PetNADTM

FCV Detection Kit

For Feline calicivirus

User Manual

For Research Use Only

Manufacturer:

GeneReach Biotechnology Corporation

TEL: 886-4-24639869 FAX: 886-4-24638255

No. 19, Keyuan 2nd Rd., Central Taiwan Science Park, Taichung City, Taiwan 407

Web Site: www.petnad.com

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INTENDED USE

PetNADTM FCV Detection Kit is intended for *in vitro* detection of FCV RNA based on insulated isothermal polymerase chain reaction (iiPCR) technology. This kit is designed specially to be used with an iiPCR-compatible instrument, **POCKIT**TM Nucleic Acid Analyzer. The assay is intended for use by veterinarians or technicians with basic laboratory skills.

This kit is intended for research use only.

SUMMARY AND EXPLANATION

Feline calicivirus (FCV) is highly contagious and widespread in the general cat population. The virus is shed predominantly in oral and nasal secretions in acute phase, when cats show symptoms including fever, conjunctivitis, nasal discharge, sneezing, and oral ulceration. During recovery, many cats continue shedding, until at least 30 days post-infection, only a few for several years (Wardley, 1976). Disease prevalence is proportional to the number of cats in the household, with the highest prevalence usually seen where large groups are

accommodated together (Wardley et al., 1974).

PCR is one of the most commonly accepted methods that provide high sensitivity and specificity for FCV detection. However, conventional PCR assays take three to four hours, and require sophisticated thermocyclers and well-trained technicians to perform. GeneReach has developed **PetNAD**TM FCV Detection Kit based on iiPCR technology, which significantly reduces reaction time and offers sensitivity and specificity comparable to those of conventional nested PCR (Tsai, 2012; Chang, 2012). Furthermore, this simple and easy assay could be completed rapidly in a portable **POCKIT**TM Nucleic Acid Analyzer.

PRINCIPLES OF THE PROCEDURE

In iiPCR, hydrolysis probe-based chemistry is used to generate fluorescent signal during amplification of target RNA. The primers and probe target ORF1 gene and do not cross-react with nucleic acid from host and other feline upper respiratory pathogens.

PRODUCT DESCRIPTION

A. Materials Provided (24 tests/kit)

Component	Co	ntents or Purpose	Amount
Premix Pack	■ FCV Premix (lyophilized		24 bags (1 FCV Premix
	pell	let) containing dNTPs,	vial and desiccating
	prir	ners, probe, and enzyme	agent/bag)
	for	amplification.	
	■ Des	siccating agent pack.	
Premix Buffer B	■ Reaction buffer to re-dissolve		2 vials (1.3 ml/vial)
	the	lyophilized pellet.	
P(+) Standard	■ Dried plasmid containing		1 vial
	FC	V partial sequence.	
Standard Buffer	andard Buffer Reaction		1 vial (110 μl/vial)
	P(+) Standard.	
R-tube			1 bag (24 pieces/bag)
Cap			1 bag (24 pieces/bag)
User Manual			1 copy

B. Materials and Equipments Required, but Not Provided

- 1) **PetNAD**TM Nucleic Acid Co-prep Kit
- 2) **POCKIT**TM Nucleic Acid Analyzer: **PetNAD**TM-compatible instrument.
- 3) **cubee**TM Mini-Centrifuge (cubee)
- 4) Micropipette and tips

C. Storage and Stability

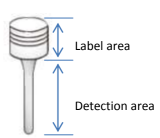
- 1) The kit should be stored at 4°C and is stable until the expiration date which is stated on the label.
- 2) Store Premix vials in sealed Premix Pack to avoid hydration of lyophilized components.
- 3) Reconstituted P (+) Standard is stable for 6 months at 4°C. Aliquot reconstituted P (+) Standard to avoid degradation of nucleic acid.

D. Sample Type

Nucleic acid extracted from swab sample (i.e. oropharyngeal, conjunctival and nasal swab)

PRECAUTIONS

- A. Do not open R-tube(s) after reaction to prevent any carryover contamination.
- B. Perform extraction and amplification in two independent spaces to minimize contamination.
- C. Do not reuse R-tube and Premix.
- D. Include the P(+) Standard to:
 - 1) Ensure **POCKIT**TM Nucleic Acid Analyzer is working normally.
 - 2) Ensure detection kit performance after storage.
- E. To get optimal fluorescence detection.
 - Wear powder-free gloves to handle R-tubes.
 - 2) Do not label in the detection area of R-tube.

