

DNA
DIAGNOSTIC

User Manual

Enterit 4 calves V

RNA extraction – cDNA synthesis - qPCR - Interpretation



USER MANUAL
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1. PURPOSE OF THE TEST

Calf diarrhea also known as scours is a major problem in dairy and beef productions because it can result in serious economic implications. The calves can die in severr cases, however calves that survive will have poorer performance compared to healthy calves. Several pathogens are associated with calf diarrhea including bacteria, viruses and parasites.

Enterit 4 calves V is a fast, reliable and easy-to-use method for the detection of the pathogenic virus and parasites (Bovine rotavirus, Bovine coronavirus, *Cryptosporidium parvum* and Eimeria species) linked to calf diarrhea. Using a simple 96 deepwell-based extraction protocol followed by a sensitive and highly specific RT-qPCR reaction, the entire protocol from RNA extraction to result can be carried out in 5 hours.

2. PRINCIPLE OF THE TEST

Enterit 4 calves V allows for the detection of Bovine rotavirus, Bovine coronavirus, *Cryptosporidium parvum* and Eimeria species in feces samples from calves. It includes the following four main steps:

The RNA extraction step uses the extraction kit for DNA/RNA extraction from 0.1 g of feces sample. The sample is dissolved in resuspension buffer and transferred to a plate containing bead solution and then mixed, spun and washed. The DNA is extracted by the lysis mix buffers at 37 °C for 20 minutes following by 95°C for 15 minutes and debris are pelleted by centrifugation. Now the DNA is in the supernatant.

The cDNA synthesis step uses 5 µL of RNA template which is added to a well in a 96 well plate containing ready to use RT cDNA master mix. The cDNA synthesis reaction runs for 1 hour and 15 minutes.

The PCR step uses 5 µL of cDNA template which is added per well in a 96 well qPCR plate containing E4CV qPCR master mix. The qPCR reaction run for approximately 52 minutes. Each qPCR reaction contains four sets of primers and fluorescence probes for specific detection of two viruses and two parasites. A fifth set of primers and probe detects an internal amplification control (IAC). Each of the four probes emit a specific light (color) enabling the identification of viruses and parasites present in the sample. Probes with CY5, ROX, HEX and FAM fluorophores are present for detection of Bovine rotavirus, Bovine coronavirus, *Cryptosporidium parvum* and Eimeria species amplicons, respectively. The tests use the standard 96 well plate format and ready-to-use reaction mixes. The test requires only pipettes, plate mixer, a vacuum aspirator, a centrifuge, incubation oven, a thermal heating block and a qPCR instrument.

The interpretation step involves inspection of the amplification plots. The functionality of the qPCR reaction is controlled by the presence of IAC amplicons resulting in an ATTO signal with a Ct value at 27-32. The CY5, ROX, HEX and FAM signals ≤ 37 show the sample is positive for Bovine rotavirus, Bovine coronavirus, *Cryptosporidium parvum* and Eimeria species, respectively. No Ct value or Ct > 37 indicate that the sample is negative.

3. KIT COMPONENTS AND STORAGE

The Enterit 4 calves V contains material for testing 384 samples.

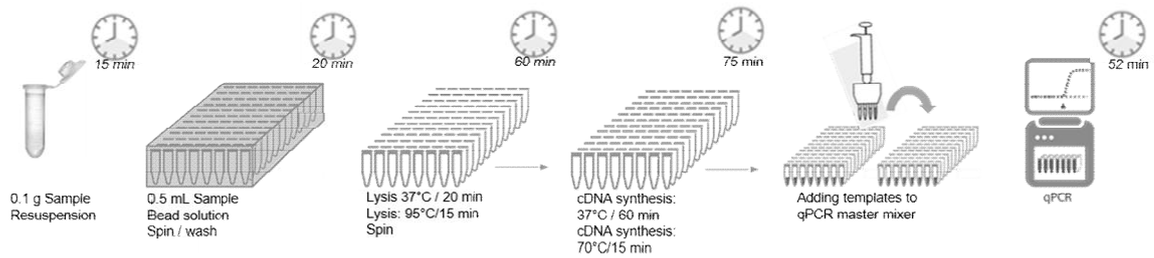
The Enterit 4 calves V contains one box for storage at room temperature (RT) and **two box** for storage at -20°C:

