SHENTEK

Mycoplasma DNA Detection Kit (2G)

User Guide

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Version: A/1 For Research Use Only Product No.: 1509841 Reagents for 50 Reactions

Huzhou Shenke Biotechnology Co., Ltd.

(IMPORTANT: Please read this document carefully before experiment.)

1. Product information

Product description

MycoSHENTEK[®] Mycoplasma DNA Detection Kit (2G) is used, together with MycoSHENTEK[®] Mycoplasma DNA Extraction Kit (2G), to qualitatively determine whether there is mycoplasma, spiroplasma, or acholeplasma contamination in master cell bank, working cell bank, virus seed stock, or cell culture-derived products, etc. Validated according to USP 63, EP 2.6.7 and JP XVIII for mycoplasma detection with a detection limit of 10 CFU/mL.

The detection kit uses real-time PCR technology to detect more than 200 species of Mycoplasma, Spiroplasma and Acholeplasma with high specificity. It has been

validated in a variety of matrices, as well as on non-mycoplasma specie

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For extraction information, please refer to MycoSHENTEK® Mycoplasma DNA

Extraction Kit (2G) User Guide (Product No. 1509840).

Kit contents and storage

WARNING: Please read the Material Safety Data Sheets (MSDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, mask and gloves.

Reagent	Part No.	Quantity	Storage
Internal Control (IC) (2G)*	NNA035	1 bottle	-20°C
Positive Control (PC) (2G)*	Positive Control (PC) (2G)*NNA0391 bottle		-20 C
My qPCR Reaction Buffer (2G)	NNB004	$400 \ \mu L \times 1 \ tube$	-20°C,
My Primer & Probe MIX (2G)	NNC065	75 μ L × 1 tube	protect from light
DNA Dilution Buffer(DDB)	NND001	$1.5 \text{ mL} \times 2 \text{ tubes}$	-20°C

Table 1. Kit components and storage

* Reconstitute lyophilized Internal control (IC) (2G) in 600 μ L of DNA dilution buffer, and lyophilized Positive control (PC) (2G) in 500 μ L DNA dilution buffer, respectively.

The kit components can be stored at appropriate conditions for up to 24 months.

Please check the expiration date on the labels.

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Lightcycler 480 Real-Time PCR System

Required materials not included in the kit

- Nonstick, RNase-free microfuge tubes, 1.5 mL, 2.0 mL
- > PCR 8-well strip tubes with caps or 96-well plates with seals
- > Low retention filter tips 1000 μ L, 100 μ L, 10 μ L
- ➢ 75% Ethanol
- > UNG enzyme (Please check the effective amount for best use.)

Related equipment

- Laminar flow cabinet or biosafety cabinet
- Benchtop microcentrifuge & PCR strip/plate centrifuge

- Microplate and micro test tube shaker
- Vortex mixer
- Real-time PCR System
- Microplats shaker
- Pipettes, 1000 μL, 100 μL, 10 μL

Workflow



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2. Methods

Experiment preparation

- 1. Wear appropriate protective eyewear, mask, clothing and gloves.
- 2. Irradiate the tabletop, pipettes and tubes with UV for 30 minutes, and disinfect with 75% ethanol.
- 3. Thaw the kit completely at 2-8°C or melt on ice, vortex and spin briefly.

qPCR MIX preparation

1. Determine the number of reaction wells according to the number of test samples and control samples with 2 replicates for each sample generally.

Number of reaction wells = (1 Positive control (PC) + 1 No Template control)

(NTC) + 1 Negative control sample (NCS) +N Unknown samples)×2

2. Thaw the kit completely at 2-8°C or melt on ice, then prepare qPCR MIX according to the following table.

Reagents	Volume/reaction	Volume for 30 reaction (includes 10% overage)
My qPCR Reaction Buffer (2G)	8 μL	264 μL
My Primer & Probe MIX (2G)	1.5 μL	49.5 μL
Internal Control (IC) (2G)*	0.5 μL	16.5 μL
Total volume	10µL	330 μL
UNG enzyme	0.1 U	3.3 U

