



# CARNIVORE PROTOPARVOVIRUS 1

## qPCR TEST HANDBOOK

For Research Use Only. Not intended for diagnostic use.

CAT NO.: YSL-qP-IC-CPPV1-100

100 reactions  
with Internal Control and Lyophilised MasterMix

VERSION 8.5



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## INTENDED USE

This qPCR test offers an efficient and user-friendly solution for the detection of Carnivore protoparvovirus 1 (CPPV1) in extracted nucleic acid samples from a variety of sources. It is intended for use by trained professionals in a suitable molecular biology laboratory.

## SPECIFICITY AND SENSITIVITY

### Specificity

The YouSeq qPCR test for detection of Carnivore protoparvovirus 1 is designed to have the broadest detection profile possible and detect all clinically relevant strains. The primers and probes typically have a  $\geq 95\%$  homology with all reference data used, from relevant, publicly available databases at the time of design.

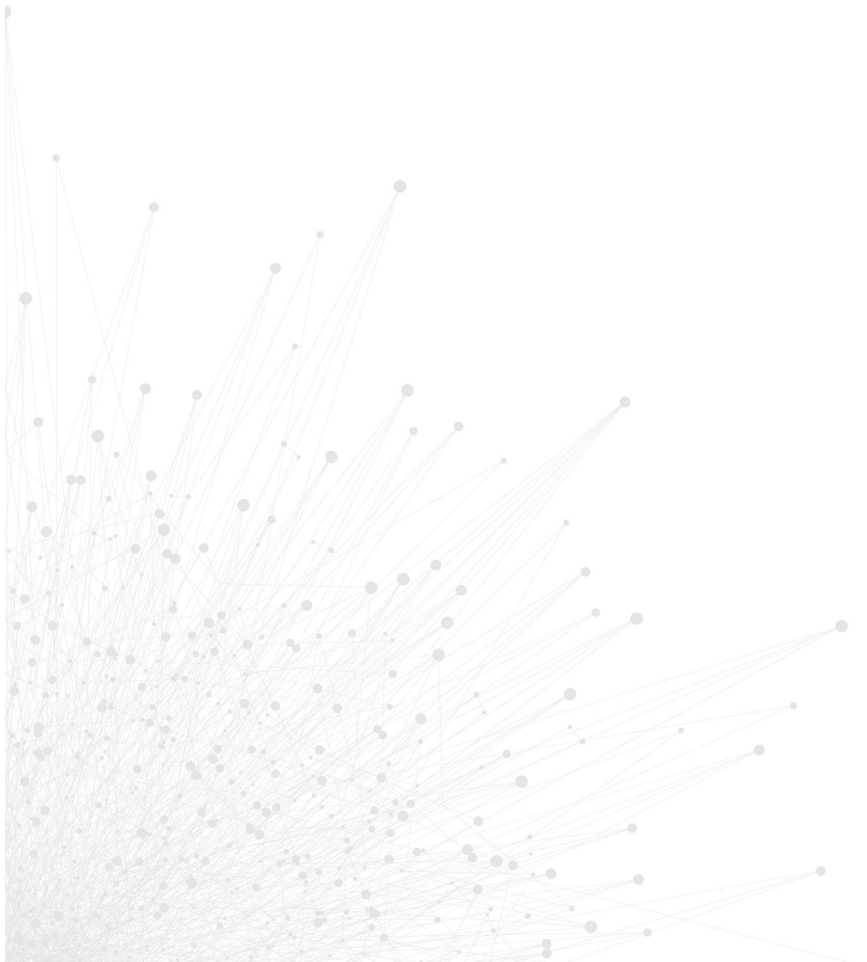
The target gene for Carnivore protoparvovirus 1 (NS1 gene) has been demonstrated to have a distinctive sequence making it an ideal target for highly specific detection.

The primers will detect the following strains: Canine parvovirus, Feline panleukopenia virus, Mink enteritis virus, Feline parvovirus, Raccoon dog parvovirus and Pangolin protoparvovirus.

For further information on the detection profile of the product, please do not hesitate to contact our team: [support@youseq.com](mailto:support@youseq.com)

### Sensitivity

The qPCR test is suitable for the detection of Carnivore protoparvovirus 1, across a wide dynamic range. Under ideal PCR conditions the assay can detect less than 100 copies of the target in the PCR reaction.





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| Component                                                   | Cap Colour                                                                          | Volume  |
|-------------------------------------------------------------|-------------------------------------------------------------------------------------|---------|
| Carnivore protoparvovirus 1 specific primer/probe mix (FAM) |  | 100 µL  |
| Internal Control primer/probe mix (VIC/HEX)                 |  | 100 µL  |
| Lyophilised Tetra™ 2X qPCR MasterMix                        |  | 1.1 mL* |
| MasterMix Resuspension Buffer (MMRB)                        |  | 1.5 mL  |
| Carnivore protoparvovirus 1 Positive Control (PTC)          |  | 500 µL* |
| Internal Control DNA Template                               |  | 500 µL* |
| Template Resuspension Buffer (TRB)                          |  | 1.5 mL  |
| DNase/RNase Free Water                                      |  | 1.5 mL  |
| ROX Passive Reference Dye                                   |  | 10 µL   |

\* Supplied dried – requires resuspension. See instructions in resuspension section.