BOX	COMPONENTS	AMOUNT	STORAGE
I	Buffer F	5 bottle of 110 mL	RT
	96 deepwell plates (2mL), empty	4 x 96 Deep Well Plate	RT
	Sealed 96 Deep Well Plates with bead solution	4 x 96 Deep Well Plate	RT
	Wash Buffer	4 bottle of 110 mL	RT
	Buffer P	2 bottles of 30 mL	RT
	96 Well Plates (0.2 mL clear tubes) with cap	4 x 96 well plate (0.2 mL clear tubes) with cap	RT
	Adhesive seals for Deep Well plates	16 pieces	RT
	Cap mats for the PCR 96 Well Plates	8x 96 cap for PCR tube	RT
II	Lysis Buffer-I	2 bottle of 12 mL	-20°C
	Mix I Additive for Lysis Buffer-I	2 tube of 1334 µL	-20°C
	Sealed 96 Well Plates (0.2 mL white tubes) with E4CV qPCR Master Mix	4 x 96 well plates	-20°C
III	Sealed 96 Well Plates (0.2 mL clear tubes) with cDNA Master Mix I	4 x 96 well plates	-20°C

4. REQUIRED EQUIPMENT AND MATERIALS NOT INCLUDED IN THE KITS

- Centrifuge for 96 well plates. Centrifuge **must be capable of running at 5000xg**.
Example: Heraeus Multifuge X3R Centrifuge with HIGHPlate 6000 rotor.
- Vacuum system or plate washer capable of aspirating supernatant from Deep Well plates.
Example: Vacuum system connected to an 8-channel adaptor (contact DNA Diagnostic for information) or vacuum system with pipette tips (35mm from tip to rim stop).
- qPCR instrument, Stratagene Mx3005 with filters for ATTO (440nm-492nm), FAM (492nm-516nm), HEX (535nm-555nm), ROX (585nm-610nm), CY5 (635nm-665nm).
- Vortexer for 96 well plates.
- Ice bucket or 0-4°C cooling block.
- Standard lab pipettes and sterile filter tips.
- Mixer for a Deep Well plate.

5. PROTOCOL



RNA extraction:

1. Add 1.2 mL Buffer F to a well in the empty 2 mL tubes.
2. Homogenize the feces sample and transfer 0.1g \pm 0.03g feces to a corresponding 2 mL tube with buffer F.
Note: If feces sample is liquid or watery, transfer 100 μ L of sample.
3. Close the tube and mix vigorously on a mixer/vortexer until samples are homogenized, otherwise prolong mixing time/intensity.
4. Spin tube at 1000xg for 1 minute. Spin can be performed in the tube directly, or by transferring of all sample to a 2 mL Deep Well plate.
5. Carefully open the lid of the 2 mL tube or the seal of 2 mL Deep Well plate containing sample. Transfer 500 μ L of supernatant to a 96 Deep Well plate containing bead solution. Mix by pipette up and down. Seal the plate with adhesive seal.
6. Spin Deep Well plate containing bead solution and samples at 5000xg for 5 minutes.
7. Remove supernatant with a vacuum system or plate washer. It is important to remove the supernatant completely. Be careful not to touch or remove the pellets.
Note: It is important to remove all supernatant from the pellet. Therefore, correct adjustment of the plate washer syringes is required.
8. Add 1 mL Wash Buffer to Deep Well plate.
9. Cover wells with adhesive seal and vortex briefly.
10. Centrifuge the plate at 5000xg for 5 minutes.
11. Remove supernatant with a vacuum system or plate washer. It is important to remove the supernatant completely. Be careful not to touch or remove the pellets.
Note: It is important to remove all supernatant from the pellet. Therefore, correct adjustment of the plate washer syringes is required.
12. Prepare fresh Lysis-I Mix by adding 6 μ L Mix I additive to 54 μ L Lysis Buffer-I and 60 μ L Buffer P.

