

cfhDNA MONODOSE dtec-qPCR Test

Genetic detection of Cell-free human DNA

Edition E09 (06/2018)

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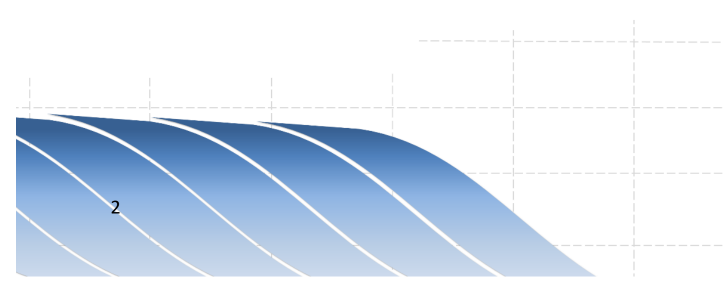


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DESCRIPTION

The **cfhDNA MONODOSE dtec-qPCR Test** contains individual ready-to-use tubes containing all the components needed for **Cell-free human DNA** detection by using qPCR. The target is a multiple-copy gene, 200 copies per genome, with a slow evolutionary rate. Due to the multiple-copy and the high degree of conservation, can be used for quantification of low amounts of circulating cell-free DNA (cfDNA) present in human blood. Cell free DNA is becoming an important clinical analyte for prenatal testing, cancer diagnosis and cancer monitoring. A multiple-copy gene provides an increase in signal amplification, allowing the detection of low DNA amounts.





PRINCIPLE OF THE METHOD

Polymerase chain reaction (PCR) allows the amplification of a target region from a DNA template by using specific oligonucleotides. In real-time PCR (qPCR), the accumulating amplified product can be detected at each cycle with fluorescent dyes. This increasing signal allows to achieve sensitive detection and quantification of pathogens.

KIT CONTENT

TargetSpecies MONODOSE dtec-qPCR (INDIVIDUAL TUBES), contains a dehydrated mixture of specific primers and labelled probe, dNTPs, BSA, polymerase and buffer at optimal concentrations. 24 or 96 rxn

[OPTIONAL] Internal Control, will include additionally in MONODOSE tubes: primers, probe and DNA template for an internal control of PCR.

DNase/RNase free water (GREEN CAP), 1.5 ml

Standard Template (RED CAP), dehydrated target copies for positive control.

Standard buffer (BLACK CAP), exclusive for resuspension of the Standard Template. 500 μl

STORAGE CONDITIONS

All the components of **cfhDNA MONODOSE dtec-qPCR Test** are stable at room temperature for transport. At arrival, if not immediately used, it should be stored at -20 °C. Individual tests are stable for one year (see expiration date on the label).

For **Standard Template** (**RED CAP**) we recommend, once dissolved, store **in an exclusive box** at -20 °C.

cfhDNA MONODOSE dtec-qPCR Test

MATERIALS REQUIRED BUT NOT PROVIDED

- DNA isolation kit (GPSpin extraction/purification kits recommended)
- DNase/RNase free water (to prepare standard curve dilution)
- Sterile pipette tips with filters
- qPCR tubes, strips or plates
- Micropipettes
- Vortex mixer
- Spinner centrifuge
- Cooling block
- Real-time PCR device

WARNINGS AND PRECAUTIONS

- (i) For research use only
- ① To avoid possible misuse, carefully read the handbook.
- ① Proper training is recommended for correct operation of the kit.
- According to good laboratory practices, always wear a suitable lab coat, disposable gloves, and protective goggles.
- ① All the instruments used must been verified and calibrated according to the manufacturer's recommendations.

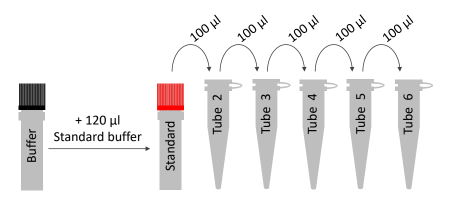
General precautions

- ▲ To prevent contamination of micropipettes, use sterile tips with filters.
- ▲ Micropipettes used to dilute the Standard Template should not be used for other PCR reagents.
- ▲ Extract, store and prepare positive materials (samples, positive controls and PCR products) in a separately laboratory environment.
- ▲ To avoid cross-contamination with the positive control, pipette it after closing reaction tubes with negative control and samples.
- ▲ Work in a row and keep components refrigerated in a cooling block.
- ▲ Protect the primer/probe from prolonged exposure to light.
- ▲ After preparing MONODOSE tubes, run reactions immediately.



PREPARATION OF STANDARD CURVE DILUTION SERIES

- 1) Pipette 900 μ l of **DNAse/RNAse free water** (not provided) into five tubes and label as 2 to 6
- Pulse-spin the Standard Template (RED CAP), reconstitute with 120 μl of Standard buffer (BLACK CAP) and vortex thoroughly, label as num. 1
- 3) Pipette 100 μ l of diluted **Standard Template** (RED CAP), into tube 2
- 4) Vortex thoroughly and pulse-spin
- 5) Change tip and pipette 100 μ l from tube 2 into tube 3
- 6) Vortex thoroughly and pulse-spin
- 7) Repeat steps 5 and 6 with the tubes 4 to 6 to complete serial dilution



Standard curve dilution series	copies/µl	copies in 5 µl
Standard Template (RED CAP)	2 x 10 ⁵	10 ⁶
Tube 2	2 x 10 ⁴	10 ⁵
Tube 3	2 x 10 ³	10 ⁴
Tube 4	2 x 10 ²	10 ³
Tube 5	2 x 10	10 ²
Tube 6	2	10

Pipette 5 μ l of template into each well for the standard curve according to your plate set-up. The final volume in each qPCR reaction well is 20 μ l.

