

### Intended Use

The CPV + CCV Antigen Combo Rapid Test Device is a combined device to differentially diagnose the presence of canine parvovirus (CPV) antigens and canine coronavirus (CCV) antigens in canine feces or vomit.

## **Principle**

The CPV + CCV Antigen Combo Rapid Test Device is based on sandwich lateral flow immunochromatographic assay. The test device has two testing windows. Each testing window has an invisible T (test) zone and C (control) zone. When sample is applied into the sample wells on the device, the liquid will laterally flow on the surface of the test strip. If there is enough CPV antigen or CCV antigen in the sample, a visible T line will appear in the corresponding testing window. The C line should always appear after a sample is applied, indicating a valid result. By this means, the device can accurately indicate the presence of CPV antigen or CCV antigen in the sample.

## Storage and Stability

Store as packaged in the sealed pouch either at room temperature or refrigerated (2-30 °C). The test is stable through the expiration date printed on the sealed pouch. The test device must remain in the sealed pouch until use. DO NOT FREEZE. Do not use beyond the expiration date.

### **Precautions**

- Do not use after expiration date.
- Handle all specimens as if they contain infectious agents.
  Observe established
- precautions against microbiological hazards throughout testing and follow the standard
- procedures for proper disposal of specimens.
- Wear disposable gloves and eye protection when specimens are being tested.
- Humidity and temperature can adversely affect results.
- Do not remove test device from its pouch until immediately before use.
- Do not reuse the test kit.
- Do not mix components from different lot and different products.

## **Additional Special Equipment**

#### **Materials Provided**

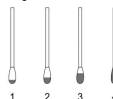
• Test devices • Droppers • Swabs

- Plastic tubes with buffer
  Package insert
  Materials Required But Not Provided
- Timer

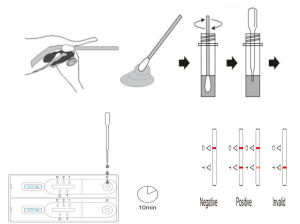
### **Direction For Use**

Allow the test device, specimen, buffer, and/or controls to equilibrate to room temperature (15-30 °C) prior to testing.

1. Collect dog's feces or vomit with the swab stick from dog's anus or on the ground. The amount of fecal swab as follows:



- 1. Not enough
- 2. Good
- 3. Good
- 4. Too Much
- 2. Insert the wet swab into the provided plastic tubes with buffer. Agitate it to assure good sample extraction.
- 3. Place the test device on a clean and level surface. Hold the dropper vertically and **transfer 3 drops of extracted sample** (approximately 120  $\mu$ I) to the specimen well (S) of the test device, then start the timer. See illustration above.
- 4. Interpret the result in **5-10 minutes**. Result **after 10 minutes** is considered as invalid.



# Interpretation of Results

**Positive:** Two lines appear. One line should always appear in the control line region(C), and another one apparent colored line should appear in the test line region.

**Negative:** One colored line appears in the control region(C).No apparent colored line appear in the test line region.

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**Invalid:** Control line fails to appear. Insufficient specimen volume or incorrect procedural techniques are the most likely reasons for control line failure. Review the procedure and repeat the test with a new test device. If the problem persists, discontinue using the test kit immediately and contact your local distributor.

### **Performance Characteristics**

Relative Sensitivity: 93.10% (95%Cl\*: 83.27% - 98.09%) Relative Specificity: 97.52% (95%Cl\*: 92.93% - 99.49%)

Accuracy:96.09% (95%CI\*: 92.11% - 98.41%)

\*CI = confidence interval compared to RT-PCR method.

### Limitations

CPV+CCV Antigen Combo Rapid Test Device is for in vitro diagnostic use only for canines. All results should be considered with other clinical information available from veterinarian. For confirming the results, confirmatory methods such as PCR are recommended.



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