Lysis-I Mixer	1 reaction	9 reaction	50 reaction	100 reactions
Lysis Buffer-I	54 µL	486 µL	2700 µL	5400 µL
Mix I additive	6 µL	54 µL	300 µL	600 µL
Buffer P	60 µL	540 µL	3000 µL	6000 µL
Total	120 µL	1080 µL	6000 µL	12000 µL

13. Add 120 µL Lysis-I Mix, to each pellet and cover with new adhesive seal. Mix briefly on a Deep Well mixer.
14. Spin down shortly to bring the liquid to the bottom (e.g. 10 seconds at 1500 x g). Transfer of 60 µL (up to 120 µL) from each sample into the corresponding tubes in the 96 well plate (clear 0.2 mL) and close the tubes with the caps. Avoid cross contamination.
15. Incubate the 96 well plate at 37°C for 20 minutes and at 95°C for 15 minutes.
16. Cool the 96 well plate on ice for 5 minutes
Note: The incubation at 37°C and 95°C can be done using a PCR instrument programmed: 37°C for 20 min. → 95°C for 15 min. → 4°C for 5 minutes.
17. Centrifuge the plate at 5000xg for 5 minutes.

If lysate sample subsequently will be used for bacteria detection, follow the step 18 on Enterit 4 calves B manual protocol.

18. Carefully transfer 5 µL of each aqueous phase directly to the corresponding well of the 96 well plate with cDNA Master Mix I in step 21.
Note: Due to the instability of RNA, it is recommended to perform the cDNA synthesis before storing samples.

cDNA Synthesis:

19. Take a cDNA master mix I plate from freezer and thaw it on ice. Briefly spin the plate to collect the cDNA master mix in the bottom.
20. Remove and discard the seal from cDNA master mix I plate.
21. Carefully transfer 5 µL supernatant from the 96 well 0.2 mL plate (step 18) to the corresponding well in the cDNA master mix I plate (clear 0.2 mL).
22. Incubate the cDNA reaction at 37°C for 60 minutes.
23. Inactivate Reverse Transcriptase enzyme by heating to 70°C for 15 minutes.
24. Cool the cDNA reaction tube on ice for 1 minute and spin it again for 1 minute to collect condensate.
Note: The cDNA samples can be kept at -20°C until use.

qPCR analysis:

25. Take a E4CV qPCR master mix plate (white 0.2 mL) from freezer and thaw it on ice. Briefly spin the qPCR plate to collect the qPCR master mix in the bottoms.

26. Remove and discard the seal from qPCR plate.
27. Carefully transfer 5 μ L supernatant from the 96 well 0.2 mL plate (step 24) to the corresponding well in the E4CV qPCR master mix plate.
Note: It is important to keep Master Mix on ice while loading template and to run the qPCR within 15 minutes.
28. Carefully cover the qPCR reaction plate with new lids and spin at 1000-1500xg for 30 seconds to 1 minute.
29. Insert qPCR plate into the qPCR instrument.
30. Run the qPCR reaction using the instrument setting and the running program below:

Instrument settings for the MX3005P qPCR machine

Filter Gain Settings:

ATTO	CY5	ROX	HEX	FAM
1x	1x	1x	1x	2x

Note: If the raw data signal (R) is lower than 5000 for a color at the cycles 7-11 then increase the filter gain setting during the next runs resulting in a raw data signal (R) of 5000.

The guidelines for setting baseline and threshold should be checked manually to ensure that threshold lines are in the exponential area of the amplification plot and above the background.

Threshold Fluorescence: Select “Background Based Threshold” to cycles 7-11 and set the “Sigma multiplier” to 10.

To avoid bias of threshold wells including amplification curve with Ct<15 is flagged out. The threshold is then looked based on the rest of the sample and the sample with Ct<15 are put into the analysis again.

Baseline Correction: Select “Adaptive baseline”.