PROTOCOL & AMPLIFICATION REGIME

Add the desired volume of sample ranging from 5 μ l up to a maximum qPCR volume of 20 μ l and, when needed, complete this final volume by adding DNase/RNase free water (GREEN CAP) (i.e., 7 μ l sample + 13 μ l water). Vortex thoroughly and pulse-spin. To determine the sample volume, please take into account the possible presence of inhibitors.

GPS[™] reagents contains BSA and are **compatible with all real-time PCR thermal cyclers, glass capillary or plate based**. Plastic of the *Generic tube* is compatible with: StepOne[™], StepOnePlus[™], ABI 7500 Fast, LightCycler[®] 96, LightCycler[®] Nano, CFX96[™], PikoReal[™] 24-well, DNA Engine[®] systems, MiniOpticon[™] 48-12 and Opticon[®] 2. For other devices, please, transfer the content of the MONODOSE (20 µl) to appropriate tubes.

Take into account that the fluorescent signal must be collected by using the FAM channel for the target. If the internal control is added use the HEX channel.

	Step	Time	Temperature
	Activation	1 min	95 °C
40 Cycles	Denaturation	10 sec	95 °C
	Hybridization / Extension and data collection ¹	1 min	60 °C

1 Fluorogenic signal should be collected during this step by using the **FAM** channel for the target and by using the **HEX** channel for the internal control.



RECOMMENDED REACTION CONTROLS

These qPCR reaction controls are recommended when considering the guidelines of ISO/IEC 17025 Standard. When setting-up your qPCR protocol, select the controls considered better suits your quality system.

Negative Control (Ctrl -): Add 20 μ l of DNase/RNase free water (GREEN CAP) to one tube. Accordingly, this reaction should be negative. A positive result may be considered as a symptom of DNA contamination in the water, making the test inconclusive. Water must be replaced.

Positive Control (Ctrl +): Prepare a standard curve dilution series as described above from the **Standard Template** (RED CAP). Add 5 μ l of the standard template dilution (i.e., 2 x 10² copies/ μ l; Tube 4) to 15 μ l of water. A positive result indicates that qPCR setup is correct and works. If negative, the test should be carefully repeated after checking the thermal protocol.

Matrix Inhibition Control (M-Inh): We recommend to run reactions in parallel to test possible inhibition effects of compounds (inhibitors) present in the sample matrix. Simply, to a reaction tube, add a known amount of **Standard Template** (RED CAP) (i.e., 2 x 10^2 copies/µl) together with the sample.

Reagent	Volume
Standard template dilution (i.e., 2×10^2 copies/µl) ¹	5 μl
DNase/RNase free water (GREEN CAP)	10 µl
Sample	5 µl
FINAL REACTION VOLUME	20 µl

1 Tube 4 of the curve dilutions series obtained from Standard Template (RED CAP)

An optimal result should show a positive signal, equal or higher (same or lower Ct) than these found for the Positive Control alone (tube 4, 10^3 copies). Inhibition may be total (negative result) or partial, observing a considerable increase in the Ct when compared to this of the Standard Template dilution added. If inhibition is observed, a sample dilution to 1/10 may be recommended (if concentration is not close to detection limit). The matrix inhibition control is external, allowing to check the inhibition on the main target of interest.

Extraction Negative Control (ExtCtrl -): Perform an extraction according to your extraction protocol without addition of sample. Add 5 μ l of extraction negative sample to 15 μ l of water. In this case, the test for DNA includes the reagents used in the extraction steps. If positive, when the **Negative Control** is negative, a contamination occurs during the extraction process. Extraction reagents must be discarded.

Extraction Positive Control (ExtCtrl +): Perform an extraction according to your extraction protocol adding the **Standard Template** (RED CAP), or DNA extracted from pure cultures into the first extraction buffer. The positive extraction control would include the effectiveness of the extraction method used. A positive result should be expected. If negative, extraction must be carefully repeated or the extraction method replaced.

INTERPRETATION OF RESULTS

The linear regression of logarithm of the copy number versus Ct gives the constants Y-intercept and slope of the standard curve (equation 1). The number of copies in the sample can be calculated based on the regression (equation 2).

$$Ct = Yinter + Slope \ x \ log(copy \ number)$$
 (1)

$$Copy \, number = 10^{\frac{(Ct-Yinter)}{Slope}}$$
(2)



To obtain the sample quantification directly from the device, the Standard dilution series must be defined in the software of your qPCR device as Standard with the specified copies for each dilution (see PREPARATION OF STANDARD CURVE DILUTION SERIES, page 5). Standard curve can be defined as total copy number or copies/µl.

To refer the values obtained with qPCR to the sample material, please take into account the elution volume after extraction, the sample volume processed and any dilution performed.

Ctrl +	Sample	Ctrl -	ExtCtrl -	IC/M-Inh	Interpretation
-	+/-	+/-	+/-	+/-	Experiment fails
		+	+/-	+/-	PCR reagents contaminated
+	+ + -	+	+/-	Possible contamination at extraction step	
		-	-	+	Positive sample
+	+	+	Negative sample		
	-	-	_	-	PCR inhibition

Key symbols + and - : amplification does or not occur, respectively

To estimate genomic copies in the sample, 200 gene copies obtained from qPCR may be considered as corresponding to a single genome.

VALIDATION METHODS

All batches are calibrated with a standard curve from 10^6 to 10 copies with our Standard Template. Diverse parameters are evaluated: Ct, slope, R² and efficiency. All this information is available in the **Quality Certification** provided to the customer by GPS^M.

CONTACT INFORMATION

For any question and technical support, contact to our address <u>support@geneticpcr.com</u>. For quotes, orders, or new target designs, please contact <u>orders@geneticpcr.com</u>.



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