Precautions

- 1. This product is only suitable for in vitro diagnostic testing, and the experimenter should have received professional training and obtained the relevant work permit.
- 2. It shall be implemented strictly in accordance with the management norms of gene amplification laboratory promulgated by the relevant competent authorities. The laboratory is divided into three areas for operation (reagent preparation area, sample preparation area and amplification area), items in each area shall not be cross-used, and special instrument and equipment shall be used in each area.
- The performance characteristics of this test kit for the sample types listed in the section "Intended use" have been identified. The performance of this test kit for other sample types has not been evaluated
- 4. The tip with filter and centrifuge tube used in the experiment should be autoclaved, and without DNase and RNase. After use, it is directly put in the disposal bottle containing 1% sodium hypochlorite.

nufactSymbols meaning

8	Do not re-use		Manufacturer	EC REP	Authorized representative in the European Community
Ť	Keep dry	М	Manufacture Date	8	Do not use if package is damaged
	Caution	arc arc	Store at 2-8 °C	IVD	In vitro diagnostic medical device
CE	CE Symbol	漱	Keep away from sunlight	LOT	Lot number
REF	Catalogue number	Ĩ	Consult instructions for use	2	Use by date

Basic Information

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techstar®

Monkeypox Nucleic Acid Detection Kit (Fluorescence PCR)

Instruction For Use

For in vitro diagnostic use only For Prescription Use only

Name	Monkeypox Nucleic Acid Detection Kit (Fluorescence PCR)		
REF	SC535482	SC535962	
Specification	48 reactions/kit	96 reactions/kit	

Intend Use

For the qualitative detection of Monkeypox virus nucleic acid in skin exudate samples, The results are only for clinical reference and should not be used as conclusive evidence for the diagnosis or exclusion of diseases.

Principle

This product uses qPCR technology to qualitatively analyze the nucleic acid of Monkeypox virus. During the amplification process, when the reaction system does not contain Monkeypox virus nucleic acid, the probe is complete, because the 3 'quenched group is close to the 5' reporter group, the fluorescence emitted by the reporter group is absorbed by the quenched group, and no fluorescence signal is emitted. When the Monkeypox virus nucleic acid was contained in the reaction system, the fluorescent probe bound to the template during primer extension is cleaved by Taq enzyme (5'-3' exonuclease activity), and the reporter group was separated from the quenched group to emits a fluorescence signal.

The real-time amplification curve can be generated by fluorescent quantitative PCR instrument automatically according to the detected fluorescence signal, so as to achieve the purpose of qualitative analysis of Monkeypox virus nucleic acid in the sample.

The assay also includes an endogenous ribonuclease P (RNaseP) as an internal control (labeled with CY5), to monitor the entire process of specimen collection, nucleic acid extraction, and PCR amplification.

Materials Provided

Constituent	Component	48 reactions/kit	96 reactions/kit
Dilution Buffer	Nuclease-free water	1.0mL/vial, 1 vial	2.0mL/vial, 1 vial
MPV-D PCR Mix	MPV-D PCR Mix Primer Probe dNTPs Enzyme		2 vial
Positive control	Plasmid containing the target gene	1 vial	1 vial
	and internal control	i viai	i viai
Negative control	Nuclease-free water	2.0mL/vial, 1 vial	2.0mL/vial, 1 vial

Note: The components in different batch are not interchangeable.

Materials and Equipment Required But Not Provided

Applied Biosystems[™] Real-Time PCR System 7500.

Alternatively, Roche LightCycler® 480 System, or QuantStudio 5 Real-Time PCR System. • QIAamp DNA Mini Kit (cat. #51304 or 51306).

Alternatively, Nucleic Acid Extraction Kit (cat. SC906). Optionally, Nucleic Acid Extraction Kit (cat. SC902/905).

