

# One-step real-time Probe RT-PCR for Arabis mosaic virus (ArMV) Detection from plant samples

Catalog	Kit Size (RXN)
TBS42014-50	50
TBS42014-100	100

## **DESCRIPTION**

Arabis mosaic virus (ArMV) RT-qPCR Detection Kit is one step real-time reverse transcription polymerase chain reaction (RT-PCR) assay intended for the detection of ArMV infection in plant samples. It combines both reverse transcription (RT) and Taqman probe-based PCR amplification to occur in a single reaction tube. The kit is optimized for the two reactions in a real-time "single step". This One-step qRT-PCR assay offers the end-users an efficient, easy to use, and reliable alternative to conventional "two-step" sequential qRT-PCR for detecting ArMV.

The ArMV RT-qPCR Detection Kit contains all components for RT-PCR reaction including RT-PCR enzymes, primer-probe, internal control, positive control, negative control, and buffers. The ArMV target gene is labeled with fluorescence Fam, and internal control is labeled with Hex.

#### **APPLICATIONS**

This kit is used for the detection of ArMV infection in plant samples.

#### **KEY FEATURES**

- One-step complete qRT-PCR reaction in a single tube.
- Reduce contamination in the operating process.
- Accurate detection and quantification of ArMVd target gene.
- 4x RT-PCR mix makes it easier to adjust the sample size in a one-tube reaction.

## KIT CONTENTS

Component	50RXN	100RXN
4X RT-PCR Master Mix	0.25mL	0.5 mL
10X ArMV Primer-probe Mix	0.1mL	0.2 mL
ArMV Positive Control	50 μL	100 μL
ArMV Negative Control	50 μL	100 μL
4x RT-PCR Buffer	0.25 mL	0.5 mL
DEPC Water	1.25 mL	1.25 mL

#### STORAGE CONDITIONS

Store all components at -20°C in a non-frost-free freezer. Shelf life is 12 months after receipt.

The kit is shipped on ice.

### PROTOCOL

Precautions: RT-PCR should be assembled in a nuclease-free environment. RNA sample preparation, reaction mixture assembly, PCR and subsequent reaction analysis should be performed in separate areas. The use of "clean", automatic pipettors designated for PCR and aerosol resistant barrier tips are recommended. Note: Nuclease Removal (TBS6013) is useful for removing RNase contamination in the working area and tools.

- 1.RNA isolation is performed with a suitable approach. We recommend our Hybrid-R RNA purification kit (catalog: 305-101) for RNA extraction from plant samples.
- 2. Prepare the following reaction mixture on ice

Components	Reaction Vol.:20 μl	Concentration
Total RNA	Variable	5 pg - 1 μg/rxn
RT-PCR Mix (4x)	5 μl	1X
Primer-probe Mix (10x)	2 μl	1x
RT-PCR Buffer (4x)	5 μl	1x
DEPC water	Adjust to the fina	l volume to 20

#### Positive or Negative Control: 4µl/well.

The ArMV target gene is labeled with Fam, and internal control is labeled with Hex

#### Note:

- 1. Gene specific primers and probe must be used for RT*qPCR* amplification.
- 2. Please ensure no salt crystals are present in the RTqPCR Mix before use. If salt crystals are observed, mix until crystals are completely dissolved and absent.
- 3. For Positive and negative control, use 2 µl of Positive or negative control to replace the RNA sample.
- 3. Gently mix and ensure all the components are at the bottom of the amplification tube. Centrifuge briefly if needed.
- 4. Program the thermal cycler so that cDNA synthesis is followed immediately by qPCR amplification.

Step	Temperature	Duration	Cycle(s)
1	50°C	20 mins	1
2	95°C	1 min	1
3	95°C	15 secs	40
	60°C	60 secs	40

### **Recommendations for Optimal Results**

- Aliquot reagents to avoid contamination and repeated freeze-thaw cycles.
- Start reaction as soon as the reaction mixture is prepared and always keep the reaction mixture chilled in an ice box prior to qRT-PCR reaction.

### **DATA ANALYSIS**

After completion of the run, save and analyze the data following the instrument instructions. Analyses should be performed separately for each fluorescence channel using a manual threshold setting. Thresholds should be adjusted to fall within exponential phase of the fluorescence curves and above any background signal. The procedure chosen for setting the threshold should be used consistently.



The results can be interpreted as below:

Positive: ArMV gene Ct value is in 12-36, and internal control

(Hex), Positive control, and Negative control are

normal.

Negative: ArMV gene Ct value is ≥37, and internal control

(Hex), Positive control, and Negative control are normal.

### **RELATED PRODUCTS**

TBS6025: Microbial DNA Magnetic Extraction

TBS42015: HLVd RT-qPCR Detection

TBS42018: Trichothecene-producing Fusarium Species

TaqProbe qPCR Detection

TB\$42019: Fusarium Species qPCR Detection

TBS42025: 4-In-1 Aspergillus qPCR

TBS42029: STEC and Salmonella Multiple qPCR

TBS42031: Listeria Monocytogenes qPCR

TBS42032: Listeria Genus qPCR TBS42033: Bacillus Cereus qPCR TBS42043: Bacillus Species qPCR

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