Genious 2X SYBR Green Fast qPCR Mix

Cat. No.: RK21204



Product components

Components	Component number	Size-1	Size-2
		5 mL	25 mL
Genious 2X SYBR Green Fast qPCR Mix *	RM21204	5 × 1 mL	25 × 1 mL
50X ROX Reference Dye I	RM21465	220 μL	5 × 220 μL
50X ROX Reference Dye II	RM21466	220 μL	5 × 220 μL

^{*} Including ABclonal Genious Hot Start Taq DNA polymerase, Mg²⁺, dNTPs, SYBR* Green I, etc.

Product Description

Real-time Quantitative PCR is a technique using DNA double-stranded dyes, usually SYBR® Green I, to quantitative detect the initial amount of DNA by amplification. ABclonal Genious 2X SYBR Green Fast qPCR Mix is a fluorescence reagent for qPCR reactions using the SYBR® Green I. This product provides real-time data on DNA amplification during PCR by performing quantitative fluorescence signal detection on SYBR®/FAM channels. This product uses the antibody-based hot-start Taq enzyme for amplification, which greatly improves the specificity of the product while ensuring the amplification effect. At the same time, by optimizing the qPCR Mix Buffer system, the product is suitable for multiple species, providing a powerful tool for multidisciplinary experimental needs.

This product is a 2X pre-mixed enzyme, contains all the components required for qPCR except primers and templates, providing great convenience for experimental manipulation.

Instruments

Rox types	qPCR Machines
No ROX	Bio-Rad iCycler serious, Roche Light Cycler serious, Qiagen/Corbett serious and others
ROX Reference Dye I	ABI 7000/7300/7700/7900, ABI StepOne/StepOnePlus, Eppendorf and others
ROX Reference Dye II	ABI 7500, ABI ViiATM7, ABI QuantaStudio serious, Stratagene serious, Corbett Rotor Gene 3000 and others

Note: The ROX Reference Dye is different for different instruments. For adding or not, please refer to the above models.

Storage

- 20°C for long term storage and 4°C for a short period and Light protection is demanded.

Materials Required

- 1. EP tubes, PCR tubes and other related materials.
- 2. qPCR specific primers and templates.
- 3. qPCR plates and seal membrane.

Usage Notes

- 1. Before using Genious 2X SYBR Green Fast qPCR Mix, please make sure that the mix is thawed completely and then placed it on ice for use.
- 2. Mix other qPCR components with Mix thoroughly and gently by pipetting or vortexing. After usage, place the mix back to -20°C for long time storage or 4°C for short period usage.
- 3. Genious 2X SYBR Green Fast qPCR Mix contains Taq polymerase, all operation should be performed on ice.
- 4. Genious 2X SYBR Green Fast qPCR Mix is equipped with a specially designed ROX reference dye. ROX reference dye is added according to the qPCR instrument model.
- 5. To avoid contamination, pipette tips with filters is suggested.
- 6. To guarantee better qPCR results, DNA template in good quality is suggested.

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Protocol

Before Use

- 1. Specificity of primers should be checked and a final concentration of 0.2 μM is suitable for most of primers.
- 2. The length of amplification products is usually range from 70 bp to 200 bp.
- 3. Dilute the template in gradient.
- 4. Add 1 pg-50 ng DNA as PCR templates and a "No Template Control sample" is suggested.
- 5. To ensure the confidence of experiment, at least 2 repeats of each samples is suggested.

Experiment procedure

1. Prepare the following reaction systems on ice:

Components	Volume
Genious 2X SYBR Green Fast qPCR Mix	10 μL
Forward Primer (10 µM)	0.4 μL
Reverse Primer (10 µM)	0.4 μL
gDNA or cDNA (<50 ng)	2 μL
ROX	0.4 μL
ddH_2O	to 20 µL

- (1) Dissolve Genious 2X SYBR Green Fast qPCR Mix at room temperature then placed it on ice for further usage. Before using, agitate the Mix thoroughly and centrifuge to collect all solution.
- (2) Calculate the amount of mix need, generally a 10% extra amount is suggested.
- (3) Dispense solution in sterile PCR or EP tubes in case of any contamination.
- (4) Add all components listed in above table, agitate the tubes gently to mix thoroughly (avoid bubbles) and centrifuge it.
- (5) Dispense the reaction solution into qPCR plates and seal the plates with optical membrane.
- (6) 2500 rpm centrifuge the qPCR plates to collect all solution.
- 2. Program qPCR reaction as follows:

Stage 1	Denaturation	Reps: 1	95 ℃	3 min
Stage 2	Cylces	Reps: 40-45	95 ℃	5 s
			60 ℃	30-34 s *
Stage 3	Melt Curve	Reps: 1	default	

^{*} Confirm there is a signal collection step after each extending step. The extending time is varied according to different machines: 30 s for StepOne Plus, 31 s for 7300 and 34 s for 7500.