## RECOMMENDED ADDITIONAL REAGENTS & MATERIALS

Nucleic Acid extraction kit – Internal Control DNA is to be included in the sample extraction. See ‘Use of Internal Control DNA’ section below.

General laboratory equipment (pipettes, pipette tips, (micro)centrifuge tubes, compatible strip tubes/plates, plate seals, etc.).

qPCR instrument with channels to detect FAM and VIC/HEX.

## BEST PRACTICE

### Decontamination:

Before beginning laboratory work, thoroughly decontaminate any work surfaces and pipettes being used, to eliminate potential contamination.

### General use and set-up:

All components should be fully defrosted with contents at the bottom of the tube before opening. To ensure contents are at the bottom, centrifuge or gently tap the tube. After use, reagents should be returned to the freezer.

Once any reagents are resuspended, mark the tick box on the tube for future reference. After this, or after combining reagents, the tube should be pulse vortexed to ensure it is mixed well.

It is advised to set up the tubes/plate and reaction mix on ice to minimise artefact formation, which may reduce sensitivity.

When preparing the qPCR reaction mix, it is recommended to incorporate an overage when calculating the total number of reactions to compensate for potential volume losses incurred during pipetting.

### Set-up environments:

It is best practice to set up qPCR tubes/plates in two different environments - a clean (no template) lab and PCR (template) lab.

No Template Control(s) (NTC) and Positive Control(s) (PTC) should be included in every run. To reduce contamination, NTCs and samples can be set up and sealed in a clean lab before moving to the PCR lab.

## BENCH SIDE PROTOCOL

### RESUSPENSION

Before first use, resuspend the designated component with the correct reagent and specified volume, as per the table below:

1. Add the resuspension reagent and pulse vortex the tube to ensure each is mixed well.

| Component                                          | Reagent | Volume | Location       |
|----------------------------------------------------|---------|--------|----------------|
| Internal Control DNA Template                      | TRB     | 500 µL | Extraction lab |
| Lyophilised Tetra™ 2X qPCR MasterMix               | MMRB    | 1.1 mL | Clean lab      |
| Carnivore protoparvovirus 1 Positive Control (PTC) | TRB     | 500 µL | PCR lab        |

### ROX (INSTRUMENT DEPENDENT)

ROX is required for instruments that use ROX as a passive reference. The table below outlines the qPCR instruments that require the addition of ROX.

1. If ROX is required:
  - a. Dilute the ROX supplied according to the table below based on the intended qPCR instrument to be used.
  - b. Directly transfer 5µL of prepared ROX to the resuspended Tetra™ MasterMix.

| Level of ROX      | Instruments                                                            | Step 1: Volume of water to add to ROX tube | Step 2: Add to MasterMix vial |
|-------------------|------------------------------------------------------------------------|--------------------------------------------|-------------------------------|
| High ROX:         | Applied Biosystems 7000, 7300, 7700, 7900, StepOne, StepOne Plus       | No Dilution Required                       | 5 µL                          |
| Low ROX:          | Applied Biosystems 7500 & 7500 FAST, ViiA7, Quantstudio, Stratagene MX | 130 µL                                     | 5 µL                          |
| ROX Not Required: | All Other Instruments                                                  | Not Required                               | Not Required                  |

### USE OF INTERNAL CONTROL DNA

1. At the relevant step of the extraction protocol, pause and add 5 µL of the resuspended Internal Control DNA into the extraction/lysis buffer for each sample that is to be extracted.

**Please note:** Do not add this Internal Control DNA directly into the biological sample as this may cause degradation of the control DNA.

2. Continue nucleic acid extraction as per the manufacturer's instructions.

## qPCR REACTION SET-UP

1. Retrieve the required components and appropriate plasticware for qPCR reaction set-up.
2. In an appropriately sized (micro)centrifuge tube, combine the following reagents to create a reaction mix that will cover all required tube/wells (e.g. **samples**, **NTC** and **PTC**).

**Please note:** When calculating required reactions, include an overage to allow for volume loss during pipetting

| Component                                             | Volume (per reaction) |
|-------------------------------------------------------|-----------------------|
| Tetra™ 2X qPCR MasterMix                              | 10 µL                 |
| Carnivore protoparvovirus 1 specific primer/probe mix | 1 µL                  |
| Internal Control primer/probe mix                     | 1 µL                  |
| Reaction mix volume                                   | 12 µL                 |

3. Mix the combined reagents by briefly vortexing or inverting.
4. Dispense 12 µL of the reaction mix into all required tubes/wells.
5. For the **NTC(s)**, add 8 µL of DNase/RNase Free Water into required tube/well(s).
6. For each extracted **sample**, add 8 µL into required tube/well(s).

