

Distribuito in ITALIA da Li StarFish S.r.l. Via Cavour, 35 20063 Cernusco S/N (MI) telefono 02-92150794 fax 02-92157285 www.listarfish.it

For research use only Cat. No. IP11073 | **48 Tests**

Canine Herpesvirus Detection Kit

Test for the detection of Canine Herpesvirus by one-step PCR

User Manual

REV.2.2

Canine Herpesvirus Detection Kit

Table of Contents

1.		otion	
2.	Storage	9	1
3.	Conten	ts ·····	1
4.	Specim	en	1
5.	Additio	nal required materials	2
6.	Proced	ure ·····	2
	6.1	DNA preparation	2
	6.2	Amplification ·····	2
	6.3	Detection of amplification product ······	
	6.4	Interpretation	3
	6.5	Elimination of carry-over contamination	3
7.	Notice		4
8.	Trouble	shooting	4
9.	Orderin	ng information	4

Canine Herpesvirus Detection Kit

■ NOTE :	

7. NOTICE

- This product was designed to detect more than 100 copies of target gene(or gene segment). When the copy number of target present in the test reaction is less than 100, a false-negative(a negative test result when the attribute for which the subject is being tested actually exists in that subject) may occur. Use this product For Research Use Only.
- Do not use any reagent after the expiration date.
- Do not use together with reagents of other products.
- Follow the instructions.
- Take care in handling of specimen to minimize risk of infection.
- The PCR process is covered by patents issued and applicable in certain countries. iNtRON Biotechnology, Inc. does not encourage or support the unauthorized or unlicensed use of the PCR process. Use of this product is recommended for persons that either have a license to perform PCR or are not required to obtain a license.

8. TROUBLE SHOOTING

- 1 In the case of difficult to interpret results due to non-specific bands.
 - Reduce amount of template by 1/10 dilution and reacts again.
- 2 Preparation of PCR reaction at room temperature may cause the non-specific band.
- 3 All procedure should be carried out on ice.

9. ORDERING INFORMATION

Product	Catalog No.
Canine Herpesvirus Detection Kit	IP11073
Viral Gene-spin™ Viral DNA/RNA Extraction kit	17151
SiZer ™ 100 DNA Marker	24073

1. DESCRIPTION

Canine herpesvirus (CHV) can cause fading puppy syndrome, upper respiratory tract disease (kennel cough) and abortion/ stillbirths in dogs. The main route of transmission appears to be oronasal from infected puppies or from nasal or vaginal excretions of adults.

The virus spreads rapidly through kennels but usually only causes disease in very young puppies. Infection of newborn puppies commonly results in death. Puppies infected with CHV at the time of birth will generally start to show clinical signs of infection at four to six days of age. Infected puppies will exhibit persistent cry, a diminished suckling response, yellow green diarrhea and abdominal pain. Fever is usually not present. Death frequently occurs within 48 hours after clinical signs are noted. One or all pups in a litter infected at birth may show signs of herpesvirus infection. Infection of adults or puppies over 3 weeks old results in replication in the respiratory tract without clinical disease. The virus can undergo latent infection and reactivation, and further shedding can be induced by immunosuppression or stress.

Canine Herpesvirus Detection Kit is direct detection of canine herpesvirus on the basis of a genetic database, so it can diagnose very fast and accurately. It can amplify only specific gene using the PCR (Polymerase Chain Reaction) method, and take only 2~3 hours for detection. Therefore, it is a very fast accurate, reliable technique.

2. STORAGE

The components of *Canine Herpesvirus Detection Kit* should be stored at -20 $^{\circ}$ C, under this condition, the kit is stable until expiration date stated on the label.

3. CONTENTS

Canine Herpesvirus PCR Pre-mixture	es
DNase/RNase-free water (white cap) · · · · · · 1 via	al
CHV positive control (Yellow cap)	al

Component in 20 ^{µℓ} reaction	
i-StarTaq [™] DNA Polymerase dNTPs PCR Reaction buffer Chemical stabilizer Gel loading buffer 8-MOP (dissolved in DMSO) Primers for CHV	

4. SPECIMEN

Performs the test with whole blood, tracheal wash, lung, liver vesicular, throat, nasopharyngeal or conjunctival swab. The specimen should be stored at -20° C prior to use.

5. ADDITIONAL REQUIRED MATERIALS

- Disposable gloves
- DNA extraction kit (see 6.1 DNA preparation method)
- Pipettes
- Sterile pipette tip
- Vortex mixer
- Centrifuge for microcentrifuge tubes
- Thermal cycler
- Electrophoresis kit
- UV transilluminator

6. PROCEDURE

Please read through the entire procedure before starting.

6.1 DNA Preparation

Various manufacturers offer DNA isolation kits. Please carry out the DNA isolation according to the manufacturer's instructions. The following standard extraction kit is recommended.

Product	Catalog No. Manufacturer	
Viral gene-spin [™] Viral DNA/RNA Extraction Kit	17151	iNtRON Biotechnology, Inc.

6.2 Amplification

- Prepare appropriate PCR premix tubes and label. And one PCR premix tube for positive control.
- ② Add $2\mu\ell$ of template DNA into the PCR premix tube.
- 3 Add 18 μ l of DNase/RNase-free water into the PCR premix tube to total volume as 20 μ l.
- 4 Add 2μℓ of positive control and 18μℓ of RNase-free water into a PCR premix tube for monitoring of amplification and easy interpretation.
- (§) Dissolve the blue pellet by pipetting. Note: The pellet is easily dissolved, by letting the mixture stand at R.T. for 1-2minutes after adding water.
- ⑥ (Optional) Add mineral oil. This step is unnecessary when using a thermal cycler that employs a top heating method (general methods).
- 7 Perform PCR reaction of samples as the below process using PCR machine.

PCR cycle		Temp.	Time
1 Cycle	Initial Denaturation	94℃	5 min.
	Denaturation	94℃	30 sec.
40 Cycles	Annealing	50℃	30 sec.
	Extension	72℃	40 sec.
1 Cycle	Final extension	72℃	5 min.

6.3 Detection of Amplified Products

- Prepare 1.5% agarose gel containing RedSafe[™] Nucleic Acid Staining Solution. (Cat. No. 21141)
- ② Load $7\mu\ell$ of PCR product and positive control on agarose gel without adding a loading-dye buffer and perform electrophoresis.
- 3 Run electrophoresis by 100V (required about 30~40 minutes).
- 4 Identify the result on ultra-violet (UV) transilluminator.

6.4 Interpretation

Expected PCR product size : 369 bp

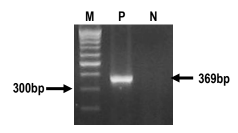


Fig 1. Electrophoresis of PCR product by Canine Herpesvirus Detection Kit

Lane M: 100bp Molecular ladder (iNtRON Biotechnology)

Lane P : CHV positive sample Lane N : Negative control

6.5 Elimination of carry-over contamination

- Each PCR/RT-PCR Pre-mixture contains 8-methoxypsoralen (8-MOP) for preventing of carry-over contamination.
- All PCR products should be discarded after UV irradiation (10 min/365nm) for preventing from carry-over contamination.