kit is a ready-to-use premix for PCR that can be used to detect the *PJ.D6D* gene and the *NC.FAD3* gene that has been introduced into genetically modified (GM) soybeans to generate Stearidonic acid (SDA).

Genetically modified organisms (GMOs) are organisms whose genetic material has been artificially modified to obtain desired characteristics, such as increased yield and resistance to pests and pathogens. Only those that have the approval of a regulatory agency, such as the Ministry of Food and Drug Safety in South Korea can be sold. Nowadays, many countries/regions require manufacturers to label products that include genetically modified products, leading to an increase in the demand for GMO detection technology.

This product contains vacuum-dried components specific to MON87769 GMO soybean including DNA polymerase, primers, dNTPs, and reaction buffer required for PCR. This ready-to-use kit simplifies preparation of PCR mixture as the user only has to add template DNA and nuclease-free water. After the reaction, since tracking dye is included, the samples can be applied directly on agarose gel for analysis without adding extra solution.

Features & Benefits

- Convenience & Reproducibility: All reactants necessary for PCR including primers are lyophilized in each PCR tube, providing reproducible results in a convenient way.
- Sensitivity: By applying the patented PyroHotStart (Enzymemediated HotStart) technology that minimizes non-specific reactions and maximizes reaction efficiency, only the target gene can be effectively amplified even with a trace amount of template DNA.
- Stability: Included stabilizer in the PCR reaction mixture provides increased stability compared to solution-type products.

Composition

Composition	25 µl reaction
Top DNA Polymerase	1 U
dNTPs (dATP, dCTP, dGTP, dTTP)	Each 250 µM
Reaction buffer with 1.5 mM MgCl ₂	1X
Stabilizer and tracking dye	0
MON87769 Forward primer	0.5 µM
MON87769 Reverse primer	0.5 µM

* Note: For research use only. Not for use in diagnostic or therapeutic procedures.

Specifications

Top DNA Polymerase				
5' \rightarrow 3' exonuclease activity	No			
$3' \rightarrow 5'$ exonuclease activity	No			
3'–A overhang	Yes			
Fragment size	111 bp			

Storage

Store at -20°C. If stored in the recommended temperature, this product will be stable until the expiration date printed out on the label.

Online Resources



Visit our **product page** for additional information and protocols

Ordering Information

Description	Cat. No.
AccuPower [®] MON87769 PCR Kit, 0.2 ml thin-wall 8- tube strips with attached cap / 96 tubes	K-2962

Notice

BIONEER corporation reserves the right to make corrections, modifications, improvements and other changes to its products, services, specifications or product descriptions at any time without notice.

Explanation of Symbols



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Experimental Procedures

Steps		Procedure Details				
1	Add template DNA	1. After preparing the template DNA and nuclease-free water, add the template DNA to the <i>AccuPower</i> [®] MON87769 PCR Kit.				
2	Preparation of reaction mixture	 Add nuclease-free water into PCR tubes to make a total volume of 25 µl. (Do not include the volume of the dried premix in the PCR tubes.) Completely dissolve the vacuum-dried pellet by vortexing, and briefly spin down. 				
		 Place PCR tubes on the thermal cycler. Perform the reaction under the following conditions. 				
	and and	Step	Temperature	Time	Cycles	
		Pre-denaturation	95°C	10 min	1 cycle	
3		Denaturation	95°C	30 sec		
	han a ha ta maa ti'a ma in a	Annealing	60°C	30 sec	40 cycles	
	Incubate reactions in a thermal cycler	Extension	72°C	30 sec		
		Final extension	72°C	7 min	1 cycle	
		* Note: Users can adjust the protocol according to their instrument and template sequences to get optimal results.				
4	Analyze with gel electrophoresis	 6. After the reaction, maintain the reaction mixture at 4-8°C. 7. Load samples on agarose gel without adding a loading-dye mixture, and perform gel electrophoresis for analysis. 				

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