

SinaProbe HS-qPCR Master Mix, 2X

REF MM2161

Σ 1250μl/100 reaction



Wet or Dry Ice

RUO

Store at: -20°C

(not more than 50 thawing-freezing cycles)

Component

SinaProbe HS-qPCR Master Mix, 2X	1250μl
----------------------------------	--------

Description

SinaProbe HS-qPCR Master Mix, (2x) is developed for quantitative real-time PCR with fluorescent probes. It includes all the components necessary for PCR (Highly processive recombinant HS-Taq DNA polymerase, Deoxy nucleoside triphosphate mix, PCR buffer, Mg²⁺, ROX dye except for DNA template, primers and probe). The mix is optimized for consistent and efficient real-time hot-start PCR of genomic, plasmid and viral DNA samples. It is supplemented with additives that increase half-life and processivity of HS-Taq DNA polymerase by enhancing its stability during PCR. SinaProbe HS-qPCR Master Mix, 2x does not contain substances affecting primer annealing temperature and characteristics of template melting. DNA polymerase included in the SinaProbe HS-qPCR Master Mix, 2x is inactive at room temperature, and its activation requires preheating of the reaction solution at 95°C for 5min. **The master mix is ideally compatible for all PCR platforms/devices as well as platform using ROX passive dye as a reference guide: Life Technologies (ABI) 7000, 7300, 7700, 7900HT, StepOne Plus.**

(PMH-047-00/01) (1)

SinaProbe HS-qPCR Master Mix, 2x contains:

100mM Tris-HCl (pH 8.5 at 25°C), 100mM KCl, 0.4mM each deoxy nucleoside triphosphate, 10mM MgCl₂, 0.1U/μl HS-Taq DNA polymerase, 0.025% Tween 20, stabilizers of HS-Taq DNA polymerase, 0.9μM ROX fluorescent dye.

Applications:

- Real-time PCR with fluorescently labeled probes
- Conventional PCR
- High-throughput PCR
- Multiplex PCR
- Genotyping

Benefits of use

- The enzyme with hot start capability enhances reaction specificity
- Activation of HS-Taq DNA polymerase requires not more than 5 min heating
- High selectivity and reaction yield
- Reduced preparation time
- Low chance of contamination during preparation of PCR solution
- Standardized conditions of the same-type reactions (reduce pipetting error during mixing PCR components in a series of experiments)
- Minimized efforts

Limits of use

- Not recommended to use for real-time PCR with intercalating dyes.

Reaction mixes features

- Recombinant HS-Taq DNA polymerase has 5'→3' DNA-dependent polymerase activity and 5'→3' exonuclease activity.
- Allow normalization to ROX reference dye (The presence of ROX does not affect the course of PCR and shift in fluorescence signal in case if the mix is used with other PCR platforms).
- prevents re-amplification of extraneous PCR products

(2)

Recommended qPCR reaction mix:

1. Unfreeze the reaction mixture and stir gently.
2. Add the following components into thin-well PCR tubes considering the final volume of a reaction mixture equal to 25µl:

Component	Volume	Final concentration
SinaProbe HS-qPCR Master Mix, 2x	12.5	1X
Forward primer	variable	0.1 – 600nM
Reverse primer	variable	0.1 – 600nM
Probe	variable	0.1–300nM
DNA template	Variable	1pg – 1µg
Sterile water	up to 25µl	-

Recommended qPCR cycles:





Step	Temperature, °C	Incubation time	Number of cycles
Preliminary denaturation	95	5-7 min	1
Denaturation	95	15 sec	25- 50
Annealing	50- 68	10-30 sec	
Elongation	58- 72	30-60 sec	

Or:


Step	Temperature, °C	Incubation time	Number of cycles
Preliminary denaturation	95	5-7 min	1
Denaturation	95	15 sec	30- 50
Annealing/ Elongation	50- 68	1min	




(3)

Signs

Signs	Definitions
	For Research Use Only
	Name and address of the manufacturer of the product
	Product technical code
	Product shipping conditions



 Unit 9, Rouyesh building, Science and Technology Park,
Tarbiat Modares University, Pajouhesh Blvd, Tehran, Iran

 +982191082111
 hi@sinaclon.com
 www.sinaclon.com

(4)