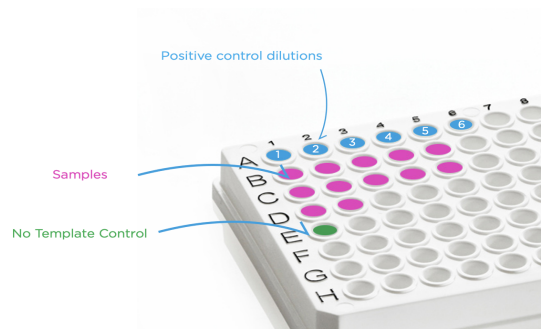
**Please note:** It is best practice to seal the **NTC(s)** and **sample** tubes/wells before proceeding to the positive control steps.

7. For the **positive controls(s)**, follow either step a. or b. below:
  - a. For **qualitative** results, add 5 µL of the resuspended **PTC** (tube 1) and 3 µL of DNase/RNase Free Water, into each designated positive control tube/well. Proceed to step 8.
  - b. For **quantitative** results, perform a serial dilution of the resuspended **PTC** (tube 1) to create a six-point standard curve:
    - I. Add 45 µL of Template Resuspension Buffer into 5 fresh microcentrifuge tubes and label them 2, 3, 4, 5 and 6.
    - II. Pipette 5 µL of **PTC** into tube 2.
    - III. Mix by pipetting up and down 10 times.
    - IV. Change pipette tip and pipette 5 µL from tube 2 into tube 3.
    - V. Mix by pipetting up and down 10 times.
    - VI. Repeat steps 'IV' and 'V' to complete the dilution process for tubes 4, 5 and 6.
    - VII. Add 3 µL of DNase/RNase Free Water into each designated positive control tube/well.
    - VIII. Add 5 µL of the required **PTC** dilution (tubes 1 – 6), into each designated positive control tube/well.

**Please note:** The described standard curve provides a dynamic range as outlined in the table below.

| Tube No.                   | Copies of Target/reaction |
|----------------------------|---------------------------|
| 1                          | 1,000,000                 |
| 2                          | 100,000                   |
| 3                          | 10,000                    |
| 4                          | 1,000                     |
| 5                          | 100                       |
| 6                          | 10                        |
| 8. Seal the tube(s)/plate. |                           |

9. Briefly vortex the sealed tube(s)/plate, then spin in a centrifuge to ensure all reagents are fully resuspended and at the bottom of the tubes/wells before proceeding.



## qPCR AMPLIFICATION PROTOCOL

1. Load the tubes/plate onto the qPCR instrument and set up the qPCR protocol following the table below.
2. Set the total reaction volume to 20 µL.

| Temperature                                                                            | Time       | Number of Cycles |
|----------------------------------------------------------------------------------------|------------|------------------|
| 95°C                                                                                   | 3 minutes  | -                |
| 95°C                                                                                   | 15 seconds | x 45             |
| 60°C  | 60 seconds |                  |

 Collect fluorogenic data through FAM and VIC/HEX channels during this step.

3. Start the run.

## INTERPRETATION OF RESULTS - OVERVIEW

If using single threshold analysis - YouSeq recommends setting the threshold at 10% of the End Point Fluorescence (EPF) for each channel.

- For the Target channel, use the [PTC](#) (tube 1) EPF to set the threshold.
- For the Internal Control, use the average EPF from [samples](#) to set the threshold.

### Results interpretation:

| Reaction Type             |                             | qPCR Signal |          |
|---------------------------|-----------------------------|-------------|----------|
| Positive control (tube 1) |                             | 18.5 ± 2    | 18.5 ± 2 |
| No template control       |                             | -           | -        |
| Sample                    | Internal Control: (VIC/HEX) | ≤ 31        | ≤ 31     |
|                           | Target: (FAM)               | +           | -        |

Result

Positive result

Negative result



## INTERPRETATION OF RESULTS – CONTROLS EXPLAINED

### Positive control

The **PTC** (tube 1) should amplify in a Cq range of  $18.5 \pm 2$  for the target. If this Cq range is not achieved, the sample test result is invalid and should be repeated.

To achieve the most accurate quantitative result from the positive control standard curve, an efficiency between 90% to 110% is desirable. If it falls outside of this range, preparing a fresh standard curve and repeating the run may improve the efficiency value.

**Please note:** The positive control is a sequence representative of the target region and does not contain the organism's entire genome. The positive control does not include the Internal Control sequence and should not be expected to amplify in the VIC/HEX channel.