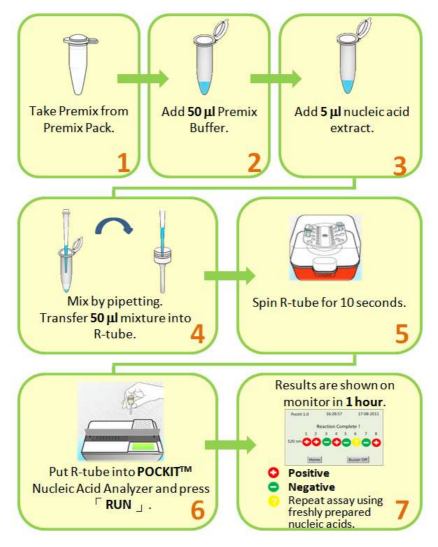


LIMITATIONS

- A. The test should be used only for testing nucleic acid extracted from animal specimen. Do not add specimen (i.e. whole blood) directly into Premix.
- B. **PetNAD**TM Nucleic Acid Co-prep Kit is recommended for nucleic acid extraction.
- C. Any deviation from recommended procedure may not achieve the optimal results and should be validated by the users.
- D. It is strongly recommended to use freshly prepared nucleic acid (within 1 hour after extraction) to achieve optimal results with **PetNAD**TM FCV Detection Kit.
- E. Vaccination with a modified-live FCV vaccine may result in positive PCR results for a few weeks after vaccination. Killed or vectored-recombinant vaccines will not interfere with PCR testing. PetNADTM is recommended in sick animals with clinical signs and/or laboratory abnormalities consistent with infection or in an animal with a suspected subclinical infection as based upon history, physical examination and clinical laboratory findings.

PROCEDURE

A. PetNADTM FCV Detection Kit Quick Guide



B. Procedure

Note: Before using for the first time, add 100 μ l Standard Buffer to P(+) Standard. Store reconstituted P(+) Standard at 4°C.

- 1) Label R-tube(s) in the label area.
- 2) Prepare one Premix for each sample. (Premix tube is in Premix Pack. Each Premix Pack contains one Premix.)

Note: If the pellet is not found at the bottom of the tube, spin tube briefly to bring it down.

- 3) Add 50 µl Premix Buffer B to each Premix tube.
- 4) Add 5 μl nucleic acid extract or P(+) Standard to each Premix tube. Mix by pipetting up and down.
- 5) Transfer 50 µl Premix/sample mixture into R-tube.
- 6) Seal top of each R-tube with a cap. Make sure R-tube is capped tightly.
- 7) Place R-tube into the holder of **POCKIT**TM.
- 8) Spin tube briefly in **cubee**TM to make sure all solution is collected at the bottom of R-tube.

Note: Start reaction within 1 hour to prevent nucleic acid degradation.

Note: Make sure there are no bubbles in the tube.

9) **POCKIT**TM reaction:

Note: Please see the user manual of POCKITTM for details.

a) Turn on **POCKIT**TM, which should complete

- self-testing within 5 minutes.
- b) Select "520 nm".
- c) When "System READY" is displayed, place the holder with R-tube(s) into the reaction chamber.
- d) Tap cap of each R-tube to make sure the tube is positioned properly.
- 10) Close lid and press "Run" to start reaction program.
- 11) Test results are shown on the monitor after reaction is completed.

DATA INTERPRETATION

* One example of results shown on the monitor.



520nm	Interpretation
0	FCV Positive
0	FCV Negative
3	Repeat reaction with freshly prepared nucleic acid.

ANYLYTICAL SENSITIVITY

The detection limit of **PetNAD**TM FCV Detection Kit is about 10 copies/ reaction.

TROUBLESHOOTING

Problems	Possible causes	Solutions
False Positive	1) Reuse of micro-	■ Micro-centrifuge tubes, tips,
	centrifuge tubes,	R-tubes and Premix are for
	tips, R-tubes and	single-use only. Reusing these
	Premix.	accessories would cause cross-
		contamination.
		■ Used micro-centrifuge tubes, tips,
		R-tubes and Premix should be
		collected and discarded according
		to local regulation. Do not place the
		waste close to the working area to
		prevent cross-contamination.
	2) Contaminated	■ Disassemble and clean up
	micropipette	micropipette.
		■ Use aerosol-free tips.
	3) Contaminated	■ Consult with a GeneReach
	reagent	technical support representative or
		local distributor.
	4) Contaminated	■ Consult with a GeneReach
	working area	technical support representative on
		how to clean up working.

Problems	Possible causes	Solutions
False	1) Nucleic acid	■ Consult manual of nucleic acid
Negative	extraction failed.	extraction kit.
	Bad nucleic acid quality or nucleic acid concentration too high	 Check sample storage condition. Please refer to Troubleshooting section of PetNADTM Nucleic Acid Co-prep Kit. If a spectrophotometer is available, check OD 260/280 ratio. This ratio should be between 1.4 and 2.0.
	3) PCR inhibition	■ Do not overload nucleic acid. ■ Spike nucleic acid sample into P(+) Standard reaction for a parallel PCR reaction. Negative results indicate the presence of inhibitors in the nucleic acid. In that case, prepare another nucleic acid extract.
Heavy contamination of amplicons in reaction chamber of	Leakage or spill of reaction from R-tube into reaction chamber of POCKIT TM .	■ Consult with a GeneReach technical support representative or local distributor.
POCKIT TM .		

REFERENCE

- Chang, H.F. G., Tsai, Y.L., Tsai, C.F., Lin, C.K., Lee, P.Y., Teng, P.-H., Su, C. and Jeng, C.C., (2012) A thermally baffled device for highly stabilized convective PCR. *Biotechnology Journal* 7(5): 662-666, doi: 10.1002/biot.201100453
- 2. Tsai Y.L., Wang H.T. T., Chang H.F. G., Tsai C.F., Lin C.K., Teng P.H., Su C. and Jeng C.C., (2012). Development of TaqMan probe-based insulated isothermal PCR (iiPCR) for sensitive and specific on-site pathogen detection. *PLoS ONE* 7(9): e45278. doi: 10.1371/journal.pone. 0045278
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