- Vortex mixer.
- Microcentrifuge.
- Micropipettes (2 or 10 $\mu L,$ 200 μL and 1000 $\mu L).$
- Racks for 1.5 mL microcentrifuge tubes.
- 10% bleach (1:10 dilution of commercial 5.25-6.0% hypochlorite bleach).
- Disposable powder-free gloves and surgical gowns.
- 1.5 mL microcentrifuge tubes (DNase/RNase free).
- + 96-well 0.2 mL PCR reaction plates or appropriate optical reaction tube.

Storage conditions and Expiration period

Store at 2-8 $^\circ$ away from light for 12 months or 20-25 $^\circ$ away from light for 6 months. Date of manufacture: see label Expiration Date: See label

Applicable Instrument

ABI7500, ABI QuantStudio 5, CFX96 Dx Real-Time PCR Detection Systems, CFX96 Touch Real-Time PCR Detection System, CFX Opus Real-Time PCR Systems and Roche LightCycler480

Specimen Collection and Preparation

- 1. Available sample types: skin exudate swabs.
- After collection, the swab was placed in normal saline or other sample preservation solution (Hank's solution or guanidine-based virus transport media).
- Appropriate specimen collection, transport, storage and processing procedures are required for the optimal performance of this test. Improper collection, storage, or transport of specimens may lead to false negative results.
- 4. The specimen may be tested immediately after collection, or it may be stored at 2-8 C for up to 24 hours before testing. If the expected detection time exceeds 24 hours, you can choose to store the samples below -70°C.

Laboratory Procedures

1. Sample processing

- 1.1 DNA should be collected from fresh specimen to ensure suitable DNA quality and quantity. DNA should be extracted using the QIAamp DNA Mini Kit (Qiagen) or Nucleic Acid Extraction kit manually (Art.No.SC902/SC905/SC906, Wuxi Tech-star Technology Co.,Ltd) according to the manufacturer's Instruction of Use.
- 1.2 Add 500µL Negative control or Nuclease-free water to the Positive control. Vortex the tubes to mix then centrifuge at 6000rpm for 30s. Positive Control and Negative control should be processed simultaneously alongside the specimen.
- 1.3 After extraction, the DNA should be used immediately or stored at -20 $\rm C$ for use later. When handling the Positive control, please take precautions to avoid contamination of the specimen sample.

A The Positive Control and Negative Control only need to be extracted once, and the extracted nucleic acid can be directly used in the next experiment.

2. Reagent preparation

- 2.1 Take out the kit contents and thaw thoroughly at ambient temperature. Prepare 96-well plates or appropriate optical reaction tube for real-time PCR based on the estimated number of reactions.
- 2.2~ Adding 960 $\!\mu$ l Dilution Buffer to the MPV-D PCR Mix for redissolve and mix it.
- 2.3 Pipette 20 μ L of PCR-Mix into each well.

▲ The remaining PCR-Mix, Positive control and Negative control must be stored at under -18°C immediately, and it should be used within 20 days, repeated freezing and thawing times should not exceed 7 times.

3. Sample Addition

Add 5µL extracted sample DNA, Negative control or Positive control, and close the 96-well reaction plate with appropriate lids or optical adhesive film, and centrifuge at 6000rpm for 30s.

A Make sure that at least one Positive Control and one Negative Control is used per run.

4. Amplification(ABI7500, for example)

- Double-click 7500 software or select Start>>All Programs>>Applied Biosystems>>7500 Software.
- 4.2 Click "New Experiment" to enter Experiment menu. In the Experiment Properties interface, enter identifying information for the experiment; you can leave other fields empty.
- 4.3 Select "7500 (96 Wells)", "Quantitation-Standard Curve" (for the experiment type); "TaqMan Reagents" (for reagent); and "Standard" (for ramp speed).
- 4.4 Go to Plate Setup > Define Targets and Samples > Define targets > Add New Target > set Target Name and Reporter as shown below:
- Target 1. Monkeypox: Reporter FAM; Quencher None Target 2. IC: Reporter CY5; Quencher None
- $4.5 \ \ \, {\rm Go\ to\ Define\ Samples\ >\ Add\ New\ Sample\ >\ Input\ PC,\ NC\ and\ Sample\ (Test\ Specimen)}$
- 4.6 Click "Assign Target and Samples" to set targets and well positions for PC, NC and Samples to be analyzed. in the "View Plate Layout" interface, enter the name of samples and controls to include in the reaction plate in corresponding well, and select the sample/target reactions to set up. Select "None" for passive reference.

