

Minute Virus of Mice probe qPCR Detection Kit

Product Number: DTK960

Shipping and Storage

Low temperature transportation, stored at -20°C, with a shelf life of one year. The positive control should be placed separately and should not contaminate other reagents.

Component

Component	50T
2 × Probe qPCR Mix	100μL
DEPC-H ₂ O	1mL
Fluorescent template diluent	1mL
Mouse parvovirus qPCR mixture	260μL
Mouse parvovirus qPCR positive control (1 × 10E8 copy/μL)	50μL

Application

This kit is designed for the detection of mouse parvovirus.

Features

Developed based on the principle of fluorescence quantitative PCR using probe technology, it features the following characteristics:

1. Ready to use, users only need to provide a sample DNA template.
2. Primers and other components have been optimized for high sensitivity.
3. Provide positive controls to distinguish false negative samples.
4. High specificity, primers are designed based on highly conserved regions of mouse parvovirus DNA sequences, and do not cross react with DNA from other viruses.
5. It can be used for both qualitative and quantitative testing. When used for quantitative detection, the linear range should be at least 5 orders of magnitude.
6. This product is sufficient for 50 fluorescent quantitative PCR reactions using a 20 μ L probe system.
7. This product can only be used for scientific research.

Protocol

1. DNA extraction (sample preparation area)

- 1.1. If there are N samples to be extracted, it is best to set N+2 extractions, with the additional being PC (positive control for sample preparation) and NC (negative control for sample preparation). You can take 10μL of 1000 fold dilution of the positive control and add a certain amount of water to make the total volume consistent with the specified volume of the sample to be extracted, which can be used as PC. In addition, water can be used as NC.
- 1.2. Extract and purify sample DNA using a self selected method, and this kit is compatible with most nucleic acid extraction kits on the market.

2. Dilute standard curve sample (sample preparation area)

Due to the high concentration of positive control, the following dilution operations must be performed in a separate area to avoid contaminating the sample or other components of this kit.

- 2.1. Mark 6 centrifuge tubes, namely 7, 6, 5, 4, 3, and 2.
- 2.2. Add 45μL of fluorescent template diluent separately using a core gun tip (preferably using a core gun tip, the same below).

- 2.3. Add 5µL of 1×10^8 copy/µL positive control (provided by the reagent kit) to tube 7, shake thoroughly for 1 minute, and obtain 1×10^7 copy/µL standard curve sample. Put it on ice for later use.
- 2.4. Change the gun head and add 5µL of 1×10^7 copy/µL positive control (obtained from the previous dilution) to tube 6. Shake thoroughly for 1 minute to obtain a standard curve sample of 1×10^6 copy/µL. Put it on ice for later use.
- 2.5. Change the gun head and add 5µL of 1×10^6 copy/µL positive control (obtained from the previous dilution) to tube 5. Shake thoroughly for 1 minute to obtain a standard curve sample of 1×10^5 copy/µL. Put it on ice for later use.
- 2.6. Repeat the above operation until obtaining standard curve samples with 6 dilutions. Put it on ice for later use. If no standard curve is required, dilute the positive control to 1×10^5 copies/µL.

3. Reagent preparation (reagent preparation area)

Prepare sufficient qPCR tubes (sample tube, negative control tube, positive control tube) and add the following components to each qPCR tube.

Component	N sample tubes	qPCR negative control	qPCR positive control
2 × Probe qPCR Mix	each 10µL	10µL	10µL
Mouse parvovirus qPCR mixture	each 5µL	5µL	5µL

Transfer to the template addition area.

4. Add Template (Template Add Area)

Add 5µL of template to each qPCR tube, in the order of negative control (DEPC-H₂O), test sample template, and mouse parvovirus qPCR positive control. Centrifuge for 30 seconds and immediately perform amplification reaction.

5. Amplification reaction (amplification and product analysis area)

Place the qPCR tube in the corresponding position of the qPCR amplification instrument sample slot for amplification. The amplification procedure is as follows:

Process	Temperature	Time
Pre denaturation	95°C	3min
qPCR reaction	95°C	15sec
(45 cycles)	60°C	20sec
Signal channel	FAM channel collects fluorescence signals	

6. Result Analysis

- 6.1. If creating a standard curve, plot the standard curve with the log value of positive control concentration as the horizontal axis and Ct value as the vertical axis. Calculate the log value of the DNA concentration of the sample from the standard curve based on the Ct value of the sample to be tested, and determine its concentration.
- 6.2. If no standard curve has been created, the results shall be judged according to the following criteria:
 - Positive control (1×10^5 copies/µL) result: Ct value < 30, with significant exponential growth, showing a typical S-shaped curve.
 - Negative control result: Ct value > 40 or no Ct value, no significant exponential growth period or plateau period.
 - Sample testing results: Ct value < 38, with a significant exponential increase, indicating the detection of mouse parvovirus in the sample, and the result is positive; A Ct value greater than 40 or no Ct value indicates that no mouse parvovirus was detected in the sample, and the result is negative; If the Ct value is within the range of 38-40, the sample should be retested. If the Ct value of the repeated experiment is still within the range of 38-40 and there is a significant exponential increase, it is judged as positive. Otherwise, it is judged as negative.