## **FIASHTEST**

[Product Name]
Pseudorabies virus gB /gE PCR Test Kit (Lyophilized)

[Package Specifica ns]

[Intended Use]
The kit uses fluorescence PCR method to identify two pathogenic genes of Pseudorabies virus gB /gE. This product requires fluorescence quantitative PCR instrument operation and can achieve rapid detection of POCT.

Testing Principle]

The test kit uses nucleic acid extraction reagents to extract the nucleic acid (DNA/RNA) from the sample.

Under the action of a high-efficiency reverse transcriptase, cDNA complementary to the RNA template is synthesized in a one-step reactior using RNA as the template. Under the action of Taq enzyme, the copy number of the specific target fragment is amplified through cycles of high-temperature denaturation, annealing at a moderate temperature, and extension using DNA as the template. The fluorescence-labeled specific probe hybridizes with the amplified target fragment, and the 5—3 exonuclease activity of Taq polymerase separates the reporting group an equencher group of the fluorescence probe, emitting a specific fluorescence signal. The specific fluorescence PCR instrument, and the result is determined based on the Ct value of the sample and the formation of the amplification curve.

[Contents]			
Item	Quantity	Storage	
PCR master mix	16 pcs	-20°C (Away from light)	
Instructions for use	1 pcs		
Sample buffer	16 pcs	Room Temperature	
Swab	16 pcs		
Biohazard bag	16 pcs		

# [Storage conditions and shelf I 1. Shelf life: 24 months. 2. Production date and expiration

[Compatible Instruments]
This test kit is compatible with FLASHTEST refluorescence PCR instrument.

[Sample] Tissue, throat sw

- Tissue, throat swab, fresh feces

  [Sample Handling]
  1. Tissue:
   Collect 1g of sample from each part of the bird to be tested (ex: Lung, tonsil, kidney, liver, brain tissue).
   Cut and mix sample. Take tig sample from the mixed tissue. Add 1ml of saline to the sample and run homogenization.
   Centrifugle homogenates at 3000 rpm for 2 minutes.
  2. Swab:
   Throat swab: Use a swab to moderately wipe the oral.
   Fresh feces swab: Use a swab to collect an appropriate amount.
   With the swab in the sample buffer, shake it thoroughly to fully dissolve the pathogen on the swab head into the buffer.
  3. Add 200 μL of mixed buffer to the nucleic acid extraction cartridge for extraction.

[Specimen storage]
Samples used for nucleic acid extraction and detection should be tested as soon as possible.
Samples to be tested within 24 hours can be stored at 4°C.
Samples that can not be tested within 24 hours should be stored at -20°C for up to 10 days.
Avoid repeated freezing and thawing of samples.

[Instructions for Use]
1. Add Elution
1.1 Add 20 µL of elution from magnetic bead extraction, to each PCR tube.
Close the lid flightly.
1.2 Shake all the liquid to the bottom of the PCR tube. Use the vortex mixer to mix the PCR tube horoughly, for 5 seconds. After mixing, make sure all liquid is at the bottom of the PCR tube, by shaking the tube again (optional: use a small centrifuge for 3 seconds to shift all liquids to the bottom.)

### 2. PCR Ar

2.1 Set the parameters as follows:			
Step	Temperature	Time	Cycle
1	55°C	3min	1
2	94°C	30s	1
3	94°C 58°C	5s 20s	×40

2.2 The reaction volume is 20µL. Fluorescence channels:				
Channel	FAM	VIC	ROX	Cy5
Target	PRV-gB	Internal reference		PRV-gE

erpr

Parameter	Reference Range	Result Interpretation
Internal Control	Ct ≤ 37 and there is a clear exponential amplification curve	Valid
Control	Ct > 37 or No Ct	Invalid
Pathogen	Ct ≤ 37 and there is a clear exponential amplification curve	Positive
	Ct > 37 or No Ct	Negative

3.2 Test Result Interpretation			
Pathogen Result	Internal Control Result	Test Result Interpretation	
Positive	Valid	Pathogen Positive	
Negative	Valid	Pathogen Negative	
Any Result	Invalid	Test invalid, please retest	

[Test Limitations]

1. The test results of this kit should be comprehensively analyzed in conjunction with other relevant physical examination results and should not be used as the sole basis for diagnosis.

2. Improper sample collection, transportation, storage, handling, and inadequate laboratory conditions may lead to inaccurate results.

3. Other unconfirmed interferences or PCR inhibitors may lead to false negative results.

4. Sequence variations caused by mutations or other factors in the targ gene of the virus being tested may lead to false negative results.

- [Product Performance]

  [Product Performance]

- Sample is 50%.

  [Notes]

  I. Before using a PCR kit, check the lyophilized PCR mix at the bottom of the tube is in good condition (white and clumped). Liquiffied lyophilized PCR mix an not be used. After opening, it should be used as soon as possible or stored away from light.

  2. This product is only for in vitro testing (for animals). All operations mus strictly follow the instructions.

  3. Overloading samples may result in false negatives. Retest is recommended.

  4. Avoid bubbles in PCR tubes. Keep the tube cap firmly closed.

  5. Use disposable tips, gloves, and laboratory coats.

  6. After tests, disinfect the workbench with 10% hypochlorous acid, 75% ethanol, or UV light.

  7. All items in the kit should be treated as biowaste and handled in accordance with local laboratory regulations.