Table 2.	qPCR	MIX	pre	paration
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If IC has been added during sample extraction, an equal volume of DNA dilution

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Table 3. qPCR Reaction MIX preparation in each well

(Positive control)	10 μL qPCR MIX + 20 μL PC
NTC (No Template Control)	10 μ L qPCR MIX + 20 μ L DNA Dilution Buffer
NCS (Negative control sample)	10 μ L qPCR MIX + 20 μ L of extracted NCS
Unknown sample	10 μ L qPCR MIX + 20 μ L of extracted unknown sample

					1		-		-			
NTC	NCS				S1	S1					PC	А
NTC	NCS				S2	S2					PC	В
					S3	S3						С
					S4	S4						D
					S5	S5						Е
					S6	S6						F
					S7	S7						G
					S8	S8						Н
1	2	3	4	5	6	7	8	9	10	11	12	

Table 4. Example of 96-well plate layout

• This example shows one Positive control (PC), one No-template control (NTC),

one Negative control sample (NCS), and eight unknown samples, with duplicate

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qPCR program setting

Please refer to the program setting as follows:

- 1. Run a new method program and select the quantitative PCR assay template.
- 2. Run a new Probe template, and type the name "Mycoplasma-2G". Select FAM in the Reporter Dye drop-down list and select (none) in the Quencher Dye drop-down list. Select VIC in the Reporter Dye drop-down list and select (none) in the Quencher Dye drop-down list, then click OK. Select the detection reference fluorescence as ROX (optional).
- 3. Set PCR cycling conditions:
 - a. Set the cycling reaction volume to $30 \ \mu$ L.
 - b. Set the temperature and the time as following:

	_		
Step	Temp.	Time(mm:sec)	Cycles
UNG treatment	25°C	10:00	1
Activation	95°C	10:00	1
Denaturation	95°C	00:15	
Annealing	62°C	00:30	45
Extension	72°C*	01:30	

Table 5. qPCR	running temperate	ure and time
- 1	0 1	

*Instrument will read the fluorescence signal during this step.

Result analysis

The following instructions apply only to the 7500 instrument with SDS v1.4.

1. Select the Results tab, then click Amplification Plot. In the Analysis Settings

window, enter the following settings:

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by clicking on the Task Column drop-down list and then click

a. NTC wells: target DNA detector task = NTC.

b. NCS, unknown samples wells: target DNA detector task = Unknown.

3. Select File >> Export >> Results. In the Save as type drop-down list, select

Results Export Files, then click Save.

NOTE: If you use a different instrument or software, refer to the applicable instrument or software documentation. Usually real-time PCR instrument software automatically export the data report.

■ Acceptance Criteria

1. Guidance for the control samples as in the following table.

Quality control	FAM	VIC	
samples			
	Ct≥40.00 or absence of specific	Ct<35.00 and normal	
NTC	amplification for duplicate runs	amplification for duplicate runs	
	Ct≥40.00 or absence of specific	Ct<35.00 and normal	
NCS	amplification for duplicate runs	amplification for duplicate runs	
	Ct<35.00 and normal	Ct<35.00 and normal	
PC	amplification for duplicate runs	amplification for duplicate runs	

Table 6. Control sample result analysis

* The QC sample analysis shall be considered in relationship to LOD (Limit of

Detection) based on method validation data.

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able 7 Onknown sample result a

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amplification	amplification for duplicate runs	and presence of PCR
	amprineation for duplicate runs	inhibitors
	Ct<40.00 and normal	Negative
Ct≥40.00 or	amplification for duplicate runs	negative
non-specific	Ct>10.00 or observes of gradifie	Not conclusive, and
amplification	Ct≥40.00 or absence of specific amplification for duplicate runs	presence of PCR
	amprineation for duplicate runs	inhibitors

* If the VIC signal is inhibited, it is necessary to repeat the sample preparation or appropriately remove the inhibitors during the sample preparation and repeat the assay.

Note: In an event that the sample is special, or some abnormalities occur, and

results difficult to determine, please contact us for technical support.

Effective date: 03 Jul. 2024

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