5X PrimePath™ Probe qPCR Kit, No ROX, GPR Protocol-At-A-Glance

I. Introduction

5X PrimePathTM Probe qPCR Kit, No ROX, GPR (Cat No. 638347 & 638349) enables the user to perform accurate qPCR on DNA samples. The highly concentrated master mix allows for flexibility in handling larger sample volumes. The entire protocol from setup to detection takes less than one hour, enabling fast results.

II. Required Materials

This protocol applies to the following Takara Bio products:

• 5X PrimePath Probe qPCR Kit, No ROX, GPR (Cat No. 638347 & 638349)

Additional materials required:

- Primers and probes
- Micropipette and tips (with hydrophobic filters)
- Vortex mixer
- Benchtop centrifuge for tubes or plates
- 1.5 ml Eppendorf tubes, 200 μl PCR tubes, or 200 μl PCR plates for sample preparation
- Tubes or plates for real-time PCR
- A real-time PCR machine (real-time systems that do not require ROX reference dye, such as CFX96 Real-Time PCR Detection System, Bio-Rad, etc.)

III. Protocol

- 1. Mix the tube of 5X PrimePath Probe qPCR Mix by inverting several times then spin down for 10 sec; the mix may appear cloudy after storage, but this does not affect its performance.
- 2. On ice and using the table below, combine everything for the qPCR reaction mix EXCEPT the DNA sample for all planned reactions, plus 10% of the total reaction mix volume for overage.

NOTES:

- If the mixture is used immediately, it can be combined at room temperature (no ice).
- Do not add the DNA sample at this time. It is listed in the table to indicate what the final mixture will
 contain.

qPCR reaction mix (per 25 μl* reaction)

Reagent	Final	Singleplex	Multiplex
	concentration		(N [†] targets)
5X PrimePath Probe qPCR Mix, No ROX, GPR	1X	5 µl	5 µl
PCR Forward Primer (10 μM)	0.2 µM	0.5 µl	N^{\dagger} x 0.5 μ l
PCR Reverse Primer (10 µM)	0.2 µM	0.5 µl	N^{\dagger} x 0.5 μ l
Probe (10 µM)	0.2 µM	0.5 µl	N^{\dagger} x 0.5 μ l
DNA sample [‡]	_	2 µl§	2 µI [§]
RNase Free H₂O	_	Up to 25 μl**	Up to 25 μl**
Total volume per reaction		25 µl	25 µl

^{*}Mix, as written, can also be used per 20 µl reaction.

[†]Where *N* represents the total number of targets of interest. The total reaction volume will still be 25 µl, while 0.5 µl of each primer and probe will be added per target.

[‡]Do not add to the reaction mix; this will be added to the mix in Step 4.

[§]Most common sample volume of Takara Bio PCR products. It is possible to add more; however, depending on the chemicals contained in the sample solution, higher sample volume may have adverse effects.

^{**}Adjust the reaction volume according to the recommendations for the real-time PCR instrument used.

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- 3. Add [25 µl Volume of DNA Sample] of the qPCR reaction mixture into a PCR tube or a 96-well PCR plate.
- 4. Dispense the volume of DNA sample into the PCR tube or plate well. Spin down for 10 sec.
- 5. Set up the thermal cycler and run the assay using the reaction conditions below:

95°C 20 sec 40 cycles: ____ 95°C 1 sec 60°C 20 sec

NOTE: Please follow the instruction manual of the real-time qPCR machine used. If the default setting does not work, perform manual analysis per the instruction manual.

6. After the reaction is complete, check the amplification curve. Confirm that the analytical parameter is correct and that the Ct value has been calculated.

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