### No template control

The **NTC** tube/well(s) should be negative, with no amplification.

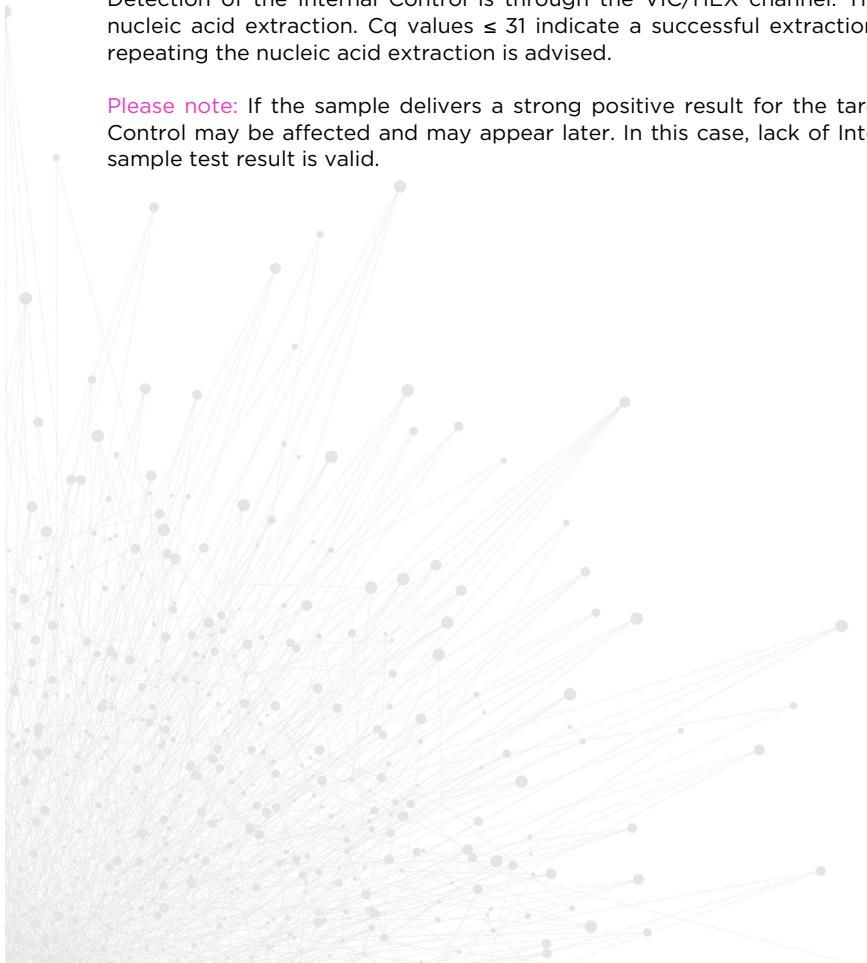
**Please note:** Background laboratory contamination can result in a very late signal in **NTC** tube/well(s). If the **NTC** has amplification, comparison to the sample test result is necessary:

- If the **NTC** is  $\geq 5$  Cq later than the sample signal, it can be considered negative, and the sample test result is valid.
- If the **NTC** is  $< 5$  Cq later, the sample test result is invalid – the test should be repeated after potential sources of contamination have been removed.

### Internal control

Detection of the Internal Control is through the VIC/HEX channel. This gives information about the efficiency of the nucleic acid extraction. Cq values  $\leq 31$  indicate a successful extraction has taken place. If the signal is later than this, repeating the nucleic acid extraction is advised.

**Please note:** If the sample delivers a strong positive result for the target of interest, then amplification of the Internal Control may be affected and may appear later. In this case, lack of Internal Control amplification is acceptable, and the sample test result is valid.





## PRODUCT SPECIFICATIONS

### Storage

Store at -20°C from arrival. The product's shelf life is outlined as an expiry date on the pouch label.

### Suitable input material

This qPCR test will work well with any source of good quality nucleic acid. Good quality is defined as nucleic acid with high integrity (not degraded). Poor quality input nucleic acid is a leading cause of test failure.

### Regulatory status

This product has been developed for Research Use Only (RUO) and is not intended for diagnostic use. It should not be used for diagnosis of disease or infection unless specifically approved by the regulatory authorities in the country of use.

### Quality Control

In accordance with the YouSeq Ltd ISO EN 13485-certified Quality Management System, each lot of Carnivore protoparvovirus 1 qPCR Test Kit is tested against predetermined specifications to ensure consistent product quality. The primers/probe(s) typically demonstrate  $\geq 95\%$  in silico specificity to their intended target and periodically checked against newly available sequence information to maintain their detection profile.

### Technical Assistance

For customer support, please contact:

e-mail: [support@youseq.com](mailto:support@youseq.com)

phone: +44 (0)333 577 6697

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