Step	Temperature		Fluorescence measured	Cycle	
1	95°C	3min	No	1	
2	95°C	15sec	No	45	
3	58°C	30sec	Yes	45	

4.8 Click "Run". In the Run interface, save the experiment. Click "START". 4.9 After the run completes, take out the sample andproceed to data analysis.

5. Analysis of result

The results are automatically saved after the reaction, and the Baseline Starting Cycle, Baseline End Cycle and Threshold of the Baseline are adjusted according to the analyzed images. Generally, the Baseline Starting point is set to 3--15. The baseline endpoint should be set at 5-20, and the threshold should be in the exponential phase of the amplification curve, and the curve for negative quality control should be flat or below the threshold line. Click Analysis to automatically analyze the results.

Interpretation of result

1. Quality control standards

- 1.1 Negative control:No typical S-type amplification curve or no Ct for FAM and CY5 channels.
- 1.2 Positive control: FAM and CY5 channels showed a typical S-type a amplification curve.
- If both meet the requirements of 1.1 and 1.2, the experiment is effective, otherwise it is invalid

2. Result interpretation

When the above quality control conditions are met, analysis as follows:

- 2.1 **Positive:** The FAM channel has a S-type amplification curve, and Ct≤38, it can be determined as Monkeypox virus nucleic acid positive;
- 2.2 Negative: The FAM channels to be tested have no Ct value, and the CY5 channel showed a typical S-type amplification curve, which was judged to be Monkeypox virus nucleic acid negative.
- 2.3 Suspected: If FAM channels to be tested has S-type amplification curve, and Ct>38, the sample needs to be extracted and tested again. If the retest results are consistent, it can be judged as Monkeypox virus nucleic acid positive; if the retest results are negative, it can be judged as Monkeypox virus nucleic acid negative.
- 2.4 If there is no typical S-type amplification curve in the FAM and CY5 channels, it indicates that the system is inhibited or mis operated, and the test is invalid. Therefore, the samples should be retested.

Limitations

- Negative result cannot completely exclude pathogen infection, and the concentration of target genes in sample below the detection limit or the mutation of target sequences to be tested can also result in negative result.
- False negative or false positive result may be caused by improper sample collection, transport and handling, as well as improper experimental operation and experimental environment.
- 3. The samples collected from the patients who used drug therapy may result in false negative results.
- 4. Diseases caused by other bacterial or viral pathogens cannot be ruled out.
- 5. The testing result of this product are only for clinical reference and should not be used as the sole basis for clinical diagnosis and treatment, and clinical management of patients should be considered in combination with their symptoms/signs, medical history, other laboratory tests and treatment responses.
- 6. Thenucleic acid may remain in the body for a long period of time, which is not related to virus activity. A positive test result does not necessarily mean that the virus is infectious, or that the virus is a pathogen that causes clinical symptoms.,

Performance Characteristics

1. Limit of Detection

The LoD of the Monkeypox Nucleic Acid Detection Kit was estimated by testing the standardized dilutions of pseudo-virus (n = 20 each). The lowest target level at which \geq 19/20 produced positive results was 500 Copies/mL. This value was then confirmed by testing 20 replicates.

2. Analysis specificity

The weak positive sample and medium/strong positive sample were tested with three batches of Monkeypox Nucleic Acid Detection Kit. Each batch of reagents is tested by 2 people per day, and each person test once in the morning and afternoon. Each sample was tested 2 times in parallel and is tested continuously for 20 days. The results met the following criteria.

- 2.1 Weak positive sample: the positive detection rate should be 100%, the Coefficient of variation of Ct of FAM channel should ≤5.0%.
- 2.2 Medium/strong positive sample: the positive detection rate should be 100%, the Coefficient of variation of Ct of FAM channels should ≤5.0%.