- 1. Draw a standard curve according to Ct values of endogenous gene. The value of R2 should be more than 0.98 and the slope of curve should be in the range of -3 to -3.5 which means the PCR amplification efficiency is in the range of 90% to 120%.
- 2. The standard deviation (STD) of Ct values should be less than 0.2 and the variation of Ct value for different experiment should be less than 0.5 (the threshold value of different experiments should be same when comparing Ct values).
- 3. The single melt curve indicate the no non-specific amplification products or primer dimmers, and the Tm value in melt curve is usually in the range of 80 to 95°C.

Troubleshooting

Melt Curve Show Multiple Peaks

- a. Primer Design: Design the primer following basic primer design protocols.
- b. Primer Concentration Too High: lower down the concentration of primers.

Unusual Amplification Curves

Genious 2X SYBR Green Fast qPCR Mix



- a. Amplification Curve Not Smooth: Too low amplification signal, increase the template input and make sure the qPCR Mix is stored properly.
- b. Inconsistent Amplification Curve: Bubbles causes abnormal qPCR results, centrifuge the plate prior to running it.
- c. Abnormal Amplification Curves: the default baseline value of machine is set to be from 3 to 15, the baseline setting can be changed according actual amplification conditions. Besides, the degradation of template may affect the curve.

No Amplification Curves after Reaction

- a. Not Enough PCR Cycles: the PCR cycle number is usually set to be 40. It should be noted a higher cycle number may increase the background signal.
- b. Primer Degradation: Use electrophoresis to confirm the integrity of primers.
- c. Confirm the Signal Collection Step: the signal collection step are usually set to be after the annealing-extending step for two-step qPCR and after extending step for Three-step qCPR.
- d. Template Input Too Low: Increase template concentration or add extra repetition.
- e. Template Degradation: Use freshly prepared template (Use electrophoresis to confirm integrity of template).
- f. Not Enough Initial Denaturation Time: Genious 2X SYBR Green Fast qPCR Mix uses Hot-Start Taq polymerase, the pre-denaturation time should be at least 3 min.

Ct Value Too Late

- a. Low Amplification Efficiency: Optimize reaction condition or change primer.
- b. Template Input Too Low: Increase template concentration or add extra repeat.
- c. Template Degradation: Use freshly prepared template (Use electrophoresis to confirm Integrity of template).
- d. Too Long PCR Products: The length of amplification products is usually in the range of 70 bp-200 bp.
- e. PCR Inhabitation Reagent: use new template or dilute the template.
- f. Too Short Pre-denaturation Time: Genious 2X SYBR Green Fast qPCR Mix contains Hot-Start Taq polymerase, the pre-denaturation time should be at least 3 min.

NTC Shows Amplification

- a. Contamination: Use sterile water to conduct experiment and the all operation is suggested to be done in clean room to avoid aerosol contamination.
- b. Non-Specific PCR Products: analyze with melt curve.

Inconsistent Results

- a. Inconsistent Sample Added: Use proper pipetting techniques.
- b. Inconsistent Temperature in qPCR Machine: ensure periodic machine calibration.
- c. Template Concentration Too Low: the lower template input, the poorer qPCR result is. Increase the template concentration.
- d. Inconsistent Threshold Set: when comparing the qPCR results in different plates, make sure the threshold value of each experiments is same.

Related Products

Name	Catolog	Size
ABScript Neo RT Master Mix for qPCR	RK20432	10 RXN / 100 RXN
ABScript Neo RT Master Mix for qPCR with gDNA Remover	RK20433	10 RXN / 100 RXN
ABScript II cDNA First-Strand Synthesis Kit	RK20400	50 RXN / 100 RXN

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