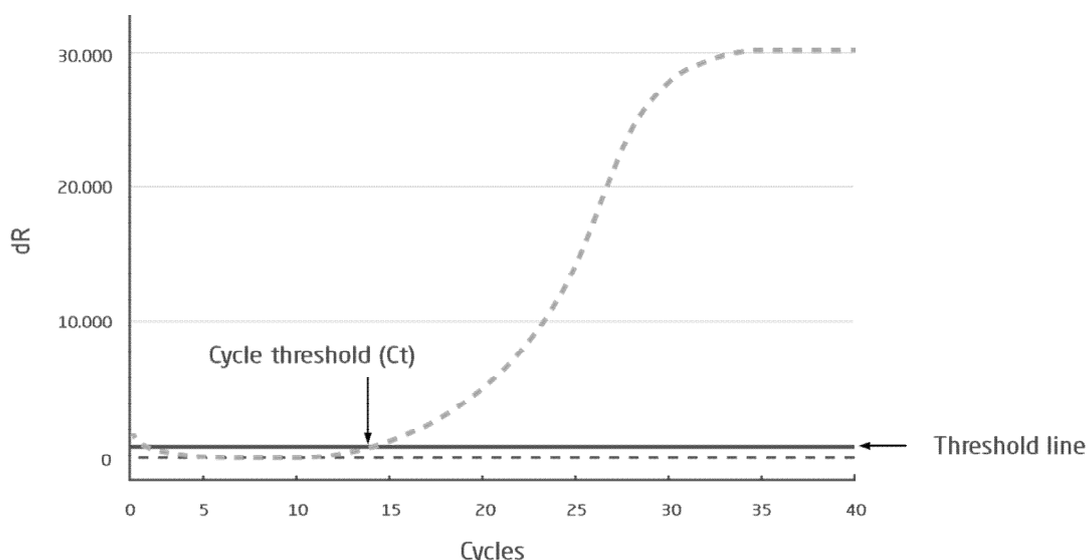
Activate the following filters: ATTO, CY5, ROX, HEX, FAM.

qPCR running program

Condition	Temperature	Duration	Number of cycles	Stage
Heat activation	95°C	1 min	1	Pre-incubation
Denaturation	95°C	5 sec	40	Amplification
Annealing/extension	60°C	25 sec*		

* Read fluorescence for ATTO, FAM, HEX, ROX, CY5 at the end of each annealing/extension step.

Interpretation of analysis:



Amplification plot with the number of PCR cycles is shown on the x-axis, and the background subtracted fluorescence from the amplification reaction is shown on the y-axis. The threshold line is used to find the Ct value, where it intersects the amplification curve. Ct is optimally determined in the early exponential phase, when the fluorescence starts increasing due to PCR amplification. The threshold line is calculated automatically by the instrument giving in this example a Ct value of 14.

31. Check amplification plot for ATTO signals and the signals should be present in all reactions. The Ct must be in the interval 27-32 to confirm the qPCR reaction is functional.
*Note: ATTO Ct can be above 32 or missing if the Ct for one or more of FAM, HEX, ROX, CY5 is low. Then the DNA sample should be diluted 5-10 times and a new qPCR reaction should be analyzed using this diluted DNA. A dilution of 10 times will theoretically move Ct values 3.3 higher. Reactions with no FAM, HEX, ROX, CY5 signal and ATTO signal absent or outside the interval 27-32 **must be retested**.*
32. Check amplification plot for CY5, ROX, HEX, FAM signals and interpret the results as described in the table below.

Target	Filter	Positive	Negative	Retest
		Ct	Ct	Ct
Bovine rotavirus	CY5	≤37	>37 or absent	
Bovine coronavirus	ROX	≤37	>37 or absent	
<i>Cryptosporidium parvum</i>	HEX	≤37	>37 or absent	
Eimeria species	FAM	≤37	>37 or absent	
IACV	ATTO	≤32		>32 or absent

*Note: Sample with Ct values below 37 are considered as true positive.
Sample with Ct values above 37 should be considered negative.*

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DNA Diagnostic A/S was established in 1992. DNA Diagnostic A/S is an ISO 13485 certified developer, manufacturer, and worldwide supplier of PCR based in vitro diagnostic kits.

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